

Brief Communication

An omission procedure reorganizes the microstructure of sign-tracking while preserving incentive salience

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Appetitive sign-tracking, in which reward-paired cues elicit approach that can result in cue interaction, demonstrates how cues acquire motivational value. For example, rats will approach and subsequently interact with a lever insertion cue that signals food delivery upon its retraction. However, lever deflections are rapidly reduced once rats are trained on an omission schedule in which lever interactions cancel food delivery. Here we evaluated the change in sign-tracking response topography in rats exposed to such an omission procedure. Lever deflections dropped precipitously when they canceled reward. However, rats that were on an omission schedule continued to approach, sniff, and contact the lever without pressing it, and did so at comparable rates to rats that were not under an omission schedule. Thus, sign-tracking was maintained, albeit in a different manner, following omission. Such findings show that the motivational attraction to reward cues can be expressed with remarkable persistence and flexibility.

[Supplemental material is available for this article.]

Appetitive sign-tracking is a phenomenon in which a reward-paired cue elicits approach that can result in cue interaction (Brown and Jenkins 1968; Jenkins and Moore 1973; Hearst and Jenkins 1974; Boakes 1977). For example, rats will acquire a conditioned response (CR) in which they will approach, contact, and bite a lever conditioned stimulus (CS) that signals the delivery of a food unconditioned stimulus (US). Sign-tracking is a key model for studying behavioral and neural mechanisms of normal and excessive motivational attraction to reward-paired stimuli (Lajoie and Bindra 1976; Berridge 2004; Tomie et al. 2008; Flagel et al. 2010; Robinson and Berridge 2013; Huys et al. 2014; Robinson et al. 2014).

Negative automaintenance, in which a lever press cancels reward, has been used to show that sign-tracking can be markedly sensitive to instrumental contingency changes (Williams and Williams 1969; Stiers and Silberberg 1974; Locurto et al. 1976), suggesting a sensitivity to response–reward associations (Skinner 1992). For example, Locurto et al. (1976) have found that lever contacts after sign-tracking are markedly reduced in rats moved to an omission schedule, with similar rates of lever contacts compared with rats exposed to extinction or random cue/reward delivery. However, typically, sign-tracking does not decline to zero (Atnip 1977; Eldridge and Pear 1987; Schwartz and Williams 1972a; Stiers and Silberberg 1974; Woodard et al. 1974), suggesting some motivational persistence as well. Thus, sign-tracking CRs may be partly sensitive to contingency and partly under control of motivational forces that promote its persistence.

Generally, it remains unclear to what quantitative extent the motivational attraction to the CS actually declines along with the declining CR when it cancels reward. Is the incentive value of the CS retained but masked by reduced lever pressing measures, or is it reduced as well?

To clarify this issue, we studied the response topography of sign-tracking CRs under negative automaintenance, as lever depressions are only one aspect of sign-tracking (Hearst and Jenkins 1974). Indeed, sign-tracking components can be ex-

pressed differently in sign-tracking and negative automaintenance conditions (Schwartz and Williams 1972b; Stiers and Silberberg 1974; Locurto et al. 1976; Davey et al. 1981), as well as when motivational processes are enhanced by brain manipulations (Mahler and Berridge 2009). Thus, we evaluated how sign-tracking may persist through differential expression of CRs.

Rats ($n = 10$) that had acquired a discriminative sign-tracking response were exposed to conditions in which lever depression led to reward cancellation (Supplemental Methods). These rats were the “sham” group used in a published experiment on the effects of accumbens lesions on sign-tracking acquisition (Chang and Holland 2013). The rats were restricted to 85% of pretask weights throughout the study. They were first trained on a task in which insertion of a CS+ lever for 10-sec predicted reward (0.1 mL of 8% sucrose) in a food cup following its retraction. A second CS– lever was inserted similarly on different trials, but did not predict reward. Within each 64-min session, there were 25 CS+ and 25 CS– trials (mean ITI = 77 sec; randomized order; lever assignment counterbalanced). Over the course of 12 training sessions, rats developed a robust sign-tracking CR to the CS+. Rats in Group Omission ($n = 5$) were then exposed to negative automaintenance conditions for 12 more days, during which CS+ lever depression canceled reward delivery. Rats in Group Control ($n = 5$) were yoked to the reward schedule of paired rats in Group Omission, thus equating reward exposure.

