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Violation of eating expectancies does not reduce conditioned desires for chocolate

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ABSTRACT

Although eating desires can be easily learned, their extinction appears more difficult. The present two-session study aimed to investigate the role of eating expectancies in the short and longer-term extinction of eating desires. In addition, the relationship between eating desires and conditioned evaluations was examined to test whether they might share a similar mechanism. It was hypothesized that the short-term extinction of eating desires would be more successful after the disconfirmation of eating expectancies (instructed extinction or IE), while resulting in worse longer-term extinction because omission of the food reward during extinction is not surprising. In contrast to the hypotheses, it was found that IE had no effect on the short-term and longer-term extinction of eating desires. Eating desires correlated with conditioned evaluations only to some extent. It is concluded that eating expectancies do not mediate the short-term extinction of conditioned eating desires. In addition, their longer-term extinction does not appear to be facilitated by a greater violation of eating expectancies. This suggests that it might not be necessary to focus on expectancy violation in cue exposure therapy to reduce eating desires.

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1. Introduction

Obesity prevalence has increased substantially over the last decades, most western countries now reporting approximately 10–35% of their adult population to be obese (Berghofer et al., 2008; Ogden, Carroll, Kit, & Flegal, 2014). One important contributor to obesity is the “obesogenic” environment, in which omnipresent food cues signal the availability of palatable, high-calorie foods. Exposure to these cues can elicit physiological and psychological reactions, including an increased desire to consume the food (Jansen, 1998). These cue-elicited eating desires are thought to stimulate overeating and weight gain (Fedoroff, Polivy, & Herman, 2003; Jansen, Havermans, & Nederkoorn, 2011), and appear heightened in overweight individuals and in those with eating psychopathology (Ferriday & Brunstrom, 2011; Jansen et al., 2003; Karhunen, Lappalainen, Tammela, Turpeinen, & Uusitupa, 1997). This highlights the need for investigating the etiology of cued eating desires, as well as finding ways to effectively tackle them.

It is thought that learning processes play an important role in cue-elicited eating desires. For instance, the sight and smell of food are thought to have become conditioned stimuli (CSs) predictive for food intake (unconditioned stimulus or US) through repeated CS-US pairings (Jansen, 1998). As a result, a CS (food cue) can elicit conditioned appetitive responses (CRs) such as a heightened desire to eat, increased salivation, and an explicit eating expectancy. Moreover, learning theory predicts that these CRs extinguish after repeated exposure to the CS alone. Thus, theoretically, after repeated exposure to the sight and smell of palatable food without consumption, conditioned responses such as desires to eat should decline. Such successful extinction of responses to food cues is thought to increase one’s ability to abstain from eating, and ultimately, result in improved weight loss success (Jansen, 1998; Jansen et al., 2011; Jansen, Stegerman, Roefs, Nederkoorn, & Havermans, 2010; Wardle, 1990). In support, the few studies conducted on cue exposure therapy (CET), in which overweight individuals or those with eating psychopathology are repeatedly exposed to food cues without eating, indeed suggest CET to be effective in reducing cue-elicited cravings and eating binges (e.g., Boutelle et al., 2014; Jansen, Broekmate, & Heymans, 1992; Jansen, Van Den Hout, De Loof, Zandbergen, & Griez, 1989; Martinez-Mallén et al., 2007; Schyns, Roefs, Mulkins, & Jansen, 2016; Torro et al., 2003). Despite
these very promising findings, the evidence for a superiority of CET over control treatments at follow-up is mixed (Boutelle et al., 2014; Jansen et al., 1992). Since CET is rooted in learning theory, it might be optimized by studying the mechanisms that underlie the (long-term) extinction of appetitive responses to food cues.

