Chapter 2

Stimulant Self-Administration

Leigh V. Panlilio

Abstract

Stimulants such as cocaine and the amphetamines are widely abused due to their rewarding effects. Much of what we know about drug abuse and drug reward comes from research involving stimulants, and much of this research involves using drug self-administration as an animal model of drug abuse. In this chapter, the example of stimulant self-administration is used to illustrate: (1) the basic methodology of drug self-administration procedures and (2) the behavioral principles that apply to addiction and animal models of addiction. Many variations of the self-administration procedure have been developed to model specific aspects of drug abuse, to assess the rewarding effects of drugs, and to assess the effects of treatments. The chapter describes how these variations are devised by stipulating the behavioral requirements for receiving the drug (i.e., the schedule of reinforcement) and incorporating drug-related environmental cues analogous to those that occur in the human drug-abuse environment.

Key words: Animal models, Drug abuse, Methodology, Stimulus, Reinforcement, Behavior, Learning

1. Introduction

Drug abuse is a behavioral phenomenon that has much in common with other behaviors, but also has unique aspects that set it apart. Addiction research is largely concerned with uncovering and explaining these unique aspects. Similarly, addictive drugs share certain properties with each other, but they also have differences that allow them to be divided into general classes. Drugs that give the user a feeling of alertness and energy are referred to as stimulants. This loose classification includes a range of drugs that vary with respect to their pharmacological actions and potential for abuse. Cocaine, amphetamine, and methamphetamine are the most widely abused drugs in this class, and also the most intensively studied. Along with opioids, such as heroin, these stimulants represent the de facto standard against which other addictive drugs are compared.
Drug self-administration is the primary animal model used to study drug abuse. In this chapter, the example of stimulant self-administration will be used to illustrate: (1) the basic methodology of drug self-administration procedures, and (2) the behavioral principles that apply to addiction and animal models of addiction. Examples will be given to illustrate certain points, but the chapter is not intended as a review of the literature. It is intended to provide a brief overview of the basic techniques used in this active area of research, to introduce the central concepts behind these techniques, and to provide a background for the other chapters in the book.

1.1. The Problem of Stimulant Abuse

In the United States alone, over two million people meet the criteria for dependence or abuse of cocaine or other stimulants (1). This abuse has high economic and social costs related to health care, crime, and loss of productivity (2). In the individual, stimulants can produce a variety of adverse effects, including insomnia, anorexia, tremors, teeth grinding, dizziness, repetitive movements (stereotypy), schizophrenia-like (psychotomimetic) symptoms, hyperthermia, a variety of cardiovascular effects, muscle rigidity, intracerebral hemorrhage, convulsions, respiratory depression, and sudden death (3, 4). Amphetamines and amphetamine derivatives such as methylenedioxymethamphetamine (“ecstasy”) can also have lasting neurotoxic effects (5, 6). The fact that addicted individuals continue to seek and consume stimulants despite these adverse consequences attests to the power of the drugs’ rewarding effects.

1.2. Drug Reward

The scientific consensus is that stimulants and other addictive drugs are abused (at least initially) because they increase neurotransmission in the mesolimbic dopamine pathway, a part of the reward system of the brain (7–9). Cocaine elevates dopamine levels by blocking the reuptake of dopamine after it has been released into the synaptic cleft (10). Amphetamine and methamphetamine elevate dopamine levels by several mechanisms, including stimulating dopamine synthesis and inverting the direction of dopamine uptake, causing a massive release of dopamine into the synaptic cleft (11, 12). While non-stimulant drugs of abuse also increase mesolimbic dopamine transmission, they do so less directly, generally by altering the activity of neurons “upstream” of the neurons that release dopamine.

The normal function of the reward system is to increase contact with biologically meaningful substances, stimuli, and events in the natural environment, such as food, water, and sex. This normal function is “hijacked” when addictive drugs pharmacologically activate the system. The effects of the drug reinforce drug-taking behavior, making it more likely to be repeated in the future. As drug use continues, the people, places, and things associated with
the rewarding effects of the drug come to function as cues that have rewarding effects of their own. These cues motivate the individual to seek the drug, and they guide the complex sequences of actions involved in obtaining, preparing, and administering the drug. Thus, the behavior of abusing drugs is shaped and maintained by the effects of both the drug itself and drug-related features of the environment.

To understand the etiology of addiction and to develop therapeutic treatments, it is necessary to study how drugs and drug-related cues affect behavior and the brain. Drug self-administration provides an animal model of drug abuse that is ideal for studying the rewarding effects of drugs and drug-related cues. These procedures typically involve implanting an intravenous catheter that allows a rat or monkey to self-inject a drug by performing a simple response (e.g., pressing a lever) that activates a motor-driven syringe pump. Laboratory animals will self-administer most of the same drugs that are abused by humans (13), and stimulants are readily self-administered by rodents and nonhuman primates. The basic drug self-administration procedure is highly flexible, and many variations have been developed to focus on specific aspects of drug reward, abuse, and addiction. Due to its flexibility, face validity, and close correspondence to human drug-taking behavior, drug self-administration is the “gold standard” among animal models of drug abuse.

Drug self-administration procedures do have some disadvantages compared to other animal models of drug abuse, such as drug-discrimination and place-conditioning procedures (see Chap. 6). For example, self-administration procedures generally require many training sessions per subject, and these sessions are relatively long. This limits the number of subjects that can be studied, reducing the statistical power of these studies. However, in most cases, small group size can be compensated for by using a within-subject experimental design that increases power by studying each subject under multiple conditions. Self-administration procedures can also be technically difficult to implement, requiring specialized equipment and skills. For example, the experimenter must monitor each animal’s progress closely and sometimes adjust the training parameters for specific subjects to obtain consistency of performance across subjects (see (14)). The experimenter must be skilled in shaping animal behavior, programming the experimental events, and implanting catheters. Catheter failure represents one of the most significant obstacles to this research, often causing subjects to be dropped from a study before it is finished. In rats, catheters typically last about 3–6 months. In monkeys, a catheter usually lasts a year or longer, and once it fails a new catheter can be implanted in the same vein or a different vein. So, individual monkeys can be studied for many years and are well
suited for long-term studies. Rats are better suited than monkeys for short-term studies, such as those focusing on the initial acquisition of drug use or involving irreversible treatments.

