1. Introduction

Alcohol consumption leads to several cognitive deficits, either temporary, due to the acute intoxication or permanent, following a chronic use. According to the dual process theory (Wiers et al., 2007; Stacy and Wiers, 2010), chronic alcohol use induces a sensitization of the appetitive system in combination with an increase of the salience of alcohol-related cues (Robinson and Berridge, 1993) as well as a progressive decline of the regulatory executive system (Parsons, 1998), as evidenced particularly by a lack of executive resources required to inhibit inappropriately salient responses to alcohol. Indeed, several studies reported that alcohol abusers show an attentional bias (Field et al., 2007; Townshend and Duka, 2001) as well as automatic approach behaviors towards alcohol-related cues (Field et al., 2008; Wiers et al., 2007). These two processes (i.e., altered inhibition capacities and increased attentional bias for alcohol) seem to be involved in the development of alcohol-seeking behaviors (Fadardi and Cox, 2008; Nigg et al., 2006) and in alcohol relapse (Field and Cox, 2008).

Inhibition capacities are often studied with a Go/No-go task. In this task, participants are required to respond by pressing a key when frequent Go stimuli are presented and to withhold their responses for infrequent No-go stimuli. This paradigm typically gives rise to larger N200 and P300 amplitudes during No-go trials as compared to Go trials (Bruin and Wijers, 2002; Smith et al., 2008). While the frontal N200 would reflect a general control process during No-go trials (Nieuwenhuis et al., 2003), there are debates over its possible role in the inhibition of a premature response (e.g., Falkenstein et al., 1999; Lavric et al., 2004), in the conflict monitoring between execution and inhibition of a single response (Nieuwenhuis et al., 2003; Yeung and Cohen, 2006) or in neither of these (Smith et al., 2007). The fronto-central P300 would index the late decision process to inhibit the motor response (Smith et al., 2010; Gajewski & Falkenstein, 2013; Huster et al., 2013).

In alcohol-dependent patients, reduced N200 peak amplitude for Go as well as No-go trials has been observed in comparison to controls in an equiprobable Go/No-go task (Pandey et al., 2012). Moreover, in line with the lack of inhibition observed following chronic alcohol consumption using behavioral tasks (Fillmore, 2003; Noël et al., 2010), several studies have found smaller P300 amplitude during No-go trials in alcohol-dependent patients (Cohen et al., 1997; Kamarajan et al., 2005). Similar to alcohol-dependent patients, heavy social drinkers exhibit a reduction of the No-go P300 amplitude compared to light drinkers (Oddy and Barry, 2009). Lack of inhibitory skills have also been associated with compensatory neuronal mechanisms allowing drinkers to achieve performance levels similar to those in controls with increased No-go P3 amplitude and higher prefrontal activation in...
binge drinkers, as compared to controls, when successful inhibition responses were analyzed (López-Caneda et al., 2012).

Interestingly, many studies have reported specific ERP patterns in chronic alcohol users confronted to alcohol-related cues. Most studies conducted on individuals with low sensitivity to alcohol, on heavy drinkers (i.e., more than 4 drinks on any day or 14 per week and usually classified as score higher than 11 at the Alcohol Use Disorders Identification Test, AUDIT; Fleming et al., 1991) and on alcohol-dependent patients have shown larger P300 amplitudes and shorter P300 latencies elicited by alcohol-related pictures or alcohol-related words as compared to neutral stimuli (Bartholow et al., 2007, 2010; Hansen et al., 2003; Herrmann et al., 2000, 2001; Namoong et al., 2004). These results suggest that alcohol-related cues are more salient for chronic alcohol users. In contrast, college binge drinkers showed similar P300 amplitudes elicited by alcohol-related pictures during the first stage of development of an abusive use of alcohol as compared to non-binge drinkers (Petit et al., 2012b). However, greater P100 amplitudes to alcoholic stimuli as compared to neutral stimuli in young social drinkers were found in this study, suggesting enhanced perceptual processing toward alcohol cues.

