Motivational factors underlying aversive Pavlovian-instrumental transfer

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While interest in active avoidance has recently been resurgent, many concerns relating to the nature of this form of learning remain unresolved. By separating stimulus and response acquisition, aversive Pavlovian-instrumental transfer can be used to measure the effect of avoidance learning on threat processing with more control than typical avoidance procedures. However, the motivational substrates that contribute to the aversive transfer effect have not been thoroughly examined. In three studies using rodents, the impact of a variety of aversive signals on shock-avoidance responding (i.e., two-way shuttling) was evaluated. Fox urine, as well as a tone paired with the delivery of the predator odor were insufficient modulatory stimuli for the avoidance response. Similarly, a signal for the absence of food did not generate appropriate aversive motivation to enhance shutting. Only conditioned Pavlovian stimuli that had been paired with unconditioned threats were capable of augmenting shock-avoidance responding. This was true whether the signaled outcome was the same (e.g., shock) or different (e.g., klaxon) from the avoidance outcome (i.e., shock). These findings help to characterize the aversive transfer effect and provide a more thorough analysis of its generalization to warning signals for different kinds of threats. This feature of aversive motivation has not been demonstrated using conventional avoidance procedures and could be potentially useful for applying avoidance in treatment settings.

Aversive Pavlovian conditioning, also commonly called defensive (or fear) conditioning has been utilized as a model of human emotional dysregulation for decades. By repeatedly pairing a neutral stimulus (e.g., tone) with an aversive outcome (e.g., shock), the tone eventually produces anticipation of shock and elicits conditioned responding (e.g., freezing, changes in heart rate, endogenous analgesia). Many studies using this behavioral procedure across a variety of species have allowed for the identification of important structures and pathways in the central nervous system crucial for the acquisition and expression of this form of learning (Maren and Fanselow 1996; Herry and Johansen 2014; Krabbe et al. 2018). As this understanding grew, identifying ways to reduce conditioned responses using stimulus exposure became a focus of the field because of its high clinical relevance to treating humans suffering from disorders of fear and anxiety (Bouton 2004; Rauch et al. 2006; Milad and Quirk 2012; Kim and Jung 2018; Meulders 2020). While this approach does reduce defensive behaviors over the course of treatment, recovery effects based on context, time, and internal state make extinction less appealing as a long-term treatment option for humans (Bouton et al. 2006; Delgado et al. 2006; Hofmann 2007; Beckers et al. 2013; Markowitz and Fanselow 2020). In response, interest in active avoidance and aversive instrumental motivation began to reemerge as an alternative approach to reducing defensive responding (Bravo-Rivera et al. 2014; Gillan et al. 2014; Krypotos et al. 2015; LeDoux et al. 2016).

In contrast to Pavlovian conditioning, where the subject passively experiences tone-shock pairings, active avoidance uses a response (e.g., lever press or shuttling) that can terminate trials and prevent harm. As the subject transitions from passive to active response modes, defensive behaviors (i.e., freezing) are gradually reduced. While this may seem an encouraging analog to extinction for understanding how defensive reactivity can be attenuated over time, active avoidance studies are often difficult to interpret and have a number of conceptual and control problems that can complicate the issue further (Sidman 1953; Rescorla and Solomon 1967; Bolles 1972; Fanselow and Lester 1988; Fanselow 1997; Kim et al. 2006; LeDoux et al. 2016; Cain 2019). Use of an aversive Pavlovian-instrumental transfer procedure offers analysis of aversive motivation without many of the problems normally associated with active avoidance studies (Campese et al. 2013, 2014). However, while a handful of studies have shown the capacity for an aversive conditioned stimulus to modulate avoidance behavior, a systematic analysis of this phenomenon is, as of yet, lacking in the literature. The purpose of the studies below is to provide a better understanding of the motivational factors that contribute to the ability of an aversive conditioned cue to control active avoidance responses. This was done by examining whether motivational control of avoidance extends to (1) nonspecific aversive conditioned stimuli, (2) predator odor, (3) cues for predator odor, and (4) associatively and motivationally opposite, yet functionally equivalent cues (i.e., an appetitive inhibitor). The studies reported below suggest that aversive motivational control over avoidance is...
limited to previously neutral cues that have acquired aversive properties via pairings with an aversive outcome.