Figure 1A presents the mean number of lever presses per minute to the CS+ and CS– over the last day of sign-tracking acquisition (Last Acq) and over the course of negative automaintenance (Sessions 1–12). For automated measures of lever pressing (and food cup entries), we analyzed the entire 10 sec of CS+ and CS– presentations. Both Omission and Control rats showed similarly high levels of lever pressing on the last day of sign-tracking

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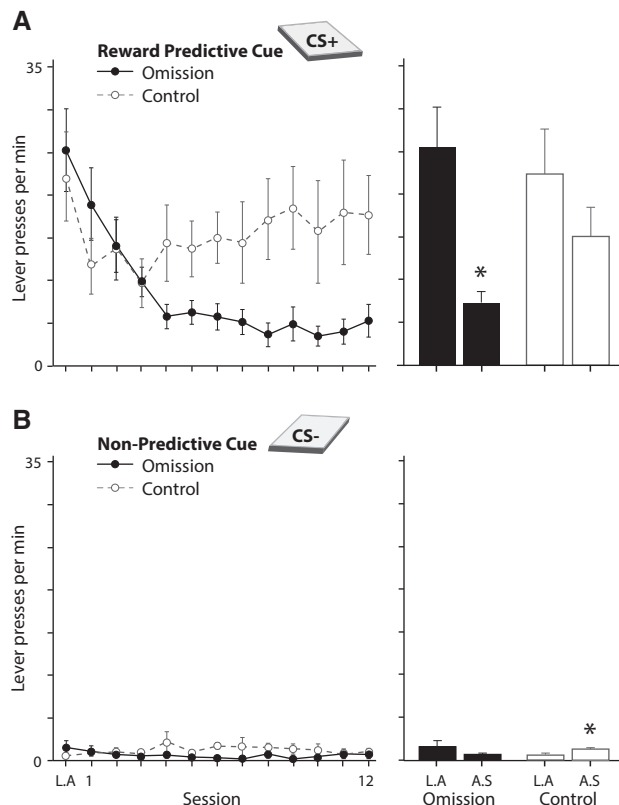


Figure 1. Negative automaintenance of sign-tracking. Lever presses per minute during, (A) the reward predictive CS+ lever insertion and, (B) during the nonpredictive CS- lever insertion. Line graphs at left show data from the last acquisition day (LA) followed by the 12 procedure sessions for Omission rats (black lines and circles) and yoked Control rats (dashed lines and open circles). Shown at right are the last acquisition compared with the average of the 12 sessions (AS). Rats in the Omission group markedly reduced CS+ lever pressing, but pressing still remained elevated above zero presses and above the minimal CS- presses. Control rats did not significantly change pressing, with the exception of a small elevation in CS- presses. (*) $P < 0.05$ (LA to AS within group).

acquisition (Cue: $F_{(1,8)} = 47.69$, $P < 0.01$; Group $F_{(1,8)} = 0.30$, $P = 0.60$; interaction: $F_{(1,8)} = 0.11$, $P = 0.75$).

Over the course of negative automaintenance, Omission rats rapidly reduced behaviors that led to depression of the CS+ lever once the negative contingency was imposed (Fig. 1A). CS+ pressing in yoked Control rats, however, remained statistically unchanged (Fig. 1A). Still, despite the negative contingency, Omission rats maintained higher levels of pressing to the CS+ than to the CS- throughout training (Fig. 1A,B). A 2 (Cue: CS+ versus CS-) \times 2 (Group: Omission versus Control) \times 12 (Session) ANOVA confirmed a main effect of Cue ($F_{(1,8)} = 36.59$, $P < 0.01$) and a marginal effect of Group ($F_{(1,8)} = 5.03$, $P = 0.06$), but no effect of Session ($F_{(11,88)} = 1.15$, $P = 0.34$). In addition, there were Group \times Session ($F_{(11,88)} = 4.46$, $P < 0.01$) and Cue \times Group \times Session interactions ($F_{(11,88)} = 4.14$, $P < 0.01$). The Cue \times Group and Cue \times Session interactions did not reach statistical significance (largest $F_{(1,8)} = 4.37$, $P = 0.07$). Concerning the nature of the Cue \times Group \times Session interaction, ANOVAs run over four-session blocks of negative automaintenance (Sessions 1–4, 5–8, and 9–12) confirmed that Omission and Control rats showed comparable levels of discriminative pressing to the CS+ and CS- over the first block (Sessions 1–4) (Cue \times Group: $F_{(1,8)} < 0.01$, $P = 0.95$). However, Control rats showed higher lev-

els of pressing than Omission rats to the CS+ over the second (Sessions 5–8) and third (Sessions 9–12) blocks of training (Sessions 5–8, Cue \times Group: $F_{(1,8)} = 8.07$, $P = 0.02$; Sessions 9–12, Cue \times Group: $F_{(1,8)} = 5.65$, $P = 0.045$).

