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Amphetamine-Induced Sensitization and Reward Uncertainty Similarly Enhance Incentive Salience for Conditioned Cues

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CITATION
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Amphetamine and stress can sensitize mesolimbic dopamine-related systems. In Pavlovian autoshaping, repeated exposure to uncertainty of reward prediction can enhance motivated sign-tracking or attraction to a discrete reward-predicting cue (lever-conditioned stimulus; CS+), as well as produce cross-sensitization to amphetamine. However, it remains unknown how amphetamine sensitization or repeated restraint stress interact with uncertainty in controlling CS+ incentive salience attribution reflected in sign-tracking. Here rats were tested in 3 successive phases. First, different groups underwent either induction of amphetamine sensitization or repeated restraint stress, or else were not sensitized or stressed as control groups (either saline injections only, or no stress or injection at all). All next received Pavlovian autoshaping training under either certainty conditions (100% CS–UCS association) or uncertainty conditions (50% CS–UCS association and uncertain reward magnitude). During training, rats were assessed for sign-tracking to the CS+ lever versus goal-tracking to the sucrose dish. Finally, all groups were tested for psychomotor sensitization of locomotion revealed by an amphetamine challenge. Our results confirm that reward uncertainty enhanced sign-tracking attraction toward the predictive CS+ lever, at the expense of goal-tracking. We also reported that amphetamine sensitization promoted sign-tracking even in rats trained under CS–UCS certainty conditions, raising them to sign-tracking levels equivalent to the uncertainty group. Combining amphetamine sensitization and uncertainty conditions did not add together to elevate sign-tracking further above the relatively high levels induced by either manipulation alone. In contrast, repeated restraint stress enhanced subsequent amphetamine-elicited locomotion, but did not enhance CS+ attraction.

Keywords: motivation and reward, uncertainty, incentive salience, sensitization, amphetamine and stress

Rewards (unconditioned stimuli or UCSs) and the Pavlovian conditioned stimuli (CSs+) or cues that predict them can take on motivational magnet properties that make them ‘wanted’ and approached. The incentive-salience hypothesis suggests that mesolimbic dopamine released in the nucleus accumbens and other brain limbic structures in response to encountering a cue for reward, is part of the mechanism that triggers surges of cue-triggered ‘wanting’ for rewards, and produces motivated attraction toward their associated cues that become ‘wanted’ CSs+ (Berridge & Robinson, 1998). Repeated exposure to drugs of abuse such as amphetamine are known to sensitize mesolimbic dopamine-related systems and to enhance incentive salience (Bradberry, Barrett-Larimore, Jatlow, & Rubino, 2000; Giorgetti, Hotsenpiller, Ward, Teppen, & Wolf, 2001; Vezina, 2004; Wyvoll & Berridge, 2001). Incentive sensitization has been suggested to be an important mechanism of addiction (M. J. F. Robinson, Robinson, & Berridge, 1994; T. E. Robinson & Berridge, 1993).

Similarly, recent evidence suggests that uncertainty of reward delivery in CS–UCS relations, creating an approach toward a gambling-like scenario, can also raise dopamine levels and similarly contribute to the motivational attraction to cues for uncertain rewards, which could be potentially relevant to addiction-type pursuit of gambling by some individuals (Boileau et al., 2013; Hart, Clark, & Phillips, 2015; Linnet et al., 2012; M. J. F. Robinson, Anselme, Fischer, & Berridge, 2014; Zack, Featherstone, Mathewson, & Fletcher, 2014). We have previously demonstrated the incentive salience amplification as enhancement of sign-tracking, or higher motivated attraction toward a reward-predictive CS+ when the CS–UCS relationship is uncertain (e.g., a 50% probability that sucrose UCS will follow the CS+) than when it is certain (i.e., 100%). A sign-tracking response occurs when an
animal approaches, sniffs, and nibbles its CS+ (insertion of a metal lever accompanied by an auditory label) that predicts subsequent sucrose reward, whereas a goal-tracking response involves a Pavlovian approach instead toward the goal or sucrose dish where the UCS will appear, also triggered by the CS+ presentation (Boakes, 1977; Hearst & Jenkins, 1974). Higher attraction to an uncertain cue is paradoxical in one sense, in that it contradicts the idea that the motivation value of a reward cue should always be linearly proportional to the predictive value of that CS+. But the dissociation between motivational attraction and predictive certainty is consistent with the incentive salience hypothesis that cue ‘wanting’ (CS attraction) is separable from cue learning (CS–UCS association). Enhanced attraction to an uncertain reward CS+ is also compatible with the idea that gambling-related cues become potent triggers of incentive motivation in part because of the uncertainty of their reward association.

