Pavlovian-to-Instrumental-Transfer in Alcohol Dependence – a pilot study

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Abstract

Background: Pavlovian processes are thought to play an important role in the development, maintenance and relapse of alcohol dependence, possibly by influencing and usurping ongoing thought and behavior. The influence of Pavlovian stimuli on on-going behavior is paradigmatically measured by Pavlovian-to-instrumental-transfer (PIT) tasks. These involve multiple stages and are complex. Whether increased PIT is involved in human alcohol dependence is uncertain. We therefore aimed to establish and validate a modified PIT paradigm that would be robust, consistent, and tolerated by healthy controls as well as by patients suffering from alcohol dependence, and to explore whether alcohol dependence is associated with enhanced Pavlovian-Instrumental transfer.

Methods: 32 recently detoxified alcohol-dependent patients and 32 age and gender matched healthy controls performed a PIT task with instrumental go/no-go approach behaviours. The task involved both Pavlovian stimuli associated with monetary rewards and losses, and images of drinks.

Results: Both patients and healthy controls showed a robust and temporally stable PIT effect. Strengths of PIT effects to drug-related and monetary conditioned stimuli were highly correlated. Patients more frequently showed a PIT effect and the effect was stronger in response to aversively conditioned CSs (conditioned suppression), but there was no group difference in response to appetitive CSs.

Conclusion: The implementation of PIT has favorably robust properties in chronic alcohol-dependent patients and in healthy controls. It shows internal consistency between monetary and drug-related cues. The findings support an association of alcohol dependence with an increased propensity towards PIT.

Keywords: Pavlovian-to-instrumental-transfer, alcohol dependence, human
Introduction

Between 75 to 85 % of alcohol-dependent patients relapse after detoxification [1,2]. This is despite their stated desire to remain abstinent, and even in the face of severe consequences during relapse [3,4]. Contextual cues may be particularly important in this process [5-7].

Pavlovian conditioning describes the process by which neutral cues acquire value by predicting the occurrence of a rewarding or punishing event. Such cues often occur in drug-taking environments, and previous studies on both animals and humans have described their role in development, maintenance and relapse of alcohol dependence [8-10].

The dopamine system, which is influenced by most (if not all) drugs of abuse, is known to be key to one type of Pavlovian conditioning termed ‘habitual’ or ‘model-free’ learning [11-14]. Model-free learning depends on iteratively updating expectations through prediction errors (discrepancies between expected and experienced rewards). Phasic dopamine cell firing is known to report such reward prediction errors [15,16] and to be causally involved in Pavlovian conditioning [17]. Recently, work on sign-tracking highlighted the importance of individual variations in this type of learning for addiction, with sign-trackers, who rely more on ‘model-free’ learning based on phasic dopamine teaching signals [18,19] being at increased risk for dependence [20,21]. Hence, variability in how subjects learn about Pavlovian conditioned stimuli is becoming a neurobiologically, clinically and theoretically coherent account for one risk factor for dependence.

One paradigm formalizing the influence of Pavlovian Conditioned Stimuli (CSs) on ongoing instrumental behavior is Pavlovian-to-instrumental-transfer (PIT; e.g. [22]). In general PIT, positive Pavlovian cues enhance instrumental responses while negative Pavlovian cues inhibit instrumental behavior (conditioned suppression) independent of specific outcomes while in specific PIT Pavlovian cues associated with a specific outcome influences ongoing behavior with the same outcome only [23,24]. Animal studies on general PIT have shown the involvement of dopamine and the nucleus accumbens [23,25]; PIT effects are reduced under dopamine antagonists [26-28]; and drug-related cues enhance both behavioral [29-32] and neuronal [33] PIT effects in drug-treated groups. In humans, PIT effects are also measurable and involve a similar set of neural structures (including the amygdala and nucleus accumbens; [34-40]. General PIT thus has important parallels with sign-tracking.