We further evaluated the degree of sign-tracking within each group. For Control rats, the ANOVA confirmed a main effect of Cue ($F_{(1,4)} = 18.68$, $P = 0.01$) but no effect of Session ($F_{(11,44)} = 0.91$, $P = 0.54$) or Cue \times Session interaction ($F_{(11,44)} = 0.83$, $P = 0.61$). For Omission rats, there were main effects of Cue ($F_{(1,4)} = 34.54$, $P < 0.01$) and Session ($F_{(11,44)} = 8.41$, $P < 0.01$), as well as a Cue \times Session interaction ($F_{(11,44)} = 9.59$, $P < 0.01$). In addition, separate ANOVAs over four-session blocks of negative automaintenance for each group confirmed main effects of Cue for all three blocks (Control: smallest $F_{(1,4)} = 10.37$, $P = 0.03$; Omission: smallest $F_{(1,4)} = 9.11$, $P = 0.04$).

These data indicate that rats under the yoked Control condition maintained stable levels of sign-tracking to the CS+ across the 12 d under the new reward contingency. In contrast, rats under the Omission condition starkly reduced lever pressing, but still retained it at a low level. This conclusion was supported further by findings from comparing, within-subjects, the last sign-tracking acquisition day (LA) to the average of sessions (AS) of negative automaintenance (Fig. 1A, right): the Omission group showed a significant reduction in CS+ lever presses ($P = 0.01$), but the Control group did not ($P = 0.21$). CS- lever pressing was low throughout for both groups, though there was a marginal but significant rise in CS- pressing in Control rats ($P = 0.04$; Fig. 1B, right).

However, lever depression measures do not capture the full repertoire of a sign-tracking CR, and our analyses of the CR topography suggested a different story. Stimulus approach and consummatory behaviors were video-scored during every fifth CS+ and CS- presentation (Mahler and Berridge 2009) for the last acquisition day and sessions 4, 8, and 12. Behaviors were sampled every 2 sec for the duration of each CS presentation. Stimulus consummatory behaviors included lever grasps (grabs of the lever with forepaws) and lever bites (active bites or nibbles of the lever). Stimulus approach included lever orienting (head movement in close proximity toward the lever), lever sniffs (at least one sniff action in close proximity of the lever), and nonconsummatory lever contacts (bodily touching of the lever without biting or grasping). Because rats have to approach the lever in order to show a consummatory behavior, an approach behavior was also counted if a rat was grasping or biting the lever. Behavior counts were then totaled for each session and added by category.

On the last acquisition day, Omission and Control rats displayed similarly high levels of approach behaviors during the CS+ but not the CS- [Cue ($F_{(1,8)} = 133.56$, $P < 0.01$), no effect of Group ($F_{(1,8)} = 1.28$, $P = 0.29$), or interaction ($F_{(1,8)} = 2.46$, $P = 0.16$)]. The groups were also similar in stimulus consummatory behaviors on that day, which were more frequent on the CS+ than the CS- lever, due to the high levels of consummatory responses to the CS+ and minimal response generally to the CS- (Cue: $F_{(1,8)} = 24.77$, $P < 0.01$; Group: $F_{(1,8)} = 0.51$, $P = 0.50$; interaction: $F_{(1,8)} = 0.51$, $P = 0.50$).

Despite a dramatic decrease in lever presses over the course of negative automaintenance, Omission rats continued to approach the sucrose-paired lever at comparable rates to Control rats. Approach behaviors remained high for the CS+ for both groups, while approach to the CS- remained low [Cue: $F_{(1,8)} = 34.15$, $P < 0.01$; no effect of Group: $F_{(1,8)} = 0.46$, $P = 0.52$, Session $F_{(2,16)} = 1.55$, $P = 0.24$, Cue \times Group: $F_{(1,8)} = 1.73$, $P = 0.23$, or other interactions: largest $F = 1.74$, $P = 0.21$]. In comparing behaviors within each group, Omission rats showed a greater number of approach behaviors to the CS+ than CS- ($F_{(1,4)} = 36.63$, $P < 0.01$), as did Control rats ($F_{(1,4)} = 7.89$, $P = 0.048$).

