Persistence of a hyperthermic sign-reversal during nitrous oxide inhalation despite cue-exposure treatment with and without a drug-onset cue

Karl J Kaiyala1,*, Stephen C Woods2, and Douglas S Ramsay1,3,4

1Department of Oral Health Sciences; University of Washington; Seattle, WA USA; 2Department of Psychiatry and Behavioral Neuroscience; University of Cincinnati; Cincinnati, OH USA; 3Department of Orthodontics; University of Washington; Seattle, WA, USA; 4Pediatric Dentistry; University of Washington; Seattle, WA USA

Abbreviations: CEXP, cue exposure; CR, conditioned response; CS, conditioned stimulus; DHL, dry heat loss; DOC, drug onset cue; EHL, evaporative heat loss; HC, home cage; HL, heat loss; HP, heat production; N2O, nitrous oxide; Tc, core temperature

Keywords: allostatic, drug addiction, extinction, homeostasis

We asked whether chronic tolerance and the hyperthermic sign-reversal induced by repeated 60% N2O exposures could be extinguished using a cue-exposure paradigm. Rats received 18 N2O administrations in a total calorimetry system that simultaneously measures core temperature (Tc), metabolic heat production (HP), and body heat loss (HL). Each exposure entailed a 2-h baseline period followed by a 1.5-h N2O exposure. The 18 drug exposures induced a robust intra-administration hyperthermia in which the initial hypothermic effect of N2O inverted to a significant hyperthermic sign-reversal during N2O inhalation due primarily to an acquired robust increase in HP. The rats were then randomized to one of 3 extinction procedures (n = 8/procedure) over a 20-d interval: 1) a N2O-abstinent home-cage group (HC) that received only the usual animal care; 2) a cue-exposure group (CEXP) in which the animals were placed in the calorimeter 8 times but received no N2O; and 3) a drug-onset-cue group (DOC) in which animals received a brief N2O exposure in the calorimeter that mimicked the first 3 min of an actual 60% N2O trial. Following the extinction sessions, all rats received a 60% N2O test trial and Tc, HP and HL were assessed. The hyperthermic sign-reversal remained fully intact during the test trial, with no significant differences observed among groups in any post-baseline change in any thermal outcome. These data suggest that cue exposure may not be an efficacious strategy to reduce sign-reversals that develop with chronic drug use.

Introduction

Drug use fosters a web of biopsychosocial phenomena that can promote a transition to drug addiction in some individuals and under some conditions.1 Among the processes thought to contribute to this progression is associative learning, whereby drug-associated cues acquire the power to elicit motivationally salient symptoms such as drug craving and withdrawal-like effects that encourage further drug taking.2,3 Cues associated with repeated drug administration have been found to acquire the ability to elicit responses that counter the effects of the drugs, and when these responses are elicited in the presence of the drug, they consequently lessen the drug’s impact and thereby contribute to drug tolerance.2,4,5 These cues can include obvious stimuli such as the specific drug environment or drug-taking paraphernalia, or else arbitrary stimuli such as novel odors. Repeated presentation of drug-relevant conditioned stimuli (CS), in the absence of the drug itself, can extinguish the ability of the CS to elicit compensatory conditioned responses (CR), thereby attenuating the ability of the relevant cues to elicit responses that contribute to tolerance and withdrawal effects.2 For example, if rats tolerant to the hypothermic effects of ethanol are given repeated presentations of ethanol-paired stimuli, but now in the absence of ethanol, the learned compensatory responses mediating tolerance are extinguished and the hypothermic effect of ethanol is reinstated.6,9 This extinction strategy, often called cue-exposure therapy, remains a focus of interest among researchers and clinicians seeking effective treatments for drug addiction.10,11

The evidence for the efficacy of cue-exposure therapy has been inconsistent, and one possibility for this is that principles of learning may not always have been appropriately applied.1,2 A vital consideration in this regard involves the identity of the critical drug-associated CSs that elicit CRs. Most approaches to extinction-based therapy have focused on traditional exteroceptive cues (e.g., environmental cues such as drug paraphernalia), whereas evidence suggests that interoceptive cues exerted by the drug itself during a drug administration can play dominant roles.
as CSs, especially in situations where the onset of the drug effect is sufficiently gradual so that synchronous build-up of interoceptive cues provides a reliable and salient predictor of the upcoming larger drug effect. Failures of cue-exposure therapy may consequently reflect failures to target and extinguish the most salient interoceptive cues in the form of the effect of the drug itself, which reliably signals an impending greater drug effect. In the context of drug taking, these kinds of interoceptive CSs are called drug-onset cues (DOCs).