However, how individual differences in PIT are associated with alcohol dependence in humans is not known. Such paradigms involve multiple stages, and are thus both lengthy and complex. Their reliability has not been examined in patient populations. Thus, there is a need for tasks that measure the individual subjects’ tendency towards model-free Pavlovian learning. We here present a PIT task (adapted from Huys, et al. [36] and Geurts el al. [39]) that measures Pavlovian influences on instrumental approach/no approach behavior. The paradigm carefully matches the instrumental expectations of the actions themselves. It includes both Pavlovian stimuli conditioned to predict monetary outcomes, and drug-associated vs neutral stimuli. We expected to see stronger PIT effects both for stimuli predicting monetary outcomes, and for alcohol-related pictures in patients suffering from Alcohol Use Disorder (AUD).
Methods

Participants

In a bi-centre study, we tested 32 recently detoxified alcohol-dependent patients (M_{age}=42.13, SD_{age}=9.78; 29 male) and 32 age and gender matched healthy controls (M_{age}=42.34, SD_{age}=10.29, 29 male). Exclusion criteria were: major psychiatric or neurologic disorders, a history of any dependence syndrome or current substance abuse (assessed by drug urine testing) except nicotine dependence in healthy controls and nicotine and alcohol dependence in patients, intake of medications or drugs known to interact with the CNS within at least four half-lives post last intake (including detoxification treatment with benzodiazepines or chlomethiazole). We included right-handed subjects only. Alcohol dependent patients had a minimum of three years AUD, 72 hours to 21 days of abstinence, as well as a low to moderate severity of withdrawal symptoms (CIWA-Ár <3, [41]). To ascertain inclusion and exclusion criteria we used a computer-based clinical interview (Composite International Diagnostic Instrument, CIDI, [42]). All participants gave written informed consent to participate; ethical approval for the study was obtained from the ethics committee of Charité Universitätsmedizin, Berlin and Universitätsklinikum Dresden. Participants received a monetary compensation for study participation.

Setting

The task was programmed using Matlab with the Psychophysics Toolbox Version 3 (PTB-3, [43,44]) extension. It was presented on a Dell laptop screen (instrumental training, forced choice) and on a projector via a mirror system (Pavlovian training and PIT). Participants performed the PIT task in a functional magnetic resonance imaging (fMRI) setting as part of a larger study examining neural correlates of learning mechanisms in alcohol dependence (see www.lead-studie.de). We here report behavioral data only. The instrumental training was conducted before the scanner session; the Pavlovian and PIT part inside the MRI scanner; and the forced choice task after the scanning session. Participants wore MR compatible SIEMENS headphones; the volume was individually adapted. Responses were made on a 1x4 Current Design MR compatible response box button using the right index finger (instrumental response in training and transfer) or two buttons using the left and the right index finger (forced choice).
Pavlovian-to-instrumental-transfer (PIT) paradigm

The PIT paradigm (adapted from Huys, et al. [36] and Geurts et al. [39]) consisted of four parts: 1) instrumental training, 2) Pavlovian training, 3) Pavlovian-to-instrumental-transfer and 4) a forced choice task (see Figure 1).

Instrumental Training

During instrumental training, subjects were instructed to collect shells by button presses. One instrumental stimulus appeared on the left or right side of the screen (counterbalanced). The subject’s task was to move a dot toward the stimulus by repeated button presses in order to collect it or to do nothing within 2s. These two instrumental choices resulted in monetary wins or losses, presented immediately after each trial via a picture of a 20 Eurocents coin for 1.5s. Feedback was probabilistic. A “good” shell was rewarded in 80% and punished in 20% of trials if collected (go) and vice versa if not collected. A “bad” shell was rewarded in 80% and punished in 20% of the trials if not collected (no-go) and vice versa if collected. The classification of a shell as “good” or “bad” and the order of stimulus presentation were randomized over participants. All six shells consisted of two different colors, were highly

Figure 1. The PIT paradigm consisted of four parts: 1) Instrumental Training: go (1A) or no-go (1B) responses, reinforced by probabilistic outcomes. 2) Pavlovian Training. Audiovisual compound cues (‘fractal CSs’) were associated with one of five outcomes. 3) Pavlovian-to-instrumental-transfer: subjects performed the instrumental task in nominal extinction, i.e. no explicit outcomes were presented. The background was tiled with either drink Cs (top) or fractal CSs (bottom). 4) At the end, subjects performed forced choices between two fractal CSs (4A), a fractal and a drink CS (4B) or two drink CSs (4C).
discriminable with respect to color and shape, and had comparable visual features (such as size and resolution).

The instrumental training comprised a minimum of 60 trials. To match for instrumental performance we established a learning criterion (80% correct choices over 16 trials). Instrumental training ended once participants reached the learning criterion or at a maximum of 120 trials.

