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Craving modulates attentional bias toward alcohol in severe alcohol use disorder: An eye tracking study.

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1 **ABSTRACT**

2 **Background and Aims:** Competing models disagree on three theoretical questions regarding
3 alcohol-related attentional bias (AB), a key process in severe alcohol use disorder (SAUD): (1)
4 is AB more of a trait (fixed, associated with alcohol use severity) or state (fluid, associated with
5 momentary craving states) characteristic of SAUD?; (2) is AB purely reflecting the over-
6 activation of the reflexive/reward system or is it also influenced by the activity of the
7 reflective/control system?; (3) does AB rely on early or later processing stages? We addressed
8 these issues by investigating the time course of AB and its modulation by subjective craving
9 and cognitive load in SAUD.

10 **Design and Setting:** A free viewing eye-tracking task in a laboratory setting, presenting
11 pictures of alcoholic and non-alcoholic beverages, combined with a concurrent cognitive task
12 with three difficulty levels.

13 **Participants:** We included 60 SAUD patients (30 self-reporting craving at testing time; 30
14 reporting a total absence of craving) and 30 controls matched on sex and age.

15 **Measurements:** We assessed AB through early and late eye-tracking indices. We evaluated
16 the modulation of AB by craving (comparison between patients with/without craving) and
17 cognitive load (variation of AB with the difficulty level of the concurrent task).

18 **Findings:** Measures of late AB processes indicated that patients with craving allocated more
19 attention towards alcohol-related stimuli than patients without craving ($p < .001$, $d = 1.093$),
20 resulting in opposite approach/avoidance AB according to craving presence/absence. Patients
21 without craving even showed a stronger avoidance AB than controls ($p = .003$, $d = .806$). AB did
22 not vary according to cognitive load ($p = .962$, $\eta_p^2 = .004$).

23 **Conclusions:** The direction of alcohol-related AB (approach/avoidance) is determined by
24 patients' subjective craving at testing time and does not function as a stable SAUD trait. While
25 relying on later/controlled attentional stages, AB is not modulated by the saturation of the
26 reflective/control system.

- 1 **Keywords:** alcohol use disorder, attentional bias, avoidance bias, eye-tracking, cognitive load,
- 2 alcohol

1 INTRODUCTION

2 Alcohol-related attentional bias (AB) is the preferential allocation of attention toward
3 alcohol-related stimuli. Prominent theoretical models assume that AB plays a causal role in the
4 onset and persistence of severe alcohol use disorder (SAUD) [1-3]. The incentive-sensitization
5 theory [4] postulates that repeated alcohol exposures sensitize the reflexive/reward system,
6 enhancing the incentive properties of alcohol-related cues through conditioning. Becoming
7 more salient, these cues capture attention (i.e., generate AB) and guide individuals towards
8 alcohol consumption. Hence, interventions targeting AB emerged, postulating that reducing
9 AB through attentional retraining would reduce consumption and relapse risk. These
10 interventions, while increasingly implemented in clinical settings with some promising effect on
11 clinical outcomes, led to inconsistent results regarding their impact on AB [5-6]. Such
12 discrepancies might result from the fact that several theoretical questions remain to be clarified
13 in this research field, namely is AB: (1) mostly a trait (fixed, associated with SAUD severity) or
14 state (fluid, associated with momentary motivational states) characteristic of SAUD; (2) purely
15 reflecting the over-activation of the reflexive/reward brain system or also influenced by the
16 activity of the reflective/control system?; (3) characterized by an early/automatic hijacking of
17 attention by alcohol-related stimuli or rather relying on later and more controlled processing
18 stages?

19 Regarding the first question, traditional models assume that AB progressively develops
20 through associative learning and reflexive/reward system over-sensitization, finally constituting
21 an enduring and potentially permanent SAUD characteristic [4,7]. These models thus
22 understated the sensitivity of AB to momentary states (e.g., motivational component of current
23 craving) compared to the influence of stable factors related to SAUD (e.g., duration, severity).
24 In the last decade, there has been more emphasis [8] on how fluctuating factors would
25 moderate the behavioural expression of the reflexive/reward system (i.e., AB). Taking a step
26 further, Field and colleagues suggested that AB is partly driven by temporary changes in
27 appetitive and/or aversive states [9]. According to their theoretical account, AB would result

1 from momentary motivational evaluations of alcohol-related stimuli, hence constituting a state
2 rather than trait SAUD marker. The subjective evaluation (positive, negative, ambivalent) of
3 alcohol-related cues would lead individuals to maintain their attention on it or conversely ignore
4 it, resulting in different AB patterns [9]. The reported [10-11] intra-individual AB fluctuation
5 according to current motivational value of alcohol (e.g., motivational component of current
6 craving [12], drinking status [13]) supports this proposal. Patients with SAUD might present an
7 AB strongly affected by their current states, which would hence not constitute a key causal
8 factor for SAUD persistence, raising doubt on the rationale of AB retraining. However, the very
9 few studies exploring AB in SAUD used unreliable measures [14], applied on recently-
10 detoxified patients (known to frequently present aversive/ambivalent alcohol evaluation and
11 low craving), which might explain the inconsistent results [15]. The only study [16] using reliable
12 eye-tracking measures showed both an avoidance bias in recently-detoxified patients with
13 SAUD and a positive correlation between AB and craving. These results call for directly
14 addressing the inconsistent theoretical assumptions regarding AB fluctuations [4,9].