Results

Shock-avoidance motivation generalizes to signals for other threats

Studies of Pavlovian-instrumental transfer (PIT) have predominantly been done using appetitive motivation (Estes 1943, 1948; Rescorla 1994; Holland 2004; Corbit et al. 2007; Corbit and Balleine 2016). The findings from these studies have contributed to an immensely rich understanding of the psychological and neural structures underlying various forms of appetitive or reward-based motivation. These studies benefit from the wide variety of appetitive reinforcers available and the readiness with which many distinct behaviors can be shaped when rewarded with different kinds of foods (Kruse et al. 1983; Dickinson et al. 1996; Ranaldi et al. 2009). Using these response-outcome options, appetitive PIT studies have demonstrated that different forms of motivation (i.e., sensory specific as opposed to general motivation) depend on different pathways in the brain (Hall et al. 2001; Holland and Gallagher 2003; Corbit et al. 2007; Shiflett and Balleine 2010). These findings have provided valuable insight toward our understanding of appetitive motivation and instances of its dysregulation (Dickinson et al. 2000; Wyvell and Berdidge 2001; Holland 2004; Wise and Koob 2014).

On the other hand, aversive motivation has not been approached from this perspective. Because it is more limited in the range of response options, and fewer perceptually distinct outcomes are studied, it can be difficult to isolate sensory specific encoding. However, some studies have shown that other kinds of aversive outcomes can potentially be used in a similar manner to more thoroughly probe the underlying psychology of aversive motivation (Rescorla 1974; Diaz-Mataix et al. 2011; Campese et al. 2019). For example, loud klaxon horns have been used as an unconditioned stimulus in previous studies of appetitive PIT (Campese et al. 2017a). These studies analyzed the effects of Pavlovian extinction on transfer using a within-subjects design where a second Pavlovian cue was paired with shock. Thus, the possibility remains that the capacity of the cue associatively linked to the klaxon to motivate avoidance can be attributed to generalization from the shock cue.

To address this, we trained two separate groups of rats to associate tone with shock, or tone with klaxon. In one subset of these animals, Pavlovian freezing tests were conducted the following day, whereas the remainder underwent Sidman avoidance for shock and were subsequently tested for transfer (see Fig. 1). This study found that freezing to the tone in both the klaxon and shock groups was statistically equal (F(14)=0.06, P=0.95) (see Fig. 1B). Both groups also showed (see Fig. 1C) increased avoidance responding over training blocks (F(3,56)=12.79, P<0.001), the rates of which were comparable (F(Group,1)=4.86, P=0.36). Finally, transfer testing revealed more responding during the stimulus compared with the baseline period to a similar degree for both groups (F(Interval,1)=5.45, P=0.04, F(Group,1)=1.97, P=0.18, F(Interval*Group,1)=1.30, P=0.27) (see Fig. 1D).

Fox urine does not influence rates of two-way shuttle box shock-avoidance

In order to determine whether the ability of conditioned threat signals to augment active shock-avoidance extends to unlearned threat signals, a group of subjects were exposed to fox urine in the avoidance apparatus after successful shock avoidance training (see Fig. 2). Another group of subjects had odor paired with tone and were tested for the tone’s ability to support transfer. Shuttle rates in these groups were compared with control groups exposed to water instead of urine either during Pavlovian training or transfer testing. All groups acquired avoidance comparably over training blocks (F(Block,4,68)=72.85, P<0.001, F(Group,3)=0.13, P=0.94). Transfer testing showed that neither a cue for odor, nor the odor itself had any influence over shuttle rates (F(Interval,1)=4.12, P=0.06, F(Group,3)=1.24, P=0.33, F(Interval*Group,3)=0.53, P=0.67) (see Fig. 2C). A follow up test in a nonavoidance context showed that the fox urine produced significantly more freezing than water when avoidance responding was not available (t(15)=6.44, P<0.001) (see Fig. 2D).