The self-administration of drugs by animals is presumed to be directly analogous to drug use by humans. Although similar drug self-administration procedures can be used in a laboratory setting with either human or animal subjects, there can be distinct advantages to using animals. The scientific method requires that researchers have the ability to control and manipulate the conditions that produce addiction. But, it would be unethical to induce addiction in humans for research purposes, and drug-experienced human volunteers have varied and complex histories that can interact with the experimentally imposed conditions being studied. In contrast, the environment and life experiences of laboratory animals can be carefully monitored and controlled. In addition, many techniques for studying the brain mechanisms involved in drug self-administration are not feasible or appropriate for use in humans and can only be implemented in animals.

The main drawback to studying drug self-administration in animals rather than humans is that species sometimes differ with respect to the reward-related effects of drugs. For example, rodents seem to differ from humans and other primates with respect to the rewarding effects of THC, the main psychoactive component of marijuana (15). But, even though the details can differ between species, the basic principles and phenomena do generalize across species, and the overall similarities between animals and humans support the validity of using animal models. In fact, a productive and efficient use of resources for medication development is to use rodents to test hypotheses generated from in vitro findings, then use primates to verify the most promising findings from testing in rodents, and finally to use human volunteers to evaluate the treatments that were found to be effective in primates.

Historically, drug self-administration procedures were directly adapted from the techniques used by experimental psychologists to study behavior maintained by natural rewards, such as food. Focusing on the rewarding effects of drugs in this manner places the study of addiction within a comprehensive approach to the study of behavior and how it is shaped by its interactions with the environment. Using this approach, researchers have created many variations of the self-administration procedure that focus on specific addiction-related aspects of the behavior, such as how much the drug is valued as a reward (see Sect. 3), how drug intake is
regulated (see Sect. 2.1, and Chap. 10), and what factors can cause or prevent relapse (see Chap. 17).

Variations of the self-administration procedure can be created by modifying the schedule of reinforcement, the set of contingencies that define the relationship between behavior and the delivery of reinforcement (16). The schedule specifies requirements such as what response produces the reinforcer, the number of times the response must occur, the amount of time that must pass before another reinforcer can be obtained, and whether any cues are provided to signal the availability of the reinforcer. With drug and nondrug reinforcers, in the laboratory and in the natural environment, the schedule of reinforcement is a powerful determinant of the pattern of responding. The sections below briefly describe the most commonly used schedules of drug self-administration and the specialized purposes they serve.

### 2.1. Continuous Reinforcement

Continuous reinforcement is the simplest schedule of reinforcement: only one response is required for each injection of the drug. However, it should be noted that in practice, not every response will produce an injection; since the injection takes some time to deliver, there is typically a timeout period of at least a few seconds following each injection under continuous reinforcement as well as most other schedules of reinforcement. Any responses that occur during the timeout period do not produce additional drug. In many studies, longer timeouts (typically 30–60 s) are instituted to prevent the animal from self-administering another injection before the previous injection has been delivered and adequately distributed to the brain.

One way that continuous reinforcement is used is to study the acquisition of drug self-administration (see Chap. 9). Even when an experiment will involve a more complex schedule later in training, a continuous-reinforcement schedule is used during initial training because consistently and repeatedly associating the response with the reinforcing effects of the drug facilitates acquisition of the response. Once responding has been established, more complex response requirements (such as those described in the sections below) can be gradually introduced.

The simple, direct relationship between responding and receiving the drug under continuous reinforcement makes this schedule well suited for studying rates and patterns of drug intake. Under this schedule, self-administration tends to occur in a specific pattern. Early in the session there is a period known as the loading phase, in which several injections are taken in relatively rapid succession. The remainder of the session is known as the maintenance phase, in which there is a pause of fairly constant duration following each injection. If the dose is changed, the duration of pausing changes, with higher doses producing longer pauses.
Over the course of the session, the level of drug in the animal’s body also follows a predictable pattern. During the loading phase, when injections occur rapidly, the drug level steadily increases. During the maintenance phase, when injections are evenly spaced in time, the drug level rises quickly after each injection, then gradually decreases as the drug is eliminated from the body during the post-injection pause. Most interestingly, the drug level where the next response occurs tends to be about the same with each successive injection. Thus, the animal appears to regulate its drug intake in such a way that the level of effect is not allowed to fall below a certain minimum (17–21).

This phenomenon of regulated drug intake is a quintessential aspect of drug self-administration, occurring reliably across many different laboratories, species, and drug classes. Consequently, understanding why it occurs might provide insight into the unique nature of drug reward. There are four basic mechanisms that might contribute to the highly regular post-injection pausing that underlies regulated drug intake (21–23). First, the drug may produce behavioral “side effects” (i.e., effects not related to reinforcement) that cause responding to slow down or cease temporarily. Second, the drug may have aversive, punishing effects when a high drug level is reached (see Sect. 2.8). Third, achieving a certain level of drug effect may produce satiation, such that the animal is no longer motivated to obtain reward. Fourth, even though the animal is still motivated, taking more drug may have no effect when the reward system becomes saturated by a high level of drug, so the animal learns to detect when this happens and pause until it detects that the drug level has dropped (24).

Understanding the mechanisms underlying regulated drug intake may also provide insight into the dysregulation of drug intake that is often associated with severe addiction. It has long been known that animals given access to stimulants 24 h per day develop excessive levels of intake (25–27). In recent years, it has been shown that when rats are provided with extended access to the drug (e.g., sessions lasting 6 h or more), they develop many of the hallmarks of addiction, such as escalated intake (18; see Chap. 10) and increased susceptibility to relapse (28). Thus, extended-access conditions can be used to provide an animal model of drug abuse that more closely approximates addiction, as opposed to casual, controlled use.