15 The second question relates to the dual-process models [1,17], postulating that SAUD
16 emerge from (1) the under-activation of the “reflective/control system”, responsible for
17 deliberative and controlled responses, and (2) the over-activation of the “reflexive/reward
18 system”, initiating automatic and appetitive behaviours. In this view, AB results from the
19 overactivation of the reflexive/reward system, but the role played by the reflective/control
20 system in its occurrence remains unclear. Indeed, dual-process models stated that situational
21 factors such as cognitive load could selectively impair the reflective/control system, leading the
22 reflexive/reward system to take the lead (therefore assuming a continuous interaction between
23 systems). Nevertheless, they simultaneously stated that reflexive/reward processes operate in
24 an effortless manner, independently from the availability of cognitive resources [8,18]. Previous
25 studies also suggested that AB is not an artefact of patients’ impaired cognitive/executive
26 functioning [19] but rather a genuine consequence of the reflexive/reward system’s
27 overactivation [20]. Still, studies in other psychopathological states showed that AB might be
28 increased by executive dysfunction [21-23], suggesting that AB is affected by the activity of the

reflective/control system. In line with this proposal, the paradigms classically used to measure AB cannot rule out the possibility that participants use their executive functions to voluntarily modify their AB, for example by using oculomotor inhibition to actively avoid saccades towards alcohol-related stimuli. An experimental way to assess this role of the reflective/control system in AB would be to saturate this system through a concurrent cognitive task that places high demands on cognitive resources, thus hampering their ability to modulate AB. In other words, the temporary reduction of available cognitive resources would reduce the ability of the reflective/control system to modulate AB. Conversely, if AB is independent of the reflective/control system, saturating this system should have no influence on AB.

The third question is whether AB relies on early and automatic attentional processes (generating an uncontrolled capture of attention towards alcohol [1,17]), or on later and more controlled processes (being related to longer processing time for alcohol-related cues and/or to a difficulty to disengage attention from them [24]). When simultaneously presenting alcohol-related and non-alcohol-related stimuli, eye-tracking allows the dissociation, with high temporal/spatial resolution, of (1) the initial attentional capture quickly following the appearance of alcohol-related cues, by measuring early indices like first Area of Interest (AOI) visited (i.e., percentage of trials in which alcohol versus non-alcohol AOIs were fixated first), from (2) the controlled maintenance of attention towards alcohol, by measuring late indices like dwell time (i.e., sum of fixation times on alcohol versus non-alcohol AOIs during the whole trial). An early/automatic AB is thus inferred by modification of the early indices (e.g., higher percentage of first fixations towards alcohol-related stimuli) while a late/controlled AB is inferred by modification of late indices (e.g., higher dwell time for alcohol-related stimuli). Eye-tracking studies showed that AB mostly appears at the late processing stages in subclinical and clinical populations [25-27], thus questioning its early/automatic nature.

Here, we directly address these three conceptual questions, as we (1) clarify whether AB is stable or affected by motivational states by comparing recently-detoxified patients with SAUD with or without craving in a free viewing eye-tracking task assessing AB. If AB is modulated by temporary appetitive states such as the motivational component of craving,

1 patients reporting craving at testing time (defined here as the intense and irrepressible desire
2 to consume alcohol right now) will tend to show an AB towards alcohol-related stimuli (i.e.,
3 approach bias), while patients without craving will tend to avoid these stimuli (i.e., avoidance
4 AB); (2) investigate whether reducing the capacity of the reflective/control system (by placing
5 a temporary load on cognitive resources) would increase AB. If the reflective/control system
6 plays a role in AB, its saturation in the high cognitive load condition of the concurrent task (i.e.,
7 the reduction of cognitive resources available to inhibit AB) will lead to a higher AB towards
8 alcohol-related stimuli compared to the low cognitive load condition (where cognitive resources
9 are available to voluntarily modulate AB); (3) determine the temporal dynamics of AB by
10 distinguishing early/automatic and late/controlled processing steps. If AB is related to late
11 processing stages rather than early/automatic ones, it will only be observed for the indices of
12 controlled attentional maintenance (i.e., dwell time, namely higher total time spent on alcohol-
13 related stimuli than on non-alcohol-related stimuli) and not for the indices related to the early
14 attentional capture (i.e., first AOI visited, namely higher number of first fixations towards
15 alcohol-related stimuli than towards non-alcohol-related stimuli).