Aversive motivation alone is not sufficient to generate PIT

Some conceptualizations of motivation envision mutually antagonistic appetitive and aversive systems where positive stimuli engage an appetitive center and suppress an aversive counterpart, vice versa for negative valence cues (Konorski 1948). This view has received empirical support from studies of motivational interactions...
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where cues from opposing motivational/associative classes are shown to be functionally equivalent (Dickinson and Pearce 1977; Dickinson and Dearing 1979; Nasser and McNally 2012). According to this view, a cue that predicts the absence of food (i.e., an appetitive inhibitor) and one that predicts the presence of shock (i.e., an aversive excitor) should both engage “aversiveness” and should, therefore, support similar kinds of motivational effects. For example, Dickinson and Dearing (1979) showed that both appetitive inhibitors and aversive excitors can facilitate (or super condition) learning to a novel cue paired with an outcome from the opposite motivational class. Specifically, an appetitive inhibitor facilitated fear learning, while an aversive excitor enhanced appetitive conditioning. To determine whether appetive PIT and active avoidance are sensitive to these kinds of motivational interactions, we evaluated the capacity of an appetitive inhibitor (i.e., a cue predicting the absence of food) to enhance shuttle rates for subjects trained to avoid shock.

Following magazine training, subjects underwent discrimination training where a tone was trained as a conditioned inhibitor for three-session blocks during the Sidman avoidance training phase. (C) Responses per minute during the pre-CS and CS periods for each group tested for aversive transfer. (Furine) fox urine, (CSW) CS water, (CSFx) CS fox urine. (D) Percent time freezing in a nonavoidance context (i.e., a standard Pavlovian chamber) during the baseline (BL) and odor presentation period (OD). Starting with samples of six rats per group, one animal was eliminated from each condition for poor performance with the exception of the water exposed control group, in which all six rats successfully acquired avoidance responding. Asterisks represent statistical significance at the 5% level.

Figure 2. (A) The experimental design used to evaluate predator odor effects of freezing and active avoidance. (B) Avoidance responding for three-session blocks during the Sidman avoidance training phase. Responses per minute during the pre-CS and CS periods for each group tested for aversive transfer. (Furine) fox urine, (CSW) CS water, (CSFx) CS fox urine. (D) Percent time freezing in a nonavoidance context (i.e., a standard Pavlovian chamber) during the baseline (BL) and odor presentation period (OD). Starting with samples of six rats per group, one animal was eliminated from each condition for poor performance with the exception of the water exposed control group, in which all six rats successfully acquired avoidance responding. Asterisks represent statistical significance at the 5% level.

Discussion

The studies reported above replicate and extend the aversive Pavlovian-instrumental transfer phenomenon in a number of ways. A shock-paired cue was shown capable of enhancing active avoidance as was a cue paired with a different aversive event, a klaxon-horn. While previous studies (Campese et al. 2017a) showed that subjects trained to avoid shock demonstrated transfer to a shock paired cue, as well as a klaxon-paired cue, the current findings were observed using a between-subjects design, making it less likely that this result depends on generalization. This is an encouraging step toward using using PIT to isolate different forms of aversive motivation (i.e., sensory specific versus general).