Potential interaction between sensitization and uncertainty is suggested by findings that exposure to uncertainty in reward conditions can promote cross-sensitization to subsequent amphetamine (Berridge & Robinson, 1998; Boileau et al., 2013; Linnet et al., 2012; Singer, Scott-Raillton, & Vezina, 2012; Zack et al., 2014). Also, the excitatory effects of reward uncertainty on behavior are relatively persistent in time, similar to sensitization (Bradberry et al., 2000; Giorgetti et al., 2001; Vezina, 2004; Wyvell & Berridge, 2001), even after the uncertainty conditions are reduced (M. J. F. Robinson, Anselme, et al., 2014). However, it is not yet known how reward uncertainty and amphetamine sensitization would compare or interact regarding the motivational attraction to reward CSs+. Finally, stress interacts with drug use, cue-induced craving, and reinstatement of drug taking, and exhibits cross-sensitization to amphetamine (Berridge & Robinson, 1998; Buffalari & See, 2009; Logrip & Zorrilla, 2012; Stewart, 2003; Zorrilla, Logrip, & Koob, 2014). It is therefore of interest to know how amphetamine sensitization or repeated restraint stress might interact subsequently with reward uncertainty and sign-tracking.

Here rats were initially sensitized to amphetamine or exposed to repeated restraint stress, or else were not sensitized or stressed as controls. Then all rats received Pavlovian conditioning (autoshaping), with a lever insertion as the CS+ and sucrose pellet delivery as the UCS, to assess sign-tracking and goal-tracking responses in an undrugged state. Locomotor activity and sensitization were then assessed by administering a single amphetamine challenge. Based on previous work, it was predicted that (a) reward uncertainty would enhance sign-tracking, and that (b) previous amphetamine exposure or stress exposure would similarly enhance sign-tracking toward a reward-predictive CS+.

Materials and Methods

Animals and Housing Conditions

Female Sprague–Dawley rats (N = 48; 200–280 g) were bred and reared by the research group from animals purchased from Harlan (Indianapolis, IN), similarly to our previous studies. Rats were weaned at 21 days of age and housed (41 cm × 25.4 cm × 20.3 cm) in groups of two–three animals with possible litter effects controlled for during group assignment. Animals had ad libitum access to water and chow, and were food restricted to 90% of free-feeding body weight during autoshaping procedures. All rats were handled for a minimum of 4 days before the start of any of the procedures. For additional details, see M. J. F. Robinson, Anselme, et al. (2014). All experimental procedures were approved by the University Committee on the Use and Care of Animals at the University of Michigan.

Apparatus

**Locomotor activity chambers.** Sixteen locomotion chambers (actometers) controlled by cross-break software were used to measure the activity level of the rats. Locomotor activity was monitored in Plexiglas chambers (41 cm × 25.4 cm × 20.3 cm) that contained a clear plastic insert in the cage center (23 × 6.3 × 20.3 cm). Locomotor activity was measured by the total number of photocell beam breaks in a photo-beam array.

**Sensory stimulation and restraint-stress chambers.** In clear Plexiglas cages (41 × 25.4 × 20.3 cm), rats experienced restraint stress by being placed in clear cylindrical restraints equipped with air holes, a tail slot, and Velcro straps and being exposed to a light source (1000–1300 lux) and loud sounds supplied by a continuous hard-rock album (by Iggy Pop & The Stooges; 80–86 dB).

**Autoshaping.** Each rat was trained and tested in one of eight Med Associates (St. Albans City, VT) autoshaping chambers (30.5 cm × 24.1 cm × 21 cm) with Plexiglas floor, ceiling, and walls. Each chamber was equipped with a magazine dish (3 cm × 2 cm × 1 cm), two levers (one acting as a CS and the other as a control stimulus) on either side of the magazine, and two video cameras. For more details, see M. J. F. Robinson, Anselme, et al. (2014).

Procedure

**Phase 1: Sensitization induction.**

**Amphetamine sensitization.** On Day 0, all rats (amphetamine-sensitization and saline-control groups) received an intraperitoneal injection of saline (1 ml/kg) and were allowed to habituate in the novel locomotion chambers for 45 min to measure their baseline locomotor activity. Starting on Day 1, depending on group assignment, rats received either amphetamine injections (N = 16; 2 mg/kg, IP) or saline (N = 16) each day for 7 days, and each time were immediately afterward placed in the locomotion chamber. To assess locomotion, photo-beam breaks were measured in 5-min bins for the span of 45 min. Following the 7 days of injections, rats were returned to their home cage to allow sensitization to incubate for an additional period of 7 days.