16 **METHODS**

17 **Participants**

18 Before starting data acquisition, we recruited and allocated thirty patients (15 women)
19 who self-reported craving at testing time (i.e., scored higher than zero at the baseline craving
20 Visual Analogue Scale, VAS: 0 = no desire at all to consume alcohol right now, to 100 = terrible
21 desire to consume alcohol right now) to the experimental *craving* group, and thirty patients (10
22 women) who did not report craving at baseline (i.e., scored zero at the VAS) to the *non-craving*
23 group. All patients fulfilled DSM-5 criteria for SAUD and were tested during their detoxification
24 treatment in three Belgian hospitals. They had all been abstinent for at least 7 days and were
25 free of other psychiatric comorbidities (except tobacco use disorder). We recruited thirty
26 healthy controls (15 women) through social networks and e-mails. Controls were free of any
27 past/present psychiatric disorder and personal/parental SAUD history. They consumed less

than ten standard alcohol units (10gr of pure ethanol per unit) per week and never exceeded three units per day. They scored lower than 8 at the AUDIT [28] (score from 0 to 40; $\alpha=.960$) and had to refrain from consuming alcohol the day before testing. Exclusion criteria for both groups included polysubstance use disorder and major past/present neurological trauma and/or disorder. They all had normal or lens corrected vision and were fluent in French.

Procedure

Participants provided written informed consent and were tested individually. Before starting data acquisition, they filled out questionnaires assessing state anxiety (STAI-S; score from 20 to 80; $\alpha=.960$) and current alcohol craving. We used the Alcohol Craving Questionnaire Short Form Revised (ACQ-SF-R [29]; score from 12 to 84; $\alpha= .874$) for a multi-dimensional assessment of craving and the VAS single-item (score from 0 to 100) to obtain a quick and specific measure of the motivational component of craving. The computerized experimental task comprised three parts and lasted 20-30 minutes. We re-assessed craving through the VAS after each part.

We performed a standard 9-point eye-gaze calibration at the beginning of each experiment part. Between them, participants filled out questionnaires measuring psychopathological variables, namely depression (BDI-II [30]; score from 0 to 39; $\alpha=.914$), trait anxiety (STAI-T [31]; score from 20 to 80; $\alpha=.949$) and impulsivity (UPPS-P, [32]; score from 20 to 80; $\alpha=.898$). The study protocol followed the Declaration of Helsinki and was approved by the Ethics Committee of Saint-Luc-UCLouvain Clinics (reference number: 2019/26MAR/141). After the experiment, we debriefed participants and controls received financial compensation. As this study was not pre-registered, its results should be considered as exploratory in line with the Addiction journal guidelines.

Apparatus

Participants were seated at a desk, facing an eye-tracker camera and an Asus Display Laptop PC with a 17.3-inch FHD screen (resolution 1080x1920; refresh rate 120Hz, placed

60cm away from the eyes). We controlled the presentation of the task and its synchronization with eye-tracking using OpenSesame [33]. We recorded eye movements using an EyeLink Portable Duo (SR Research, Canada; sampling rate 1000Hz; average accuracy range 0.25°-0.5°, gaze tracking range 32° horizontally, 25° vertically).

Free viewing eye-tracking task

The AB task was replicated from [34]. In each trial, participants had to first fixate a central fixation dot appearing on the background screen for at least 100ms. We used this dot as a drift check for eye-gaze calibration, and to ensure that participants focused their attention at the centre of the screen. Once the eye-tracking device detected the eyes at the centre of the screen, a 4x4 matrix replaced the dot for 6000ms. The matrix presented 16 full color 250x250 images of eight alcoholic and eight non-alcoholic beverages without context. The four inner pictures always consisted of two alcohol and two non-alcohol pictures, while we randomized the 12 outer pictures. Participants were asked to freely look at the pictures. To support participants' task engagement, we presented three types of stimuli: bottle, bottle with empty glass, bottle with filled glass. A total of 218 pictures were extracted from the ABPS battery [35], the selected stimuli being culturally relevant for the Belgian population.

Participants completed three experimental conditions, each containing 54 trials: one presenting solely the free viewing task (i.e., *Baseline*, see Figure 1) without any concurrent task, and two presenting the free viewing task alongside a concurrent cognitive task of increasing difficulty (i.e. low cognitive load, *Level 1* and high cognitive load, *Level 2*). *Levels 1* and *2* were used to saturate the reflective/control system and hence to explore the impact of depleting cognitive resources on AB, always measured through the free viewing task performed simultaneously. The baseline free viewing task was always performed first to ensure a valid measure of AB (i.e. uncontaminated by the concurrent cognitive task). Then, we presented *Levels 1* and *2* in a counterbalanced order between participants, to control for potential learning or fatigue effects between these two conditions.