Furthermore, the class of stimulus capable of harnessing this kind of motivation was more specifically tied to acquired Pavlovian motivation. Inmate threat stimuli such as fox urine, which caused significant freezing responses, were not capable of augmenting avoidance. One would expect that the predator odor might suppress avoidance by increasing freezing, however, this was not observed. Previous studies with avoidance have been interpreted as suggesting that reductions in passive behaviors like freezing are replaced with some sense of “agency” or control over emotional reactivity (Moscarello and Hartley 2017). Indeed, studies using aversive PIT have shown that much like extinction, conditioned freezing behaviors are reduced following avoidance. However, unlike extinction, these reductions in defensive responding survive the passage of time and changes to test contexts (Campese et al. 2017b). It is possible that the failure of predator odor to suppress avoidance behavior reflects this change in how aversive threats are processed following avoidance. However, it could also be argued that relative to shock, predator odor is simply a less aversive outcome. Exposure to the more potent stimulus in the avoidance context could have produced a contrast effect rendering it not threatening in that environment (Crespi 1942; Flaherty 1982). Similarly, a cue paired with predator odor was not successful at modulating avoidance responding. Comparable null findings have been reported with traditional freezing measures (Wallace and Rosen 2000; McGregor et al. 2002; Blanchard et al. 2003).

Moreover, it was shown that general aversive motivation is not sufficient to drive aversive PIT. A signal for the absence of

P < 0.001) (see Fig. 3B). Control subjects, comparatively, showed flat responding across training (F(day,15) = 1.51, P = 0.11, F(stimulus,1) = 4.88, P = 0.054, F(day*stimulus,15) = 0.83, P = 0.64) (see Fig. 3C). All subjects were then trained on shock-avoidance in the shuttle boxes where both groups acquired shuffling over sessions comparably (F(block,4) = 36.04, P = 0.01, F(group,1) = 0.08, P = 0.78, F(block*group,4,72) = 0.42, P = 0.79) (see Fig. 3D). After a reminder discrimination session, subjects underwent transfer tests in the shuttle boxes. These tests found that the ability of an appetitive inhibitor to generate aversive transfer was no stronger than that of the untrained tone in the control group (F(interval,1) = 1.9, P = 0.19, F(group,1) = 0.91, P = 0.35, F(interval*group,1) = 0.33, P = 0.58) (see Fig. 3E).

To confirm that the tone had acquired true inhibition in discrimination subjects, these subjects were randomly split into two groups that underwent summation and retardation testing independently (Rescorla 1969). Summation testing found that the tone effectively reduced magazine approach to a novel appetitive excitor (t(9) = 2.66, P = 0.03) (see Fig. 3F). Additionally, when trained as an appetitive excitor relative to another, untrained cue (i.e., flashing light), the untrained cues acquired excitation significantly more quickly while the tone’s acquisition was retarded by the accrued inhibition (t(9) = 4.43, P = 0.01) (see Fig. 3G).
食品在减少逃避反应方面并不具备抑制的特性。尽管被证明具有真正的抑制性质，但应该注意，这并不是一种可以轻易识别的抑制性质。在出现逃避反应时，声音被证明能够有效地减少 appetitive 激励。正因为如此，这种方法被用作开发在计算机游戏中模拟厌恶刺激的一种非常有用的对比。在计算机游戏中模拟厌恶刺激的研究结果表明，人们显示厌恶转移的现象。然而，这些研究还存在一些限制，包括对厌恶处理的范围和对激励区别的限制。在厌恶处理的背景下，这种现象和应用到人类和啮齿动物研究中。

## Materials and Methods

### Subjects

80只斯普拉格-道利特鼠被用于下面的实验。这些实验主要在条件室中进行，条件室由Coulbourn Instruments制造。这些实验使用了与人类相似的程序，但使用的限制和这些人类程序有很大不同（例如，厌恶处理和威胁处理）。在厌恶处理中，这种程序和行为的研究有共同点（例如，厌恶处理和威胁处理）。使用这种准备可以产生令人鼓舞的结果。使用这种程序，成像和其他限制可以使用这种准备。

### Apparatus

研究主要在条件室中进行，条件室由Coulbourn Instruments制造。厌恶性条件性行为的训练和测试session是在标准条件下进行的（模型编号H10-11R-TC）而逃避行为的训练和测试session是在两道穿梭室中进行的（模型编号H110-11R-SC）。听觉刺激（例如，5 kHz的音）和0.5 mA的正脉波脚刺激用于
these sessions were presented using a programmable audio generator (model no. A12–33) and a precision animal shock (model no. H13–15), respectively, both manufactured by Coulbourn. Klaxon horns (114 dB) were manufactured by Wolof (model no. 330).