Extinction can be studied using conditioning paradigms, in which cue-elicited eating desires and eating expectancies are first established by repeatedly pairing a cue (CS) with a US (food) (e.g., Bongers, van den Akker, Havermans, & Jansen, 2015; Van den Akker, Havermans, Bouton, & Jansen, 2014; Van Gucht, Vansteenwegen, Van den Bergh, & Beckers, 2008b). This acquisition phase is followed by an extinction phase, during which repeated CS - no US pairings occur. Findings suggest that eating desires can be quickly acquired, but only when a participant is consciously aware of the CS-US contingency (i.e., when reporting heightened US expectancies upon CS presentation) (Van den Akker, Jansen, Frenzt, & Havermans, 2013). Thus, during acquisition, US expectancies are likely essential for developing cued eating desires. During extinction however, the two responses can diverge: even when eating expectancies reduce, eating desires can remain heightened (Van Gucht, Vansteenwegen, Beckers, & Van den Bergh, 2008b). These divergent extinction patterns suggest an involvement of separate response systems that are differentially affected by extinction. Specifically, it may be that eating expectancies relate more to an anticipatory response system, preparing the organism for an incoming stimulus (Van Gucht, Vansteenwegen, Van den Bergh, & Beckers, 2008b). In contrast, regarding the desire to eat, it has been noted that its resistance to extinction resembles extinction in evaluative conditioning: conditioned evaluations do not extinguish easily as well (Baeyens, Crombez, Van den Bergh, & Eelen, 1988; Van Gucht et al., 2008b). It might be that CS evaluations are based on the mere activation of the US representation in memory, i.e., reflecting a mere referential learning in which the CS “makes one think of” the US in the absence of actual eating expectancies (Hermans, Vansteenwegen, Crombez, Baeyens, & Eelen, 2002; Baeyens, Eelen, Crombez, & Van den Bergh, 1992). This activation of the US representation could also be sufficient for experiencing heightened eating desires (Van Gucht, Baeyens, Vansteenwegen, Hermans, & Beckers, 2010). Extinction may have more impact on response systems that prepare an individual for an incoming stimulus than on systems that are based on the mere activation of the US representation in memory (Van Gucht et al., 2008b; see also Luck & Lipp, 2015; Sevenster, Beckers, & Kindt, 2012), causing eating desires and CS evaluations to extinguish more slowly than US expectancies.

However, there is also evidence suggesting a closer relationship between expectancies and eating desires during extinction. For instance, although it indeed seems difficult to completely extinguish cue-elicited eating desires, their extinction can be achieved to a certain extent (Van den Akker et al., 2014; Van den Akker et al., 2015). This pattern is, in fact, very similar to the extinction of US expectancies: although expectancies diminish, they usually do not show complete extinction either (Van den Akker et al., 2014; Van den Akker et al., 2015; Van Gucht et al., 2008b). In addition, in a conditioning study conducted in smokers, explicit disconfirmation of acquired US expectancies (i.e., “instructed extinction” or IE) immediately eliminated cue-elicited craving for cigarettes, suggesting that expectancies acted as a mediator for cue-elicited cigarette craving (Field & Duka, 2001). Thus, it seems possible that the lack of complete extinction of eating desires during extinction in previous studies was caused by a lack of complete extinction of eating expectancies, rather than by the involvement of separate response systems. A complete and immediate elimination of acquired eating expectancies by IE can help determine how closely food cravings and US expectancies are related: if conditioned cravings were to disappear immediately after CS-US disconfirmation, this would provide evidence for a mediating role of expectancies in the extinction of eating desires, and against the involvement of different response systems. Conversely, if conditioned cravings remain heightened despite an elimination of expectancies, this would provide evidence against a mediating role of expectancies in the extinction of eating desires, and would provide additional support for the idea of different response systems underlying the extinction of US expectancies and conditioned desires.

Extinction performance during one experimental session does not need to be predictive for longer-term extinction learning. New memories require time to consolidate (McGaugh, 2000), which is why actual learning is best tested at a later point in time. In case of IE, one could expect worse longer-term extinction learning, despite a possibly better short-term extinction performance. One reason for this might be the altered degree of “surprise” of non-occurrence of the US during extinction. Surprise, or violation of US expectancies, is thought to play a major role in (extinction) learning: a smaller discrepancy between expected and actual occurrence of the US should result in poorer (inhibitory) learning (Craske, Treanor, Conway, Zbozinek, & Vervliet, 2014; Rescorla & Wagner, 1972).