1 **Concurrent auditory cognitive task**

2 In *Level 1*, we presented a series of digits orally through headphones and participants
3 had to detect the appearance of a target digit (“5”) by mouse-clicking (low cognitive load, as
4 participants just had to pay attention to the presented digit without further cognitive
5 processing). In *Level 2*, we presented another series of digits orally and participants had to
6 mouse-click each time the sum of the two last digits was 10 (high cognitive load, as participants
7 had to store digits in their working memory and perform additions throughout the task). A male
8 French voice [36] pronounced each digit with the same pace. We used Audacity® software to
9 mark the onset/offset of each digit and then compressed the sampled period in an OGG file.
10 The duration of enunciation and silent periods for each digit was set to 2000ms. In both levels,
11 we presented the series of digits in a continuous way to keep the difficulty level constant. The
12 dependent variables for this concurrent cognitive task were correct responses (i.e., percentage
13 of target digits for which the participant correctly mouse-clicked), false alarms (i.e., number of
14 non-target digit for which the participant mouse-clicked) and delayed responses (i.e., number
15 of target digits for which the participant mouse-clicked after the onset of the following digit).

16 **Data analysis**

17 No data-reduction procedure was performed as the free-viewing task did not measure
18 any behavioural performance. Seven individuals (2 patients with craving, 3 patients without
19 craving, 2 control participants) were not able to participate in our experiment due to poor eye-
20 tracking calibration. We set our sample size to 30 participants per group based on previous
21 work exploring AB in SAUD [12,16,37-39]. We performed a simulation-approach power
22 analysis for exploratory three-way ANOVA designs [40] assuming an alpha of 0.05, a sample
23 size of 30 per group and a Bonferroni adjustment for multiple comparisons, which indicated a
24 power of 0.65 and 0.95 for detecting a small to medium effect size ($\eta_p^2=.022$) in our 3x2 and
25 3x2x3 within-between interactions respectively. We defined AOI for the free viewing task as
26 the zone in pixels covered by each image, leading to 16 AOIs per trial (corresponding to the
27 number of stimuli shown per trial in the free viewing task). We assessed early AB processes

through the *first AOI visited* index, and late AB processes through *dwell time* index. EyeLink algorithms qualified gaze samples as fixations or saccades. For the concurrent task, we performed a manipulation check to check that the level of difficulty was higher in Level 2 than Level 1 by contrasting the percentage of correct responses, and the number of false alarms and delayed responses in both levels.

We performed all statistical analyses using SPSS 27.0. We performed between-group comparisons (independent t-tests; chi-square tests) on demographic, psychopathological and alcohol-related variables. As an estimate of reliability, we computed Cronbach's alpha for the first AOI visited and dwell time measures at baseline¹. Following a well-established procedure [41,42], we calculated AB scores (difference scores or percentages for alcohol vs. non-alcohol stimuli) separately for each matrix, leading to 54 AB scores for each AB measure. Considering the established left-gaze bias effect on the first AOI visited (i.e., left hemifield preference related to reading direction [43]), we also estimated the reliability of this measure when comparing AOIs on the left vs. right of the screen. We performed a manipulation check of the cognitive load and the increased difficulty from Level 1 to Level 2 by conducting 3x2 ANOVAs on data from the concurrent task (correct responses, false alarms, delayed responses) with GROUP (craving/non-craving/controls) as between-subjects factor and LEVEL (1/2) as within-subject factor. We performed 3x3x2 ANOVAs on AB indices (first AOI visited/dwell time, to test our third hypothesis) with GROUP (craving/non-craving/controls, to test our first hypothesis) as between-subjects factor, LEVEL (baseline/1/2, to test our second hypothesis) and STIMULI (alcohol/non-alcohol) as within-subject factors. We rerun these analyses by adding education level, age at first consumption, state and trait anxiety, depression and impulsivity as covariates (since at least two groups differed on these potentially confounding variables, see Results section) and reported the results in Supplementary Material. We conducted Post-Hoc tests with a Bonferroni-corrected p-value of $\alpha_{\text{altered}}=.05/3=0.017$. Finally, we performed Pearson's

¹ We dismissed the inclusion of two supplementary eye-tracking measures from our analyses (i.e., first saccade latency, number of AOIs visited) because of their redundancy with the main measures of early and late AB processes, but also in view of the poor reliability for the latter ($\alpha=.385$), and inability to test the reliability of the former (as the number of observations was too small when separated by type of stimuli, trial and participant).

correlations to explore the influence of all demographic, psychological and alcohol-related variables on AB magnitude (indexed by dwell time) and on craving at baseline.

RESULTS

Demographic, psychopathological and alcohol-related measures

Both groups of SAUD patients had fewer years of education, earlier age at first consumption (for craving SAUD patients only), and higher state anxiety, trait anxiety, depression and impulsivity than controls (Table 1). Craving and non-craving SAUD patients did not differ except for trait anxiety.

