

Please cite the Published Version

Harkki, Juliana, Tuovinen, Pauli, Jousimäki, Veikko, Barreto, Goncalo, Rapeli, Pekka, Palomäki, Jussi, Annevirta, Jonne, Puisto, Anna-Helena, McGlone, Francis P, Nieminen, Heikki and Alho, Hannu (2024) CT-optimal touch modulates alcohol-cue-elicited heart rate variability in Alcohol Use Disorder patients during early abstinence: a randomized controlled study. Alcohol. ISSN 0741-8329

DOI: https://doi.org/10.1016/j.alcohol.2024.11.004

Publisher: Elsevier

Version: Accepted Version

Downloaded from: https://e-space.mmu.ac.uk/637563/

Usage rights:

CC BY

Creative Commons: Attribution 4.0

Additional Information: This is an open access article published in Alcohol, by Elsevier.

Enquiries:

If you have questions about this document, contact openresearch@mmu.ac.uk. Please include the URL of the record in e-space. If you believe that your, or a third party's rights have been compromised through this document please see our Take Down policy (available from https://www.mmu.ac.uk/library/using-the-library/policies-and-guidelines)

CT-optimal touch modulates alcohol-cue-elicited heart rate variability in Alcohol Use Disorder patients during early abstinence: a randomized controlled study

Juliana Harkki, Pauli Tuovinen, Veikko Jousimäki, Goncalo Barreto, Pekka Rapeli, Jussi Palomäki, Jonne Annevirta, Anna-Helena Puisto, Francis McGlone, Heikki Nieminen, Hannu Alho

PII: S0741-8329(24)00177-0

DOI: https://doi.org/10.1016/j.alcohol.2024.11.004

Reference: ALC 7337

To appear in: Alcohol

Received Date: 26 July 2024

Revised Date: 26 October 2024

Accepted Date: 21 November 2024

Please cite this article as: Harkki J., Tuovinen P., Jousimäki V., Barreto G., Rapeli P., Palomäki J., Annevirta J., Puisto A.-H., McGlone F., Nieminen H. & Alho H., CT-optimal touch modulates alcohol-cue-elicited heart rate variability in Alcohol Use Disorder patients during early abstinence: a randomized controlled study *Alcohol*, https://doi.org/10.1016/j.alcohol.2024.11.004.

This is a PDF file of an article that has undergone enhancements after acceptance, such as the addition of a cover page and metadata, and formatting for readability, but it is not yet the definitive version of record. This version will undergo additional copyediting, typesetting and review before it is published in its final form, but we are providing this version to give early visibility of the article. Please note that, during the production process, errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

© 2024 The Author(s). Published by Elsevier Inc.



CT-optimal touch modulates alcohol-cue-elicited heart rate variability in Alcohol Use Disorder patients during early abstinence: a randomized controlled study

<u>Juliana Harkki</u> 1,*,@, Pauli Tuovinen 1,@, Veikko Jousimäki 1,@, Goncalo Barreto 1,2,@, Pekka Rapeli 3,@, Jussi Palomäki 4,@, Jonne Annevirta 1,@, Anna-Helena Puisto 1,@, Francis McGlone 1,5@, Heikki Nieminen 1@, Hannu Alho 2,@

1: Department of Neuroscience and Biomedical Engineering, Aalto University, Finland

2: Clinicum, Faculty of Medicine, University of Helsinki and Helsinki University Hospital, Finland

3: Department of Psychiatry, Helsinki University Central Hospital, Finland

4: Department of Public Health and Welfare, Finnish Institute for Health and Welfare, Finland

5: Faculty of Science & Engineering, School of Life Sciences, Manchester Metropolitan University, UK

* : Corresponding author

Corresponding author: Juliana Harkki Juliana.harkki@aalto.fi +358405609757 Agricolankatu 11 C 33 00530 Helsinki Finland

Declaration of interest: JH and PT hold a patent for a mechanism for a CT-optimal haptic device (IPID3248). Other authors have no interests to declear.

Abstract

Alcohol Use Disorder (AUD) is a chronic brain disorder associated with a high risk of relapse and a limited treatment efficacy. Relapses may occur even after long periods of abstinence and are often triggered by stress or cue induced alcohol craving. C-tactile afferents (CT) are cutaneous nerve fibers postulated to encode pleasant affective touch and known to modulate physiological stress responses. However, their translational potential has not yet been explored extensively in controlled clinical trials. This randomized controlled study aimed to investigate the potential of CT stimulation in modulating relapse predicting biomarkers, physiological cue-reactivity, and subjective alcohol craving in AUD patients in early abstinence.

Twenty-one participants meeting DSM-5 criteria for mild to moderate AUD received CToptimal touch or a non-CT-optimal control treatment while exposed to neutral, stressinducing, and alcohol-related visual stimuli. The tactile treatment was provided with a robotic device, eliminating the social elements of touch. Heart rate variability (HRV), salivary cortisol, and subjective craving were assessed at the baseline, during and after the treatment and stimuli exposure.

The results showed that CT-optimal touch significantly reduced alcohol-cue-elicited standard deviation of normal-to-normal intervals (SDNN) HRV compared to the control group, shifting the HRV reactivity to the direction known to indicate lower relapse susceptibility. Cortisol levels showed no significant differences between the groups, and subjective alcohol craving increased after alcohol cue exposure in both groups.

This study found that CT-optimal touch modulates autonomic cue-reactivity in AUD patients, encouraging further research on the therapeutic potential of affective touch. Future research should explore the long-term effects and real-world clinical relevance of CT-optimal touch in alcohol relapse prevention.

Keywords: Affective touch, Alcohol Use Disorder, C-tactile afferents, cue-reactivity, craving, heart rate variability, stress, randomized controlled trial.

1 Introduction

2

Alcohol use disorder (AUD) is a chronic brain disorder with a high risk of relapse even after
long periods of abstinence. Eventual relapse occurs in up to 75% of the patients attempting to
quit drinking with current pharmacological and psychosocial treatments (Boothby & Doering,
2005). The high relapse rate indicates the critical need for novel interventions. However, few
relapse-preventing therapies have emerged since the approval of opioid receptor antagonists,
such as naltrexone, nearly 30 years ago (Center for Substance Abuse Treatment, 2009).