2.2. Ratio Schedules

Ratio schedules specify the number of responses that are required for each injection. Under a fixed-ratio schedule, the same number of responses is required for each injection. This schedule is usually designated by the abbreviation “FR” followed by the response requirement. For example, under FR 10 an injection is delivered for every tenth response. Continuous reinforcement is sometimes referred to as FR 1. With reinforcing drugs and with nondrug
reinforcers such as food, FR schedules that require a substantial number of responses for each reinforcer typically produce a specific pattern of responding known as “break and run”; there is an initial period of nonresponding (i.e., a break) after delivery of the reinforcer, followed by a period of rapid responding (i.e., a run) that continues until the required number has been reached.

In many cases, FR schedules of drug self-administration are used as baselines in studies that involve the testing of potential treatments for addiction (see Sect. 4). In other cases, fixed-ratio schedules are used to assess whether a novel drug has reinforcing effects. For example, when a medication is developed for a purpose such as treating pain or obesity, it is important to determine whether it also has rewarding effects that make it liable to be abused. One way to evaluate the rewarding effects of the new drug is to train animals to self-administer a drug with known abuse liability, such as cocaine, then see if the self-administration response is maintained when the syringe is filled with the new drug instead of cocaine. Ratio schedules are advantageous for these purposes because they engender substantial rates of responding if the new drug is an effective reinforcer (see Sect. 3). In contrast, the rate of responding maintained by a reinforcing drug under a continuous-reinforcement schedule can be quite low, especially if the drug has long-lasting effects, and can sometimes be difficult to distinguish from the rate that occurs when the drug is replaced by a placebo such as saline solution.

Continuous reinforcement is used to determine when the animal will take the drug if it is freely available. In contrast, a progressive-ratio schedule is used to determine how much the animal will “work for the drug.” Under a progressive-ratio schedule, the required number of responses is increased with each successive injection. The increases usually occur in steps according to an exponential progression. For example, a commonly used progression is 1, 2, 4, 6, 9, 12, 15, 20, 25, 32, 40, 50, 62, 77, 95, 118, 145, 178, 219, 268, 328, 402, 492, and 603. This kind of schedule is designed to determine the point at which the response requirement becomes so high that the drug’s reinforcing effects no longer maintain responding. Typically, the pattern of responding will become less consistent and long pauses will begin to appear as the requirement increases. Once a designated criterion is reached, for example, 1 h without a response, the final response requirement that was successfully completed is taken as the “breaking point.” The breaking point is a relatively direct measure of the strength of the drug as a reinforcer at the tested dose (see Sect. 3.3). The criterion chosen for determining the breaking point should be chosen to be substantially longer than the post-injection pause that would occur under continuous reinforcement or a low-requirement fixed-ratio schedule. The progression listed above was designed to start with a ratio of 1 and escalate quickly
enough that the animal will cease responding within a 5 h session (29). If the criterion is not met within a session, researchers sometimes simply use the highest ratio completed within the allotted session time as the datum. Although this may give orderly results, it is not clear whether these results would be comparable to “true” breaking points, defined by a nonresponse criterion. An alternative procedure when the criterion is not met within a single session is to start the next day’s session with a response requirement that is the same as (or one or two steps lower than) the highest step already reached, rather than starting again at the first step of the full progression.

Second-order schedules are one kind of procedure that incorporates the effects of drug-related environmental cues into the drug self-administration procedure (30). These schedules represent a combination of two simpler schedules, such as a fixed-ratio schedule and a fixed-interval schedule (the latter of which specifies the amount of time that must pass before the reinforcer becomes available). Under such a schedule, completing the requirements of the fixed-ratio schedule produces a brief stimulus presentation (e.g., a colored light for 2 s). Once the amount of time specified by the fixed-interval schedule elapses, the drug is given along with the next brief stimulus presentation. For example, Katz and Goldberg (31) used a second-order schedule of cocaine self-administration with squirrel monkeys. The brief stimulus was presented on a fixed-ratio 10 schedule (i.e., every tenth response produced the stimulus), and the first stimulus presented after 5 min was accompanied by an injection of cocaine. For comparison, a simple fixed-interval schedule was also studied, in which cocaine was delivered for the first response that occurred after 5 min, and the stimulus was only presented during the injection. As is typical of this kind of schedule, the second-order schedule generated much higher rates of responding than the simple fixed-interval schedule.

Responding occurs at higher rates under second-order schedules because the brief stimulus comes to have reinforcing effects of its own. These reinforcing effects result from classical conditioning (32), in which the stimulus associated with the effects of cocaine becomes capable of eliciting responses similar to those elicited by cocaine itself. The conditioned-reinforcing effects of the drug-associated stimulus can be seen most clearly when delivery of the drug is discontinued; even though the response produces only the stimulus, responding can be maintained at high levels for many sessions. These conditioned-reinforcing effects of the brief stimulus model the effects of cues that influence human behavior in the drug-abuse environment. For example, in order to experience the effects of cocaine, a person must come into contact with a series of specific cues: money, the drug supplier, the
packaging and physical properties of the drug, the place where the drug will be used, the injection paraphernalia, the sensation of the needle puncturing the skin. Each of these cues acts as a conditioned reinforcer that maintains part of the sequence of behavior.

Second-order schedules are valuable because they incorporate the effects of drug-related cues into an elegant animal model that begins to approach the complexity of drug abuse in the natural environment. However, it can also be valuable to have a simpler model of conditioned reinforcement. This can be accomplished with procedures in which a stimulus is first associated with the effects of a drug, then used to reinforce a novel response. During the conditioning phase, the drug can either be delivered automatically by the experimenter or self-administered by the animal (33). For example, Di Ciano and Everitt (34) trained rats to self-administer cocaine by poking their nose into a small hole in the wall of the training apparatus. Each cocaine injection was paired with a 20-s presentation of a cue light. Then, during a test session, the rat was presented for the first time with two levers, one of which produced the cue light for 1 s for every third response, and one of which had no effect. The rats responded more on the lever that produced the light, indicating that this drug-paired stimulus had become a conditioned reinforcer capable of establishing a new response.