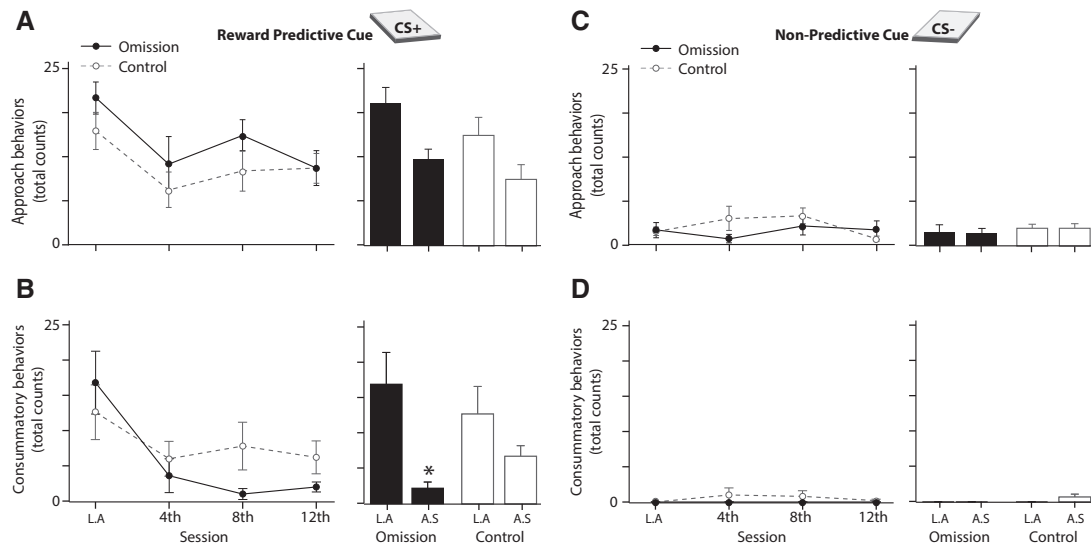


Figure 2. Omission procedure dissociates approach versus consummatory features of sign-tracking. (A) Approach behaviors directed to the CS+ lever (orient, sniff, contact) on the last acquisition session and sessions 4, 8, and 12 of negative automaintenance for Omission and Control rats. Bar graph shows averaged sessions 4, 8, and 12 (AS) compared with the last acquisition session (LA). (B) Consummatory behaviors directed to the CS+ lever (grasp and bite). Omission rats increased CS+ approach behaviors and decreased consummatory behaviors that could lead to lever depression and reward cancellation, thus maintaining a fairly high level of CS+ engagement relative to Control rats but reallocating it to mainly approach behaviors. (*) $P < 0.05$ (LA to AS within group for CS+). (C) Approach behaviors directed to the CS- lever. (D) Consummatory behaviors directed to the CS- lever. No comparisons between LA and AS were significant for CS-.

These trends were further supported by repeated-measures ANOVAs comparing the last acquisition day to the average of negative automaintenance behavior: Omission rats did not significantly change their CS+ approach ($P = 0.07$; Fig. 2A), nor did Control rats ($P = 0.10$; Fig. 2A), and neither Group changed their CS- approach (P 's > 0.05) (Fig. 2C).

Stimulus consummatory behaviors followed a similar trend (Fig. 2B). Overall, rats showed a greater number of consummatory behaviors to the CS+ than CS-, but there were moderate group differences (Cue: $F_{(1,8)} = 7.82$, $P = 0.02$; Group: $F_{(1,8)} = 4.11$, $P = 0.08$; Session: $F_{(2,16)} = 0.25$, $P = 0.78$; all interactions: largest $F = 1.68$, $P = 0.23$). Nevertheless, under direct repeated-measures comparison to the last acquisition day, Omission rats showed significantly reduced consummatory actions ($P = 0.027$) on average during negative automaintenance, which were 87% reduced. Control rats' consummatory behaviors did not change significantly ($P = 0.20$; 47% reduced). Neither group changed their consummatory response to the CS- (P 's > 0.05) (Fig. 2).