**Repeated restraint stress.** Stress rats were exposed daily to restraint and sensory-stress stimulation (N = 16; exposure to bright light and loud discordant soundtrack), for a period of 45 min each day for 7 days. Restraint has previously been shown to produce a robust activation of the hypothalamic–pituitary–adrenal axis (Evanson & Herman, 2015; Maghsoudi et al., 2014), while sensory-stress stimulation potentiates mesolimbic elicitation of defensive treading and is avoided when given a choice (Reynolds & Berridge, 2008). No-stress control animals were singly placed in a separate cage for 45 min each day for 7 days and allowed to freely explore. All animals were then returned to their home cage and allowed to incubate for 7 days.

Prior to Pavlovian training, rats were exposed to sucrose pellets in their home cage, and received 1 day of magazine training (25
pellets were dropped one by one into the magazine dish in the absence of CS lever presentation, 30–90 s intertrial interval (ITI)) 7 days after the last day of sensitization treatments.

**Phase 2: Pavlovian autoshaping.** Rats in the amphetamine, saline, stress, and no-stress groups were further subdivided into either CS–UCS certainty (C, 100%-1) or uncertainty (U, 50%-1–2–3) autoshaping conditions. In the certainty condition, 100% of CS lever presentations were immediately followed by the delivery of a single sucrose pellet. In the uncertainty condition, CS lever presentations were either followed by no sucrose pellet 50% of the time or by either 1, 2, or 3 pellets with an equal probability (16.7%) for the remaining 50% of CS+ presentations; rewarded and nonrewarded trials were intermixed on a random basis. We have shown elsewhere that, in itself, the variability in food amounts (1–2–3) used here generated similar performance to that obtained with fixed food amounts (100%-1), but that it contributed to performance enhancement when associated with a 50% probability of reward (Anselme, Robinson, & Berridge, 2013). All groups underwent 8 autoshaping sessions over 8 days. In every condition, each rat received a total of 36 CS+ presentations per session for a total of 36 UCS sucrose pellets delivered. Each CS+ presentation consisted of the insertion of a lever into the chamber for 8 s, accompanied by a 2.9 kHz tone and a light at the base of the lever. Sucrose pellet delivery to the magazine dish occurred immediately after retraction of the lever. Seven groups of eight individuals were obtained: amphetamine + certainty (Amph + C), amphetamine + uncertainty (Amph + U), saline + certainty (Saline + C), saline + uncertainty (Saline + U), stress + certainty (Stress + C), stress + uncertainty (Stress + U), and no stress + uncertainty (No stress + U). The saline groups were used as controls for comparison to both the amphetamine and stress groups. Indeed, based on the results of the No stress + U groups, we see that the injection of saline neither impacted performance in autoshaping or performance during the test of locomotor sensitization.

**Sign-tracking versus goal-tracking assessment.** An animal would be classified as a sign-tracker if it was to sniff, nibble, bite, and grasp the CS+ lever, resulting in lever presses three times more frequently than it did the sucrose dish during CS+ presentations on the final training day. The criterion for classification as goal-tracker was to sniff, nibble, bite and grasp the dish, resulting in magazine entries at least three times more frequently than the lever during CS+ presentations on the last day of training. An individual could be classified as a mixed responder if it directed between 33% and 66% of its total number of responses to CS+, and the remaining responses to the goal dish (Flagel, Watson, Robinson, & Akil, 2007). All animals developed some form of conditioned response across testing, whether sign-tracking, goal-tracking or some mixture of both.

**Phase 3: Test of locomotor sensitization.** Following the last day of autoshaping, all animals received an amphetamine injection (0.75 mg/kg, IP) and were placed in locomotion chambers (the same as in Phase 1) for 45 min. Activity levels were measured in 5-min bins. Animals in the stress and no-stress conditions first received a saline injection and were placed in locomotion chambers for 45 min (to control for habituation and provide a baseline recording of general activity) prior to receiving an amphetamine injection.

**Statistical Analysis**

Mixed ANOVAs were used for most between- and within-subject comparisons. Planned comparisons allowed us to assess one data set relative to another. One-way ANOVAs were used as appropriate. Two-tailed tests were used, and the null hypothesis was rejected at p < .05. All measurements are indicated as mean ± SE. Due to a recording malfunction, data for one animal in the Saline + C group was removed from the analysis on Day 5 of autoshaping.