Our laboratory has developed a rat model for investigating drug tolerance, using nitrous oxide (N₂O) and assessing thermal parameters. This model permits us to deliver N₂O, a pharmacologically active gas that promotes significant hypothermia upon initial administration, while synchronously measuring core temperature (Tc), heat production (HP) and heat loss (HL). Using this approach, we have demonstrated that serial administration of N₂O to rats causes tolerance to develop in the form of a lessened hypothermia over trials resulting primarily from the growth of a HP response during repeated N₂O administrations. Further, if N₂O administrations are continued after full tolerance has developed, rats eventually exhibit a significant hyperthermic sign-reversal during repeated N₂O administrations. This hyperthermic sign-reversal during N₂O inhalation, and to a lesser extent by adaptations that slightly impede HL, has been suggested to have motivational consequences because rats will oppose their own dysregulated hypothermia by moving to a cooler ambient temperature if one is available. This finding is compatible with an allostatic interpretation of addiction, which proposes that withdrawal-like sign-reversal states may motivate behaviors that oppose the sign-reversal state. For example, making a behavioral response that increases the amount of drug in the body (i.e., drug taking) could cause a greater pharmacological effect that opposes, and thereby reduces, the aversive sign-reversal.

The primary goal of the present study was to determine whether the responses mediating the hyperthermic sign-reversal that eventually develops with repeated N₂O administrations might be acquired through an associative mechanism and therefore be eliminated using extinction procedures. If extinction could inhibit the elicitation of HP responses during N₂O administration, the hyperthermic sign-reversal of Tc should diminish, and the degree of tolerance development should be reduced as well. The present study was designed to induce thermal tolerance, including a hyperthermic sign-reversal of Tc during repeated N₂O administrations in rats. After the sign-reversal was well-established, rats were randomly assigned to one of 3 groups: 1) a N₂O-abstinent home-cage control group, 2) a cue-exposure group that received extinction trials to the environmental cues associated with the N₂O delivery chamber, or 3) a drug-onset cue group that received extinction trials that provided both the environmental cues plus a brief interoceptive drug-onset cue of N₂O. Following the extinction trials, a final N₂O administration determined whether the responses mediating tolerance or the hyperthermic sign-reversal of Tc were diminished by the extinction intervention.

Materials and Methods

Subjects
Male Long-Evans rats (Charles River, N = 24, 25–28 d of age upon arrival) were maintained on a 12-h:12-h light/dark cycle (lights on at 0700 h) at an ambient temperature of 22 ± 1°C. Rats were group-housed in polycarbonate tubs with free access to tap water and pelleted chow (5053 Picolab Rodent Diet 20, Animal Specialties and Provisions, Quakertown, PA). All animal procedures were approved by the University of Washington Institutional Animal Care and Use Committee.