Taken together, these results indicate that poor inhibition responses and automatic processes of alcohol-related cues in heavy drinkers could be reflected by modulations of the N200 and P300 amplitudes. However, only a few studies have investigated the effects of inhibition deficits towards alcohol-related cues, most researches to date mainly focused on the appetitive or attractive value of alcohol-related cues. Nevertheless, from a clinical point of view, being able to inhibit the consumption of an alcoholic beverage when exposed to alcohol-related cues is very important for alcohol-dependent patients to maintain abstinence, and also for heavy drinkers who wish to regulate their consumption. Indeed, it could be assumed, similarly to alcohol-dependent patients, that heavy drinkers are less able to inhibit a response toward alcohol-related cues that capture attention automatically. However, two studies did not observe such a deficit in heavy social drinkers (Adams et al., 2013; Nederkoorn et al., 2009), whereas detoxified alcohol-dependent patients showed impaired response inhibition towards alcohol-related cues after 19.7 ± 2.7 days of abstinence (Noël et al., 2007). However, several ERP studies revealed early cerebral dysfunctions related to alcohol consumption before any detectable behavioral impairment in social drinkers (Bijl et al., 2005; Nichols and Martin, 1996; Oddy and Barry, 2009), and in binge drinkers (López-Caneda et al., 2013, 2014; Maurage et al., 2009, 2012; Watson et al., 2014). To our knowledge, only one electrophysiological study assessed the inhibition abilities toward alcohol-related information (Petit et al., 2012a). The authors used a letter Go/No-go task in which the letters were superimposed on a background picture displaying three different emotional contexts (neutral, alcohol-related, and non-alcohol-related) among heavy and light drinkers. Results showed higher false alarm (FA) errors (i.e., erroneously pressing the key) and a delayed P300 latency in heavy drinkers during No-go trials associated with an alcohol-related context as compared to a non-alcohol context. This suggests less attentional allocation to the Go/No-go task in favor of alcohol-related background information.

The aim of the present study was to assess inhibition capacity and alcohol-cue reactivity among heavy and light drinkers with a classic letter Go/No-go task. In addition, the participants performed a modified Go/No-go task in which they had to inhibit a response towards neutral or alcohol-related pictures. We first hypothesized that heavy drinkers, by comparison to light drinkers, would exhibit inhibition deficits revealed by more FA during No-go trials involving letters as well as reduced N200 and P300 amplitudes (Kamarajan et al., 2005; Montgomery et al., 2013; Pandey et al., 2012). Second, in the modified Go/No-go task, we expected reduced inhibition performances towards alcohol-related cues (Noël et al., 2007; Petit et al., 2012a) combined with delayed P300 latencies (Petit et al., 2012a) in heavy drinkers as compared to light drinkers. Third, we assumed that heavy drinkers would display larger P300 amplitudes than light drinkers to alcohol-related pictures, reflecting higher emotional salience of these stimuli for alcohol abusers (Herrmann et al., 2001). As the present study assesses response inhibition towards alcohol stimuli, approach and avoidance to alcohol, subjective craving and impulsivity were assessed.

2. Materials and methods

2.1. Participants

Participants were recruited at the Faculty of Psychology of the University of Liège (Belgium) through interviews and online advertisements. They were pre-screened with the Alcohol Use Disorder Identification Test (AUDIT) and either discarded or assigned into one of two groups according to their reported consumption of alcoholic beverages: light drinkers (AUDIT score ≤ 6; see Saunders et al., 1993) or heavy drinkers (AUDIT scores ≥ 11, see Fleming et al., 1991). Participants whose AUDIT scores were comprised between the two limits were excluded from the study as well as alcohol abstainers. Participants included had no major medical problem, no history of central nervous system disorders (including epilepsy and brain trauma), no regular use of drugs other than alcohol and nicotine, no uncorrected serious vision issue (e.g. color blindness), and had French as native language. The final sample comprised 30 participants aged between 18 and 33 years old. The two groups (N = 15) were matched on age and gender (eight females in each group). The groups’ characteristics are shown in Table 1. No differences between groups were found regarding levels of anxiety and depression, which is of importance as some studies emphasized relationships between inhibitory control and negative affect states (Kaiser et al., 2003). The study was approved by the Ethic Committee of the Faculty of Psychology of the University of Liège and all participants gave their signed informed consent before taking part in the experiment.