Even when conditioned reinforcement or environmental cues are not the focus of a drug self-administration study, it is common practice to present a visual or auditory stimulus each time the drug is delivered. And, even when such stimuli are not intentionally provided by the experimenter, it is still likely that there are injection-related cues, such as the sound of the motorized syringe pump and the feeling of room-temperature fluid being injected. Whether incidental or intentionally programmed by the experimenter, these injection-related cues can have an important influence on drug self-administration. Such cues provide immediate feedback that the drug is being delivered, bridging the delay between the response and the onset of the drug’s effects, and thereby making the drug more effective at reinforcing the response. This situation is comparable to the standard procedure for studying food reinforcement, in which a stimulus such as a feeder click is immediately provided when the response occurs, but there is a delay between the pressing of the lever and the food pellet being taken into the mouth. Providing a drug-paired stimulus also makes the laboratory model more comparable to the natural drug-abuse environment, where ingestion of a drug is almost always accompanied by some kind of cue. Interestingly, one effect that stimulant drugs can have is to enhance the effectiveness of conditioned reinforcers. For example, amphetamine
and other stimulants can increase the conditioned-reinforcing effects of stimuli associated with food or water (35).

The conditioned-reinforcing effects of stimuli associated with drug delivery can induce relapse to drug use in humans, and this phenomenon can be modeled in laboratory animals with cue-induced reinstatement procedures (e.g., see (36); see Sect. 2.7, and Chap. 17). It should also be noted that conditioned reinforcement provides the basis for place-conditioning procedures (see Chap. 10), an animal model of drug reward in which the drug’s effects are associated with the features of an environmental context.

2.5. Multiple Schedules

As described above, drug-related cues act as conditioned reinforcers to establish and maintain the sequences of behavior that ultimately lead to experiencing the effects of the drug. An equally important role of environmental cues is to guide behavior by indicating what response is required at each step of the sequence. Cues that provide this guiding effect – by signaling when a specific response can produce a specific reinforcer or conditioned reinforcer – are known as discriminative stimuli. Discriminative stimuli can be incorporated into drug self-administration procedures by using a multiple schedule in which the experimenter presents tones or lights to signal when the drug is available. Under these conditions, the animal’s response will readily come under control of the discriminative stimulus, increasing in frequency when the stimulus is present and decreasing when it is absent (14, 37). In both the laboratory and the human drug-abuse environment, these cues exert considerable power over drug-related behavior, determining when and where specific responses will occur. For example, like conditioned reinforcers (which are produced by the animal’s response), discriminative stimuli (which are presented automatically by the experimenter) can reinstate drug seeking in an animal model of cue-induced relapse (e.g., (38)).

Although drug-related cues are often studied in the laboratory by focusing on the effects of a single cue, the natural environment is composed of a vast number of cues that can be encountered in various configurations. Multiple schedules can be used to study how the individual cues that make up these configurations interact to influence drug self-administration. For example, a tone and a light can be presented separately to signal the availability of cocaine during training, then these cues can be presented together for the first time during a test session. The combined power of these cues can cause rats to double their intake of cocaine (37). In contrast, if a tone signals when cocaine is available and a light signals when cocaine is not available during training, the light acquires inhibitory properties that can substantially decrease the amount of responding controlled by the tone during a subsequent test in which the cues are combined (39). Thus, individual
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Sixty cues can increase or decrease drug seeking depending on their relationship to the drug, and when multiple cues are encountered they interact to influence behavior. The effects of combined cues depend on both the discriminative control exerted by each cue (i.e., whether it occasions a habitual increase or decrease in responding) and the conditioned-incentive effects of the cue (i.e., whether it is associated with an increase or decrease in the rate of reinforcement; see 40).

Another way that multiple schedules are used is to evaluate whether a potentially therapeutic treatment can selectively decrease drug self-administration. For this purpose, a multiple schedule is used in which responding produces food pellets in the presence of one stimulus and drug injections in the presence of another stimulus (e.g., 41). The goal is to discover treatments that decrease drug-reinforced responding while leaving food-reinforced responding intact. If a treatment decreases both food and drug responding, it might be producing general sedative or motor-depressant effects rather than altering the reinforcing effects of the drug. Or, it might be blocking the reinforcing effects of both the drug and natural reinforcers, an effect that could make it less desirable as a treatment.

2.6. Chained Schedules

Chained schedules are related to second-order and multiple schedules in that (1) they are complex schedules composed of simpler components and (2) they incorporate the effects of environmental cues. The difference is that in chained schedules, the responding in one component of the schedule has the effect of advancing the schedule to the next component, with the drug only received in the final component. For example, Olmstead et al. (42) used a chained schedule of cocaine self-administration in which rats’ “drug-seeking” and “drug-taking” responses occurred on separate, retractable levers. At the beginning of a reinforcement cycle, the seeking lever was inserted and a light was presented to indicate that responses on that lever could provide access to the taking lever. Once the rat had responded on the seeking lever for about 30 s, the next response on the seeking lever caused the taking lever to be inserted. The first response on the taking lever produced an injection of cocaine, and the lights were turned off to signal a timeout period. In this schedule, the retractable levers not only provided a means of responding, but served as cues. That is, insertion of either lever functioned as a discriminative stimulus to respond on that lever, and insertion of the taking lever functioned as a conditioned reinforcer for responding on the seeking lever.