Experimental design and procedures
Figure 1 presents an overview of the experimental design. Rats were placed individually in a calorimeter for 18 exposure sessions to 60% N₂O so as to induce thermal tolerance followed by a hyperthermic sign-reversal as observed in previous research. Exposures were conducted on Monday, Wednesday and Friday for 6 consecutive weeks. Each exposure entailed a 2-h baseline pre-exposure period followed by a 1.5-h 60% N₂O administration. Exposures began at either 0900 h (morning) or 1300 h (afternoon) and each rat was consistently tested at the same time of day. The first exposure session began >7 d after surgery and recovery [body mass was 157.2 ± 25.0 g (mean ± SD)]. After the 18 trials, the 24 rats were randomized (counterbalancing for morning and afternoon exposure sessions) into 3 extinction groups: 1) a home-cage group (HC), 2) a cue-exposure group (CEXP), and 3) a drug-onset cue group (DOC). The 3 extinction procedures were conducted over a 20-d interval. During this time, the CEXP and DOC groups were provided with 8 calorimetry sessions that occurred on the same days of the week and with the same starting times and durations as in the induction period. Except for a brief 3-min delivery of N₂O in the DOC group (described below), the control gas was delivered for the entire 3.5-h extinction session. HC rats were maintained in their home cages during the entirety of the extinction period.

Total calorimetry and N₂O administration systems
Total calorimetry simultaneously measures the rates of total HL and metabolic HP, the 2 underlying determinants of Tc that reflect the influence of control mechanisms involved in regulating Tc. This system is described in detail elsewhere (see Part I of the online supplement). In brief, 6 total-calorimetry systems equipped for telemetric Tc assessment served as gas-tight exposure chambers for N₂O. All gas mixtures were delivered to each chamber at a constant flow rate of 1.5 L/min. In the 60% N₂O gas condition, the drug was administered such that the target concentration of 60% N₂O was achieved rapidly. This goal was accomplished by a 2-stage administration in which 72% N₂O, 21% O₂, and 7% N₂ was delivered for the first 6.5 minutes. Subsequently, the gas blend was switched to 60% N₂O, 21% O₂, and 19% N₂ for the remainder of the delivery of the 60% N₂O gas condition. The drug concentration in the chamber did not exceed 60% N₂O using this procedure. The control gas condition consisted of a blend of 0% N₂O, 21% Oxygen (O₂), and 79% nitrogen (N₂). A drug-onset cue was made by
delivering the first 3 minutes of the 60% \( \text{N}_2\text{O} \) gas condition followed by delivery of the control gas condition. This procedure for the drug-onset cue produced a maximum 39% \( \text{N}_2\text{O} \) concentration in the chamber, whereupon the incurred gas was switched to control gas such that the percent \( \text{N}_2\text{O} \) in the chamber decayed exponentially to 0% over \(~15\) min (graphically depicted in Fig. 1).\(^{21}\)

The DOC and CEXP protocols were identical except for the initial period of \( \text{N}_2\text{O} \) inhalation in the DOC group.

Telemetric measurement of \( T_c \), data acquisition and instrument control

Telemetric measurement of \( T_c \) was accomplished using a commercial system from Data Sciences International (Saint Paul, MN) that consists of a Data-Exchange Matrix, Physio-Tel Receiver (Model RPC-1), Dataquest ART 4.2 software, and an implantable battery-powered temperature sensor (model TA-F40) implanted in the rat’s peritoneal cavity. The antenna system used in each direct calorimeter consists of 2 radio ferrite coils oriented perpendicularly to each other and epoxied underneath a Plexiglas platform that holds them 2 mm above the floor of the direct calorimeter. The antennae wires exit the calorimeter through a gas-tight port and are connected to the RPC-1 receiver base. All other instrument control and data acquisition were performed using custom programs written in LabVIEW 6.8 (National Instruments, Austin, Texas).

Surgical placement of the telemetric temperature sensor

A telemetric temperature sensor was implanted surgically into each rat’s peritoneal cavity using isoflurane anesthesia (3–5% for induction and 1–3% for maintenance) while the rat was on a 39 °C heating pad. Meloxicam (an NSAID) was provided in the drinking water (0.02 mg/ml \( \text{H}_2\text{O} \)) from 1 day before to 2 days after surgery.