After IE, the US omission that occurs during extinction is not very surprising, therefore possibly resulting in worse longer-term extinction learning. This would have implications for cue exposure therapy. If a reduction in US expectancies prior exposure (extinction) sessions results in worse extinction learning, cue exposure therapy might benefit from maximizing these expectancies prior to an exposure session — i.e., heightening expectancies for the US (eating) to occur.

The primary aim of the present study was to evaluate the role of US expectancies in the short and longer-term extinction of conditioned eating desires. To examine this, extinction performance in an IE condition (i.e., receiving explicit disconfirmation of the acquired CS-US contingencies before extinction) was compared with a “normal” extinction condition (no extinction instructions) on two subsequent days. It was expected that IE would speed up the short-term extinction of eating desires relative to a normal extinction procedure, while resulting in worse extinction learning, as reflected by a greater spontaneous recovery (the recovery of responding that occurs after the mere passage of time) (Pavlov, 1927; Rescorla, 2004) and slower re-extinction during test after a 24 h delay. Alternatively, it may be that eating desires are more closely related to evaluations of the CS rather than to US expectancies because they may share a similar mechanism. To examine this possibility conditioned CS evaluations were measured before and after each extinction session, and correlations between US expectancies, eating desires, and CS evaluations were assessed.

### 2. Methods and materials

#### 2.1. Participants

48 participants took part in the study, of which two participants were replaced by additional participants because they did not show awareness of the CS-US contingency (i.e., they did not report clear differential US expectancies towards the end of acquisition), and one other participant because she did not show up for the second session. Participants were eligible to participate in the study if they were 1st or 2nd year undergraduate female students, between 17 and 25 years old, proficient in Dutch, and had indicated to like chocolate. Further, care was taken that no participant had previously participated in an appetitive conditioning study. To standardize hunger, participants were instructed to have a small meal two hours prior to each session and to refrain from consumption thereafter (only the drinking of water was allowed). As a cover
story, participants were told the study was about the memory of taste. Participants received either a voucher worth €12.50 or course credit for participation. The study was approved by the local ethical committee.

2.2. Measurements

2.2.1. US expectancy and desire to eat

100 mm-Visual Analogue Scales (VAS) were used to assess expectancy to receive chocolate (‘To what extent do you expect this box to contain chocolate at this moment?’) and subjective desire for chocolate (‘When looking at this box, how strong is your desire for chocolate at this moment?’) (in this order). Ratings ranged from 0 (‘certainly expect this box not to contain chocolate/no desire at all’) to 100 (‘certainly expect this box to contain chocolate/very strong desire’).

2.2.2. CS evaluations

Evaluations for the CS+ and CS− were assessed using two VAS (‘How pleasant do you find the box depicting the elephant/fish?’) (in this order). Ratings ranged from 0 (‘not pleasant at all’) to 100 (‘extremely pleasant’).

2.2.3. Hunger

To be able to control for possible group differences in hunger, participants filled in a VAS (‘How hungry are you at this moment?’) ranging from 0 (‘not hungry at all’) to 100 (‘extremely hungry’).

2.2.4. US liking

To be able to control for possible group differences in the liking of the chocolate (the US) used, participants filled in a VAS (‘How much did you like the chocolate?’), ranging from 0 (‘not at all’) to 100 (‘extremely’).

2.2.5. Dutch Eating Behaviour Questionnaire (DEBQ; Van Strien, Frijters, Bergers, & Defares, 1986)

The DEBQ was administered to be able to control for possible group differences in eating styles. This 33-item questionnaire provides a measure of external eating (10 items), emotional eating (13 items), and restraint (10 items). Each item is scored on a 5-point scale, ranging from “never” to “very often”, higher scores indicating a higher degree of emotional or external eating, or restraint.

All questionnaires were administered in Dutch.

2.3. Stimuli

US: A small piece of Belgian milk chocolate (approximately 1.3–1.5 g, Rousseau) served in a small cup served as US. The USs were kept out of sight of the participants.