**Pavlovian Conditioning**

At the beginning of each trial, a compound conditioned stimulus (CS) consisting of a multicolored fractal-like distorted image on the right or left side of the screen (counterbalanced across subjects, fixed within subjects) paired with a pure tone via headphones was presented for 3s. Henceforth, the combined audiovisual CSs involving fractals will be referred to as ‘fractal CSs’. After a delay of 3s (presenting two fixation crosses at the two potential CS locations), a coin (unconditioned stimulus, US) was presented for a further 3s on the opposite side. Subjects were instructed to observe the CSs and USs and to memorize the pairings. The CS-US association was deterministic and fixed within, but randomized across subjects. The set of stimulus pairings consisted of two positive CSs paired with +2 and +1 Euros, one neutral CS paired with 0 Euros (picture of a zero) and two negative CSs paired with -1 and -2 Euros (coins with superimposed red cross). Subjects completed 80 trials.

To ensure that the Pavlovian stimuli had comparable subjective ratings and visual features, we created a set of 20 pictures by distorting and recoloring food photographs with the GIMP software. All pictures had equal mean luminance values, and equal root-mean-square contrasts of the luminance values [45]. An independent sample of 75 people (42 female, M_age = 29.8 and SD_age = 12.2) rated the pictures on a 7-point-Likert scale according to the 4 dimensions: pleasure, arousal, beauty, and alcohol craving. We chose five highly discriminable CSs with respect to color and shape out of those pictures with equal ratings on all subjective rating dimensions.

**Pavlovian-to- Instrumental-Transfer**

Subjects performed the instrumental task again with either fractal or drink CSs tiling the background. No outcomes were presented, but subjects were instructed that their choices still counted towards the final monetary outcome. There were four drink CSs: two were alcoholic (photographs of the participant’s favourite alcoholic drink) and two neutral (photographs of a water glass; all with homogeneous white background). Drink stimuli were paired with the sound of pouring a drink into a glass. Participants completed 162 trials whereby 9 different background stimuli (5 CSs previously paired with money, 4 pictures of beverages) were shown 18 times each in a pseudorandom order. The response window was 3s with 2-6s inter-stimulus-interval (individually exponentially distributed jitter).

**Forced choice task**

Finally, participants completed a forced choice task. Subjects had to choose one of two compound CSs. All possible CS pairings were presented three times in an interleaved, randomized order and stimuli were presented one at a time for 2s. Slow responses led to a reminder requesting faster responses. We used these data to verify acquisition of Pavlovian expectations.

**Data analysis**

Data were analyzed using Matlab 2011a [46] and IBM SPSS Statistics 20 [47]. When performing ANOVAs, we report Greenhouse-Geisser corrected statistics where appropriate.
(after testing for homogeneity and sphericity using Box’s test of equality of covariance and Mauchly’s test of sphericity).

**Instrumental performance:** we computed a 2x2 repeated measures ANOVA for the number of button presses, both for the instrumental training and in the PIT task. There was one within-subjects factor for approach go vs approach no-go and one between-subjects factor group (healthy controls vs. patients).

**Pavlovian Training:** Individual acquisition of Pavlovian associations was assessed by the fraction of correct answers on the forced choice task using a $\chi^2$-test. Group comparisons were performed with two-sample t-tests or signed rank tests, as appropriate.

PIT data for fractal CSs and drink CSs were assessed separately. The number of button presses was averaged for each instrumental – Pavlovian stimulus combination. We used repeated measures ANOVAs with the number of button presses as dependent variable and two factors (CS and group). The group factor was between-subject. The CSs were coded as within-subjects factors. For fractal CSs there were five levels corresponding to each outcome, for drink CSs there were two levels corresponding to alcohol or non-alcohol. Individual PIT effects were quantified by regressing the mean number of button presses on the value of the CSs and retaining the slope. This was done across all CSs to estimate the overall PIT effect; across neutral and negative CSs for conditioned suppression, and across neutral and positive stimuli for the positive PIT effect. Simple group comparisons were performed by Wilcoxon signed rank test if conditions for t-tests were violated.

To estimate the stability of the PIT effect, we computed the individual PIT effect slopes for the first and second half of the experiment separately, and then correlated these (Pearson linear correlation for Gaussian and Spearman Rank coefficient for non-Gaussian data). Finally, we conducted correlation analyses between the individual regression coefficients calculated for fractal and drink CSs for each subject.
Results

Instrumental behavior

Figure 2. Instrumental performance. Mean number of button presses: subjects learned to collect a “good” shell (reward when collected, go condition) and to leave a “bad” shell (reward when not collected, no-go condition). Results are shown for the instrumental learning task (A) and the PIT task (B). In the PIT task the subjects had to perform the instrumental response they learned in the instrumental training without direct feedback. Black bars represent standard errors of the mean. *** - p<0.001.