As expected, both groups of SAUD patients reported consuming more alcohol per week and had higher AUDIT score than controls, and non-craving SAUD patients reported lower craving than craving SAUD patients (for all VAS and ACQ-SF-R) and controls (for all VAS).

Manipulation check: Concurrent cognitive task

Correct responses, false alarms and delayed responses. We found Level effects showing lower percentages of correct responses [$F(1,73)=38.380$, $p<.001$, $\eta_p^2=.345$], more false alarms [$F(1,73)=9.880$, $p=.002$, $\eta_p^2=.119$] and more delayed responses [$F(1,73)=9.175$, $p=.003$, $\eta_p^2=.112$] in Level 2 than Level 1. Other main effects and interactions were inconclusive (all p -values $\geq.078$).

These results show that the concurrent cognitive task successfully intensified the saturation of the reflective/control system by increasingly recruiting cognitive resources between Levels 1 and 2.

Free viewing AB task

AB reliability

Internal consistency of AB scores was high for dwell time ($\alpha=.976$). It was low for the first AOI visited when comparing alcohol vs. non-alcohol AOIs (regardless of the position of those AOIs; $\alpha=.047$) but conversely high when comparing AOIs on the left vs. right (regardless of their alcohol/non-alcohol content; $\alpha=.908$).

This latter result suggests that the low reliability of the first fixation towards alcohol/non-alcohol AOIs is mostly due to the predominance of the left-gaze bias [participants more frequently directed their first fixation on the left AOI compared to the right AOI; $t(89)=11.412$, $p<.001$, $d=1.203$] rather than the poor psychometric quality of this measurement.

Early AB processing stages

First AOI visited. We found a TYPE effect showing that participants performed first fixations more frequently towards non-alcohol-related than alcohol-related stimuli. Other main effects and interactions were inconclusive (Table 2).

Late AB processing stages

Dwell Time. We found a TYPE effect showing longer dwell time for non-alcohol-related than alcohol-related stimuli. We also found a STIMULI*GROUP interaction (Figure 2): SAUD patients with craving presented longer dwell time on alcohol [$t(58)=4.234$, $p<.001$, $d=1.093$], and shorter dwell time on non-alcohol [$t(58)=3.586$, $p<.001$, $d=.926$] than SAUD patients without craving (Table 3). Moreover, both SAUD patients without craving and controls showed longer dwell time on non-alcohol than alcohol-related stimuli [non-craving: $t(29)=5.635$, $p<.001$, $d=1.029$; controls: $t(29)=2.775$, $p=.010$, $d=.507$], but this difference was higher in non-craving SAUD patients than controls [alcohol: $t(58)=3.122$, $p=.003$, $d=.806$; non-alcohol: $t(58)=2.826$, $p=.007$, $d=.730$]. Other main effects and interactions were inconclusive (Table 2)².

² The stimuli*group interaction for dwell time remained significant in the supplementary analyses that added as covariates all the demographic, psychological and alcohol-related variables differing between groups. Other main effects and interactions were inconclusive for both AB indices (see Table S1 in Supplementary material for the detailed results).

Note that the LEVEL effect did not lead to any significant main effect or interaction on indices of AB, suggesting that the results of the free viewing task and the related AB were not modulated by the intensity of the cognitive load generated by the concurrent task.

Correlations

Dwell Time. In SAUD patients, we found positive correlations between dwell time at baseline and craving before (VAS: $r=.443$, $p<.001$; ACQ: $r=.546$, $p<.001$) and after (VAS: $r=.486$, $p<.001$) the task. All other correlations in SAUD patients and controls between AB and demographic, psychological or alcohol-related variables were inconclusive (all $p\text{-values}\geq.060$).

Craving. In SAUD patients, we found a negative correlation between craving VAS score at baseline and age ($r=-.289$, $p=.025$). All other correlations in SAUD patients and controls were inconclusive (all $p\text{-values}\geq.197$).

DISCUSSION

Addiction models postulate that AB is a major index of the overactivation of the reflexive/reward system [4,17], causally implicated in SAUD persistence. In the last decade, concurrent models placed greater emphasis on the moderating role of situational factors (e.g., craving, cognitive load) on the links between alcohol use severity and AB [8-9]. Nevertheless, a theoretical blur persists on the nature and role played by AB in SAUD. We thus experimentally addressed three remaining questions on AB, namely how is it affected by current motivational state and cognitive load, and what is its time course.