One of the characteristic symptoms of AUD and an important contributor to relapses is 10 11 alcohol craving (Koob & Volkow, 2016; Sinha, 2013b). Craving may be triggered by 12 alcohol-related cues, i.e. incentive saliences that precipitate the abnormally high 13 dopaminergic activity in the nucleus accumbens (NAc) associated with the hedonic state of 14 alcohol intake. Another equally important trigger for craving is negative mental stress, 15 involving the hypothalamic pituitary adrenal (HPA) axis, amygdala and peripheral stress 16 responses (Koob, 2013; Koob & Volkow, 2016; Sinha, 2013b). Exposure to chronic stressors 17 or aversive life events is known to increase the risk of harmful use of alcohol and the severity 18 of AUD (Sinha, 2022). Moreover, aversive stress-related symptoms, such as anxiety, 19 negative mood and aggressive behaviour, are typical in early abstinence from alcohol, which 20 may further increase craving and the risk of a relapse (Koob & Volkow, 2016; Sinha, 2013a). 21 In short, stress has a bidirectional impact on alcohol craving and the occurrence of relapses; it 22 may trigger craving, leading to the harmful use of alcohol, whereas the harmful use of 23 alcohol may generate the symptoms of stress contributing to the vicious cycle of addiction. 24

Neural adaptations resulting from a chronic use of alcohol alter both peripheral and central
stress systems, manifesting, for instance, as dysregulation of the autonomic nervous system
(ANS) (Blaine et al., 2017; Koob, 2013; Sinha, 2009; Sinha et al., 2009). This neural
dysregulation is illustrated clearly by heart rate variability (HRV), an established biomarker
of ANS activity. Generally, in a healthy population, a higher resting state HRV is considered
to reflect higher activity of the parasympathetic division of ANS, thus indicating a lower
stress level. Correspondingly, stress exposure typically leads to a decrease in HRV. However,

32 numerous studies have shown that AUD patients have an abnormally low resting state HRV,

33 yet it increases when exposed to stressors or alcohol-cues (for reviews see Cheng et al., 2019;

Ralevski et al., 2019). Furthermore, heightened high-frequency HRV in response to alcohol-

35 cues predicts susceptibility to relapse in abstinent AUD patients, suggesting that HRV may

36 be used as a relapse predicting biomarker (Garland et al., 2012).

37

38 As stress has been recognized as one of the key factors in alcohol craving and relapses,

39 treatments targeting stress-related mechanisms have attracted research interest in recent years.

40 However, to the best of our knowledge, no previous studies in this field have addressed the

41 potential stress regulating properties of somatosensory systems, namely C-tactile afferents

42 (CT).

43

44 CTs are slowly conducting unmyelinated low-threshold mechanoreceptor nerve fibers that 45 are postulated to code for the pleasant (rewarding) properties of social and affective touch 46 (Löken et al., 2009; McGlone et al., 2014; Olausson et al., 2016; Pawling et al., 2017). The 47 optimal range of tactile stimulus velocity for CT firing is from 1 to 10 cm / s; both slower and faster speeds resulting in a weaker response in the firing rate of CTs during 48 microneurography experiments (Löken et al., 2009). This velocity is also typical for gentle 49 50 social touch, such as a caress. For instance, Püschel et al (2022) showed that mothers who 51 were asked to stroke their preterm infants, did so in the CT-optimal velocity range. Further 52 microneurography studies have shown that the firing frequency of CTs positively correlates 53 with perceived pleasantness of skin stroking (Essick et al., 1999; Essick et al., 2010; Löken et 54 al., 2009). These observations are supported by a study by Morrison et al. (2011), which 55 showed that patients with a genetic deficit of C-fibers rated CT optimal skin stroking as less 56 pleasant than did the control group of healthy individuals.

57

58 CTs interact with several neurobiological mechanisms associated with AUD. Firstly, they 59 project to the insular cortex (IC), a region of high importance in interoception and affective processes, also known to be a crucial brain site in modulation of cue-induced substance 60 61 craving (Björnsdotter et al., 2009; Davidovic et al., 2019; Löken et al., 2009; Naqvi et al., 2014; H. Olausson et al., 2002; Olausson et al., 2016; Pawling et al., 2017). The IC has 62 functional connectivity with brain regions central to reward, motivation, and addictive 63 64 processes, influencing dopaminergic activity within the ventral tegmental area and NAc 65 (Girven et al., 2020). The elevated dopamine levels in the NAc also occur as a result of tactile

skin stimulation, further highlighting the effect of affective touch on reward pathways

67 (Maruyama et al., 2012). Moreover, affective social touch, typically applied at CT-optimal

68 velocity and force, modulates the endogenous μ -opioid system, known to have a pivotal role

69 in alcohol-induced dopamine increase and the pleasurable effects of alcohol (Gilpin & Koob,

- 70 2008; Korpi et al., 2015; Nummenmaa et al., 2016).
- 71

72 Crucially, CT-optimal touch seems to effectively modulate stress responses. Hence the 73 affective touch system has been suggested as an indispensable component of our physiological stress regulation (Morrison, 2016; Blaine et al., 2017; Pawling et al., 2017b). 74 75 One of the most intriguing examples of this is a study by Walker et al (2020), which 76 demonstrated that a mere 10 minutes of slow stroking touch per day had a striking effect on 77 stress-resilience in rats, nearly abolishing the anxiety-related behaviour and corticosterone 78 increase in a rodent model of chronic mild stress. Human studies have shown that CT-optimal 79 and affective touch have an ability to both increase oxytocin and to lower cortisol level, as well as to reduce peripheral stress system activity manifesting, for instance, as a lower heart 80 rate (Ditzen et al., 2007; Eckstein et al., 2020; King & Becker, 2019; Püschel et al., 2022; 81 82 Uvnäs-Moberg et al., 2014; Walker et al., 2017). Importantly, CT-optimal touch has been identified as a potent modulator of HRV. For instance, a study by Triscoli et al. (2017) 83 84 showed that HRV of healthy subjects increases with robotic CT-optimal stroking touch. 85 Moreover, a recent study demonstrated that dynamic touch at CT-optimal velocity, increased 86 the HRV of preterm infants, whereas static touch had no such effect (Manzotti et al., 2023). However, to date, no published data exists on the potential effects of CT-optimal touch on 87 88 HRV and ANS regulation in the AUD patient population.