2.2. Questionnaires

2.2.1. Alcohol consumption

Alcohol consumption was assessed with a self-reported measure based on the timeline follow-back method (Sobell and Sobell, 1990). Participants reported how many standard alcohol drinks (i.e., 10 g) they had consumed during the previous week (Wiers et al., 1997). Furthermore, the number of days they drank more than six drinks of alcohol in one occasion during the past 2 weeks was recorded (Wiers et al., 1997).

Table 1

<table>
<thead>
<tr>
<th>Measure</th>
<th>Description</th>
<th>Light drinkers</th>
<th>Heavy drinkers</th>
<th>t</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td></td>
<td>21.2 (3.9)</td>
<td>21.7 (1.8)</td>
<td>0.5</td>
<td>0.64</td>
</tr>
<tr>
<td>Educational level</td>
<td>14.1 (2.9)</td>
<td>14.8 (1.8)</td>
<td>0.8</td>
<td>0.41</td>
<td></td>
</tr>
<tr>
<td>AUDIT score</td>
<td>4.0 (2.1)</td>
<td>19.5 (6.1)</td>
<td>9.3</td>
<td>&lt;0.0001</td>
<td></td>
</tr>
<tr>
<td>≥6 units in one occasion/last 2 weeks</td>
<td>0.1 (0.3)</td>
<td>1.5 (1.3)</td>
<td>3.9</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Mean number of consumption per week</td>
<td>2.3 (2.6)</td>
<td>18.5 (15.1)</td>
<td>4.1</td>
<td>&lt;0.001</td>
<td></td>
</tr>
</tbody>
</table>

The table shows the mean (standard deviations) for each demographic, psychological and alcohol consumption characteristics in heavy and light drinkers, t value of the Student’s test comparison between group and the associate probability.
2.2.2. Alcohol use disorder identification test

The AUDIT (Saunders et al., 1993) includes 10 multiple-choice items measuring alcohol consumption, drinking behaviors and alcohol-related problems. This measure was used to assign the participants into the two groups (light and heavy drinkers, see above).

2.2.3. Impulsiveness

The Barrat Impulsiveness Scale-11 (BIS-11) is a 30-item self-report questionnaire measuring the behavioral construct of impulsiveness. It consists of three subscales: motor impulsiveness, cognitive impulsiveness and non-planning impulsiveness. Each item consisted of a 4-point Likert scale from “rarely/never” (1) to “almost always” (4).

2.2.4. Approach and avoidance

The Approach and Avoidance of Alcohol Questionnaire (AAAQ, McEvoy et al., 2004) is a 14-item self-report questionnaire developed to assess the compulsive urge to drink alcohol and the avoidance towards alcohol. Participants had to indicate the extent to which they agreed or not with each item on a 9-point Likert scale ranging from “not at all” (0) to “very strongly” (8).

2.2.5. Depression

The Beck Depression Inventory (BDI-II, Beck et al., 1996) is a 21-item self-report questionnaire used to screen for depressive symptoms experienced during the past 2 weeks. Extensive reliability and validity data have been reported (Beck et al., 1996).

2.2.6. Anxiety

The State-Trait Anxiety Inventory for adults form Y (STAI form Y, Spielberger, 1983) is a 40-item self-report questionnaire used to assess the state anxiety (how people feel now) (20 items) and the trait anxiety (how people usually feel, independently of the situation) (20 items).

2.3. Procedure

Participants were tested in a sound-attenuated room. They filled in the four questionnaires (AAAQ, BDI, BIS, STAI) before taking part in two different Go/No-go tasks.

2.3.1. Letter Go/No-go task

This task consisted of one training session without electroencephalographic (EEG) recording (20 trials, 15 Go trials and 5 No-go trials) and one test session (100 trials, 75 Go trials and 25 No-go trials) during which ERPs were recorded. On each trial, a fixation cross was presented on the center of the screen for 1200 ms followed by a letter for 500 ms. The Go frequent stimulus was the capital letter “O” (size of 500 × 400 mm) and the No-go infrequent stimulus was the letter “E.” The letters were displayed on a 17-in. monitor, and the participants were seated 1 m from the screen. They were instructed to press the spacebar, with their dominant hand, as quickly and accurately as possible.