Half the patients achieved the instrumental training criterion (after 82.1 trials on average) and 20 out of 32 controls (after 74.7 trials on average). The remaining subjects performed all 120 trials of the instrumental training. Overall, subjects learned to press more when approach was more rewarded (go condition) and to press less when not approaching was more rewarded (no-go condition); $F_{(1, 62)}=54.11, p<0.001$, Figure 2 A). This difference was stably maintained during PIT ($F_{(1, 62)}=33.27, p<0.001$, Figure 2 B). There was no effect of group either during instrumental training ($F_{(1, 62)}=2.42, p=0.13$), or during the PIT part of the experiment ($F_{(1, 62)}=2.26, p=0.14$), and there were no interactions between condition (go/no-go) and group in either training or PIT part ($F_{(1, 62)}=0.84, p=0.37$ and $F_{(1, 62)}=0.01, p=0.94$ respectively).
Forced choices of fractal CSs

Figure 3. Pavlovian conditioning. A: Successful Pavlovian learning is visible in the forced choice task indicating a high preference for the fractals associated with higher outcomes. Bars show mean probability of better stimulus for those subjects performing above chance (black dots). Red crosses show subjects performing at or below chance. B: Choice probabilities involving drink CSs in the forced choice task, comparing fractals CSs with drink CSs or comparing two drink CSs. Alc: alcoholic drink; Wtr: water; Alc>CS: probability of choosing drink of fractal CS; Wtr>CS: probability of choosing water over fractal CS. Alc>Wtr probability of choosing alcoholic over water CS. Black bars represent standard errors of the mean. ** - p<0.01.

To assess the acquisition of Pavlovian values, we analysed preferences in the forced choice trials involving only pairs of fractal CSs (Figure 3 A). Twenty-eight out of 32 patients and the same number, 28 out of 32 controls, preferred higher-valued fractal CSs overall (with individual p<0.05). Patients and controls chose the better of the two fractals on 88.21+/-0.02% and 89.42 +/- 0.03% of the trials, respectively. There was no significant group difference (p=0.95, z=-0.06, rank sum=987, Wilcoxon rank sum test). In one healthy control the forced choice data were missing. This and the eight subjects who failed to show preference for fractals CSs with higher associated value were excluded from the PIT analyses as Pavlovian conditioning was uncertain.

Forced choices involving beverage CSs

Subjects also performed forced choices involving pictures of their favorite drink and/or water and of drinks paired with fractal CSs (Figure 3 B). Patients chose fractals over alcoholic CSs more than controls (p<0.01, z=-2.65, rank sum=639, Wilcoxon rank sum test; Bonferroni correctable for 3 comparisons), but otherwise did not differ from controls.
PIT effects: fractal CSs

There was a significant main effect of fractal CSs ($F_{(1.301, 53)}=28.14, p<0.001$, Greenhouse-Geisser corrected), but this did not differ between groups (fractal CS x group $F_{(1.301,53)}=0.78, p=0.412$, Greenhouse-Geisser corrected, Figure 4 A). Exploratory analyses of the fractal CS effects on button press responses only during go or during no-go also failed to yield group differences. The effect of fractal CSs was present both in subjects who had ($F_{(1.267,53)}=14.76, p<0.001$, Greenhouse-Geisser corrected) and who had not ($F_{(1.33,53)}=11.87, p<0.01$, Greenhouse-Geisser corrected) reached the instrumental criterion.

As a measure of individual PIT effects, we calculated linear regression coefficients between the number of button presses and the fractal CS value for each individual separately. This was individually significant ($p$ values < 0.05) in 17 out of 28 patients (61 %, Figure 4 C) and in 10 out of 27 healthy controls (37 %, Figure 4 D). This fraction was trend wise different between groups ($\chi^2=3.08, p=0.08$).

PIT is usually only considered with appetitive conditioned stimuli, while the aversive side is typically examined in separate experimental setups in conditioned suppression. This motivates a separate analysis of the positive and negative limbs of the experiment. We therefore computed separate linear regression coefficients for the positive and the negative CSs, always including the neutral CS. Patients showed a stronger effect of negatively valued fractal CSs (conditioned suppression) than controls ($p<0.05$, $z=-2.34$, rank sum=644;
Bonferroni correctable for 2 comparisons, Figure 4 B). Groups did not differ in terms of the effect of positive fractal CSs \( (p=0.97, z=-0.04, \text{rank sum}=781) \).