First, we showed the role played by craving in the magnitude and direction of AB, offering experimental support to the theoretical proposals that AB is affected by current motivational state rather than stable. Indeed, craving SAUD patients spent more time looking at alcohol stimuli than patients without craving, while the reverse was found for non-alcohol stimuli. Moreover, both controls and non-craving SAUD patients showed an avoidance bias for alcohol-related stimuli (i.e., shorter dwell times for alcohol-related stimuli), this bias being even stronger in the latter group. These results are in line with recent findings showing an avoidance

1 bias in SAUD patients reporting very low craving and high abstinence motivation [16].
2 Altogether, our findings question the proposal of a long-lasting and potentially permanent AB
3 in SAUD, since we could not find any AB among recently-detoxified patients when using
4 reliable eye-tracking measures (i.e., dwell time). The opposite AB patterns between the two
5 groups of patients support the theoretical account that AB is driven by temporary changes in
6 appetitive/aversive motivational states regarding alcohol, and that its stability along the
7 disorder might have been overstated [9]. The subjective momentary evaluation of alcohol-
8 related cues (indexed here by the motivational component of craving and assessed through
9 VAS) constitutes an important predictor of whether individuals maintain and/or override their
10 gaze on them, resulting in avoidance/approach AB. In line with our hypotheses [15], we
11 showed that, during the detoxification process, non-craving patients present an avoidance bias
12 towards alcohol. This result makes sense as these patients, being abstinent in clinical settings
13 and motivated to avoid alcohol outside the clinic, present a negative evaluation and aversive
14 state towards alcohol. Such avoidance bias is also identified, although to a lesser extent,
15 among control participants, which was expected as this group comprised low drinkers
16 presenting reduced attraction towards alcohol. Conversely, craving patients show motivational
17 conflict (i.e., craving associated with abstinence motivation), thus not leading to a strong AB
18 towards alcohol (i.e., no significant difference with control participants) and confirming that AB
19 is not a strong and stable characteristic of patients presenting SAUD. Correlational analyses
20 further support this proposal that AB is influenced by current motivational states rather than by
21 stable consumption characteristics (e.g., SAUD duration, consumption frequency or intensity),
22 as AB was not associated with any index of SAUD severity, except VAS craving scores
23 (specifically assessing the motivational component of craving). Future research should
24 investigate whether AB might also fluctuate with other components of this multi-dimensional
25 construct (e.g., bodily sensations) and/or momentary aversive states (as postulated by [9]).

26 The concurrent cognitive task supported the proposal that AB relies on the over-
27 activation of the reflexive/reward system and might be quite independent from the
28 reflective/control system, as AB patterns were not influenced by the extent of cognitive

resources available. However, this null result cannot be interpreted as definitive proof that the reflective/control system does not influence AB. Indeed, while our manipulation check demonstrated that we efficiently increased cognitive load across conditions (i.e. worse performance in Level 2 than 1), the cognitive resources of the reflective/control system might have been insufficiently saturated to impact the reflexive/reward system and AB, and our sample size might have been too low to detect a subtle impact of cognitive load on AB. This independence of AB towards the reflective/control system should be confirmed in future studies that more strictly manipulate the saturation of cognitive resources.

Finally, we suggested that AB, regardless of its direction (approach/avoidance), is significantly underpinned by late attentional stages (assessed through dwell time measures) and is mostly characterized by a preferential maintenance of attention towards alcohol/non-alcohol stimuli. In contrast, the facilitated capture of attention could not be reliably assessed in the present study (as in previous studies [16,33,44]), since the measure of first AOI visited was contaminated by the classical dominance of the left side of the visual field related to reading/writing habits [43]. Nevertheless, we observed that the presumably strong attention-grabbing properties of alcohol-related stimuli (as postulated by dominant models [4]) did not overcome the left-gaze bias, even in participants showing stronger AB towards alcohol at later processing stages (i.e., SAUD patients with craving), and were actually contradicted by the higher percentage of first fixations towards non-alcohol-related stimuli in all our participants. This casts doubt on the postulated automatic/early nature of AB in SAUD [17], already questioned by heterogeneous findings when manipulating stimuli duration in behavioural experiments [45,46]. In contrast, our results regarding the late component are in line with eye-tracking studies in subclinical [47] and clinical [16] populations, as well as with earlier studies targeting such malleable late components in attentional retraining [6,24]. We thus highlighted the relevance of dwell time measures to investigate the preferential maintenance of attention throughout the trial and we encourage future studies to increase the reliability of early eye-tracking indexes by presenting stimuli vertically to override the predominant left-gaze bias

and/or by developing AB tasks specifically exploring the early attentional capture by alcohol-related cues [48].