The main advantage of this seeking–taking chained schedule is that it isolates drug seeking from drug taking. The ability to distinguish between drug-seeking and drug-taking responses is important because different mechanisms may underlie these
behaviors (e.g., see (43)). This kind of distinction can also be achieved with second-order schedules by either focusing on the behavior that occurs prior to the first injection of the session or by using a procedure in which the drug is only injected at the very end of the session (44).

2.7. Extinction

Extinction refers to discontinuing reinforcement. When a behavior such as lever pressing is no longer reinforced, its frequency will eventually drop to a very low level. However, when extinction is first instituted, the animal’s response rate will often increase for a short time before it decreases; this temporary increase in responding is known as an “extinction burst.” Furthermore, once the response has decreased to a low, stable level, it may increase again (i.e., show spontaneous recovery) if the animal is reexposed to the training apparatus after a hiatus. Thus, even though a response no longer produces the drug and does not occur for extended periods of time, it does not disappear completely.

Extinction is most widely used in addiction research as a phase of training in the reinstatement model of relapse ((45); see Chap. 17). In this procedure, extinction is used to parallel abstinence from drug use. For example, in a typical reinstatement procedure, rats are trained to self-administer cocaine, then an extinction phase is instituted in which drug delivery is discontinued until responding drops to a low level. Finally, a reinstatement test is performed in which the animal is given a treatment and allowed to respond, but the response still does not produce the drug. If the treatment increases the frequency of responding (i.e., responding is reinstated), this is considered to be analogous to a relapse of drug-seeking behavior. Three general kinds of treatment are effective in producing reinstatement in the laboratory: exposure to drugs (either the training drug or a different drug), exposure to stress (usually a series of mild footshocks), or exposure to drug-related cues (discriminative stimuli or response-produced conditioned reinforcers). These treatments correspond to the triggers that are known to induce relapse to drug use in humans.

Extinction can also be used as one part of a complex schedule of reinforcement, such as a multiple schedule that includes signaled periods in which the drug is not available. As mentioned above (see Sect. 2.5), stimuli that signal a period of extinction can produce inhibitory effects when combined with other drug-related stimuli. Extinction can also be used to assess the persistence of drug seeking. For example, there is evidence that addiction-like states in animals may cause resistance to extinction in heroin-trained rats (46) but not cocaine-trained rats (28, 47, 48). It has long been known that a history of intermittent reinforcement (i.e., training with schedules other than continuous reinforcement) makes behavior resistant to extinction, and that certain cues and contexts can maintain responding during extinction.
Understanding the factors that contribute to the persistence of behavior in the face of changes such as extinction could provide new avenues for the prevention and treatment of addiction (49).

2.8. Punishment

Continuation of drug use despite adverse consequences is a primary symptom of drug addiction. Such consequences can be modeled in the laboratory by using punishment procedures. Punishment occurs when a response produces something aversive that decreases the likelihood of the response occurring in the future. For example, if a brief footshock is delivered whenever the rat presses the lever in a drug self-administration procedure, the rate of self-administration will usually decrease. However, resistance to punishment, like resistance to extinction and resistance to conditioned suppression (see Sect. 2.9), may reveal increased compulsivity (see Chap. 13) or addiction-like increases in the motivation to receive the drug (48, 50).

Punishment can occur in the human drug-abuse environment in several different ways. Often, it is imposed by other people (e.g., employers, law enforcement) to decrease drug use. But, there can also be an inherently aversive, punishing component to the effects of the abused drug itself. For example, self-administered cocaine can produce both reinforcing and punishing effects in the same animal (51). Unfortunately, both the punitive measures meted out by society and the inherently aversive effects of abused drugs tend to be delayed relative to the rewarding effects, and this delay reduces their ability to decrease drug use. Nevertheless, adverse consequences can be one factor that limits drug use or promotes abstinence in addicted individuals.

It is worth noting that a number of abused drugs (e.g., benzodiazepines, barbiturates, ethanol) have rewarding effects but can also reverse the effects of punishment. The anti-punishment and anti-anxiety effects of these drugs might promote their coadministration with stimulants (52). For example, drugs such as diazepam and buspirone may counteract the inherent aversive effects of self-administered cocaine (53). Furthermore, anti-punishment drugs might directly induce relapse to drug use when abstinence has been achieved through punishment (54).

2.9. Conditioned Suppression

Conditioned suppression is a phenomenon in which ongoing behavior is disrupted by a conditioned stimulus. It is used in two ways in addiction research. First, like punishment, it can be used to assess the persistence of self-administration behavior; the main difference is that the aversive event is produced by the response in punishment procedures, but the aversive event occurs regardless of the animal’s behavior in conditioned suppression. For example, rats that are self-administering cocaine can be periodically presented with a visual stimulus signaling that an unavoidable footshock will be delivered. Responding will normally decrease
during presentation of the conditioned stimulus, but resistance to this suppression may be an indicator of addiction (55).

Second, conditioned-suppression procedures can be used as a model of conditioned drug effects (see (56)). For example, rats that are responding on a schedule of food reinforcement can be presented with a visual stimulus signaling that an intravenous injection of cocaine will be automatically delivered. The cocaine-associated stimulus will disrupt responding (57). This suppression induced by cocaine-associated stimuli in rats may be analogous to the cue-induced effects that humans describe as drug craving. Since conditioned suppression can be produced by either hedonically negative events (e.g., footshock) or hedonically positive events (e.g., delivery of food; (58)), the suppression induced by cocaine-paired cues could be due to either reward-related or aversive effects.

Reinforcement is said to occur when a response has an effect on the environment that makes the response more likely to be repeated in the future. The reinforcement discussed in the chapter thus far is positive reinforcement, which occurs when a response has the effect of producing something, like a drug injection or food pellet. In contrast, negative reinforcement occurs when the response becomes more likely because it eliminates something aversive. Some drugs can produce negative reinforcement by providing relief from pain, stress, or anxiety. Since withdrawal from chronically administered drugs of abuse is usually unpleasant, avoiding or escaping from this state can be negatively reinforcing. For example, morphine-dependent monkeys will press a lever that prevents or terminates injections of opioid antagonists that precipitate withdrawal symptoms (59). Although the symptoms produced by withdrawal from cocaine and other stimulants are not as severe as those produced by opioid withdrawal, stimulant withdrawal can produce depression-like effects ((60, 61); see Chap. 1). Avoiding or escaping from these unpleasant states might contribute to the persistence of drug use ((62); see Chap. 16).