One milliliter of fox urine (Minnesota Trapline Products) was delivered using syringes mounted in Razell Pumps (Model R-E) with the attached lines leading to a cup in the waste tray below each of the Coulbourn chambers. Urine was loaded into the syringes, followed by 1-mL volume air bubble and an additional 1 mL of water at the end of the line in order to mask odor. Urine and the water control were delivered at a rate of 4 mL/min.

Appetitive conditioning sessions were done in Med Associates chambers (model no. ENV-008) that were equipped with 5-V light bulbs and an audio generator card (model no. ANL 926) for stimulus delivery. Standard 45-mg pellets (Test Diet AIN-76A) were delivered to the food cup via a pellet dispenser at specific times and approach behavior was recorded using infrared sensors at the threshold of the food magazine. Follow up tests for conditioned inhibition were done in these chambers and utilized a flashing 30-sec panel light and a white noise (80 dB) auditory cue generated using this apparatus.

Procedure

Pavlovian conditioning

Three 30-sec presentations of the tone stimulus coterminated with shock (1 sec), or klaxon (5 sec) in a 15-min training session with a 5-min baseline and fixed 3-min intertrial interval (ITI). Test sessions had the same parameters, but without footshock or klaxon presentations. For sessions involving predator odor, the urine was infused into the chamber after a 10-min baseline period or beginning with the presentation of the CS in the cued preparation.

Appetitive discrimination training

Subjects were first given baseline avoidance training to eliminate poor performers. Good performers were food restricted and then trained to retrieve food pellets from the food cup over 2 d. In each of these two 20-min sessions, two pellets were delivered to the magazine on a VI60 schedule. Over the next 16 sessions subjects were trained on an appetitive discrimination task where a 30-sec house light was paired with food using a delay conditioning arrangement. A compound stimulus consisting of the house light with tone was presented without food to generate inhibition to the tone. Each 105 min session included 16 light, and 8 light + tone trials, following a 5-min baseline with a variable 3-min ITI ranging from 1 to 5 min. Summation tests were done by training a flashing light to signal food and then combining this cue with tone to determine the degree of control over magazine responding by the tone. Retardation tests were done by comparing the acquisition of magazine responses to the tone when it was paired with food relative to a novel cue (white noise).

Unsigned Sidman active avoidance

Avoidance training was done in the two-way shuttle boxes and consisted of fifteen 25-min sessions. Subjects were placed into the chamber with the house light on and the following contingencies in effect. A response-shock (R-S) interval of 30-sec was applied over a shock-shock (S-S) interval of 5 sec. If subjects failed to shuttle by the end of these intervals, a 1-sec shock was delivered. Shuttle responding produced a period of safety (i.e., the R-S interval) lasting 30-sec. Additionally, the house light was turned off for 0.3 sec as a feedback cue for each shuttle response. Shuttle responses that occurred in the absence of shock were scored as avoidance responses (ARs) and those that were initiated while a shock was present were counted as escape responses (ERs). While performance of ARs terminated ongoing footshock, both ARs and ERs initiated the R-S interval. At the end of the session the house lights were turned off and responses produced no further feedback. Subjects that failed to register two consecutive sessions with at least 20 ARs prior to day 10 of training were classified as poor performers and were removed from the study (see Lazarro-Munoz et al. 2010). Sessions were run 5–6 d per week.

Pavlovian–instrumental transfer (PIT) testing

PIT testing was conducted over two consecutive days in the shuttle box chambers and included no shocks at all. Following placement in the chamber, subjects shuttled in extinction for 15 min, receiving the blinking house light feedback cue for each shuttle. After 15 min, responses were monitored by the computer running the study to identify the point at which the subject’s response rate was at two responses per minute. At this point the tone was presented and it remained on until the subject performed 10 shuttle responses, after which the house light turned off and the session ended. Subjects were removed from the boxes and tested again the next day.