**Results**

**Phase 1: Sensitization Induction**

Saline and amphetamine groups did not differ initially in locomotion following a saline injection on Day 0 (F<sub>1,28</sub> = 0.109, p = .744). During initial induction of sensitization (Figure 1A), amphetamine administration elevated locomotor responses (beam breaks) compared with all rats receiving saline injections (group: F<sub>1,28</sub> = 114.077, p = .000). Further, amphetamine-exposed rats showed signs of an incrementally sensitized locomotor response with increased activity across the 7 days of sensitization induction (day: F<sub>7,196</sub> = 22.574, p = .000; Group × Day interaction: F<sub>7,196</sub> = 27.491, p = .000), which became increasingly pronounced (Days 1–7: F<sub>1,28</sub> = 43.675, p = .000). By contrast, locomotion levels remained unchanged across the 7 days in the saline group (Days 1–7: F<sub>1,28</sub> = 0.446, p = .510; see Figure 1A). Overall, these results suggest that repeated amphetamine administration produced detectable sensitization over the 7-day period, reflected as increased locomotion over initial basal activity level. The amphetamine sensitization regimen used here did not significantly reduce body weight below saline group levels, which might have otherwise influenced subsequent sign-tracking and goal-tracking behavior (F<sub>1,30</sub> = 1.719, p = .200).

**Phase 2: Pavlovian Sign-Tracking (Uncertainty, Prior Sensitization, and Prior Stress Effects)**

Uncertainty versus certainty of CS–UCS prediction. Following an additional week to allow for incubation of any sensitization treatment, autoshaping training was conducted for all rats, and individuals were assessed for sign-tracking versus goal-tracking phenotypes. Exposure to reward uncertainty during Pavlovian autoshaping in saline-treated rats (Saline + U group) led to significantly greater sign-tracking attraction toward the CS+ lever over 8 days, as measured by grasps and/or consumatory nibble and bite responses on the metal object, resulting in lever presses, compared with similar saline-treated rats that were trained under certainty conditions (Saline + C group; see Figure 1B). There was an effect of group (F<sub>1,14</sub> = 6.373, p = .024), and day (F<sub>7,98</sub> = 22.548, p = .000), but no interaction (F<sub>7,98</sub> = 0.544, p = .799).

Rats in Group Saline + U pressed the CS lever 167% more than rats in Group Saline + C on Day 8. As noted in Figure 1B, a significant difference between Groups Saline + C and Saline + U was shown for six of the training days (F<sub>1,14</sub> ≥ 4.683, p ≤ .048). Both saline-treated groups gradually increased response rates across 8 days of training (Days 1–8, C: F<sub>1,14</sub> = 24.226, p = .000; U: F<sub>1,14</sub> = 40.637, p = .000).
Conversely, the number of goal-tracking responses (Figure 1C), reflected by magazine-beam breaks caused by nose entries into the sucrose dish during CS+ presentations, declined under uncertainty conditions from on average 75 on the first day to approximately 10 by the final few days ($F_{1,14} = 8.136, p = .013$). By contrast, goal-tracking responses for rats in the certainty conditions remained relatively stable over the week, and did not change significantly from Day 1 to Day 8 ($F_{1,14} = 0.001, p = .973$). Consequently, the uncertainty group of saline-treated rats emitted markedly fewer goal-tracking responses than the certainty group by Day 8 ($F_{1,14} = 6.608, p = .022$). Among the sign-trackers of the two saline groups, 100% of individuals emitted at least some goal-tracking responses, but the amount of those responses remained limited (±41 on Day 8). Figure 1C indicates an overall goal-tracking difference between Groups Saline + U and Saline + C ($F_{1,14} = 7.519, p = .016$) as well as an effect of Day ($F_{7,98} = 5.576, p = .000$), and a Group × Day interaction ($F_{7,98} = 2.313, p = .032$). Consistent with the evidence shown above that uncertainty enhanced sign-tracking responses, uncertainty also lowered goal-tracking responses (probably due to response competition as rats can only perform one behavior at a time). As noted, a significant difference was shown on all but the first day ($F_{1,14} s = 5.396, ps < .036$). Here also, 100% of goal-trackers (18.7% of saline-treated individuals) emitted a small number of sign-tracking responses (±75 on Day 8).

Incidence of goal-tracker versus sign-tracker phenotypes. A direct comparison between certainty and uncertainty groups indicated that uncertainty produced more sign-trackers (Saline + U = 100%; Saline + C = 50%), and fewer goal-trackers (Saline + U = 0%; Saline + C = 37.5%). For these groups, uncertainty actually abolished the existence of any individuals who met criterion for classification as a goal-tracker phenotype (i.e., three times more approaches to the goal dish than to the lever during CS+ lever presentations; although we cannot rule out the possibility that the uncertainty group contained more exclusive sign-trackers by chance).