As our study was not pre-registered, its results should be considered as exploratory. Although none of the assessed demographic and psychopathological variables correlated with AB (including the educational level for which groups were not matched), it should also be acknowledged that other unmeasured biasing variables might have modulated AB (e.g., alcohol-related motivations, subclinical psychiatric comorbidities). Finally, while we followed the current guidelines to reliably explore AB in alcohol-use disorders, notably by using eye-tracking measures and the recommended free viewing task [49], other paradigms that require more active engagement of participants could be adapted to reliably measure AB (e.g., visual search task, [50]). Despite these limits, our findings should lead researchers and clinicians to reconsider the role of AB in SAUD and the conditions in which AB modification programs should be conducted. Some patients with high craving and/or low abstinence motivation might present genuine AB and could thus benefit from attentional training [6]. Indeed, since AB is more easily triggered by specific motivational states (i.e., high craving, positive alcohol evaluation), interventions could have stronger effects by administering attentional training when patients are currently in this state, or by using other interventions directly modifying this state [e.g., mindfulness or visual cognitive interference; 51]. However, most recently-detoxified patients already avoid alcohol-related cues, raising doubts regarding the usefulness of generalized attention training in this population. Importantly, the increasing accessibility of reliable AB measures by using low-cost eye-tracker or newly developed AB paradigms [52] helps clinicians to identify patients who will benefit most from attention training. Finally, the strong relationship between AB and craving observed here and previously [53] highlights the need to identify and target psychological factors triggering craving in SAUD to break the vicious circle between craving, AB and alcohol-seeking behaviour, traditionally described as the three pathways to relapse [54].

CONCLUSION

1 We used eye-tracking measures to clarify three major theoretical questions on AB in
2 SAUD, namely whether AB is stable, independent of the reflective/control system and
3 early/automatic. We showed that AB is not stable in detoxified patients with SAUD, but is rather
4 determined by the presence of craving, patients with/without craving presenting opposite AB
5 patterns. The absence of craving is associated with a strong avoidance AB for alcohol-related
6 cues, thus questioning most theoretical frameworks proposing that AB constitutes a central
7 and long-lasting SAUD feature [4]. We thus argue, in line with alternative theoretical proposals,
8 that AB rather expresses momentary changes in appetitive/aversive evaluation of alcohol-
9 related cues [9]. We also offered preliminary data to suggest that AB might not be influenced
10 by increased cognitive load (and might thus be quite independent from reflective/control
11 system's activity) and is mostly related to later and more controlled stages of attentional
12 processing, thus not being related to early/automatic attentional capture.

1 **Data availability**

2 The data that support the findings of this study are available on request from the
3 corresponding author.

4 **Author Contributions**

5 **Zoé Bollen:** Conceptualization (lead); methodology (lead); formal analysis (lead);
6 investigation (lead); project administration (lead); software (lead); visualization (equal); writing
7 – original draft (lead). **Arthur Pabst:** Formal analysis (supporting); visualization (equal); writing
8 – review & editing (equal). **Nicolas Masson:** Conceptualization (supporting); methodology
9 (supporting); writing – review & editing (equal). **Reinout W. Wiers:** Writing – review & editing
10 (equal). **Matt Field:** Writing – review & editing (equal). **Pierre Maurage:** Conceptualization
11 (supporting); methodology (supporting); project administration (supporting); supervision (lead);
12 writing – original draft (supporting); writing – review & editing (equal).

13

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1 Tables

2

3 **Table 1.** Demographic, psychopathological and alcohol consumption measures [mean
4 (standard deviation)] and independent samples t-test or chi square test (t or χ^2 and p-values)
5 comparing SAUD patients with craving, SAUD patients without craving and control
6 participants.

	Craving (N=30)	Non-craving (N=30)	Controls (N=30)	Craving vs Non-craving		Craving vs controls		Non-craving vs controls	
				<i>t or χ^2</i>	<i>p</i>	<i>t or χ^2</i>	<i>p</i>	<i>t or χ^2</i>	<i>p</i>
Demographic measures									
Sex ratio (M/F)	20/10	15/15	15/15	1.714	.190	1.714	.190	.000	1
Age	42.90 (10.66)	48.07 (9.35)	47.87 (10.39)	1.996	.051	1.827	.073	.078	.938
Years of education	12.47 (2.85)	13.07 (4.23)	16.07 (2.83)	.644	.522	4.873	<.001	3.192	.002
Psychopathological measures									
BDI-II	11.56 (7.03)	8.68 (8.29)	2.93 (3.69)	1.324	.192	5.778	<.001	3.449	.001
STAI-S	40.47 (15.28)	36.12 (16.76)	28.83 (7.56)	1.050	.298	3.737	<.001	2.170	.034
STAI-T	52.24 (9.18)	45.52 (12.5)	32.30 (11.99)	2.145	.037	6.617	<.001	4.109	<.001
UPPS-P	48.79 (8.28)	44.39 (9.33)	37.43 (7.51)	1.773	.082	5.274	<.001	3.114	.003
Alcohol consumption measures									
AUDIT	33.50 (5.43)	30.75 (6.68)	3.30 (1.70)	1.516	.136	28.532	<.001	21.769	<.001
First consumption (age)	13.85 (3.16)	15.72 (4.89)	15.28 (1.89)	1.756	.084	2.092	.041	.454	.652
Doses per week	32.12 (24.22)	21.90 (13.15)	0.47 (0.43)	1.802	.077	6.833	<.001	8.919	<.001
Years of SAUD	13.40 (9.70)	9.67 (8.58)	N/A	1.580	.119	N/A	N/A	N/A	N/A
Previous detoxification	2.28 (2.88)	3.00 (4.21)	N/A	.770	.445	N/A	N/A	N/A	N/A
Days of abstinence	35.50 (39.51)	39.07 (43.86)	N/A	.331	.742	N/A	N/A	N/A	N/A
Baseline craving (VAS)	22.73 (23.40)	0.00 (0.00)	2.30 (4.84)	5.322	<.001	4.684	<.001	2.601	.012
Post-Level 0 craving (VAS)	29.27 (25.98)	0.00 (0.00)	4.53 (11.20)	6.171	<.001	4.789	<.001	2.217	.031
Post-Level 1 craving (VAS)	28.03 (26.32)	0.00 (0.00)	4.67 (11.28)	5.833	<.001	4.469	<.001	2.266	.027
Post-Level 2 craving (VAS)	28.20 (27.00)	0.00 (0.00)	4.70 (11.36)	5.721	<.001	4.394	<.001	2.266	.027
Baseline craving (ACQ-SF-R)	35.70 (14.13)	17.97 (7.78)	18.17 (5.07)	6.019	<.001	6.396	<.001	.118	.907