Data reduction for total calorimetry and \( T_c \)

Dependent variables were \( T_c \), HP, DHL (dry HL) and EHL (evaporative HL). \( T_c \) was recorded at 15-s intervals while HP and HL data were recorded at 10-s intervals. Average \( T_c \), HP and HL were calculated for each 6-min bin. Data were also analyzed in terms of change (\( \Delta \)) from baseline values that were defined as the means in the final 12 min of the baseline period. Gas concentration data were recorded from each calorimeter at 1-min intervals.
**Results**

**Body mass and baseline core temperature during the induction phase.** Body mass increased markedly during the induction phase but did not differ among groups ($F_{2,21} = 0.04; P = 0.96$). At the first exposure, animals weighed 157.2 ± 5.1 g whereas by the 18th they weighed 424.3 ± 6.8 g. Baseline core temperature decreased slightly but significantly over the induction phase ($t = −7.44; P < 0.0001$; coefficient estimate $= −0.017 ± 0.002°C$ per exposure) but did not differ among groups ($F_{2,21} = 1.18; P = 0.33$). On the first exposure, baseline core temperature was 37.3 ± 0.06°C; on the 18th it was 37.0 ± 0.04°C. Thus, consistent with our use of random assignment, groups were well matched across exposures in terms of body mass and baseline core temperature, 2 variables with potentially important effects on changes of core temperature during N₂O administration.

**Development of thermal tolerance with repeated N₂O administration**

Consistent with our previous work,$^{19,21-23,29}$ repeated 90-min 60% N₂O administrations engendered the development of thermal tolerance, defined as the absence of N₂O-induced hypothermia, and a subsequent sign-reversal, defined as the occurrence of hyperthermia during drug administration (Fig. 2A, B). The thermal sign-reversal occurred reliably during exposures 6–18, as indicated by the 95% CIs in Figure 2A. The magnitude of hyperthermia during exposures 7–18 did not differ significantly among the 3 groups (adjusted for exposure number and baseline core temperature: $F_{2,28.5} = 0.71; P = 0.50$).

**Effect of drug-onset cue on thermal outcomes during extinction trials**

Comparisons between the DOC and CEXP groups indicated that the DOC had thermal consequences in the early period of measurement (min 12–24 after the t₀ presentation of the DOC; see Table 1). Specifically, transient physiological effects were elicited by the drug-onset cue’s abbreviated N₂O administration (achieving a peak value of 39% N₂O and a mean concentration of 9% N₂O during the first 20 min after time zero). These DOC-associated changes did not continue after this early period (12–24 min) and there was no evidence that DOC-associated early thermal changes increased or decreased across extinction trials ($0.21 < P < 0.97$ for within subjects analysis of exposure number for the baseline-adjusted thermal outcomes).

**Effect of extinction procedures on thermal outcomes during the test trial**

Figure 2 C and D depicts Tc and HP profiles for each extinction group during the post-extinction-period test session. Unexpectedly, the home-cage group had reliably higher baseline Tc than either the CEXP or DOC groups (by 0.74 ± 0.21°C and 0.67 ± 0.21°C, respectively; $P < 0.005$). Because this result contrasts with the absence of group differences in baseline Tc during

**Table 1. Effect of DOC on thermal outcomes measured in the 3rd and 4th 6-min bins following t₀**

| Outcome (units) | DOC minus CEXP group difference in change from baseline ± SE* | P-value |
|-----------------|-----------------------------------------------------------------|---------|
| ΔTc (°C)        | 0.20 ± 0.1025                                                   | 0.07    |
| ΔHP (W)         | 0.40 ± 0.180                                                   | 0.047   |
| ΔDHL (W)        | 0.16 ± 0.032                                                   | <0.0005 |
| ΔEHL (W)        | 0.03 ± 0.024                                                   | 0.25    |

*Based on repeated measures regression analysis encompassing the 8 extinction trials. Values are adjusted for baseline thermal values.
the induction of the thermal sign-reversal, it suggests that the rats that remained in their home cages during the extinction period generated a greater stress response when returned to the calorimeter for testing than did the 2 groups that had experienced the calorimeter environment 8 times each over the 20-d extinction period.

Notably, N₂O exposure in the test session evoked hyperthermic changes in Tc (Fig. 2C) that were undiminished...
in comparison with those observed in the latter sessions of the induction protocol (F1,18.7 = 3.14, P = 0.09; test session minus session 18 difference = 0.16 ± 0.193°C).