Prior amphetamine-sensitization effects. In contrast to saline-treated rats described above, all rats that had previously been exposed to amphetamine-sensitization (Amph + C and Amph + U) displayed relatively higher sign-tracking approaches to the CS+ lever than saline-exposed rats even when trained under certainty conditions (see Figure 1D). In this respect, rats trained with CS–UCS certainty that had earlier received amphetamine exposure (Amph + C) pressed their CS+ lever more than saline-treated rats trained under certainty as described above (Saline + C; $t_{120} = 3.219, p = .002$). Adding uncertainty to amphetamine sensitization as a combination (Amph + U) did not further elevate sign-tracking responses detectably above the already elevated sensitized level (Amph + C), indicating that the two forms of sign-tracking elevation were not additive here or that the elevation had...
reached a measurement ceiling (group: F1,14 = 0.002, p = .962; interaction: F2,98 = 0.236, p = .975; see Figure 1D). Both groups showed a steady increase in lever responses across the 8 daily sessions (Day: F7,98 = 13.325, p = .000), a clear indication that both groups correctly learned the task (Days 1–8: C: F1,14 = 11.943, p = .004; U: F1,14 = 18.322, p = .001), but the two groups did not significantly differ on any day.

However, despite higher sign-tracking responses, amphetamine-sensitization rats did not necessarily make fewer goal-tracking entries into the sucrose magazine than saline-treated rats (Amph + U vs. Saline + U: F1,14 = 1.386, p = .259; Amph + C and Saline + C: F1,14 = 0.248, p = .626). It should be noted also that goal-tracking performance did not significantly change between Day 1 and Day 8 for the rats sensitized to amphetamine: (U: F1,14 = 0.653, p = .432; C: F1,14 = 0.753, p = .400), unlike the decline over days seen in the saline-treated uncertainty group above. Also, amphetamine-sensitization rats all showed similar numbers of goal-tracking responses regardless of whether they trained under uncertainty or certainty conditions: Although there seemed to be a slight trend for uncertainty rats to track the goal less (see Figure 1E), that difference did not reach statistical significance (F1,14 = 0.451, p = .513). Overall, it appears that amphetamine sensitization enhanced sign-tracking without reducing goal-tracking, and possibly even protected goal-tracking from being significantly reduced by concomitant uncertainty training (only a slight and nonsignificant trend toward reduction of goal-tracking was observed). That sensitization pattern of sign-tracking enhancement was different from the uncertainty pattern described above, which enhanced sign-tracking at the expense of reduced goal-tracking.

Incidence of goal-tracker versus sign-tracker phenotypes. How is it possible that amphetamine maintained goal-tracking responses at normal levels while enhancing sign-tracking? In terms of individual classification as phenotypes, amphetamine-sensitized rats were primarily goal-trackers (Amph + U = 75%; Amph + C = 62.5%), with only a few goal-trackers (Amph + U = 12.5%; Amph + C = 37.5%) and only one animal in the uncertainty group displaying mixed behavior. Of the animals exposed to certainty following sensitization, sign-tracking responses ranged from 230 to 5, while goal-tracking ranged from 281 to 0 (three goal-trackers: 281–171, all others = 8). Among animals sensitized and exposed to uncertainty, sign-tracking responses ranged from 205 to 34, and goal-tracking responses ranged from 354 to 0 on Day 8 (single goal-tracker = 354, all others ≤ 84). Consequently, the lack of a significant decrease in goal-tracking in sensitized animals under uncertainty may have been largely driven by a single goal-tracker and explained by a high standard error (Amph + U: SE = 42.8). In fact, 354 magazine entries is the largest number of goal-tracking responses done by any animal across all groups and days, which may suggest that, in the few cases where goal-tracking behavior prevails under conditions of uncertainty, the addition of amphetamine sensitization produces extreme levels of attraction to the goal.

Prior stress effects. Previous stress exposure did not elevate sign-tracking under certainty conditions compared with non-stressed rats, unlike amphetamine sensitization. For example, the no-stress/no-injection control group (No stress + U) did not differ from the stressed group (Stress + U) under uncertainty conditions (group: F1,21 = 0.272, p = .608; see Figure 1F), nor did the saline-injected control groups differ from stressed groups under certainty conditions (Stress + C vs. Saline + C: F1,14 = 0.006, p = .937) or under uncertainty conditions (Stress + U vs. Saline + U: F1,14 = 0.833, p = .377). However, adding uncertainty to prior overstimulation restraint stress did appear to marginally raise sign-tracking in Stress + U rats compared with Stress + C rats, indicating that uncertainty can elevate sign-tracking in animals previously exposed to chronic stress (group: F1,14 = 4.274, p = .058).