However, there are several reasons to believe that addiction stems primarily from the positive reinforcement produced by drugs, rather than the negative reinforcement produced by avoidance of withdrawal symptoms or other unpleasant states (63). Neither precipitated withdrawal from heroin nor presentation of stimuli associated with withdrawal appear to motivate heroin seeking under a seeking–taking chained schedule in a way that would be consistent with negative reinforcement (64). Medicines that provide relief from unpleasant conditions such as pain but do not activate the reward system are not addictive, and many drugs that produce unpleasant effects during withdrawal (e.g., antidepressants, antihistamines; (65)) are not addictive. Perhaps most
importantly, escape from withdrawal symptoms cannot account for relapse to drug use in individuals who have been abstinent for a long period of time. This suggests that treatments that alleviate withdrawal symptoms might assist in achieving abstinence, but by themselves are unlikely to prevent relapse.

3. Assessing Reinforcing Effects

The assumption that the reinforcing effects of drugs underlie their potential for abuse and addiction is central to the drug self-administration model of human drug abuse. It is worth noting that these reinforcing effects are not an immutable property of the drug, but depend on a number of factors such as the animal’s reinforcement history, drug history, current state, access to alternative sources of reinforcement, and genetic makeup (see Chap. 11). As a result, being able to assess the reinforcing effects of a drug under various conditions is critical to conducting drug self-administration research. The process of assessing reinforcing effects essentially involves making comparisons. Does any dose of drug X have a reinforcing effect compared to a saline solution? How do various doses of drug X compare to each other? How do the reinforcing effects of drug X over a range of doses compare to those of drug Y? Does a potentially therapeutic treatment change these effects? Does inactivating a certain brain area change these effects?

3.1. Control Procedures

The most basic of these questions is whether a certain dose of a drug is having a reinforcing effect. For example, a rat may repeatedly press a lever that produces intravenous injections of a drug, but this in itself does not demonstrate that the drug’s effects are reinforcing. By definition, reinforcement is evident when the response that produces the drug becomes more likely to be repeated. This increase in likelihood must be measured relative to some control condition. A commonly used control procedure is to provide the rat with a second lever (an “inactive” lever) that does not produce the drug. If the rat presses the active lever more than the inactive lever, this indicates that the drug is having a reinforcing effect. This two-lever control procedure is an efficient and valid way to verify that the injections are reinforcing.

However, it should be noted that responding on the inactive lever in this two-lever control procedure cannot be considered a truly neutral, nonreinforced behavior. There can be generalization (66) of responding from the reinforced lever to the inactive lever due to their physical similarity. Furthermore, sometimes by chance, the drug may be delivered when the rat responds on the active lever soon after responding on the inactive lever, which can lead to “superstitious” responding on the inactive lever (67).
As a result, responding on the inactive lever should not be considered a measure of general locomotor activity independent of the reinforced responding that occurs on the active lever.

An informative but less commonly used control procedure is to have additional, independent groups of rats that have access to a lever but never receive the drug, or that only receive the drug passively, regardless of whether they press the lever. Even if the experimenter never programs anything to occur when the lever is pressed, rats will occasionally press it incidentally as they explore their surroundings. If the experimenter simply programs the lever to turn stimuli such as lights on and off, these stimuli changes can have an inherently reinforcing effect that maintains a certain level of responding. Importantly, passively received drugs can sometimes enhance the inherently reinforcing effects of these stimulus changes, causing substantial rates of responding ((68); see Chap. 4). Thus, depending on how they are implemented, independent-group control procedures can provide a more interpretable test for reinforcement than a two-lever procedure, and they can also provide unique insight into how drugs affect behavior.

The most important control procedure for confirming that a drug is having a reinforcing effect is to discontinue drug delivery and use the drug’s vehicle as a placebo. If the response is maintained by the reinforcing effects of the drug, responding should cease or at least decrease due to extinction when all conditions are kept the same except that the drug is no longer delivered. This demonstration of reinforcement is even clearer if the drug and extinction conditions are then repeated and the behavior increases and decreases accordingly. It is important to point out that, typically, responding only decreases to low levels after a number of extinction sessions. However, if animals are repeatedly tested with drug and extinction conditions, the extinction-induced decreases in responding will occur more rapidly. This kind of training can be valuable when the animals will be used to test the effects of treatments expected to block the reinforcing effects of the drug; animals that have learned to abruptly stop responding when the injections are not having a reinforcing effect may provide a more sensitive and valid test for blockade.

The procedure of comparing responding under drug and placebo conditions can (and usually should) be extended to comparing several different doses of the drug within each animal. Dose-dependence is an expected characteristic of any pharmacological effect. In the case of drug self-administration, demonstrating dose-dependence helps confirm that the drug is having a reinforcing effect and allows an accurate description of how a treatment changes the drug’s effects on behavior (see Sect. 4.1). When comparing doses, ideally each dose should be studied for several sessions, until the rate of responding stabilizes. The order in
which the doses are tested should be counterbalanced across animals. When each dose has been tested, the data are used to construct a dose–effect curve for each animal, showing the response rate (or some other measure) as a function of dose. These individual curves can be averaged together into a group curve to facilitate presentation of the results. But, it only makes sense to do this if the individual curves are generally consistent with each other.