89

90 Building on this foundation, we hypothesize that the rewarding and stress-regulating effects 91 of CT-optimal touch may influence physiological cue-reactivity, subsequently reducing 92 alcohol craving and preventing relapses. To test this, we investigated the impact of acute CT-93 optimal touch on HRV reactivity, salivary cortisol, and self-reported craving of early 94 abstinent AUD patients in stress- and alcohol-cue exposure. Comparable protocols have previously been used in drug research (e.g. Fox et al., 2012), however, instead of 95 pharmacological intervention, in this body of work, we explored the effect of CT-optimal 96 tactile stimulation. 97

99 Materials and methods

100

101 Experimental design

This study was a randomized single-blinded controlled trial with two parallel treatment
groups (CT-optimal treatment and non-CT-optimal control treatment). The study was
conducted in accordance with the Declaration of Helsinki and European Union's General
Data Protection Regulation. The research plan was pre-evaluated and approved by the
Helsinki University Hospital Regional Committee on Medical Research Ethics
(HUS/11938/2022).

108 Participants

109

Twenty-one participants (10 female) of ages 27 to 60 (mean 48) took part in the study. The 110 111 participants were randomly assigned to active treatment (CT-optimal touch, n=11) and 112 control treatment (non-CT-optimal touch, n=10) groups. All participants met the Diagnostic and Statistical Manual of Mental Disorders, 5th Edition (DSM-5) criteria for mild to 113 114 moderate Alcohol Use Disorder (AUD) and had maintained a minimum of 14 days without engaging in heavy alcohol consumption. Heavy alcohol consumption was defined as ≥ 60 g 115 116 of absolute alcohol per day for males and ≥ 40 g per day for females. The participants were 117 recruited via social media.

118

The participants were instructed to avoid caffeinated drinks for 24 hours, physical exercise 119 120 for 3 hours and eating, drinking and smoking for 1 hour before the experiment. On attending 121 the test facility, the participants signed the informed consent and were familiarized with the 122 protocol. A breathalyser test was performed to confirm the breath alcohol concentration was at the undetectable level and absence of withdrawal symptoms was ensured with Clinical 123 124 Institute Withdrawal assessment for Alcohol questioner (Ciwa-ar) (Sullivan et al., 1989). The 125 heavy drinking days of the past 6 weeks were recorded in the Timeline Follow Back Calendar (TLFB) (Sobell & Sobell, 1990) and Obsessive Compulsive Drinking Scale (OCDS) (Anton 126 127 et al., 1996) was used to assess the overall recent alcohol craving (mean total score 18.05)

129 The exclusion criteria included diagnosis of concomitant psychiatric disorders, current use of

addictive substances other than alcohol (excluding nicotine), mood-regulating medications,

131 recent participation in a treatment program for alcohol use disorders, autism spectrum

132 disorder (score \geq 26 in Autism-Spectrum Quotient test, (Baron-Cohen et al., 2001)),

133 dermatological conditions and cardiovascular or other clinically relevant unstable or

134 untreated illnesses.

135

136 Treatments

137

To eliminate the social aspect present in human touch, we used a robotic device to provide 138 the CT-optimal dynamic touch. Both CT-optimal touch treatment and the non-CT-optimal 139 control treatment were performed with the same robotic device that provided dynamic back-140 141 and forth strokes with a soft cosmetic brush on the skin of the left (non-dominant) forearm of the participants. The CT-optimal treatment was performed at a velocity of 2.5 to 3 cm/s and a 142 143 force of 200 to 300 mN. The non-CT-optimal control treatment was performed with velocity of 0.1 to 0.2 cm/s and a force similar to the CT-optimal treatment. The device was visible to 144 the participants during the experiment. 145

146

147 The custom-made robot used for producing the haptic stimulation was designed in a similar way as in the study by Eriksson Hagberg et al (2019) in which a similar stimulator was used 148 149 to study activation of skin nerve fibers and related brain activity. Two pneumatic muscles were used to ensure the controlled distance, force, and velocity. The pneumatic muscles were 150 151 operated with compressed air that was adjusted with a pneumatic control system located at a 152 safe distance from the participant. The muscles were reinforced and supported with glass-153 fiber composite tubes and plastic polypropylene connectors. The stimulator was equipped with an optoswitch to measure the velocity of the brush, and a load cell to measure the 154 155 applied force on the skin and to identify the skin contact. A photograph of the robot is shown in Supplement 1. 156

157

158 Stimuli

159

160 To expose participants to triggers known to induce alcohol craving, the subjects were

161 presented with alcohol-related and stress-inducing images. Neutral images that typically do

162 not induce craving, were shown as a control condition. The images for the neutral and stress stimuli were chosen from International Affective Picture System database (IAPS)(Lang, P.J. 163 164 Bradley, M.M., & Cuthbert, 2008). Each set of images contained 30 pictures of one type of stimulus, each image being visible for 10 s. The images were presented on a 27 inch monitor 165 (ThinkVision E27q-20, Lenovo, Quarry Bay, Hong Kong) located approximately 60 cm 166 away from the participant. All participants were exposed to all three types of stimuli on 167 168 separate consecutive sessions in the following order: neutral, stress-inducing, alcohol-related. 169 The mean valence/arousal ratings for the selected neutral IAPS images were 6.4/3.3 and the 170 images contained, for instance, landscapes and details of nature. The mean valence/arousal ratings for the selected stress-inducing images were 2.3/6.3 (range 1 to 9) and the images 171 172 included, for instance, violence, mutilated or severely damaged human bodies and attacking animals. The alcohol-related images were chosen from The Geneva Appetitive Alcohol 173 174 Picture database (GAAP) (Billieux et al., 2011), which contain images of alcoholic drinks, 175 bars, night clubs, and people drinking. The alcohol-related visual material was adapted to suit 176 the participant's personal preference by ensuring that the material contains images of their preferred type of alcoholic beverage. In addition, the material was modified to be 177 recognisable for the Finnish AUD patients, for example by replacing an image of a Swiss 178 179 beer bottle with a Finnish equivalent sourced from the internet.

180 HRV reactivity

To assess autonomic functioning, HRV was derived from blood volume pulse (BVP) 181 182 measured with Empatica E4 wrist band device (Empatica Inc., Milan, Italy) throughout each session excluding a 15-minute relaxation at the end of the session (Fig. 1). Of many HRV 183 184 parameters, standard deviation of normal-to-normal intervals (SDNN) was selected, as CToptimal touch has been previously show to modulate SDNN (Triscoli et al., 2017). Kubios 185 186 HRV Scientific software (Kubios Oy, Kuopio, Finland) was used to analyse HRV from the 187 BVP data using automatic artefact correction. We analysed the mean SDNN HRV of four 5-188 minute blocks from the following time points of each session: baseline, treatment minutes 5 to 10, visual stimulation, treatment minutes 20 to 25 (recovery). To determine the HRV 189 190 reactivity of each participant, the change of the SDNN HRV was calculated for each session by subtracting the baseline value from the stimulation value and for the whole experiment by 191 192 subtracting the baseline value of the first session from the values of each time point of the 193 whole experiment.