We further explored these effects by assessing the contribution of each single component of the CR to the total number of behaviors exhibited using nonparametric Kruskal-Wallis tests (Fig. 3). For this, we simply counted each observed instance of a behavior. Omission rats showed an elevation in both their proportion of lever sniffing ($P = 0.03$), which increased from 0% to 27% of the total CR, and orienting ($P = 0.03$), which increased from 5% to 25% of the total CR (Fig. 3A,B). Conversely, the proportion of the CR that was lever bites declined dramatically from 77% to 17% ($P = 0.01$) (Fig. 3A,B). These changes were at the heart of a robust restructuring of the approach versus consummatory sign-tracking CRs made to the CS+ that occurred from the last acquisition day through the negative automaintenance sessions (Fig. 3B). Overall, on the last acquisition day, 82% of the CR took the form of approach-and-consume behaviors (mainly biting). During negative automaintenance, this proportion reversed, with a nearly identical 82.5% of the CR being composed instead of approach-only behaviors. In comparison, Control rats did not show a change in their proportion of any CS+ sign-tracking CR,

instead exhibiting mostly lever bites throughout testing (Fig. 3A,B). What this suggests is that Omission rats did not cease responding once they exhibited an initial approach CR. Instead, they continued to exhibit approach-specific behaviors like orientations and sniffs, while not subsequently emitting a consummatory response. Thus, the frequency of approach-only behaviors rose for this group. In contrast, Control rats continued emitting consummatory responses, and thus approach-only counts like orientations and sniffs were comparatively low.

Percent of time in the food cup during CS presentations was also assessed under the possibility that Omission rats may have engaged more with the food source (i.e., goal-tracking) when lever presses canceled reward. This was not the case: both Omission and Control rats maintained fairly consistent and low levels of food cup time (ANOVA on Cue \times Group \times Session: largest main effect $F_{(1,8)} = 2.54$, $P = 0.15$; largest interaction $F_{(11,88)} = 1.55$, $P = 0.13$). When comparing the last acquisition day to the average of the negative automaintenance sessions, Omission rats did show a moderate increase in food cup time during the CS+ ($P = 0.036$), which appeared related to a low level of food cup time on the last acquisition day (Supplemental Fig. 1). Still, food cup entry time during negative automaintenance accounted for only 12.46% of their time during the CS+, and food cup time during the CS- was comparable at 12.82%. No group changed food cup entry time behavior during the CS- (P 's > 0.05). Hand-scored food cup orientations during the CS+ and CS- were also unchanged and similar between groups (P 's > 0.05) (Supplemental Fig. 1).

Collectively, these findings support the conclusion that sign-tracking rats exposed to an omission procedure reduce behaviors that lead to reward cancellation (lever depression) including consummatory behaviors like biting, but they concomitantly maintain approach behaviors like orientations and sniffs. Indeed, when aggregating hand-scored consummatory and approach behaviors, Omission rats showed a relatively moderate 41% decline in sign-tracking from the last acquisition day that was essentially identical to the decline in Control rats (40%). This equivalence