CS: Two children’s jewelry boxes served as Cs, and contained the US in case of a CS+ trial. One box depicted elephants (‘Elmer the Patchwork Elephant’) and had coloured squares and yellow linings, and the other depicted fish (‘The Rainbow Fish’) and was blue. Which box served as CS+ and CS− was counterbalanced across conditions.

2.4. Design and procedure

Participants were randomly assigned to one of two conditions: instructed extinction (IE) or normal extinction (NE), which differed only in instructions that were given after the acquisition phase. Participants were individually seen on two sessions, the second session always taking place exactly 24 h after the first session. Testing took place between 11 AM and 6 PM.

2.4.1. Session 1

After arrival, participants gave written consent for participation and rated their hunger. Next, they were shown the boxes and were instructed that one of the boxes would sometimes contain something they would eat, whereas the other box would never contain anything. Participants repeated this information once. After this, the conditioning procedure started.

Acquisition: During acquisition training, all participants received five CS+ and five CS− trials. A trial proceeded as follows: a box (CS+ or CS−) was placed in front of the participant for 10 s and she was instructed to look at the box. Next, she rated her US expectancy and desire to eat, after which she opened the box. In case of a CS+ trial, the box contained the US which the participant consumed. In case of a CS− trial, the box did not contain anything. The participant closed the box and an inter-trial interval of 20 s started, after which the next trial began. All instructions given during the trials were pre-recorded. Trials were presented in a random order, with the restriction that no more than two consecutive trials were of the same trial type (CS+ or CS−).

Pilot testing suggested that the interruption in the conditioning procedure after acquisition (which was necessary for instructed extinction and to measure conditioned evaluations) unwarrantedly diminished responses on the first extinction trial in the NE condition, likely because some participants suspected the “rules” had changed. Therefore, two VAS (US liking and US creaminess) were given to the participant after the first two CS+ trials had been completed, in order for participants to learn that an interruption would not be predictive for a discontinuation of CS-US pairings. These VAS were introduced in a manner similar to the interruption in the conditioning procedure after acquisition (see below).

CS evaluations and manipulation: After acquisition, CS evaluations were assessed in both conditions. The VASs were introduced as follows: “We now briefly interrupt the study so you can fill in this question.” Only in case of IE, this was followed by the instruction: “Please pay close attention; you will now receive a very important instruction. From now on, none of the boxes will contain chocolate. It is important for you to understand and remember this. Could you repeat this instruction?” Next, in both conditions, the participant was told: “Thank you. The study will now continue”. After this, the extinction phase started.

Extinction: All participants received a similar extinction procedure, during which 12 CS+ and 12 CS− trials were presented. Trials proceeded as during acquisition, with the exception that no USs were provided. Whether the first extinction trial was a CS+ or CS− was counterbalanced across conditions.

After extinction, CS evaluations, a bogus memory questionnaire, and the participant’s time of pre-experimental food intake were assessed.

2.4.2. Session 2

After arrival in the laboratory, participants rated their current hunger, and CS evaluations were assessed. Next, participants received a second extinction phase.

Re-extinction: Participants received 12 CS+ and 12 CS− trials (no USs were given). Trials proceeded as during the previous phases. Whether the first extinction trial was a CS+ or CS− was counterbalanced across conditions.

After re-extinction, CS evaluations, the participant’s suspicion about the study’s hypothesis, their time of pre-experimental food intake, the DEBQ, and the participant’s height and weight were assessed.