Figure 1G shows that, here also, goal-tracking responses for the Stress + U and No-stress + U groups were very similar (F1,21 ≤ 0.669, ps ≥ .422), except on Day 1 (F1,21 = 12.691, p = .002). Although there was a trend, no overall difference was shown between the two uncertainty conditions (Stress + U and No stress + U) and Stress + C rats (F2,21 = 3.226, p = .060). The Stress + U rats alone did however show lower levels of magazine entries when compared with Stress + C (F1,14 = 13.031, p = .003), particularly during the first 4 days of training (F1,21’s ≤ 5.935, ps ≤ .024).

Incidence of goal-tracker versus sign-tracker phenotypes. Again, uncertainty groups had a higher proportion of sign-trackers than certainty groups, regardless of whether they were previously stressed: Stress + U (ST = 87.5%; GT = 0%), No stress + U (ST = 87.5%; GT = 12.5%), and Stress + C (ST = 50%; GT = 25%). As with all groups across all conditions, there was a general trend for more sign-trackers than goal-trackers.

The main result emerging from this analysis is that current CS–UCS uncertainty for both nonsensitized rats and prior amphetamine sensitization produces similar increases in sign-tracking performance, but reduces goal-tracking performance only in saline-treated rats. Current uncertainty and prior amphetamine-sensitization groups become similar in sign-tracking under these conditions, and show higher sign-tracking than their certainty control or unsensitized control conditions. However, uncertainty and drug sensitization did not combine additively to produce a superelevation in a combined Amph + U group above those separately elevated levels (compared with Amph + C or Saline + U groups). Finally, repeated sensory and restraint-stress exposure alone, by comparison, did not produce as robust an elevation in sign-tracking, which allowed the effects of uncertainty to still be detected, at least on some training days.

Comparing sign-tracker versus goal-tracker phenotypes across all training, sensitization, and stress conditions, uncertainty conditions generally produced a higher proportion of sign-trackers (87.50%; F1,5 = 22.857, p = .005) and fewer goal-trackers (4.17%; F1,5 = 24.143, p = .004) than certainty (sign-trackers = 54.17%; goal-trackers = 33.33%). Very few individuals showed a mixed phenotype, and the low proportion was equivalent in both uncertainty and certainty groups (F1,5 = 0.714, p = .436). The propensity of uncertainty to generate a larger proportion of sign-trackers (and hence a lesser proportion of goal-trackers) had previously been suspected (Anselme et al., 2013; M. J. F. Robinson, Anselme et al., 2014) but not confirmed until now (Figure 2A).

Phase 3: Confirmation of Psychomotor Sensitization

Following the final day of Pavlovian autoshaping testing, all rats received an amphetamine challenge (0.75 mg/kg, IP) and were placed in locomotion chambers for 45 min to confirm
whether psychomotor sensitization had been previously induced by either earlier amphetamine treatment or earlier stress exposure (Figure 2B). One-way ANOVA revealed an overall sensitization effect of increased locomotor activity for both the amphetamine-treatment group (regardless of certainty/uncertainty) and the stress-treatment group (at least in uncertainty condition) compared with saline and no stress groups (group: $F_{6,49} = 5.746, p = .000$). That is, the Stress + U group displayed higher or sensitized locomotion in comparison with the Saline + U group ($F_{1,49} = 6.059, p = .017$), and Stress + U was similarly higher than No stress + U ($F_{1,49} = 4.164, p = .047$), although Stress + C and Stress + U groups did differ ($F_{1,49} = 11.611, p = .001$). Similarly, prior amphetamine sensitization elevated locomotion in rats trained under certainty (Amph + C vs. Saline + C: $F_{1,49} = 17.254, p = .000$), but not under uncertainty (Amph + U vs. Saline + U: $F_{1,49} = 2.614, p = .112$). However, a comparison of the two amphetamine groups (C and U) with the two saline groups (C and U) indicated amphetamine-induced sensitization ($F_{1,30} = 11.026, p = .002$). The saline groups should be relevant controls for the stress groups, as control groups did not differ from each other: neither between Saline + U and No stress + U ($F_{1,49} = 0.177, p = .676$) nor between Saline + U and Saline + C ($F_{1,49} = 0.144, p = .706$).

**Discussion**

Here we have confirmed that uncertainty about the prediction triggered by a CS+ for sucrose reward as UCS enhances sign-tracking behavior (approaches and grasping, nibbling, and biting of metal lever), leading to greater attraction toward that CS+ lever in Pavlovian autoshaping, compared with certainty conditions. The sign-tracking enhancement by uncertainty occurred at the reciprocal expense of goal-tracking. Uncertainty enhancement of sign-tracking was observed for unsensitized (no previous amphetamine), and repeated restraint-stress rats. Consequently, there was a greater proportion of sign-tracker phenotypes, and lower proportion of goal-tracker phenotypes in these groups, as well as a higher number of overall sign-tracking responses in individuals when trained and tested under uncertainty conditions, compared with under CS–UCS 100% certainty conditions.