Dose–effect curves describing response rates or injection rates under drug self-administration schedules typically exhibit an inverted-U shape. The peak of the curve occurs at a dose that is high enough to be reinforcing, but not so high that it produces long post-injection pauses. Along the descending limb of the curve (i.e., at doses higher than the peak), the rate of responding decreases as the dose increases, because higher doses produce longer post-injection pauses. Along the ascending limb of the curve (i.e., at doses lower than the peak), less responding is maintained than at the peak dose, probably because doses on the ascending limb are only weakly reinforcing. Response rates on the ascending limb often represent the averaging of alternating periods of rapid responding and periods of nonresponding, rather than a steady pattern of moderate responding. In many studies an ascending limb is not obtained, and responding at doses lower than the peak dose is comparable to responding under placebo conditions. Although the determinants of whether an ascending limb is obtained have not been studied extensively, likely factors include the schedule of reinforcement, the pharmacokinetic properties of the self-administered drug (i.e., how quickly and for how long it acts), and the animal’s training history.

Beyond the question of whether a drug is functioning as a reinforcer, there is the question of reinforcing efficacy, or how effective the drug is at maintaining the response. Reinforcing efficacy is measured by comparing the effects of different doses, different drugs, or even drug and nondrug reinforcers. However, these comparisons cannot be performed by simply measuring the response rate maintained by a reinforcer; response rates are influenced by too many factors. So, three specialized procedures have been developed for comparing reinforcing effects: progressive-ratio schedules, choice schedules, and behavioral economics analyses.

As described earlier (see Progressive ratio, above), breaking points under progressive-ratio schedules provide a direct assessment of how effectively a reinforcer maintains a response. Typically, the higher the dose, the higher the breaking point. However, in some studies there is a peak in the dose–effect curve above which the breaking point starts to decrease, possibly indicating that aversive side effects are beginning to appear, or perhaps that the
nonresponse criterion used to determine the breaking point is too short. An advantage of progressive-ratio schedules is that the breaking-point measure is relatively independent of the animal’s response rate. This independence is important because most drugs of abuse have behavioral side effects that can alter ongoing responding even when the drug is passively received. Depending on the drug, dose, schedule of reinforcement, and the animal’s history, such effects might involve either an increase or a decrease in response rate, and the size and direction of this effect may have nothing to do with the drug’s effectiveness as a reinforcer.

*Choice procedures* provide another means of comparing the efficacy of various reinforcers. Like progressive-ratio procedures, the measure of reinforcing efficacy obtained with choice procedures is independent of response rate. Choice procedures typically involve providing the animal with two levers: one lever produces a drug, and – depending on the study – the other lever produces the same drug at a different dose, produces a different drug, or produces a nondrug reinforcer such as food. If one lever is consistently chosen over the other, the reinforcer associated with that lever presumably has a higher reinforcing efficacy. However, it should be noted that the availability of two different reinforcers in the same session can sometimes alter their efficacies, such that the results obtained with the choice procedure do not agree with those obtained with a progressive-ratio or behavioral-economics procedure in which the reinforcers are studied separately. As with other self-administration procedures, it is important to assess choice over a range of doses. In addition, control procedures should be used to ensure that an apparent preference for a specific dose is not just due to a side bias (e.g., the animal habitually choosing the left lever); this usually entails switching the outcomes associated with the two levers and verifying that the behavior shifts appropriately.

A *behavioral-economics* approach to studying drug self-administration combines principles of psychology and microeconomics (69). Drugs are viewed as commodities, and the animal’s response output is viewed as the allocation of a resource. Typically, this kind of research involves using fixed-ratio schedules of drug self-administration and comparing various combinations of response requirements and doses. The response requirement and dose are converted to a *unit price*. For example, under a FR 10 schedule in which each injection contains 1 mg of cocaine, the unit price of cocaine would be 10 responses per mg; the same unit price (10 responses per mg) could be studied by giving 0.1 mg of cocaine in each injection under a FR 1 schedule. A *demand curve* is generated, depicting the amount of drug consumed as a function of unit price. These curves usually descend from left to right, with consumption decreasing as the price increases. The steepness of this curve indicates the *elasticity of demand*, how sensitive
consumption is to changes in price. If consumption drops off quickly as price is increased, the demand is elastic. If consumption remains stable even when the price is increased, the demand is inelastic, which indicates that the drug is highly valued and treated as a necessity.

As a general approach to studying the allocation of behavior, a behavioral-economics analysis can be applied to many situations, including progressive-ratio and choice procedures. The decreased consumption seen under high costs in a demand-curve analysis is presumably related to the cessation of responding that defines the breaking point under a progressive-ratio schedule. With regard to choice schedules, the behavioral-economics approach provides an established theoretical framework for analyzing interactions between different commodities. Such interactions are important to consider, since the availability of one commodity may decrease or increase the demand for another. For example, food and water have a complementary relationship: consuming food makes water more reinforcing. Certain commodities can also “substitute” for each other, suggesting that they satisfy the same demand. For example, when both cocaine and the short-acting opioid remifentanil were made available under a choice schedule, and the price of one drug was manipulated while holding the price of the other drug constant, monkeys increased their consumption of the fixed-price drug when the cost of the variable-price drug went up, indicating that these drugs substitute for each other as commodities (70). In contrast, in an experiment where monkeys were allowed to choose between responses that delivered ethanol or water to drinking spouts, when the price of both fluids was increased the monkeys maintained their ethanol intake by increasing their responding but did not maintain their water intake; this indicates that the demand for ethanol was less elastic than the demand for water (71).

4. Assessing Treatment Effects

4.1. Interpreting Shifts in the Dose–Effect Curve

Studying a range of doses is essential for evaluating whether a drug is having reinforcing effects. It is equally essential when evaluating whether an experimental treatment is altering the drug’s effects. For example, measuring the effects of a treatment on cocaine self-administration under a single dose of cocaine would not provide a complete picture. If the cocaine dose is near the peak of the dose–effect curve, rates of self-administration might be decreased equally by treatments that increase the effects of cocaine (i.e., make it function like a dose on the ascending limb) or decrease the effects of cocaine (making it function like a dose on the descending limb). Therefore, the effects of treatments that
alter self-administration responding should be analyzed as *shifts in the dose–effect curve*.