2.3.2. Alcohol modified Go/No-go task

The procedure was exactly the same as in the letter Go/No-go task except that the stimuli were 25 colored pictures of alcohol drinks and 25 colored pictures of neutral objects (100 trials, 75 Go trials and 25 No-go trials). The alcohol pictures included bottles of beer and wine, glasses of beer, wine, liquor and cocktails, and the neutral object pictures included objects typically used in the office, such as pen, ruler, stapler, computer mouse, and USB stick (Kreusch et al., 2013). Each individual picture was presented three times during Go trials. We used two versions of this task: one in which the alcohol-related pictures were assigned to Go trial and neutral pictures to No-go trial, and one version in which the stimuli had the reverse status. The order of the two versions was counterbalanced across participants. The pictures were displayed fully on a 17-in. monitor, and the participants were placed 1 m before the screen.

2.4. EEG recording

ERG recording, stimulus presentation and waveform analyses were performed with an ANT system (eeprobe, eevoke and eemagine EEG, respectively). EEG activity was recorded at nine sites according to the 10–20 System (Fp1, Fp2, F3, Fz, F4, C3, Cz, C4 and Pz), using Ag/AgCl electrodes, earlobes for reference and forehead for ground. All sites were cleaned with acetone and abraded to maintain impedances below 10 kΩ. Electrooculogram (EOG) activity was recorded from above the left eye. If 50% or more of the epochs of a participant contained artifacts, this participant was excluded from ERP analyses.

As a result, four participants were rejected from ERP analyses, two heavy drinkers and two light drinkers. Then, approximately 15% of trials were contaminated by eye movements or muscular artifacts and were discarded. A two-way ANOVA with Stimulus type (alcohol-related vs. neutral) as within-subject variable and Group (light vs. heavy drinkers) as between-subject variable showed that the number of rejected trials was similar in each group and condition (Stimulus type: F(1,50) = 2.4, p = 0.12; Group: F(1,50) = 2.3; p = 0.13, Stimulus type × Group: F(1,50) = 0.2, p = 0.65). The EEG was amplified by battery-operated amplifiers with a gain of 50,000 and a band pass of 0.16–30 Hz (Advanced Neuro Technology-ANT Ltd., Enschede, the Netherlands). The EEG was digitized at 256 samples for 900 ms with a 100 ms prestimulus baseline. Trials on which the EEG or EOG exceeded 50 μV were rejected automatically by the system. N200 and P300 components were defined as the maximum negative or positive peaks within the latency windows of 200–300 and 270–650 ms, respectively, from the onset of each stimulus. The detection of the peaks was performed by individual visual inspection with no automatic detection procedure. After considering artifacts and FA from the initial 25 No-go trials, approximately 21 trials were included on average for each subject (mean: 21.74, SD: 1.05).

2.5. Statistical analyses

We carried out separate analyses for the letter Go/No-go and the alcohol modified Go/No-go tasks. For the letter Go/No-go task, the percentage of FA on No-go trials and the average reaction times (RT) on Go trials of the two groups (heavy and light drinkers) were analyzed with Student’s t-test. N200 and P300 amplitudes and latencies were analyzed with four separate three-way mixed ANOVAs including Group (heavy vs. light drinkers), Trial type (Go and No-go) and Electrodes (F3-Fz-F4-C3-Cz-C4-Pz; Oddy and Barry, 2009; Petit et al., 2012a) as factors, with Trial type and Electrode as within-subject variables.

To examine whether the level of alcohol avoidance had an influence on behavioral performance and ERP components within the modify alcohol Go/No-go task, we additionally divided participants (n = 26) into two groups, split at the median (2.2) of the alcohol avoidance mean score: high alcohol avoiders (n = 13, 7 heavy drinkers and 6 light drinkers) and low alcohol avoiders (n = 13, 6 heavy drinkers and 7 light drinkers). The average percentage of FA on No-go trials and the average RT on Go trials were analyzed with three-way mixed ANOVAs including Stimulus type (alcohol-related vs. neutral picture) as within subject factor and Group (heavy vs. light drinker) and Alcohol avoidance (high alcohol avoiders and low alcohol) as between subject factors. Five-way mixed ANOVAs with Group (heavy vs. light drinkers), Alcohol avoidance (high alcohol vs. low alcohol avoiders), Stimulus type (alcohol-related vs. neutral picture), Trial type (Go and No-go), and Electrode (F3-Fz-F4-C3-Cz-C4-Pz; Oddy and Barry, 2009; Petit et al., 2012a) were conducted separately with N200 and P300 amplitudes and latencies as dependent variables.