194 Subjective alcohol craving

- 195 To assess the subjective alcohol craving, we asked the participants to rate their current
- 196 craving on a 10-point horizontal visual analogue scale (VAS). The rating was performed at
- 197 three timepoints per session; the baseline, after the visual stimuli and at the end of the
- treatment (Fig. 1.). The lowest value (point 1) on the VAS equalled the statement "If alcohol
- 199 *was now available, I would not want to drink it at all"*, whereas the highest value (the point
- 200 10) equalled the statement "If alcohol was now available, I would not be able to resist
- 201 *drinking even if I tried*". The VAS was presented to the participant on the same screen as the
- visual stimuli and the participant gave the rating by saying aloud the number corresponding
- 203 the current state of their subjective alcohol craving.

204 Salivary cortisol level

To assess the salivary cortisol levels, saliva samples were collected 5 times during each 205 206 session (total 15 samples per participant) (Fig. 1.). The time points of the saliva collection 207 were the baseline, after the first 10 minutes of the treatment, after the visual stimuli, at the end of the treatment and after the 15-minute relaxation period. The saliva was collected by 208 209 placing a synthetic swab (Salivette Cortisol, Sarstedt Inc. Nümbrecht, Germany) between the 210 tongue and the cheek of the participants for minimum 2 minutes after which it was placed in a plastic tube (Salivette) and stored in -20 °C until the analysis. Once the samples were 211 thawed, they were centrifuged in 1000 x g for 2 minutes. The samples were analysed with a 212 213 competitive enzyme-linked immunosorbent assay (Cortisol free in Saliva ELISA DES6611, 214 Demeditec Diagnostics GmbH, Kiel, Germany) and the optical density was determined at 450 nm with a well-plate reader (FLUOstar Omega, BMG Labtech, Ortenberg, Germany). If 215 216 the sample had less than 50 µl of saliva it was diluted to 1:2 or 1:5 ratio with Calibrator 0. 217 The concentrations were interpolated from the absorbance values with GraphPad Prism v. 218 10.1.1 software (GraphPad Software LLC, Boston, United States) using 4PL curve fit.

219 Procedure

220 The experiment included three consecutive sessions in which the participants were exposed

- to neutral, stress-inducing and alcohol-cue images with one type of stimuli per session while
- they received either the CT-optimal or the control treatment. The structure of one session is
- illustrated in Fig. 1.

224 At the start of each session, the participants were instructed to sit still in a comfortable position avoiding any movement. To block the noise from the pneumatic device participants 225 226 were asked to wear earplugs. Each session started with a saliva sample and a VAS rating followed by the baseline BVP measurement. After this, the treatment was started and 227 228 continued for 25 minutes including the 5-minute visual stimuli prestation in the middle. The 229 treatment was followed by a 15-minute relaxation during which the participant was allowed 230 to stand up, walk and move lightly. Total length of one session was approximately 47 231 minutes.

232 ** Fig 1. **

After the three sessions participants had a chance to participate in a voice-guided relaxation session to ensure the absence of a possible craving or stress triggered by the experiment. In addition, all participants were offered a consultation of an addiction specialist physician after their participation.

The study included a second visit that contained a cue-reactivity task test with pupillary reactivity and EEG measurements. The second experiment was followed by a two-week follow-up period during which the participants were asked to report the daily subjective craving, alcohol consumption and possible adverse events by filling an online diary. The results of the second experiment, as well as the follow-up, will be reported in separate publications.

243 Statistical analysis

244

245 Statistical analysis was performed with GraphPad Prism v. 10.1.1 software (GraphPad Software LLC, Boston, United States) software. Normality of the HRV reactivity data of 246 each session was confirmed and the data was analysed with an unpaired t-test. The change of 247 the mean HRV, the mean VAS craving ratings and the mean cortisol levels from the initial 248 249 baseline throughout the whole experiment (all three sessions and all time points) were 250 analysed independently with a repeated measures two-way analysis of variance (ANOVA). A 251 multiple linear regression analysis was conducted with HRV reactivity of the alcohol-cue 252 session as the dependent variable and treatment, initial baseline HRV, number of heavy 253 drinking days in TLFB, OCDS score, the mean change of the craving ratings in VAS in 254 alcohol-cue session and the mean initial baseline craving as independent variables.

255	
256	Results
257	
258	HRV reactivity
259	
260	The HRV of two participants from the CT-optimal treatment group could not be analysed due
261	to the irregular BVP data. Hence, the data of 9 participants from the CT-optimal group and
262	10 from the control group were included in the analysis. The results of the unpaired t-tests of
263	each sessions mean HRV change from the session's baseline to the stimulation exposure (Fig.
264	2) showed that the mean SDNN HRV of the CT-optimal treatment group was significantly
265	reduced compared to the control group in alcohol-cue exposure ($R^2 = 0.3$, 95% CI -22.71 to -
266	2.399, $p = 0.0184$). There were no differences in the mean HRV changes between the groups
267	in neutral or stress-inducing stimulation. The individual values are visualized in Supplement
268	2. The multiple linear regression analysis confirmed that treatment was the only factor having
269	a significant interaction with the alcohol-cue elicited HRV change while neither of other
270	analysed variables (the initial baseline HRV, number of heavy drinking days in TLFB, OCDS
271	score, the mean change of the craving ratings in VAS in alcohol-cue session nor the mean
272	initial baseline craving) were significantly associated with it.

273

274 ** Fig. 2 **

275

The mean HRV increased from the initial baseline of the experiment in both treatment groups 276 277 during the neutral and stress-inducing sessions (Fig. 3). In the control group the increase was seen also during the alcohol-cue session. The results of the repeated measures two-way 278 279 ANOVA showed that in the control group the chance of the HRV from the baseline was 280 statistically significant (p < 0.05) at all analysed time points excluding the baseline measurement of the alcohol-cue session. In the CT-optimal treatment group the HRV was 281 significantly increased from the baseline from the second timepoint of the stress session to 282 283 the baseline of the alcohol-cue session.