2.5. Data reduction and statistical analyses

Differential acquisition, extinction, and re-extinction of the
Desire to eat and US expectancy over time and across conditions were analyzed using repeated-measures ANOVAs for each phase of the experiment (acquisition, extinction, and re-extinction). This resulted in 2 (Condition: IE vs. NE) × 2 (CS-type: CS+ vs. CS−) × 5/12 (Acquisition Trial/Extinction Trial/Re-extinction Trial) repeated-measures ANOVAs, including CS-type (CS) and Trial (T) as within-subjects factors and Condition (C) as between-subjects factor. Spontaneous recovery was analyzed using 2 (Condition: IE vs. NE) × 2 (CS-type: CS+ vs. CS−) × 2 (Trial: EXT12 vs. RE-EXT1) RM ANOVAs. To test the immediate effects of our manipulation, 2 (Condition: IE vs. NE) × 2 (CS-type: CS+ vs. CS−) × 2 (Trial: ACQ5 vs. EXT1) ANOVAs were conducted, as well as a paired t-test for the IE condition on the differentiation on the first extinction trial. Acquisition and extinction of differential CS evaluations were assessed using repeated-measures ANOVAs as well, using CS and T (acquisition, extinction, pre-re-extinction, post-re-extinction) as within-subjects variables, and C as between-subjects variable. Finally, to assess how strongly the different responses were associated, differential responses (i.e., CS+ minus CS−) were calculated for CS evaluations, US expectancies, and eating desires, before and after each extinction phase (expectancies and desires: ACQ5; EXT12; RE-EXT1; RE-EXT12; evaluations: ACQ; EXT; RE-EXT PRE; RE-EXT POST), and correlational analyses on these responses were performed. Greenhouse—Geisser epsilon corrections are reported for all repeated-measures analyses whenever sphericity was violated.

### 3. Results

Participants characteristics are displayed in Table 1. Because baseline hunger on session 1 differed across the conditions, it was included as a centered covariate in the analyses. Since hunger did not influence patterns of conditioned responding in any of the phases, all ps > .05, in the final analyses, hunger was not included.

#### 3.1. US expectancy

**Acquisition:** Participants learned to expect to receive chocolate when presented with the CS+, as indicated by a significant CS × T interaction, F(2.89, 132.84) = 82.81, p < .001, ηp² = .64, with no differences across conditions (CS × T × C), F < 1 (see Fig. 1). This resulted in a significant CS+ vs. CS− differentiation on trial 5, F(1, 46) = 1550.71, p < .001, ηp² = .97, with a trend towards a greater differentiation in the IE condition, F(1, 46) = 3.33, p = .08, ηp² = .07.

**Effectiveness of the manipulation:** In the NE condition, differential US expectancies did not change from the last acquisition trial to the first extinction trial, as shown by a non-significant CS × T (ACQ5 vs. EXT1) interaction, F(1, 23) = 2.59, p = .12, ηp² = .10. In contrast, in the IE condition, US expectancies reduced from acquisition to extinction, F(1, 23) = 446.96, p < .001, ηp² = .95. In this condition, analyses indicated a trend towards a CS+ vs. CS− differentiation on the first extinction trial, F(1, 23) = 3.15, p = .09, ηp² = .12, this differentiation being non-significant on the second extinction trial, F(1, 23) = 1.86, p = .19, ηp² = .08. Thus, the extinction instruction was effective: it resulted in an almost-immediate and complete reduction in differential US expectancies.

**Extinction:** In the NE condition, differential expectancies diminished during extinction (CS × T), F(3.79, 87.22) = 42.56, p < .001, ηp² = .65, while unsurprisingly, this was not the case for the IE condition, F(2.05, 47.23) = 1.64, p = .20, ηp² = .07. On extinction trial 12, a significant differentiation remained in the NE condition, F(1, 23) = 11.01, p = .003, ηp² = .32, but not in the IE condition, F < 1.

**Spontaneous recovery and re-extinction:** Spontaneous recovery of US expectancies was present (EXT12 vs. RE-EXT1), F(1, 46) = 86.83, p < .001, ηp² = .65, and similar across conditions, F < 1. In line with this, on re-extinction trial 1 a significant differentiation in US expectancies was found, F(1, 46) = 150.15, p < .001, ηp² = .77, that did not differ between conditions, F < 1. Expectancies re-extinguished (CS × T), F(2.89, 132.92) = 55.76, p < .001, ηp² = .55, although a significant differentiation remained on the last re-extinction trial, F(1, 46) = 10.12, p = .003, ηp² = .18. Conditions did not differ in the course of re-extinction (CS × T × C) and in final re-extinction levels, Fs < 1. Thus, overall, US expectancies recovered after a 24 h interval and re-extinguished to a certain extent, but no differences across conditions were found.