As appropriate, Tukey’s post hoc tests and Greenhouse–Geisser corrections were applied. As the electrode effect was not the main
focus of this study, only Group × Electrode interactions are reported. Pearson’s correlations were used to test the hypothesis that N200 and P300 amplitudes and latencies (mean amplitudes averaged across seven recorded fronto-central scalp sites: F3, Fz, F4, C3, Cz, C4 and Pz) (Euser and Franken, 2012) in No-go trials for alcohol-related cues (Petit et al., 2012a) are associated with the percentage of FAs to alcohol-related cues, AUDIT scores, BIS-11 scores and the Obsessed/Compelled craving for alcohol. All statistical analyses were conducted with Statistica (version 10) for Windows.

3. Results

3.1. Behavioral data

Only RTs of trials with correct responses were analyzed (i.e., 99% in the classical Go/No-go task and 97% in the alcohol modified Go/No-go task). The t-tests computed on the RTs and the percentage of FAs in the letter Go/No-go task revealed no significant difference between heavy and light drinkers on RTs (t(28) = 1.4, p = 0.17), but a significant difference on FAs (t(28) = 2.24, p = 0.03). Heavy drinkers made more FA than light drinkers (mean = 5.6%, SD = 5.4 in heavy drinkers and mean = 2.1%, SD = 2.5 in light drinkers).

For the modified alcohol Go/No-go task, the three-way mixed ANOVA on RTs showed a significant main effect of Stimulus type (F(1,26) = 23.98, p < 0.001). Post-hoc analysis showed significant faster responses for alcohol-related pictures as compared to neutral ones (p < 0.001) for the Go trials, independently of the groups. The main effects of Group (F(1,26) = 2.79, p = 0.10) and Alcohol avoidance (F(1,26) = 0.005, p = 0.94) were not significant. There was no significant interaction. The three-way mixed ANOVA with percentage of FAs as dependent variable showed a significant main effect of Alcohol avoidance (F(1,26) = 6.25, p = 0.02). Post-hoc analysis revealed a significantly higher percentage of FAs in high alcohol avoiders compared to low alcohol avoiders (p = 0.01). No significant main effect of Group (F(1,26) = 0.06, p = 0.80) or Stimulus type was observed (F(1,26) = 0.65, p = 0.43). However, there was a significant three-way Stimulus type × Group × Alcohol avoidance interaction (F(1,26) = 4.38, p = 0.046). Heavy drinkers categorized as higher alcohol avoiders showed a non-significant tendency to perform more FAs to alcohol-related cues as compared to heavy drinkers classified as lower alcohol avoiders (p = 0.09). No such tendency was observed for FAs to neutral cues or in light drinkers.

4. ERP

4.1. Letter Go/No-go task

4.1.1. N200 component

The mixed ANOVA conducted with N200 latency as dependent variable revealed a significant main effect of Trial type (F(1,24) = 16.75, p < 0.001). The N200 latency was longer on No-go trials than on Go trials (p < 0.001). No significant main effect of Group (F(1,24) = 0.14, p = 0.70) nor significant interaction (F(2,56) = 0.008, p = 0.92) were observed.

With N200 amplitude as dependent variable, the mixed ANOVA revealed no significant main effect of Trial type (F(1,24) = 0.97, p = 0.33), no significant main effect of Group (F(1,24) = 0.08, p = 0.77), and no significant Trial type × Group interaction (F(1,24) = 0.83, p = 0.37). However, a significant Group × Electrode interaction was observed (F(6,144) = 3.31, p = 0.03). While no difference of N200 amplitude was observed in heavy drinkers as regards to electrodes, higher non-significant N200 amplitude was found in frontal electrodes (F3, Fz, F4) than parietal and central electrodes (C4, Pz) in light drinkers (p < 0.1).