By comparison, we have reported that prior sensitization to amphetamine induced an equivalent elevation of sign-tracking here, even under current certainty conditions, but did not recipro-
cally decrease in goal-tracking under any training condition. We also report that, once sign-tracking was elevated by either current uncertainty conditions or previous amphetamine sensitization, no further elevation was produced by combining both manipulations together, perhaps indicating a ceiling effect for sign-tracking enhancement. Our results suggest that prior amphetamine sensitization and current CS–UCS uncertainty are equivalent potentiators of incentive salience attributed to a discrete CS+ cue that predicts reward.

Amphetamine Sensitization

Our finding of increased sign-tracking in female rats after amphetamine sensitization is similar to the report of Doremus-Fitzwater and Spear (2011). However, we note that opposite sensitization enhancement of goal-tracking has been reported by others (Simon, Mendez, & Setlow, 2009). Although the reason for different results in our study and Simon and colleagues’ study remains unknown, we note that their study was done in male rather than female rats and involved five rather than seven daily injections of amphetamine. Also, their sensitization was conducted in the home cage whereas ours was conducted in a novel context, and home administration of amphetamine has been reported to diminish or modify sensitization induction compared with administration in a novel context (Crombag et al., 2001).

Here we found, rather unusually, that when amphetamine sensitization enhanced sign-tracking, goal-tracking still persisted at moderate levels and did not decline in amphetamine-sensitized rats, not even when uncertainty during training was added to prior amphetamine sensitization. The persistence of sensitized goal-tracking responses at control levels seems at least a step toward Simon et al.’s (2009) findings, in that goal-tracking defected from its usual reciprocal relationship to sign-tracking, suggesting that amphetamine-sensitization may have contributed to goal-tracking maintenance as a competitive alternative response. It may also reveal a potential incentive-salience component of goal-tracking, as a motivated attraction to the sucrose dish, particularly in the few animals displaying a goal-tracking phenotype. We acknowledge that sign-tracking is generally taken to more purely reflect Pavlovian conditioned approach. However, enhanced goal-tracking can occur as a motivated response in goal-tracking individuals, together with faster approach and more intense consumatory sniffs, licks, and nibbles of the metal sucrose dish, after the same brain-limbic stimulations that enhance motivated sign-tracking in sign-tracking individuals, such as μ-opioid stimulation of the central amygdala by DAMGO microinjections (DiFeliceantonio & Berridge, 2012; Mahler & Berridge, 2009). Enhanced goal-tracking as a motivated response was interpreted as indicating that a third psychological component, namely, incentive salience targeted toward the dish, can potentially contribute to goal-tracking in some individuals, especially in states of mesocorticolimbic activation. Similarly, acute amphetamine administration has been reported to increase goal-tracking while drug is on board (Holden & Peoples, 2010). Finally, either amphetamine or opioid stimulation of the nucleus accumbens can increase the relative incentive salience of a reward-proximal CS (analogous to the sucrose dish here) at the expense of a more predictive but reward-distal CS (analogous to the CS+ lever here) (Smith, Berridge, & Aldridge, 2011; Tindell, Berridge, Zhang, Peciña, & Aldridge, 2005). These considerations highlight that mesolimbic activation can produce complicated patterns of effects on incentive salience of lever versus dish as potential CSs, with consequences for sign-tracking versus goal-tracking. That complexity needs further research and explanation to better understand the conditions under which sign-tracking versus goal-tracking becomes most enhanced, but in general our results do fit a hypothesis that sensitization can enhance ‘wanting’ for a reward-predictive cue.