#### 3.2. Desire to eat chocolate

**Acquisition:** Participants acquired a differential desire to eat chocolate during acquisition (CS × T), F(2.23, 107.16) = 4.89, p = .006, ηp² = .10, and equally so for both conditions (CS × T × C), F < 1 (see Fig. 2). This resulted in a significantly higher desire to eat in response to the CS+ vs. the CS− on the last acquisition trial, F(1, 46) = 29.92, p < .001, ηp² = .39, independent of condition, F < 1.

**Immediate response to the manipulation:** The explicit CS-US disconfirmation did not appear to reduce differential desires to eat on the subsequent trial. Although differential desires diminished marginally significant from the last acquisition to the first extinction trial, F(1, 46) = 3.74, p = .06, ηp² = .08, importantly, this change did not interact with condition, F = 1.21, ns. A significant CS+ vs. CS− differentiation was still present on the first extinction trial, F(1, 46) = 17.40, p < .001, ηp² = .27, independent of condition, F < 1.

Because visual inspection of the figures suggested an overall decrease in eating desires in specifically the IE condition, additional analyses were performed on the overall change in eating desires (ACQ5 vs. EXT1). These confirmed an overall reduction in desires in the IE condition, F(1, 23) = 10.51, p = .004, ηp² = .31, but not in the NE condition, F < 1 (overall T × C interaction: F(1, 46) = 4.37, p = .04, ηp² = .09). Thus, while disconfirmation of US expectancies did not affect conditioned desires to eat, it appeared to diminish eating desires overall.

**Extinction:** Conditioned desires to eat extinguished (CS × T),

### Table 1

<table>
<thead>
<tr>
<th></th>
<th>Normal extinction (N = 24)</th>
<th>Instructed extinction (N = 24)</th>
<th>t(46)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>19.38 (1.47)</td>
<td>19.21 (1.28)</td>
<td>0.42</td>
<td>.68</td>
</tr>
<tr>
<td><strong>BMI</strong></td>
<td>23.45 (4.81)</td>
<td>22.96 (3.03)</td>
<td>0.43</td>
<td>.67</td>
</tr>
<tr>
<td><strong>Baseline hunger</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Session 1</td>
<td>36.21 (19.38)</td>
<td>48.00 (22.32)</td>
<td>1.95</td>
<td>.06</td>
</tr>
<tr>
<td>Session 2</td>
<td>41.46 (26.73)</td>
<td>48.29 (27.18)</td>
<td>0.88</td>
<td>.38</td>
</tr>
<tr>
<td><strong>US liking DEBQ</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Restricted</td>
<td>2.70 (0.73)</td>
<td>2.77 (0.69)</td>
<td>0.35</td>
<td>.73</td>
</tr>
<tr>
<td>Emotional</td>
<td>2.53 (0.65)</td>
<td>2.68 (0.53)</td>
<td>0.88</td>
<td>.38</td>
</tr>
<tr>
<td>External</td>
<td>3.26 (0.59)</td>
<td>3.21 (0.59)</td>
<td>0.27</td>
<td>.79</td>
</tr>
</tbody>
</table>
$F(3.42, 157.17) = 2.85, p = .033, \eta^2_p = .06$, and similarly so across conditions ($CS \times T \times C$), $F(3.42, 157.17) = 1.35, p = .26, \eta^2_p = .03$. Extinction of differential desires to eat was not complete: a marginally significant differentiation between the $CS_+ \text{ and } CS_- \text{ remained at the end of extinction (ext12), } F(1, 46) = 3.91, p = .054, \eta^2_p = .08$, with no differences across conditions, $F(1, 46) = 1.41, p = .24, \eta^2_p = .03$.