4.1.2. P300 component

The repeated ANOVA conducted with P300 latency as dependent variable revealed a significant main effect of Trial type (F(1,24) = 17.24, p < 0.001). The P300 latency was longer on No-go trials than on Go trials (p < 0.001). No significant Group effect (F(1,24) = 0.92, p = 0.34) or interaction (F(1,24) = 0.002, p = 0.96) were observed. The repeated ANOVA computed with P300 amplitude as dependent variable revealed a significant main effect of Trial type (F(1,24) = 62.79, p < 0.001) and no significant main effect of Group (F(1,24) = 0.14, p = 0.71) nor Trial type × Group interaction (F(1,24) = 0.87, p = 0.36). The P300 amplitude was higher during No-go trials than during Go trials (p < 0.001).

5. Alcohol modified Go/No-go task

5.1. N200 component

With N200 latency as dependent variable (see Fig. 1 and Table 2), the mixed ANOVA revealed no significant main effects or interaction. Regarding N200 amplitude (see Fig. 1 and Table 2), the ANOVA revealed a significant main effect of Trial type (F(1,22) = 10.26, p = 0.004). Higher N200 amplitude was observed during No-go trials compared to Go trials (p = 0.004). There were no other significant main effects. A significant Trial type × Alcohol avoidance interaction was observed (F(1,22) = 11.38, p = 0.002). We observed a non-significant trend showing that drinkers who experienced higher avoidance towards alcohol tended to show a higher N200 during No-go trials compared to drinkers who experienced low alcohol avoidance (p = 0.057). Moreover, the amplitude of the N200 in drinkers who experienced high avoidance towards alcohol was significantly higher in No-go trials compared to Go trials (p < 0.001) while no such difference was observed in drinkers who experienced less alcohol avoidance (p = 0.99). Finally, a significant Electrode × Group interaction was observed (F(6,132) = 5.22, p = 0.006). Post-hoc analysis revealed that while no difference of N200 amplitude was observed in heavy drinkers, N200 amplitude was higher at frontal electrodes (F3, Fz, F4) than parietal and central electrodes (C3, C4, Pz) in light drinkers (p < 0.01). A significant Electrode × Group × Trial interaction was also found (F(6,132) = 3.95, p = 0.02).

5.2. P300 component

The ANOVA conducted with P300 latency as dependent variable (see Fig. 1 and Table 3) revealed a significant main effect of Trial type (F(1,22) = 48.63, p < 0.001) and a significant main effect of Stimulus type (F(1,22) = 17.32, p < 0.001). We found delayed P300 latencies for neutral cues compared to alcohol-related cues and during No-go trials compared to Go trials. No other main effects or interaction were significant.

With P300 amplitude as dependent variable (see Fig. 1 and Table 3), the repeated ANOVA showed a main effect of Trial type (F(1,22) = 23.97, p < 0.001). Post-hoc test indicated higher P300 amplitude during No-go trials as compared to Go trials (p < 0.001). Moreover, we found a significant Stimulus type × Trial type × Alcohol avoidance interaction (F(1,22) = 4.86, p = 0.04). Higher P300 amplitude was observed in low alcohol avoiders for No-go as compared to Go trials for alcohol-related cues (p < 0.001) and no such difference was found in high alcohol avoiders (p = 0.19). No other main effects or interaction were significant.

5.3. Correlations between behavioral data and ERP data

Correlation analyses showed a positive correlation between the percentages of FA to alcohol-related cues during No-go trials with the impulsivity score (r = 0.43, p = 0.02) (see Table 4). Analyses also revealed a positive correlation between the percentage of FAs to alcohol-related cues during No-go trials.
alcohol-related cues and the P300 latency during No-go trials (r = 0.38, p = 0.05). However, AUDIT scores, obsessed craving, and impulsivity did not exhibit significant correlations with N200/P300 amplitudes or latencies.