284

285 ** Fig 3. **

287 Subjective craving

288

289	The mean ratings of subjective alcohol craving were significantly higher after alcohol-cue
290	exposure than after the neutral stimuli exposure in both treatment groups based on the
291	repeated measures two-way ANOVA (CT-optimal group CI 95% -1.964 to -0.2173, $p =$
292	0.0194, control group CI 95% -2.469 to -0.5314, $p = 0.0067$) (Fig. 4). In the CT-optimal
293	treatment group the mean rating after the presentation of alcohol-cues was also significantly
294	higher than the mean craving after the stress stimuli exposure (CI 95% -1.901 to -0.09867, p
295	= 0.0330). However, the stress stimuli did not induce significantly higher craving compared
296	to the neutral stimuli in either group. The statistical analysis of the change of the mean
297	craving ratings from the baseline showed no significant differences between the treatment
298	groups at any timepoint.
299	
300	** Fig. 4 **
301	
302	Salivary cortisol level
303	
304	Of the collected saliva samples 70 had to be discarded due to the insufficient amount of
305	saliva. The data of the analysed samples of 18 participants (8 CT-optimal, 10 controls) were
306	included in the mixed-model analysis which showed no significant differences between the
307	groups in the change of the mean cortisol level. In both groups, the mean cortisol level
308	dropped below the baseline 10 minutes after the stress exposure and remained lower than the
309	baseline until the last measurement of the experiment (Fig. 5). However, this change was
310	statistically significant only for the control group at the timepoint 25 minutes after the stress
311	exposure.
312	
313	** Fig. **
314	
315	Discussion

316

To our knowledge the present study is the first randomized controlled trial investigating theeffect of acute CT-optimal touch on cue-reactivity in AUD patients. Based on previous

evidence indicating that CT-optimal affective touch can modulate the functioning of the

autonomic nervous system and physiological stress responses, we hypothesized that CT-

optimal touch may affect the cue-reactivity of abstinent AUD patients. In line with our

322 hypothesis, our results revealed significant differences in HRV reactivity between the CT-

323 optimal touch treatment group and the non-CT-optimal touch control group during alcohol-

324 cue exposure, indicating a modulatory effect of CT-optimal touch on autonomic cue-

reactivity. Importantly, this effect was achieved with a robotic device, without the social

elements of affective human touch.

327

Contrary to the HRV patterns observed in healthy individuals, AUD patients typically exhibit 328 329 a low relaxed state HRV which elevates in response to stress or alcohol-cues. A previous study by Garland et al. (2012) showed that abstinent AUD patients with a lower cue-elicited 330 331 HRV were significantly less likely to relapse than those with a higher cue-elicited HRV, implying that HRV reactivity may serve as a physiological predictor of a relapse. Building on 332 333 this foundation, we demonstrated that CT-optimal touch lowers the cue-elicited HRV in AUD 334 patients. Our results suggest that CT activation may shift the HRV reactivity closer to that of a healthy social drinker or an AUD patient with a lower susceptibility to relapse. However, 335 336 these results require to be replicated in a larger patient group and the possible clinical relevance must be evaluated outside the laboratory to investigate whether the physiological 337 338 modulation translates into a reduction in relapses.

339

Alongside the altered HRV, the AUD-related dysregulation of physiological stress responses
manifests as a high baseline cortisol level that decreases in response to stress or cue exposure
(Junghanns et al., 2005; Sinha, 2013a). Although we observed a reduction in cortisol
concentrations after stress exposure in both groups, CT-optimal touch did not exert a notable
effect on cortisol concentrations under any condition. This finding is consistent with the
study by Triscoli et al. (2017), which demonstrated that CT-optimal robotic touch modulated
HRV but not salivary cortisol levels in healthy subjects.

347

348 Throughout the experiment, alcohol craving remained moderately low in both groups,

349 although it significantly increased after exposure to alcohol cues. Environmental factors, such

as the laboratory setting and the time of day, may have influenced these results. The type of

351 treatment did not influence the subjective experience of alcohol craving which increased

equally in both groups after the cue exposure. In addition, the differences in HRV reactivity

between the groups did not relate to the craving ratings. These results align with prior

354 observations suggesting that physiological responses to alcohol-cues do not consistently

355 correlate with conscious craving (Heinz et al., 2010). Importantly, physiological responses

356 have been identified as more reliable predictors of relapse and addiction-related behavior

than conscious substance craving (Heinz et al., 2010).

358

In contrast to our hypothesis, no significant differences in HRV reactivity were observed
between groups during the stress exposure. Additionally, neither group exhibited an increase
in craving ratings in response to stress stimuli, suggesting that the chosen stimulation
material failed to induce the intended psychosocial stress in this patient group, although
similar visual stimuli have been successfully used in studies with AUD patients before
(Garland et al., 2012).

365

Several limitations need consideration regarding this study. Firstly, our sample size was 366 367 smaller than initially intended due to challenges in recruiting patients who both met the inclusion criteria and attended the scheduled study visit. In the following studies, this 368 challenge could be mitigated by including patients who are already committed to a treatment 369 370 programme. In future investigations of stress-triggered craving, the type of stress stimuli should be reconsidered, and the stress experience should be assessed using additional tools to 371 enhance the depth of understanding. 22% of the saliva sample were discarded because of 372 373 insufficient amount of saliva, which may have affected the results of the cortisol reactivity. 374 Future studies may benefit from utilizing serum cortisol instead of salivary cortisol for 375 improved accuracy. Lastly, due to the nature of the intervention and design of the robotic 376 device, double blinding was not possible as the velocity of the skin stroking was visible to the 377 investigators.

378

In conclusion, our study provides strong preliminary evidence that acute CT-optimal touch
modulates HRV reactivity during alcohol-cue exposure in early abstinent AUD patients. The
stress-regulating effects of CT-optimal touch make it a promising translational tool and a
candidate for potential novel adjunctive therapeutic intervention in the context of AUD.
Future research should focus on investigating the long-term outcomes to establish the clinical
potential of CT-optimal touch in preventing relapses.

386 Funding

- The study was funded by Business Finland (grant 6622/31/202), Aalto University and The
- 388 Emil Aaltonen Foundation (grant 230034 N1).

389

ounding

390 Author contributions

- 391 Juliana Harkki: Conseptulization, Metholodology, Formal analysis, Investigation, Writing -
- 392 original draft, Visualization, Supervision Pauli Tuovinen: Conseptulization, Software,
- 393 Resources, Writing review & editing Veikko Jousimäki: Resources, Writing review &
- 394 editing Goncalo Barretto: Resources, Writing review & editing Pekka Rapeli:
- 395 Metholodology, Writing review & editing Jussi Palomäki: Resources, Writing review &
- 396 editing Jonne Annevirta: Investigation Anna-Helena Puisto: Investigation Francis
- 397 McGlone: Conseptulization, Metholodology, Writing review & editing Heikki Nieminen:
- 398 Conseptulization, Supervision, Writing review & editing Hannu Alho: Conseptulization,
- 399 Metholodology, Supervision, Writing review & editing
- 400

401 Acknowledgments

We want to thank Hanna Renvall and Ilkka Nissinen for support, guidance and advice
throughout the study, Veli-Matti Saarinen and Helge Kainulainen at Aalto NeuroImaging for
their help in setting up the robotic device used in the experiment, Jukka Planman for valuable
brainstorming and conversations, Biodesign Finland for planting the seed for this study and
the funders for making it possible.