Spontaneous recovery and re-extinction: Conditioned desires recovered after a 24 h delay (EXT12 vs RE-EXT1), $F(1, 46) = 6.58, p = .014, \eta^2_p = .13$, and this was similar across conditions, $F = 1.08,
ns$. This resulted in a significant $CS_+ \text{ vs } CS_- \text{ differentiation on the first re-extinction trial, } F(1, 46) = 24.13, p < .001, \eta^2_p = .34$, that was similar across conditions, $F < 1$. Conditioned desires re-extinguished to some extent ($CS \times T$), $F(5.46, 251.08) = 3.80, p = .002, \eta^2_p = .08$, with no differences across conditions ($CS \times T \times C$), $F < 1$. Again, extinction was not complete: the differential desire to eat was still significant on the last re-extinction trial, $F(1, 46) = 13.81, p = .001, \eta^2_p = .23$, and similarly for both conditions, $F < 1$.

Fig. 1. Mean US expectancy by CS-type, trial, condition, and phase of the experiment.

Fig. 2. Mean desire to eat by CS-type, trial, condition, and phase of the experiment.
3.3. CS evaluations

After acquisition, the CS+ was evaluated significantly more positively than the CS−, $F(1, 46) = 23.13, p < .001, \eta^2_p = .34$ (see Fig. 3). This differential conditioned evaluation extinguished, $F(1, 46) = 14.80, p < .001, \eta^2_p = .24$, though after extinction there was still a trend towards more positive evaluations for the CS+ vs CS−, $F(1, 46) = 2.86, p = .1, \eta^2_p = .06$. The conditions did not differ in the acquisition or extinction of conditioned evaluations, $Fs < 1$.

Differential conditioned evaluations recovered during tests at the beginning of the second session (EXT vs RE-EXT PRE), $F(1, 46) = 16.10, p < .001, \eta^2_p = .26$, and again resulting in significantly higher evaluations for the CS+ vs CS− before re-extinction, $F(1, 46) = 17.22, p < .001, \eta^2_p = .27$. Again, evaluations extinguished, $F(1, 46) = 27.57, p < .001, \eta^2_p = .38$, and this time, extinction was complete, $F < 1$. Spontaneous recovery, the course of extinction, and final extinction levels did not differ significantly across conditions, $F < 1, F(1, 46) = 2.11, p = .15, \eta^2_p = .04$; $F(1, 46) = 1.81, p = .19, \eta^2_p = .04$.

3.4. Correlational analyses

At the end of acquisition, differential evaluations correlated only with eating desires, $r(48) = .32, p = .03$. Correlations between US expectancies and evaluations, $r(48) = .03, ns$, and US expectancies and eating desires were non-significant, $r(48) = .24, p = .11$. Similarly, after extinction, conditioned evaluations correlated significantly with differential eating desires, $r(48) = .43, p = .002$, while US expectancies did not correlate with evaluations, $r(48) = -.01, ns$, nor with eating desires, $r(48) = .06, ns$. Thus, on session 1, it seems that conditioned evaluations are linked with eating desires while US expectancies are not. At the start of re-extinction, differential desires correlated with both US expectancies, $r(48) = .33, p = .02$ and conditioned evaluations, $r(48) = .38, p = .008$, which also correlated with each other, $r(48) = .28, p = .051$. After re-extinction, evaluations did not correlate with eating desires, $r(48) = .20, p = .18$, nor with US expectancies: $r(48) = -.17, p = .25$, while this time, US expectancies were inversely correlated with eating desires, $r(48) = -.37, p = .01$.

In sum, on session 1, greater positive evaluations of the CS+ (but not heightened US expectancies) appeared to consistently co-occur with increased eating desires. A different pattern was found on session 2: at the start of re-extinction, eating desires correlated with both conditioned evaluations and US expectancies. After re-extinction, differential eating desires did not correlate with conditioned evaluations and even showed an inverse correlation with US expectancies. Thus, overall, findings of correlational analyses were somewhat mixed but suggest eating desires to be more related to conditioned evaluations than to US expectancies.