6. Discussion

While the improvement of inhibition response has been suggested as a requirement to maintain alcohol consumption under control (Field et al., 2010), the specific exploration of the inhibitory capacities of alcohol-related stimuli in alcohol abuse has received limited attention in the literature. The present study investigated the inhibition processes, and more particularly toward alcohol-related cues, in heavy and light drinkers with a classical letter Go/No-go task and an alcohol modified Go/No-go task, while taking into account the level of alcohol avoidance.

We first hypothesized a general inhibition deficit and reduced N200 and P300 amplitudes in heavy drinkers during the classical letter Go/No-go task as compared to light drinkers. In line with this hypothesis, heavy drinkers made more FA in this task by comparison with light drinkers. Moreover, based on Group × Electrode interaction, while no N200 amplitude differences were observed between electrode sites in heavy drinkers, light drinkers showed a non-significant tendency to exhibit higher N200 amplitudes over frontal sites compared to central and parietal electrode. This non-significant tendency was consistent to previous findings (Enriquez-Geppert et al., 2010; Smith et al., 2007). The same pattern of anteriorization among light drinkers was significantly observed in the alcohol modified Go/No-go task. Lower N200 amplitude has previously been observed in detoxified alcohol-dependent patients who performed an equiprobable Go/No-go task as compared to a healthy control group (Pandey et al., 2012). The difference between groups was larger at frontal and central regions, and the N200 was especially affected during No-go trials. In line with these results, our data suggest that the heavy drinkers do not display the N2 anteriorization observed in the light drinkers. Chronic alcoholism may therefore be related to dysfunctional frontal activation associated with control process, particularly when suppression of a motor response is required (i.e., No-go trials).

Contrary to our hypothesis, no difference of P300 amplitude was observed between heavy and light drinkers during No-go trials. Based on the findings of Oddy and Barry (2009), we expected reduced P300 amplitudes during the letter No-go trials in heavy drinkers. Several discrepancies between the two studies could explain the absence of difference in our study. Firstly, in the study of Oddy and Barry (2009), the level of alcohol consumption was based on the number of standard alcoholic drinks per month. Here, heavy and light drinkers were classified according to the AUDIT scores. Since the AUDIT assesses the harmful effects of alcohol consumption, scores might be more influenced by negative consequences of consumption than by the frequency of consumption itself. Secondly, Oddy and Barry (2009) used an equiprobable Go/No-go task in contrast to our task that consisted of 25% of No-go trials. According to Barry and Rushby (2006), in an equiprobable Go/No-go task, the No-go stimulus is not related to response inhibition process, but rather an involuntary orientation reflex (No-go trials). In contrast, the 75%/25% distribution used here is more likely to require active inhibition of the response during No-go trials (Barry and Rushby, 2006).

Our second hypothesis concerned an impaired response inhibition towards alcohol-related cues in heavy drinkers. Unlike another study comparing detoxified alcohol-dependent patients to a control group of healthy participants (Noël et al., 2007), we showed no significant difference of inhibition performance or RT towards alcohol-related cues between heavy and light drinkers. This absence of deficit of inhibition

![Fig. 1. Illustration of the N200 and P300 on the three central electrodes (FZ, CZ, PZ) in No-go and Go trials. Waves for alcohol-related pictures and neutral pictures in heavy (red and yellow, respectively) and light drinkers (blue and dark green, respectively) are illustrated.](image-url)
null
towards alcohol-related cues in heavy drinkers could be explained by two factors. First, the level of alcohol avoidance, combined with the pattern of consumption, could play a key role in the occurrence of FA. Indeed, we found that high alcohol avoiders performed more FA than low alcohol avoiders independently of stimulus category, and that heavy drinkers who experienced high alcohol avoidance tended to make more FA for alcohol-related cues than those who experienced low alcohol avoidance. This finding is quite surprising because the self-reported level of alcohol avoidance has been found to correlate negatively with automatic approach tendencies (Barkby et al., 2012). However, recent findings report that alcohol-dependent patients inclined to avoid alcohol show higher relapse rates, suggesting that alcohol avoidance could hide approach drives (Spruyt et al., 2013). The second factor that could explain the absence of inhibition deficit towards alcohol-related cues in heavy drinkers is the low impulsivity level found in our participants. This could act as a protective factor against the difficulty to refrain a response toward alcohol (Papachristou et al., 2012). The positive correlation between the impulsivity level and the percentage of FA to alcohol-related cues found in the present study supports this hypothesis.