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References

Beckers T, Krypotos AM, Boddez Y, Effting M, Kindt M. 2013. What’s wrong with fear conditioning? Biol Psychol 92: 90–96. doi:10.1016/j.biopsycho.2011.12.015
Blanchard DC, Markham C, Yang M, Hubbard D, Madaran E, Blanchard RJ. 2003. Failure to produce conditioning with low-dose trimethylnitrosoamine or cat feces as unconditioned stimuli. Behav Neurosci 117: 360–368. doi:10.1037/0735-7044.117.2.360
Bolles RC. 1972. The avoidance learning problem. In The psychology of learning and motivation (ed. Bowar GH, Spence JT), pp. 97–145. Academic Press, Oxford.
Bouton ME. 2004. Context and behavioral processes in extinction. Learn Mem 11: 485–494. doi:10.1101/lm.78804
Bouton ME, Westbrook BE, Cocoran KA, Maren S. 2006. Contextual and temporal modulation of extinction: behavioral and biological mechanisms. Biol Psychiatry 60: 352–360. doi:10.1016/j.biopsych.2005.12.015
Bravo-Sivera C, Roman-Oritz C, Brignoni-Perez E, Sotes-Bayon F, Quirik GJ. 2014. Neural structures mediating expression and extinction of platform-mediated avoidance. J Neurosci 34: 9736–9742. doi:10.1523/JNEUROSCI.0191-14.2014
Cain CK. 2019. Avoidance problems reconsidered. Curr Opin Behav Sci 26: 9–17. doi:10.1016/j.cobeha.2018.09.002
Campese V,McCue M, Lázaro-Muñoz G, LeDoux JE, Cain CK. 2013. Development of an aversive Pavlovian-to-instrumental transfer task in rats. Front Behav Neurosci 7: 176. doi:10.3389/fnbeh.2013.00176
Campese VD, Kim J, Lázaro-Muñoz G, Pena L, LeDoux JE, Cain CK. 2014. Lesions of lateral or central amygdala abolish aversive Pavlovian-to-instrumental transfer in rats. Front Behav Neurosci 8: 161. doi:10.3389/fnbeh.2014.00161
Campese VD, Kim JT, Rojas G, LeDoux JE. 2017a. Pavlovian extinction and recovery effects in aversive Pavlovian to instrumental transfer. Front Behav Neurosci 11: 179. doi.org/10.3389/fnbeh.2017.00179
Campese VD, Soroeta JM, Vazey EM, Aston-Jones G, LeDoux JE, Sears RM. 2017b. Noradrenergic regulation of central amygdala in aversive Pavlovian-to-instrumental transfer. eNeuro 4: ENURO.0224-17.2017. doi:10.1523/E N URO.0224-17.2017
Campese VD, Kim JT, Hou M, Gupta S, Draus C, Kurpas B, Burke K, LeDoux JE. 2019. Chemogenetic inhibition reveals that processing relative but not absolute threat requires basal amygdala. J Neurosci 39: 8510–8516. doi:10.1523/JNEUROSCI.2530-18.2019
Corbit LH, Balleine BW. 2016. Learning and motivational processes contributing to Pavlovian- Instrumental Transfer and their neural bases: dopamine and beyond. Curr Top Behav Neurosci 27: 259–289. doi: 10.1007/97854_2015_388
Corbit LH, Janak PH, Balleine BW. 2007. General and outcome-specific forms of Pavlovian-instrumental transfer: the effect of shifts in motivational state and inactivation of the ventral tegmental area. Eur J Neurosci 26: 3141–3149. doi:10.1111/j.1460-9568.2007.05934.x
Crespi LP. 1942. Quantitative variation of incentive and performance in the white rat. Am J Psychol 55: 467–517. doi:10.2307/1417127

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