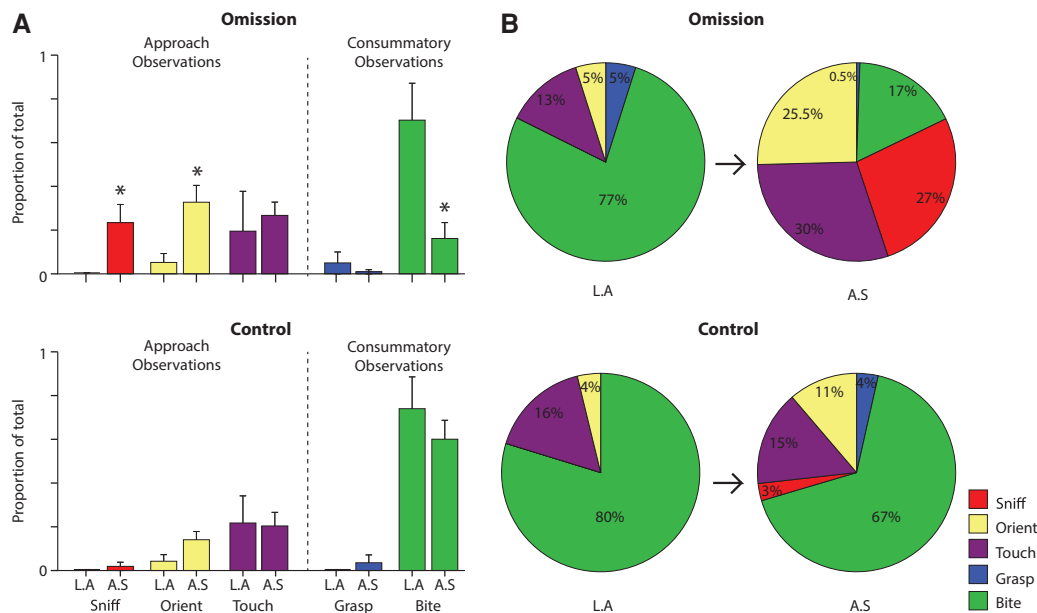


Figure 3. CS+ approach and consummatory behavior microstructure. (A) Proportion of sign-tracking responses to the CS+ on the last acquisition (LA) and negative automaintenance days (AS) that were sniffs (red), orientations (yellow), touches (purple), grasps (blue), and bites (green). *Top* graph shows the Omission group, *bottom* shows the Control group. (*) $P < 0.05$ (LA to AS within group). (B) Pie charts representing the proportion of responses on these days with approach and consummatory segments denoted.

contrasted sharply to the Omission rats' larger 71% decline in automated lever press counts, while Controls only reduced lever press counts by 22%. Thus ~60% of the acquired sign-tracking CR persisted despite the omission schedule, and the persistence was similar to Controls. This suggests that the CS+ retained much of its motivational value and that its expression in behavior was highly flexible.

Although Locurto et al. (1976) observed differences in response topography between rats placed on sign-tracking and omission schedules, these differences were not quantified. Based on a near-full reduction in lever contacts in their study, Locurto et al. (1976) argued that omission training was ineffective at maintaining responding to the lever. However, our data suggest that sign-tracking can be robustly maintained under omission training as demonstrated by a loss of terminal (consummatory) behaviors yet maintenance in approach-only behaviors.

Approach behaviors were not put under the omission contingency, and therefore we do not know what their sensitivity would be. Similarly, we recognize the potential role for stimulus-response associations in the residual lever press responses. Nevertheless, we suggest the maintenance of approach behaviors are best captured by incentive motivation concepts, which suggest that such sign-tracking CRs occur because of a motivational attraction to the CS+, a motivation that can become decoupled from trial-to-trial variations in reward outcome. The fact that rats persisted in engaging with the CS+ instead of favoring the food cup lends further support to the argument here that the incentive value of the CS+ remained high.

In the brain, limbic circuits have been implicated in sign-tracking (Cardinal et al. 2002; Mahler and Berridge 2009; Flagel et al. 2010; Chang et al. 2012a,b, 2015; Saunders and Robinson 2012; Chang and Holland 2013). A fruitful effort would be to resolve the neural mechanisms by which rats adaptively modify their responses when CS–CR–US conditions change but CS incentive salience remains strong. This question gets to the heart of an issue facing efforts to understand the brain basis of addictive behaviors, in which addicts can exhibit flexibility in their pursuit

of drug-associated stimuli. Evidence here indicates that the ability of a salient cue to pull in behavior can be rooted in a motivational process but not necessarily tied down to a specific behavioral routine. Such persistence-with-flexibility features may be useful, quantifiable targets for addiction research.

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References

- Atnip GW. 1977. Stimulus- and response-reinforcer contingencies in autoshaping, operant, classical, and omission training procedures in rats. *J Exp Anal Behav* **28**: 59–69.
- Berridge KC. 2004. Motivation concepts in behavioral neuroscience. *Physiol Behav* **81**: 179–209.
- Boakes R. 1977. Performance on learning to associate a stimulus with positive reinforcement. In *Operant-Pavlovian interactions* (ed. Davis H, Hurwitz H), pp. 67–97. Lawrence Erlbaum Associates, Hillsdale, NJ.
- Brown PL, Jenkins HM. 1968. Auto-shaping of the pigeon's key-peck. *J Exp Anal Behav* **11**: 1–8.
- Cardinal RN, Parkinson JA, Lachenal G, Halkerston KM, Rudarakanchana N, Hall J, Morrison CH, Howes SR, Robbins TW, Everitt BJ. 2002. Effects of selective excitotoxic lesions of the nucleus accumbens core, anterior cingulate cortex, and central nucleus of the amygdala on autoshaping performance in rats. *Behav Neurosci* **116**: 553–567.
- Chang SE, Holland PC. 2013. Effects of nucleus accumbens core and shell lesions on autoshaped lever-pressing. *Behav Brain Res* **256**: 36–42.
- Chang SE, Wheeler DS, Holland PC. 2012a. Effects of lesions of the amygdala central nucleus on autoshaped lever pressing. *Brain Res* **1450**: 49–56.
- Chang SE, Wheeler DS, Holland PC. 2012b. Roles of nucleus accumbens and basolateral amygdala in autoshaped lever pressing. *Neurobiol Learn Mem* **97**: 441–451.