The baseline-adjusted early and overall increases in HP, DHL or EHL during N2O exposure did not differ reliably among groups in the test session (F2,20 ≤ 2.34; P ≥ 0.12).

Discussion

The primary aim of this study was to test the hypothesis that a robust hyperthermic sign-reversal state acquired by repeated inhalation of 60% N2O might be attenuated by extinction trials involving unpaired associations with cues that were previously associated with drug inhalation. This concept is important because some models of drug addiction suggest that a sign-reversal state may contribute to the escalation of drug consumption.23,24 Thus, a procedure that would diminish or eliminate a sign-reversal could have potential therapeutic significance. In addition to evaluating extinction trials using environmental cues, the present study also evaluated the effectiveness of extinction trials that use an interoceptive drug-onset cue (DOC).15,16 This approach was founded on an extension of the traditional extinction model that holds that interoceptive cues caused by the sensory consequences of the drug itself can be critical CSs in eliciting CRs to drug administrations.4,13–17 According to this view, to successfully extinguish the effect of a DOC to elicit CRs in conditioned individuals, the extinction protocol must disassociate the initial sensory sequellae of the drug onset with the subsequent impending drug effects. It is thought that the extinction process does not erase or destroy the original learning, but rather depends on learning a new altered contingency.30–32

Our data, however, revealed that a robust acquired intra-N2O-administration hyperthermic state (i.e., the sign-reversal hyperthermia) remained fully intact in a N2O test session following a 20-d interval in which rats received extinction trials. The sign-reversal remained fully intact during the test session and, not surprisingly, the major driver of the hyperthermic state was a prompt and substantial increase in metabolic HP with the onset of N2O.

The inefficacy of the extinction procedures in our work must be interpreted with caution, given both the details of our experimental model and study design and a host of broader theoretical questions related to the mixed record of cue exposure therapy.12 An important advantage of our system is that it measures the hyperthermic sign-reversal and chronic tolerance development to N2O-induced hyperthermia.

An important advantage of our system is that it measures the underlying determinants of Tc, specifically HL and HP, via simultaneous direct gradient-layer calorimetry (HL) and indirect calorimetry (HP).20 These outcomes permit valuable insights into the behavior of the effector responses that underlie perturbations and adaptations in Tc during drug administration. We have previously documented that the initial hyperthermia evoked by N2O primarily reflects a marked increase in HL, whereas the intra-administration hyperthermic state that develops with repeated N2O treatments primarily reflects an acquired increase in HP that occurs briskly upon N2O onset.18–20 It must also be emphasized that HP and HL can exhibit substantial reciprocal changes during N2O administration such that Tc does not change, i.e., N2O’s effect to promote temperature loss via increased HL was fully obviated by a countervailing increase in HP.18 Thus, our model enables identification of agonistic pharmacological effects (HL) and antagonistic effector responses (HP) that would be masked or inferred by a model that measures only Tc. Because of this, the failure of the employed extinction procedures to attenuate changes of HP or HL demonstrates that the persistent change in the regulated variable (Tc) reflects a persistent adaptation in the behavior of its underlying determinants during N2O administration. Interestingly, Metzger et al. were unable to extinguish an acquired thermal tolerance to a cold exposure challenge despite evidence that associative processes played a role in tolerance development using the same method.37,38

Of fundamental importance to interpreting our results is the issue of whether the rats are able to sense the interoceptive DOC employed in our study. Our DOC was an initial 3-min rise in the N2O concentration with a time-concentration profile that was identical to the first 3 min of a standard 60% N2O administration. Presumably any change in sensory information or effector activity that occurred on the induction trials would also have occurred with the 3-min N2O administration. In the present study, the brief N2O DOC did in fact exert measurable physiological effects. A previous study employed a shorter duration N2O DOC that also caused significant changes in early ΔHP and early ΔDHL.21

In a previous study, we were unable to demonstrate classical conditioning of N2O-associated thermal responses using exteroceptive cues as well as interoceptive drug-onset-cues.21 As a possible explanation, we previously suggested21 that N2O’s effect as an NMDA receptor antagonist might impair the development of conditioned thermal responses owing to the important role of NMDA-glutamatergic signaling in memory acquisition.41 In light of the present failure to attenuate thermal responses using a N2O DOC in a cue-exposure paradigm, we now suggest that studies are warranted to determine whether co-administration of N2O with another drug that is known to permit conditioning (e.g., ethanol) would prevent conditioning to that drug.