407

408 **Bibliography**

- 410 Anton, R. F., Moak, D. H., & Latham, P. K. (1996). The obsessive compulsive drinking
- 411 scale: A new method of assessing outcome in alcoholism treatment studies. *Archives of*412 *General Psychiatry*, 53(3), 225–231.
- 413 https://doi.org/10.1001/archpsyc.1996.01830030047008
- 414 Baron-Cohen, S., Wheelwright, S., Skinner, R., Martin, J., & Clubley, E. (2001). The autism-
- 415 spectrum quotient (AQ): evidence from Asperger syndrome/high-functioning autism,
- 416 males and females, scientists and mathematicians. *Journal of Autism and Developmental*
- 417 *Disorders*, *31*(1), 5–17. https://doi.org/10.1023/a:1005653411471
- Billieux, J., Khazaal, Y., Oliveira, S., De Timary, P., Edel, Y., Zebouni, F., Zullino, D., &
 Van Der Linden, M. (2011). The Geneva Appetitive Alcohol Pictures (GAAP):
- 420 Development and preliminary validation. *European Addiction Research*, 17(5), 225–
- 421 230. https://doi.org/10.1159/000328046
- 422 Björnsdotter, M., Löken, L., Olausson, H., Vallbo, Å., & Wessberg, J. (2009). Somatotopic

- 423 organization of gentle touch processing in the posterior insular cortex. *Journal of*
- 424 *Neuroscience*, 29(29), 9314–9320. https://doi.org/10.1523/JNEUROSCI.0400-09.2009
- 425 Blaine, S. K., Seo, D., & Sinha, R. (2017). Peripheral and prefrontal stress system markers
- 426 and risk of relapse in alcoholism. *Addiction Biology*, 22(2), 468–478.

427 https://doi.org/10.1111/adb.12320

Boothby, L. A., & Doering, P. L. (2005). Acamprosate for the treatment of alcohol
dependence. *Clinical Therapeutics*, 27(6), 695–714.

430 https://doi.org/10.1016/j.clinthera.2005.06.015

- 431 Center for Substance Abuse Treatment. (2009). Incorporating Alcohol Pharmacotherapies
 432 Into Medical Practice. *Center for Substance Abuse Treatment*, *114*, 126.
- 433 Cheng, Y. C., Huang, Y. C., & Huang, W. L. (2019). Heart rate variability as a potential
- biomarker for alcohol use disorders: A systematic review and meta-analysis. *Drug and Alcohol Dependence*, 204. https://doi.org/10.1016/j.drugalcdep.2019.05.030
- 436 Davidovic, M., Starck, G., & Olausson, H. (2019). Processing of affective and emotionally
- 437 neutral tactile stimuli in the insular cortex. *Developmental Cognitive Neuroscience*,
 438 35(November 2017), 94–103. https://doi.org/10.1016/j.dcn.2017.12.006
- 439 Ditzen, B., Neumann, I. D., Bodenmann, G., von Dawans, B., Turner, R. A., Ehlert, U., &
- 440 Heinrichs, M. (2007). Effects of different kinds of couple interaction on cortisol and

441 heart rate responses to stress in women. *Psychoneuroendocrinology*, *32*(5), 565–574.

442 https://doi.org/10.1016/j.psyneuen.2007.03.011

- 443 Eckstein, M., Mamaev, I., Ditzen, B., & Sailer, U. (2020). Calming Effects of Touch in
- 444 Human, Animal, and Robotic Interaction—Scientific State-of-the-Art and Technical
- 445 Advances. *Frontiers in Psychiatry*, *11*(November), 1–17.
- 446 https://doi.org/10.3389/fpsyt.2020.555058
- 447 Eriksson Hagberg, E., Ackerley, R., Lundqvist, D., Schneiderman, J., Jousmäki, V., &
- 448 Wessberg, J. (2019). Spatio-temporal profile of brain activity during gentle touch
- investigated with magnetoencephalography. *NeuroImage*, 201(November 2018).
- 450 https://doi.org/10.1016/j.neuroimage.2019.116024
- 451 Essick, GK, McGlone, F., Dancer, C., Fabricant, D., Ragin, Y., Phillips, N., Jones, T., &
- 452 Guest, S. (2010). Quantitative assessment of pleasant touch. *Neurosci Biobehav Rev.*,
- 453 *Feb;34*(2), 192–203. https://doi.org/10.1016/j.neubiorev.2009.02.003.
- Essick, Greg, James, A., & McGlone, F. P. (1999). Psychophysical assessment of the
 affective components of non-painful touch. *NeuroReport*, *Jul* 13;10(10), 2083–2087.
- 456 Fox, H. C., Anderson, G. M., Tuit, K., Hansen, J., Kimmerling, A., Siedlarz, K. M., Morgan,