**PIT effects: beverage CSs**

![Figure 5. PIT effect for beverage CSs. A: Alcoholic drink CSs reduce responding compared to neutral water drink CSs. B: The mean (light grey bar) and individual (dots) difference between water and alcohol cues. Light grey dots represent individuals not showing a significant PIT effect, dark grey dots represent individuals with a significant effect \((p<0.05)\). Error bars represent standard errors of the mean.](image)

Two outliers (one patient and one control) were outside the range of 3 standard deviations around the mean and therefore removed. There was a main effect of drink CSs, with alcohol CSs reducing responses compared to water CSs \( (F_{(1,51)}=10.4, p<0.01, \text{Figure 5A}) \). The effect did not differ between groups (drink CS x group; \( F_{(1,51)}=0.28, p=0.599 \)) and was present both in patients \( (F_{(1,26)}=7.79, p<0.05) \) and marginally in controls \( (F_{(1,25)}=3.28, p=0.082) \).

We again computed individual PIT scores by fitting linear regressions. These were individually significant in 8 of the 28 patients (28.57 %; Figure 5B) and in 2 out of 27 healthy controls (7.41 %; Figure 5C). Mirroring the fractal CS result, there was a group difference in the proportion of subjects showing a drink CS effect \( (\chi^2=4.14, p=0.04) \).

**Temporal stability of PIT effects**

Individual regression coefficients for PIT effects with fractal CSs were computed separately on the first and second half of the experiment. The correlation between these was very high \( (\rho=0.79, p<0.001, \text{Spearman; Figure 6 A}) \), suggesting a temporally very stable PIT effect.
This remained true when considering patients ($\rho=0.78$, $p<0.001$; Spearman) and controls ($\rho=0.77$, $p<0.001$; Spearman) separately.

Similarly, PIT effects for drink CSs were stable between first and second halves when collapsing across groups ($\rho=0.34$, $p<0.05$, Spearman, Figure 6 B) or when considering patients ($\rho=0.52$, $p<0.01$, Spearman) alone. In contrast, there was no evidence for a stable drink CS PIT effect in controls ($\rho=0.14$, $p=0.5$, Spearman), though the group difference failed to reach significance (Fisher’s $z=1.52$, two-tailed $p=0.13$).

**PIT effects for fractal and drink CSs are correlated**

Finally, we asked whether subjects who showed a PIT effect in response to fractal CSs would also tend to show a PIT effect in response to drink CSs. Individual PIT effects in
response to drink and fractal CSs were highly correlated. This was true when collapsing across groups ($\rho=0.45$, $p<0.001$; Spearman; Figure 7), or when considering patients ($\rho=0.47$, $p<0.05$, Spearman) or controls ($\rho=0.40$, $p<0.05$, Spearman) individually.
Discussion

The current study was a pilot study to establish the feasibility of measuring the influence of Pavlovian processes on on-going behaviour in alcohol-dependent patients in comparison to healthy controls (see www.lead-studie.de). We chose Pavlovian-to-instrumental-transfer due to its substantial pre-clinical evidence base relating both to addiction and dopaminergic processes [22,48]. However, it is a complex paradigm, which has not always shown strong behavioral effects (e.g. [35], particularly inside the MR scanner), and as such it was necessary to first establish the feasibility of such a task in a challenging patient population. The three main conclusions of this pilot study are that i) PIT effects can indeed be measured behaviourally in a stable and reliable manner; ii) that PIT effects in response to drug-relevant and monetary conditioned CSs are consistent; and iii) that recently detoxified patients suffering from AUD and healthy controls differ modestly in the propensity to show PIT effects, and in the strength of conditioned suppression (negative PIT).

For task measures to have clinical validity, it is critical that they are reliable and valid within the whole cohort, and show substantial variability between individuals in a manner that is consistently related to the task construct. First, we found that individual PIT effect strength is stable over time, being highly correlated between the two halves of the experiment. Although this is very encouraging, it does not replace the need for assessing test-retest reliability with an intervening interval. The fact that the drink CS effect in controls was not stable is most likely due to poor power as only two subjects showed an individually significant effect. Overall, stability was comparable in patients and controls. Second, we attempted to measure the internal consistency of the PIT effect by correlating the effects of Pavlovian CSs predictive of monetary outcomes, and CSs representing subjects’ favourite drinks. We found that these effects correlated highly both in patients and controls, suggesting that there is one central process determining the extent to which an individual subject shows PIT effects. This correlation may also partially address issues concerning the nature of the PIT effect. Both general and specific PIT processes could contribute to the effect of the conditioned stimuli, given that instrumental behaviors were also rewarded with monetary outcomes. However, this is not possible for the drink CSs, and the strong correlation between the two effects suggests that the PIT effect in response to fractal CSs does at least contain a substantial general component. On an individual level, PIT effects varied very substantially between subjects, with around 60% in patients and 37% in healthy controls showing individually significant effects. This high variability between subjects might be useful if related to longitudinal outcomes of interest, but here led to rather small group effects. The present task design was based on Huys et al. [36] and Geurts et al. [39]. While the PIT effect strength and variability across subjects is comparable with the first, it is substantially stronger than the latter. Both previous studies employed healthy controls; but only the latter was performed inside a MRI scanner. It is as yet unclear precisely what factors in the tasks account for the variability across studies.