Thus, there is an apparent conundrum with regard to using extinction procedures in an attempt to eliminate a sign-reversal that has been suggested to encourage further drug taking. It may be that associative mechanisms are not the underlying cause of
the sign-reversed hyperthermia, such that extinction attempts are futile. This would be consistent with the NMDA-antagonistic pharmacologic action of N\textsubscript{2}O to prevent conditioning from occurring. However, that would imply that a different mechanism, one that is not reliant upon conditioning, accounts for the sign-reversal, and tolerance as well. In either instance, the promise for using extinction procedures to successfully reduce sign-reversals as a therapeutic strategy is not supported by the current data from this pre-clinical model.

Our study employed 8 extinction trials distributed across a 20-d interval. It is possible that a greater number of extinction trials is required to bring about a reduction in the sign-reversal hyperthermic state that was established during the 18 induction administrations. A major review on cue-exposure therapy noted that the number of cue-extinction trials necessary to achieve extinction had a range of 2-35 sessions.\textsuperscript{12} In addition, in our study, each extinction session involved only one DOC exposure, whereas some studies have used multiple cue exposures per trial (reviewed in\textsuperscript{12}). Some evidence indicates that the number of cue extinction trials needs to equal or exceed the number of induction trials,\textsuperscript{42}) whereas our study involved a ratio of 18 induction : 8 cue extinction trials. Accordingly, future cue exposure studies involving our paradigm should include more extinction trials and perhaps more cues per trial.

There are important implications of these findings for theories of physiological regulation. A homeostatic interpretation of N\textsubscript{2}O-elicted hypothermia and its subsequent tolerance would suggest that the initial hypothermia that begins within minutes of the first N\textsubscript{2}O session acts like a US (unconditioned stimulus) that elicits compensatory responses to counter the hypothermia. As discussed above, we have found that the initial N\textsubscript{2}O-induced hypothermia is due to a concomitant N\textsubscript{2}O-induced increase of HL. The compensatory increase of HP is initiated later, beginning from 30 min to an hour or 2, with considerable variability among animals.\textsuperscript{20,21} Thus, while the increased HL (the unconditioned drug effect) persists during the N\textsubscript{2}O administration, the subsequent increase of HP lags and ultimately raises Tc, accounting for acute tolerance.\textsuperscript{20} This is perfectly consistent with the principles of homeostasis. A perturbation (decreased Tc) elicits a UR (unconditioned response) that ultimately restores Tc to normal. Over trials, the compensatory response increases in magnitude and occurs earlier within the N\textsubscript{2}O trial, eventually completely mitigating the unconditioned hypothermia (i.e., chronic tolerance develops).\textsuperscript{19,21}

However, while these phenomena are easily explained and in fact predicted by a homeostatic adaptation model, the changes that occur with further N\textsubscript{2}O trials are not. The finding that rats develop a sign-reversal hyperthermia when administered N\textsubscript{2}O would not be predicted by theories of homeostasis and in fact is counter to its premises.\textsuperscript{22,23} It should be noted that sign-reversals that occur after the prior development of tolerance has been achieved have been described for other drugs, e.g., opioid-induced hyperalgesia.\textsuperscript{43,44}

In part because of sign-reversals and other phenomena inconsistent with homeostatic principles, many have advocated that a different model of physiological regulation be adopted, and the current front-runner is allostatia. Allostasia was in fact formulated consistent with homeostatic principles, many have advocated that an allostatic model of regulation to explain the persistent hyperthermic sign-reversal observed in this study. The failure of the sign-reversal hyperthermia, and indeed of the tolerance itself, to be extinguished using the present paradigm does not speak to either allostatic or homeostatic interpretations as both models acknowledge the importance of associative mechanisms to regulation. However, contemporary views of allostatia emphasize the important role of central stress effects and peripheral stress hormones as possible mediators of allostatic effects. Stress may play a role in mediating the sign-reversal hyperthermia observed in this study.