Uncertainty

The excitatory effect of reward uncertainty in CS+ autoshaping has long been noted (Anselme et al., 2013; Boakes, 1977; Collins, Young, Davies, & Pearce, 1983; Gibson, Farrell, Locurto, Duncan, & Terrace, 1980; Gottlieb, 2004; Kaye & Pearce, 1984; M. J. F. Robinson, Anselme et al., 2014) although its explanation also remains controversial. Some authors have suggested that increased behavioral vigor in uncertainty reflects the animal’s aversive frustration caused by intermittent nonreward (Amsel, 1958; Papini, 2003). However, we hypothesized that it was caused by increased appetitive directed to the CS+ (Anselme, 2015; Anselme et al., 2013) because the increased approach occurs under Pavlovian autoshaping conditions even when no lever-pressing response has ever been instrumentally reinforced, and because evidence suggests that uncertainty in rewards may stimulate the midbrain-mesolimbic dopamine system both in animals and humans (Anselme & Robinson, 2013; Boileau et al., 2013; L. Clark, Lawrence, Astley-Jones, & Gray, 2009; Dreher, Kohn, & Berman, 2006; Fiorillo, Tobler, & Schultz, 2003; Jousta et al., 2012; Limet et al., 2012; Singer et al., 2012; Zack et al., 2014). An appetitive interpretation may be consistent with our observation here of overlap in effect between uncertainty and prior amphetamine sensitization. Previous research has shown that amphetamine sensitization promotes mesolimbic reactivity in the form of increased dopaminergic release in areas such as the nucleus accumbens, ventral pallidum, and amygdala, and greater motivation for drug rewards (Vezina, 2004). Behaviorally, sensitization enhances incentive salience by increasing excitatory associations (Harmer & Phillips, 1999a, 1999b) and ‘wanting’ (Tindell et al., 2005) for reward associated cues, all of which can be expected to affect levels of sign-tracking which has been suggested to heavily rely on dopaminergic activity (Flagel et al., 2011; Saunders & Robinson, 2012). Our conclusion is simply that uncertainty and sensitization produced similar effects, namely with the procedures used here, to enhance sign-tracking toward the reward-predictive CS+ lever.

Ceiling Effects for Sensitization + Uncertainty Combination?

It might have been expected that prior amphetamine sensitization combined with current uncertainty together (Amph + U) would produce a greater sign-tracking attraction to the reward CS+ lever than either manipulation alone (Saline + U or Amph + C), but it did not: The combination produced an elevation over
control levels that was merely equivalent to elevations produced either by sensitization alone or by uncertainty alone. That the whole was less than the sum of its parts suggests that some sort of measurement ceiling existed which precluded further enhancement. A ceiling effect at either the behavioral level or in mesocorticolimbic circuitry may have occurred because rats pressed up to six times on average per CS+ presentation (almost one press per second) on Day 8. It is possible that a ceiling effect occurred centrally in mesolimbic circuitry, such as when dopamine released upon amphetamine administration occupies 50% of receptors and cannot occupy receptors any further (Korteekaas et al., 2004). Alternatively, perhaps the combination did indeed boost incentive salience further in a way that might be detected by other measurement procedures, but simply not detected here.

**Sensory Stress Sensitizes and Interacts With Uncertainty for Psychomotor Locomotion but not Motivation CS+ Attraction?**

Prior amphetamine pretreatment produced robust sensitization of locomotion (psychomotor sensitization) as well as of sign-tracking (incentive sensitization). Stress pretreatment did produce psychomotor sensitization, reflected in higher drug-induced locomotion here at least for rats treated under uncertainty conditions, but did not produce significant incentive sensitization. Even the psychomotor sensitization by stress that appeared only was significant in some stressed rats (i.e., stressed and then trained under uncertainty) but not others (i.e., stressed and then trained under certainty), suggesting a potential interaction between the conditions. Certainty conditions following the stress regimen could have had normalizing effects on behavior and contributed to preventing the expression of locomotor cross-sensitization. The psychomotor sensitization by stress did not transfer into incentive sensitization of CS+ motivation, suggesting either that the degree of stress sensitization was relatively weak here or, that there are differences in neural circuitry underlying psychomotor sensitization versus incentive sensitization, or both (T. E. Robinson & Berridge, 2008).

Also, we note that the relationships between stress and the reward circuit are complex and may lead to apparent contradictions, such as decreased reactivity of striatal neurons (Cabib & Puglisi-Allegra, 2012). Finally, we did not find a potentiation of locomotion by mere uncertainty exposure here, though we did find a potentiation of stress psychomotor sensitization by uncertainty, even though others have reported that uncertainty training by itself can induce psychomotor sensitization to amphetamine (Singer et al., 2012). However, our procedures were quite different from Singer et al. (2012), who used a variable-ratio schedule operant task to induce uncertainty (not Pavlovian autoshaping) and a much longer training period (55 days), making it difficult to directly compare across studies.

**Conclusion**

Our results suggest commonalities in the pathways involved in the enhancement of cue attraction by reward uncertainty and amphetamine sensitization. This further ties the behavioral effects of reward uncertainty to changes in brain dopaminergic pathways. It suggests that unpredictable reward environments may result in an increased motivation toward reward-related cues similar to psychostimulant sensitization, which may be of particular importance to the development of problem gambling.

**References**


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