P. T., & Sinha, R. (2012). Prazosin Effects on Stress- and Cue-Induced Craving and 457 Stress Response in Alcohol-Dependent Individuals: Preliminary Findings. Alcoholism: 458 459 Clinical and Experimental Research, 36(2), 351–360. https://doi.org/10.1111/j.1530-0277.2011.01628.x 460 461 Garland, E. L., Ingmar, F. H. A., & Howard, M. O. (2012). Cue-Elicited Heart Rate 462 Variability and Attentional Bias Predict Alcohol Relapse Following Treatment. 463 Psychopharmacology, 222(1), 17-26. https://doi.org/10.1007/s00213-011-2618-4.Cue-464 Elicited 465 Gilpin, N. W., & Koob, G. F. (2008). Neurobiology of alcohol dependence: Focus on 466 motivational mechanisms. In Alcohol Research and Health (Vol. 31, Issue 3, pp. 185– 467 195). National Institute on Alcohol Abuse and Alcoholism. Girven, K. S., Aroni, S., Navarrete, J., Marino, R. A. M., McKeon, P. N., Cheer, J. F., & 468 469 Sparta, D. R. (2020). Glutamatergic input from the insula to the ventral bed nucleus of the stria terminalis controls reward-related behavior. Addiction Biology, December 470 2019, 1–11. https://doi.org/10.1111/adb.12961 471 Heinz, A., Beck, A., Mir, J., Grüsser, S. M., Grace, A. A., & Wrase, J. (2010). Alcohol 472 473 craving and relapse prediction: Imaging studies. In Advances in the Neuroscience of 474 Addiction (pp. 137–161). CRC Press. https://doi.org/10.1201/9781420007350 Junghanns, K., Tietz, U., Dibbelt, L., Kuether, M., Jurth, R., Ehrenthal, D., Blank, S., & 475 476 Backhaus, J. (2005). Attenuated salivary cortisol secretion under cue exposure is 477 associated with early relapse. Alcohol and Alcoholism, 40(1), 80-85. 478 https://doi.org/10.1093/alcalc/agh107 479 King, C. E., & Becker, H. C. (2019). Oxytocin attenuates stress-induced reinstatement of 480 alcohol seeking behavior in male and female mice. *Psychopharmacology*, 236(9), 2613– 481 2622. https://doi.org/10.1007/s00213-019-05233-z 482 Koob, G. F. (2013). Addiction is a reward deficit and stress surfeit disorder. Frontiers in Psychiatry, 4(AUG), 1–18. https://doi.org/10.3389/fpsyt.2013.00072 483 Koob, G. F., & Volkow, N. D. (2016). Neurobiology of addiction: a neurocircuitry analysis. 484 485 In The Lancet Psychiatry. https://doi.org/10.1016/S2215-0366(16)00104-8 486 Korpi, E. R., den Hollander, B., Farooq, U., Vashchinkina, E., Rajkumar, R., Nutt, D. J., 487 Hyytiä, P., & Dawe, G. S. (2015). Mechanisms of action and persistent neuroplasticity 488 by drugs of abuse. *Pharmacological Reviews*, 67(4), 872–1004. 489 https://doi.org/10.1124/pr.115.010967 Lang, P.J. Bradley, M.M., & Cuthbert, B. N. (2008). International affective picture system 490

Journal Pre-proo

- 491 (IAPS): Affective ratings of pictures and instruction manual. *Technical Report A-8*.
- 492 University of Florida, Gainesville, FL.
- Löken, L. S., Wessberg, J., Morrison, I., McGlone, F., & Olausson, H. (2009). Coding of
- 494 pleasant touch by unmyelinated afferents in humans. *Nature Neuroscience*, *12*(5), 547–
 495 548. https://doi.org/10.1038/nn.2312
- 496 Manzotti, A., Cerritelli, F., Monzani, E., Savioli, L., Esteves, J. E., Lista, G., Lombardi, E.,
- 497 Rocca, S. La, Biasi, P., Galli, M., Chiera, M., & McGlone, F. P. (2023). Dynamic touch
- 498 induces autonomic changes in preterm infants as measured by changes in heart rate
- 499 variability. *Brain Research*, *1799*(August 2022), 148169.
- 500 https://doi.org/10.1016/j.brainres.2022.148169
- Maruyama, K., Shimoju, R., Ohkubo, M., Maruyama, H., & Kurosawa, M. (2012). Tactile
 skin stimulation increases dopamine release in the nucleus accumbens in rats. *Journal of Physiological Sciences*, 62(3), 259–266. https://doi.org/10.1007/s12576-012-0205-z
- 504 McGlone, F., Wessberg, J., & Olausson, H. (2014). Discriminative and Affective Touch:
- 505 Sensing and Feeling. *Neuron*, 82(4), 737–755.
- 506 https://doi.org/10.1016/j.neuron.2014.05.001
- 507 Morrison, I., Löken, L. S., Minde, J., Wessberg, J., Perini, I., Nennesmo, I., & Olausson, H.
- 508 (2011). Reduced C-afferent fibre density affects perceived pleasantness and empathy for
 509 touch. *Brain*, *134*(4), 1116–1126. https://doi.org/10.1093/brain/awr011
- 510 Naqvi, N. H., Gaznick, N., Tranel, D., & Bechara, A. (2014). The insula: A critical neural
- substrate for craving and drug seeking under conflict and risk. *Annals of the New York Academy of Sciences*, *1316*(1), 53–70. https://doi.org/10.1111/nyas.12415
- 513 Nummenmaa, L., Tuominen, L., Dunbar, R., Hirvonen, J., Manninen, S., Arponen, E.,
- 514 Machin, A., Hari, R., Jääskeläinen, I. P., & Sams, M. (2016). Social touch modulates
- endogenous μ -opioid system activity in humans. *NeuroImage*, *138*, 242–247.
- 516 https://doi.org/10.1016/j.neuroimage.2016.05.063
- 517 Olausson, H., Lamarre, Y., Backlund, H., Morin, C., Wallin, B. G., Starck, G., Ekholm, S.,
- 518 Strigo, I., Worsley, K., Vallbo, B., & Bushnell, M. C. (2002). Unmyelinated tactile
- afferents signal touch and project to insular cortex. *Nature Neuroscience*, 5(9), 900–904.
 https://doi.org/10.1038/nn896
- 521 Olausson, Håkan, Wessberg, J., Morrison, I., & McGlone, F. (2016). Affective touch and the
- 522 neurophysiology of CT afferents. In *Affective Touch and the Neurophysiology of CT*
- 523 *Afferents*. https://doi.org/10.1007/978-1-4939-6418-5
- 524 Pawling, R., Cannon, P. R., McGlone, F. P., & Walker, S. C. (2017). C-tactile afferent