In conclusion, the hyperthermic sign-reversal remained fully intact during the test trial, which suggests that cue exposure may not be an efficacious strategy to reduce sign-reversals that develop with chronic drug use.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Acknowledgments**

The authors gratefully acknowledge the contributions of Christopher W Prall and Hoang Yen Ho for their technical contributions to this study. The Helen Riaboff Whiteley Center, located at the University of Washington’s Friday Harbor Laboratories, provided an ideal environment for working on this manuscript.

**Funding**

This investigation was supported by the National Institutes of Health (NIDA grant DA023484).

**References**

1. Piazza PV, Deroche-Gamonet V. A multistep general theory of transition to addiction. Psychopharmacology (Berl) 2013; 229:387-413; PMID:23963530; http://dx.doi.org/10.1007/s00213-013-3224-4

2. Siegel S, Baptista MA, Kim JA, McDonald RV, Weise-Kelly L. Pavlovian psychopharmacology: the associative basis of tolerance. Exp Clin Psychopharmacol 2002; 10:162-83; PMID:1223979; http://dx.doi.org/10.1097/000213-013-3224-4

3. Siegel S, Ramos BM. Applying laboratory research: drug anticipation and the treatment of drug addiction. Exp Clin Psychopharmacol 2002; 10:162-83; PMID:1223979; http://dx.doi.org/10.1037/1064-1297.8.3.276

4. Dwornik BR. Learning and physiological regulation. Chicago: University of Chicago Press; 1993.

5. Ramsay DS, Woods SC. Biological consequences of drug administration: implications for acute and chronic tolerance. Psychol Rev 1997; 104:170-93; PMID:9009884; http://dx.doi.org/10.1037/0033-295X.104.1.170

6. Crowell CR, Hinson RE, Siegel S. The role of conditional drug responses in tolerance to the hypothermic effects of ethanol. Psychopharmacology (Berl) 1991; 117:3:51-4; PMID:6765789; http://dx.doi.org/10.1007/BF00841301

7. Li AD, Poulos CX, Cappell H. Conditioned tolerance to the hypothermic effect of ethyl alcohol. Science 1979; 206:1109-10; http://dx.doi.org/10.1126/science.493999
1. Kaiyala KJ, Butt S, Ramsay DS. Assessment of heat production, heat loss, and core temperature during nitrous oxide exposure: A new paradigm for studying drug effects and opponent responses. Am J Physiol Regul Intercell Physiol 2005; 288.8692-701; PMID:15563578; http://dx.doi.org/10.1152/ajpregu.00412.2004

2. Ramsay DS, Woods SC, Kaiyala KJ. Repeated nitrous oxide exposure in rats causes a thermoregulatory sign-reversal with concurrent activation of opposing thermoregulatory effectors. Temperature; 2014; 1:3

3. Ramsay DS, Woods SC, Kaiyala KJ. Drug-induced regulatory overcompensation has motivational consequences: implications for homeostatic and allostatic models of drug addiction. Temperature. 2014; 1:3

4. Feise RJ. Do multiple outcome measures require p-value adjustment? BMC Med Res Methodol 2002; 2:8; PMID:12069695; http://dx.doi.org/10.1186/1471-2288-2-8

5. Bardon (2004), Bossert and Shaham (2004), Boulanger and NM, Wasse JH. Applied Longitudinal Analysis. Hoboken, New Jersey: John Wiley & Sons, Inc; 2004.