- Chang SE, Todd TP, Bucci DJ, Smith KS. 2015. Chemogenetic manipulation of ventral pallidal neurons impairs acquisition of sign-tracking in rats. *Eur J Neurosci* **42**: 3105–3116.
- Davey GC, Oakley D, Cleland GG. 1981. Autoshaping in the rat: effects of omission on the form of the response. *J Exp Anal Behav* **36**: 75–91.
- Eldridge GD, Pear JJ. 1987. Topographical variations in behavior during autoshaping, automaintenance, and omission training. *J Exp Anal Behav* **47**: 319–333.
- Flagel SB, Clark JJ, Robinson TE, Mayo L, Czuj A, Willuhn I, Akers CA, Clinton SM, Phillips PE, Akil H. 2010. A selective role for dopamine in stimulus-reward learning. *Nature* **469**: 53–57.
- Hearst E, Jenkins H. 1974. *Sign-tracking: the stimulus-reinforcer relation and directed action*. Monograph of the Psychonomic Society, Austin.
- Huys QJ, Tobler PN, Hasler G, Flagel SB. 2014. The role of learning-related dopamine signals in addiction vulnerability. *Prog Brain Res* **211**: 31–77.
- Jenkins HM, Moore BR. 1973. The form of the auto-shaped response with food or water reinforcers. *J Exp Anal Behav* **20**: 163–181.
- Lajoie J, Bindra D. 1976. An interpretation of autoshaping and related phenomena in terms of stimulus-incentive contingencies alone. *Can J Psychol* **30**: 157–173.
- Locurto C, Terrace HS, Gibbon J. 1976. Autoshaping, random control, and omission training in the rat. *J Exp Anal Behav* **26**: 451–462.
- Mahler SV, Berridge KC. 2009. Which cue to “want?” Central amygdala opioid activation enhances and focuses incentive salience on a prepotent reward cue. *J Neurosci* **29**: 6500–6513.
- Robinson MJ, Berridge KC. 2013. Instant transformation of learned repulsion into motivational “wanting”. *Curr Biol* **23**: 282–289.
- Robinson TE, Yager LM, Cogan ES, Saunders BT. 2014. On the motivational properties of reward cues: individual differences. *Neuropharmacology* **76**(Pt B): 450–459.
- Saunders BT, Robinson TE. 2012. The role of dopamine in the accumbens core in the expression of Pavlovian-conditioned responses. *Eur J Neurosci* **36**: 2521–2532.
- Schwartz B, Williams DR. 1972a. The role of the response-reinforcer contingency in negative automaintenance. *J Exp Anal Behav* **17**: 351–357.
- Schwartz B, Williams DR. 1972b. Two different kinds of key peck in the pigeon: some properties of responses maintained by negative and positive response-reinforcer contingencies. *J Exp Anal Behav* **18**: 201–216.
- Skinner BF. 1992. ‘Superstition’ in the pigeon. 1948. *J Exp Psychol Gen* **121**: 273–274.
- Stiers M, Silberberg A. 1974. Lever-contact responses in rats: automaintenance with and without a negative response-reinforcer dependency. *J Exp Anal Behav* **22**: 497–506.
- Tomie A, Grimes KL, Pohorecky LA. 2008. Behavioral characteristics and neurobiological substrates shared by Pavlovian sign-tracking and drug abuse. *Brain Res Rev* **58**: 121–135.
- Williams DR, Williams H. 1969. Auto-maintenance in the pigeon: sustained pecking despite contingent non-reinforcement. *J Exp Anal Behav* **12**: 511–520.
- Woodard WT, Ballinger JC, Bitterman ME. 1974. Autoshaping: Further study of “negative automaintenance.” *J Exp Anal Behav* **22**: 47–51.

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