4. Discussion

The aim of the present study was to examine whether instructed extinction would affect the short-term and longer-term extinction of conditioned appetitive responses. In addition, the relationship between eating desires, eating expectancies, and CS evaluations was examined. It was found that disconfirming eating expectancies prior to extinction did not affect the short-term extinction of conditioned eating desires: in both conditions, differential desires reduced but did not extinguish completely. On session 2, spontaneous recovery and re-extinction of appetitive responses was found, while again, this was similar across the conditions. CS evaluations correlated with eating desires on session 1, but not on session 2.

The finding that on session 1, an explicit disconfirmation of eating expectancies did not affect the short-term extinction of eating desires suggests that eating expectancy does not act as mediator for the extinction of eating desires. Hence, it seems unlikely that previously observed difficulties to extinguish eating desires were caused by a lack of complete extinction of eating expectancies. This is in line with the idea that eating expectancies and eating desires stem from loosely coupled response systems that are differentially susceptible to extinction (Delamater & Westbrook, 2014). As described in the Introduction, one possibility is that eating desires and conditioned evaluations share a common mechanism, which would cause them to behave similarly throughout the different conditioning phases. Only partial evidence was found for this in the present study. On session 1, conditioned
evaluations and eating desires appeared to co-occur: both evaluations and eating desires seemed equally unaffected by instructed extinction, and differential acquisition and extinction of eating desires correlated with the differential acquisition and extinction of evaluations, but not with differential expectancies. In contrast however, on session 2, no evidence was found for eating desires to be more closely linked to evaluations than to eating expectancies. Thus, overall, our findings provide some evidence for eating desires to be associated more closely with conditioned evaluations than with US expectancies, although they also do not seem to behave in synchrony. The partial concordance between eating desires and evaluations may be explained by eating desires and CS evaluations being based on activation of the US representation in memory, reflecting a mere referential learning that is not dependent on current eating expectancies (Hermans et al., 2002). Their partial discordance however suggests that eating desires and evaluations are not homologous either. Alternatively, it may be that our self-report measure of CS evaluations did not provide accurate measures of underlying evaluative representations, but were influenced by judgment-related processes, contributing to divergences between eating desires and evaluations. Specifically, prior judgments of CS evaluations (i.e., after acquisition) may cause participants to use the most recent information rather than integrating all available information about the CS: there is evidence for (partial) revaluation. It is thought that in humans, such returns of responses can be optimized, and in which manners these techniques can be implemented in weight loss therapies. In addition, future studies may examine how successful long-term extinction of appetitive responses to food cues in humans is clearly needed.

Our findings raise the question how one might effectively extinguish conditioned eating desires. If conditioned evaluations and eating desires share a common mechanism, one approach would be to use methods that have been shown to be effective in changing conditioned evaluations. For instance, counterconditioning (pairing the CS+ with a US of opposite valence) has been shown to eliminate acquired evaluations (Baeyens, Eelen, van den Bergh, & Crombez, 1989), as well as conditioned cravings for chocolate (Van Guch et al., 2010). Other effective methods may include a more extensive extinction training (Luck & Lipp, 2015). US devaluation (decreasing the evaluative meaning of the US) (e.g., Baeyens, Eelen, Van den Bergh, & Crombez, 1992; Leer, Engelhard, Altink, & van den Hout, 2013), or cognitive reappraisal of the CS (Blechert et al., 2015). Interestingly, and underlining the potential benefit of changing conditioned evaluations, (manipulating) CS evaluations has been found to be predictive for food choice (Veling, Aarts, & Stroebe, 2013), alcohol intake (Houben, Nederkoorn, Wiers, & Jansen, 2011), and for reinstatement of conditioned fear (Zbozinek, Hermans, Prenoueau, Liao, & Craske, 2014).

In sum, it was found that eating expectancies did not mediate the extinction of eating desires, nor did greater expectancy violation during extinction enhance extinction learning. Eating desires appeared only partially linked with conditioned evaluations. One implication could be that in CET, it is not necessary to attempt maximizing the violation of eating expectancies. Future research may aim to confirm that increased violation of eating expectancies is, indeed, not effective in reducing eating desires during CET. In addition, future studies may examine how successful long-term extinction of conditioned appetitive responses such as eating desires can be optimized, and in which manners these techniques can be implemented in weight loss therapies.

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References