7

8 Note. ACQ-SF-R, Alcohol Craving Questionnaire Short Form Revised; AUDIT, Alcohol Use
9 Disorder Identification Test; BDI-II, Beck Depression Inventory-II; N/A, not applicable to this
10 group; STAI-S, State-Trait Anxiety Inventory-State; STAI-T, State-Trait Anxiety Inventory-Trait;
11 UPPS-P, Impulsive Behavior Scale; VAS, Visual Analogue Scale.

12

13

Table 2. ANOVA results for eye-tracking measures of early and late AB processing stages.

	First AOI visited				Dwell time			
	<i>df</i>	<i>F</i>	<i>p</i>	η^2_p	<i>df</i>	<i>F</i>	<i>p</i>	η^2_p
Between-subjects effects								
Group	2	1.521	.224	.035	2	.416	.661	.010
Error (Group)	85				85			
Within-subjects effects								
Stimuli	1	13.467	<.001	.137	1	18.279	<.001	.177
Stimuli * Group	2	1.029	.362	.024	2	9.688	<.001	.186
Error (Stimuli)	85				85			
Level	2	.716	.490	.008	2	2.237	.110	.026
Level * Group	4	1.885	.115	.042	4	.243	.913	.006
Error (Level)	170				170			
Stimuli * Level	2	.717	.490	.008	2	.630	.534	.007
Stimuli * Level * Group	4	1.359	.250	.031	4	.153	.962	.004
Error (Stimuli * Level)	170				170			

Note. *df* indicates degrees of freedom. η^2_p indicates partial eta-squared.

1 **Table 3.** Attentional bias eye-tracking measures for the three levels of cognitive load [mean
2 (standard deviation)] comparing SAUD patients with craving, SAUD patients without craving
3 and control participants.

	First AOI visited (in %)		Dwell time (in ms)	
	Alcohol	Non-alcohol	Alcohol	Non-alcohol
Craving (N=30)				
Baseline	25.73 (4.76)	27.27 (4.58)	2495 (631)	2355 (688)
Level 1	26.73 (3.22)	27.03 (3.28)	2387 (784)	2406 (865)
Level 2	26.14 (3.06)	27.59 (3.12)	2447 (665)	2375 (723)
Non-craving (N=30)				
Baseline	27.03 (3.37)	26.87 (3.42)	1794 (523)	3018 (531)
Level 1	25.07 (3.50)	28.87 (3.57)	1681 (735)	3054 (864)
Level 2	25.14 (3.55)	28.76 (3.52)	1796 (641)	3048 (785)
Controls (N=30)				
Baseline	25.93 (3.80)	27.93 (3.73)	2184 (437)	2617 (480)
Level 1	24.57 (4.65)	27.80 (4.83)	2137 (522)	2582 (489)
Level 2	26.07 (4.13)	27.63 (4.13)	2184 (463)	2622 (561)

4
5 Note. AOI, Area of Interest; ms, milliseconds.

6

1

2 **Figure legends**

3 **Figure 1.** Illustration of a trial from the eye-tracking free viewing task (Fig 1a) and of the
4 concurrent cognitive task with low (Level 1) and high (Level 2) cognitive load (Fig 1b).

5 **Figure 2.** Dwell time observed in patients with severe alcohol use disorder reporting craving
6 (craving), patients reporting no craving (non-craving) and control participants (controls) in the
7 free viewing task at baseline (a), level 1 (b) and level 2 (c) for alcohol and non-alcohol stimuli.