- 525 stimulating touch carries a positive affective value. *PLoS ONE*, *12*(3), 1–15.
- 526 https://doi.org/10.1371/journal.pone.0173457
- 527 Pawling, R., Trotter, P. D., McGlone, F. P., & Walker, S. C. (2017). A positive touch: C-
- 528 tactile afferent targeted skin stimulation carries an appetitive motivational value.
- 529 *Biological Psychology*, *129*(July), 186–194.
- 530 https://doi.org/10.1016/j.biopsycho.2017.08.057
- 531 Püschel, I., Reichert, J., Friedrich, Y., Bergander, J., Weidner, K., & Croy, I. (2022). Gentle
- as a mother's touch: C-tactile touch promotes autonomic regulation in preterm infants.
- 533Physiology and Behavior, 257(August). https://doi.org/10.1016/j.physbeh.2022.113991
- 534 Ralevski, E., Petrakis, I., & Altemus, M. (2019). Heart rate variability in alcohol use: A
- review. In *Pharmacology Biochemistry and Behavior* (Vol. 176, pp. 83–92). Elsevier
 Inc. https://doi.org/10.1016/j.pbb.2018.12.003
- 537 Sinha, R. (2009). Modeling stress and drug craving in the laboratory: Implications for
- addiction treatment development. *Addiction Biology*, *14*(1), 84–98.
- 539 https://doi.org/10.1111/j.1369-1600.2008.00134.x
- Sinha, R. (2013a). Modeling Relapse Situations in the Human Laboratory. *Curr Top Behav Neurosci.*, *13*, 379–402. https://doi.org/10.1007/7854
- 542 Sinha, R. (2013b). The clinical neurobiology of drug craving. *Current Opinion in*
- 543 *Neurobiology*, 23(4), 649–654. https://doi.org/10.1016/j.conb.2013.05.001
- 544 Sinha, R., Fox, H. C., Hong, K. A., Bergquist, K., Bhagwagar, Z., & Siedlarz, K. M. (2009).
- 545 Enhanced negative emotion and alcohol craving, and altered physiological responses
- 546 following stress and cue exposure in alcohol dependent individuals.
- 547 *Neuropsychopharmacology*, *34*(5), 1198–1208. https://doi.org/10.1038/npp.2008.78
- 548 Sobell, L. C., & Sobell, M. B. (1992). Timeline follow back. A technique for Assessing self-
- reported Alcohol Consumption. In R. Z. Litten (Ed.), *Measuring alcohol consumption: Psychosocial and Biological Methods* (pp. 41–72). The Humana Press Inc.
- 551 Sullivan, J., Sykora, K., Schneiderman, J., Naranjo, C., & Sellers, E. (1989). Assessment of
- alcohol withdrawal: the revised clinical institute withdrawal assessment for alcohol scale
 (CIWA-Ar). *Br J Addict.*, *Nov*;84(11), 1353-7.
- 554 Triscoli, C., Croy, I., Steudte-Schmiedgen, S., Olausson, H., & Sailer, U. (2017). Heart rate
- variability is enhanced by long-lasting pleasant touch at CT-optimized velocity.
- 556 *Biological Psychology*, *128*(December 2016), 71–81.
- 557 https://doi.org/10.1016/j.biopsycho.2017.07.007
- 558 Uvnäs-Moberg, K., Handlin, L., & Petersson, M. (2014). Self-soothing behaviors with

559 particular reference to oxytocin release induced by non-noxious sensory stimulation.

560 *Frontiers in Psychology*, 5(OCT), 1–16. https://doi.org/10.3389/fpsyg.2014.01529

- 561 Walker, S. C., Cavieres, A., Peñaloza-Sancho, V., El-Deredy, W., McGlone, F. P., &
- 562 Dagnino-Subiabre, A. (2020). C-low threshold mechanoafferent targeted dynamic touch
- 563 modulates stress resilience in rats exposed to chronic mild stress. European Journal of
- 564 Neuroscience, May, 1-14. https://doi.org/10.1111/ejn.14951
- 565 Walker, S. C., Trotter, P. D., Swaney, W. T., Marshall, A., & Mcglone, F. P. (2017). C-tactile
- afferents: Cutaneous mediators of oxytocin release during affiliative tactile interactions? 566
- Neuropeptides, 64, 27-38. https://doi.org/10.1016/j.npep.2017.01.001 567
- 568













Fig. 1 The design of a single treatment session. The experiment included three consecutive sessions, each with one type of visual stimuli. The subjects were exposed to neutral, stress-inducing or alcohol-related, images, while they received either the CT-optimal or the control treatment. 5 saliva samples were collected during each session. HRV was derived from the BVP that was measured throughout the experiment excluding the 15-minute relaxation in the end of each session. Subjective craving was assessed with VAS at the baseline, after the visual stimuli and after the 25-minute treatment. BVP = Blood volume pulse, HRV = heart rate variability, VAS = visual analog scale.



Fig. 2 The mean change of the SDNN HRV between the baseline of the session and the visual stimuli exposure. The data is expressed as mean of the change. Positive values indicate increase of the HRV, whereas negative values indicate decrease. The chance is calculated by subtracting the SDNN HRV of the five-minute epoch of the visual stimuli exposure from the SDNN HRV of a five-minute epoch at the baseline of the session. The error bar indicates SEM. A = neutral visual stimuli. B = stress-inducing visual stimuli, C = alcohol-related visual stimuli, Control = non-CT-optimal treatment, CT-opt = CT-optimal treatment, HRV = heart rate variability, SEM = standard error of means, SDNN = Standard deviation of NN-intervals * = p < 0.05.



Fig. 3 The mean change of the SDNN HRV throughout the whole experiment. The initial baseline is set as 0. The error bars indicate SEM. The asterisks indicate significant difference from the baseline. Control = non-CT-optimal treatment, CT-opt = CT-optimal treatment, bl = baseline, t 5-10 min = treatment minutes 5 to 10, stim = visual stimulus, rec = recovery, N = neutral stimuli, S = stress inducing stimuli, A = alcohol-related stimuli. SEM = standard error of means. * = p < 0.05, ** = p < 0.01.



Fig. 4 The mean self-reported craving after each type of visual stimuli exposure. The error bars indicate SEM. Stim N = neutral visual stimuli, stim S = stress-inducing visual stimuli, stim A = alcohol-related visual stimuli. Control = non-CT-optimal treatment, CT-opt = CT-optimal treatment. SEM = standard error of means. * = p < 0.05, ** = p < 0.01.



Fig. 5 The mean change of the salivary cortisol throughout the whole experiment. The initial baseline is set as 0. The error bars indicate SEM. The asterisk indicates significant difference from the baseline. Control = non-CT-optimal treatment, CT-opt = CT-optimal treatment, bl = baseline, t 5-10 min = treatment minutes 5 to 10, stim = visual stimulus, rec = recovery, N = neutral stimuli, S = stress inducing stimuli, A = alcohol-related stimuli. SEM = standard error of means. * = p < 0.05

- The stress-regulating effects of C-tactile (CT) -optimal touch are known but their therapeutic potential has not yet been investigated in translational studies.
- This study provides the first evidence of clinical relevance of CT-optimal touch in management of substance dependence.
- CT-optimal touch significantly lowers alcohol-cue-elicited heart rate variability, indicating lower relapse susceptibility in alcohol use disorder patients.
- The ability of CT-optimal touch to modulate autonomic cue-reactivity, highlights the potential therapeutic benefits of affective touch in managing alcohol use disorder.

Journal