6. Feise RJ. Do multiple outcome measures require p-value adjustment? BMC Med Res Methodol 2002; 2:8; PMID:12069695; http://dx.doi.org/10.1186/1471-2288-2-8

7. Perneger TV. What’s wrong with Bonferroni adjustments. BMJ 1998; 316:1236-8; PMID:9553006; http://dx.doi.org/10.1136/bmj.316.7139.1236

8. Rothenberg RE, and NM, Wasse JH. Applied Longitudinal Analysis. Hoboken, New Jersey: John Wiley & Sons, Inc; 2004.

9. Ramsay DS, Kaiyala KJ, Leroux BG, Woods SC. Individual differences in initial sensitivity and acute tolerance predict patterns of chronic drug tolerance to nitrous oxide-induced hyperthermia in rats. Psychopharmacology (Berl) 2005; 181:48-59

10. Ramsay DS, Kaiyala KJ. Relationship between the Development of Tolerance to Repeated Cold-Exposure in Rats. Anim Learn Behav 1995; 23:9-16; http://dx.doi.org/10.1177/009132859402300102

11. Ramsay DS, Kaiyala KJ, Leroux BG, Woods SC. Contextual and temporal modulation of extinction: behavioral and biological mechanisms. Biol Psychiatry 2006; 60:352-68; PMID:16616731; http://dx.doi.org/10.1016/j.biopsych.2005.12.015

12. Quirk GJ, Mueller D. Neural mechanisms of extinction learning and retrieval. Neuropsychopharmacology 2008; 33:56-72; PMID:17882236; http://dx.doi.org/10.1038/sj.app.300155

13. Kalant H, LeBlanc AE, Gibbins RJ. Tolerance to, and dependence on, some non-opiate psychotropic drugs. Pharmacol Rev 1971; 23:135-91; PMID:4398655

14. Lovinger DM, Crabbe JC. Laboratory models of alcoholism: treatment target identification and insight into mechanisms. Nat Neurosci 2005; 8:1471-80; PMID:16265990; http://dx.doi.org/10.1038/nn1581

15. Cunningham CL, Crabbe JC, Rigter H. Pavlovian conditioning of drug-induced changes in body temperature. Pharmacol Ther 1983; 23:365-91; PMID:1471-2288-2-8; PMID:12069695; http://dx.doi.org/10.1037/0073-274X(83)90019-0

16. Siegel S. Tolerance to the hyperthermic effect of morphine in the rat is a learned response. J Comp Physiol Psychol 1978; 92:1137-49; PMID:759560; http://dx.doi.org/10.1037/0073-274X(83)90019-0

17. Kirkkis SC, Riccio DC. Stimulus Conditions Influencing the Development of Tolerance to Repeated Cold-Exposure in Rats. Anim Learn Behav 1995; 23:9-16; http://dx.doi.org/10.1177/009132859402300102

18. Jevtic-Tozovic V, Todorovic S, Mennerick S, Powell S, Dziewian K, Beenhoff N, Zoranski CF, Olney JW. Nitrous oxide (laughing gas) is an NMBA antagonist, neuroprotectant and neurotoxin. Nat Med 1998; 4:460-63; PMID:9546794; http://dx.doi.org/10.1038/nm.487

19. Powder JR. Memory and the NMDA receptors. N Engl J Med 1996; 334:1195-8; PMID:8665707; http://dx.doi.org/10.1056/NEJMchb0902052

20. Chelonis JJ, Calton JL, Hart JA, Schachtman TR. Antenuation of the renewal effect by extinction in multiple contexts. Learn Motiv 1999; 30:1-14; http://dx.doi.org/10.1016/S0747-0744(98)80013-6

21. Ossipow MH, Liu J, King T, Vanderah TW, Porreca F. Underlying mechanisms of pronociceptive consequences of prolonged morphine exposure. Biopysci Rev 2005; 80:319-24; PMID:15799927; http://dx.doi.org/10.1002/bip.20254

22. Vanderah TW, Suena NM, Ossipow MH, Malan TP Jr, Liu J, Porreca F. Tonic descending facilitation from the rostral ventromedial medulla mediates opioid-induced abnormal pain and antinociceptive tolerance. J Neurosci 2001; 21:279-86