The second aim was to explore how AUD affected PIT. The current results suggest a heightened propensity to PIT, and a stronger negative PIT effect (conditioned suppression). Against our expectations, we were not able to show any difference in appetitive PIT. Theories of addiction put emphasis on striatal and prefrontal mechanisms [49], with the striatal component particularly involved in a shift towards habitual [50] or model-free [11,51-53] decision-making, and the pre-frontal component resulting in a concomitant impairment of goal-directed decision-making (e.g. [54]; Sebold et al., this issue). Pavlovian values are attached to stimuli, unlike instrumental values, which are attached to action or
stimulus-action pairs. However, Pavlovian values can be both model-free and model-based [14,55,56], producing general and specific PIT, respectively, and relying differentially on the nucleus accumbens core and shell [23]. In the setting of alcohol addiction, both model-based and model-free decision-making systems likely play important parts. On the one hand, recent work on sign-tracking [57] suggests a particularly important role for model-free contributions to behaviour in the nucleus accumbens core. On the basis of this, one might expect a strengthened general appetitive PIT effect to identify a certain vulnerability to drug addiction. On the other hand, work particularly involving cocaine [54-56,58] might similarly argue for a stronger outcome-specific PIT component. Indeed, in the sister paper (Sebold et al., this issue), we find on another task, that alcohol dependence results particularly in impairments of goal-directed decisions after non-rewards. We are not aware of work on conditioned suppression in AUD, and the results hence certainly need replication. However, we have recently found that dietary serotonin depletion selectively affects aversive PIT, and it is tempting to relate our current findings to emerging arguments about the involvement of serotonin in addiction [59].

Rather unsurprisingly, patients but not controls chose fractal CSs over alcoholic pictures in the forced choice tests. This may be influenced by social desirability (having undergone a recent detoxification), but also by currently active explicit motivation to abstain. Strikingly, however, alcohol-related CSs also appeared to suppress, rather than enhance responding during PIT. This finding appeared to hold for both patients and controls and also attests to the consistency between preferences and PIT effects. As we pointed out, Pavlovian CSs can derive value both through model-free and model-based mechanisms. However, an outcome-specific model-based PIT account of this is difficult as instrumental and Pavlovian CSs were not associated with or predictive of the same outcome. Accounting for this with a general PIT, model-free mechanism is equally difficult and would run counter to the long-standing view that drug-cues are appetitive. One answer to this might come from the temporal stability of the effect, comparing it right after detoxification and some time later. Some of the subjects failed to acquire the Pavlovian contingencies. This may be due to either the delay between the training and the forced choice task (approximately 25 minutes apart; see also Trick, et al. [34]). However, a caveat concerning the forced choice procedure is that it may tap explicit rather than implicit [60] processes, and that subjects might show evidence of PIT even in the absence of explicit knowledge (see e.g. [61]). However, the numbers were too small to clearly show this.

Given possible gender differences with respect to responsiveness to alcohol cues [62], we decided to focus on male participants in our pilot sample. Whereas males have a higher risk of developing alcohol dependence, lifetime prevalence numbers in females varying between 2 and 8% in Europe and the USA (e.g. [63,64]) suggest that the identification of further gender effects with respect to contextual cues is of importance. In terms of recruiting a representative sample of alcohol dependent patients we will include women in future samples; however, at this point, gender effects were beyond the scope of this pilot study.

In conclusion, these pilot results suggest that our PIT paradigm is suitable for testing alcohol dependent patients and healthy controls. We observed a stronger PIT effect in patients suffering from AUD compared to healthy controls and a high interindividual variance between subjects, which is an important factor for further studies on predicting the development and maintenance of AUD. Therefore our PIT task might allow insights into the decision-making structure underlying the disease.
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