The present results showed that all the participants were faster to process alcohol-related cues as reflected by earlier P300 latencies during Go trials and by shorter RT. Previous findings also reported faster responses to alcohol-related cues independently of participants’ alcohol consumption profile, which could mean the high salient emotional value of alcohol stimuli experienced by all participants (Kreusch et al., 2013).

When alcohol avoidance was included as a between group factor in our ERP analyses, the results showed a larger N200 amplitude tendency for alcohol-related cues in heavy drinkers as compared to Go trials among alcohol avoiders, whereas no such differences were found between No-go and Go trials for lower alcohol avoiders. In previous studies, larger N200 amplitude was reported following incongruent conditions such as the avoidance of positive stimuli (Ernst et al., 2013) or the interference by irrelevant information (Yeung and Cohen, 2006). Moreover, several studies indicated that anxious individuals seem to maintain a higher level of cognitive control to cope and to monitor the outcome of their actions reflecting by larger N200 amplitude (Righi et al., 2009; Sehlmeyer et al., 2010). As participants avoiding alcohol were constrained to process alcohol cues, they were probably in an incongruent condition, felt more anxious, leading to the recruitment of higher cognitive control resources in order to perform the task. Our results showed also that low alcohol avoiders exhibited higher P300 amplitude for alcohol-related cues during No-go trials than during Go trials, but no such difference was found in high alcohol avoiders, suggesting lesser reactivity to alcohol cues in higher alcohol avoiders. Such finding (less P300 amplitude reactivity, following by larger N200 amplitude) has previously been associated with a higher task difficulty (Gajewski & Falkenstein, 2013; Benikos et al., 2013), suggesting that inhibiting a response towards alcohol-related cue would be more difficult in high alcohol avoiders than in low avoiders. Higher alcohol avoiders could allocate more attentional resources to inhibit the response (i.e., higher N200 amplitude), leading to less reactivity to alcohol-cues as compared to low alcohol avoiders (i.e., no higher P300 amplitude for alcohol related cues). Taken together, our behavioral and ERP data suggest that the level of alcohol avoidance influences the processing of alcohol-related stimuli more than the current levels of alcohol consumption in non alcohol dependent participants.

Before concluding, some limitations of the present study should be acknowledged. The main limitation is the small sample size that may prevent the detection of some main effects, interactions, and correlations. Secondly, our sample consists of university students, and therefore our results must be extended to other populations of alcohol abusers. Thirdly, since amplitudes of some ERP components can be affected by the family history of alcoholism or substance abuse (Porejse et al., 2005; Euser et al., 2012), the absence of control over this factor in the present study constitutes a limitation. Fourthly, as the ERPs are recorded only at the median line (Fz, Cz, Pz) and fronto-central sites (F3, F4, C3, C4) in present study, further studies must be conducted with more electrode sites to allow a better source localization. Finally, we focused only on the N200 and P300 components. However, recent studies found specific modulations of earlier components like the P100 among binge drinkers. Indeed, Petit et al. (2012b) reported greater P100 amplitudes to alcoholic stimuli in comparison with neutral stimuli in young social drinkers, suggesting enhanced perceptual processing toward alcohol cues. Consequently, this study should be considered as a preliminary one and further research using a larger sample of heterogeneous alcohol consumers should be conducted to assess the ability to inhibit a response towards alcohol-related stimuli.

In conclusion, the present results indicate lower inhibition capacities in heavy drinkers as reflected by the absence of N200 anteriorization. When the inhibition capacities towards alcohol-related cues were investigated, the level of alcohol avoidance impacts the processing of such cues. Indeed, participants who experienced higher avoidance for alcohol made more FA, recruited larger attentional resources (i.e., larger N200) and weaker decisional processes (i.e., less P300 reactivity), especially for alcohol-related cues in the modified alcohol Go/No-go task. In a context involving alcohol-related cues, higher alcohol avoiders could therefore be more disturbed by the presence of alcohol-related cues.

Conflict of interest

None.

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References