If a treatment shifts the dose–effect curve to the left or right but the shape is maintained, this indicates that sensitivity to the drug has changed. If a treatment causes a rightward shift, it decreases sensitivity to the self-administered drug; this kind of effect is typically obtained when a treatment blocks the reinforcing effect of the drug, but the blockade can be overcome by increasing drug intake. A leftward shift indicates that the treatment increases sensitivity to the drug, potentiating its effects. If the curve shifts over time even when no treatments are given, leftward and rightward shifts indicate the development of sensitization or tolerance, respectively, to the self-administered drug.

Upward and downward shifts of the dose–effect curve generally indicate that a treatment alters the efficacy of the drug at producing the effect being measured. A treatment that shifts the self-administration curve upward increases the maximum rate of responding that can be maintained by the self-administered drug. A downward shift or flattening of the curve often occurs when the treatment blocks the reinforcing effects of the drug in a way that cannot be overcome by increasing drug intake. A downward shift may also occur if the treatment does not alter the reinforcing effects of the drug, but somehow interferes with performance of the self-administration response; to evaluate this possibility, separate experiments can be conducted to measure the drug’s effects on spontaneous locomotor activity or food-reinforced responding. It should be noted that for most schedules other than progressive ratio, the drug’s efficacy in maintaining high rates of responding does not necessarily correspond to its reinforcing efficacy. For example, an upward shift of the dose–effect curve might result from tolerance to response-suppressant side effects of the drug.

4.2. Detecting Shifts in the Dose–Effect Curve

Detecting shifts in a dose–effect curve usually requires many sessions of testing. For example, the following would be a thorough approach to testing the effects of a treatment on drug self-administration. The first step is to study several doses of the self-administered drug to establish a preliminary dose–effect curve. Each dose is made available for several sessions until a stable performance develops. This preliminary phase serves several purposes. It provides the animals with experience self-administering the drug under a range of doses, and it allows the experimenter to verify that the range of doses is sufficient to observe the typical inverted-U dose–effect curve. Once this preliminary curve has been established, two more curves are determined by studying each dose of the drug with and without the treatment in each subject. These dose-by-treatment combinations are studied in a mixed order, counterbalanced across subjects. The treatment and no-treatment curves are determined during the same phase of
training – rather than simply comparing a treatment curve to the preliminary curve obtained earlier – to prevent extraneous variables such as sensitization and tolerance from confounding the results. As during determination of the preliminary curve, each condition is studied for several sessions until behavior becomes stable; this allows an assessment of whether a treatment effect is consistent from day to day, takes time to fully develop, or disappears after repeated exposures.

The thorough approach described above is designed to produce the most valid and reliable results. But, it is quite time consuming, and it becomes even more time consuming when expanded to study more than one level of the treatment or to study combinations of treatments. Consequently, some alternative procedures have been developed to streamline the process. One way to quickly obtain dose–effect curves is by using a multi-dose schedule, in which the session is divided into several periods, with a different dose of the drug made available during each period (e.g., (72)). For example, a multiple schedule can be used in which there are five 30-min periods in each session, with a different stimulus presented and a different dose of cocaine made available during each period (e.g., see (73)). Once the animals have been trained with this schedule, a dose–effect curve can be obtained within each session.

Within-session dose–effect curves can also be produced with a variable-dose schedule (74). This procedure takes advantage of the fact that the duration of post-injection pausing is dose-dependent (see Sect. 2.1). In a variable-dose schedule, the dose is varied unpredictably throughout the session, with no signal to indicate which dose will be delivered by the next response. When the post-injection pauses from the session are plotted as a function of dose, the pauses are seen to be longer at higher doses. Like continuous reinforcement, variable-dose schedules are useful for studying changes in drug intake but are not appropriate for measuring reinforcing efficacy. The dose–effect curves obtained with continuous-reinforcement (when the dose is fixed within sessions but varied across sessions) and variable-dose schedules are typically congruous, except at low, nonreinforcing doses. At these doses, responding is infrequent and pauses are extremely long under continuous reinforcement. In contrast, a nonreinforcing dose will produce very short post-injection pauses under a variable-dose schedule because the animal will respond quickly until a higher dose is received.

Cumulative dosing is a within-session strategy that involves studying several doses of a treatment drug instead of several dose of the self-administered drug (e.g., see (75)). In this procedure, the session is divided into periods, and some amount of the treatment drug is injected at the beginning of each period. The cumulative amount of treatment drug affecting the animal
during each period is calculated based on the rate at which it is known to be eliminated from the body. Thus, a low dose is in effect early in the session, and the cumulative dose increases with each successive period. Since cumulative dosing usually entails removing the animal from the chamber and giving it an intraperitoneal injection before each period during test sessions, the animals must first be acclimated to being handled and receiving vehicle injections during training sessions. A disadvantage of cumulative dosing procedures is that low doses are tested early in the session and higher doses later in the session, a potentially problematic confound.

With all of these streamlined procedures, the researcher must weigh the advantages of rapid testing against disadvantages such as the lack of complete counterbalancing and the small amount of exposure to each condition. But, it should be noted that saving time is not the only potential advantage of rapid testing. Streamlined techniques may be less open to interference from extraneous variables such as gradual drifts in the baseline rate of self-administration. Or, the focus of the study may be on changes in the effectiveness of a treatment over time, making it advantageous to obtain a “snapshot” of the dose–effect curve during each daily session.

Animal models of drug abuse are easily integrated with most of the procedures currently used in neuroscience. This multidisciplinary approach has significantly advanced our understanding of the brain mechanisms that underlie reward and addiction. Unfortunately, the very ease and success of this integration can lead to the impression that drug self-administration and other behavioral methods are merely tools that serve other areas of research. But, behavior is also the clinical endpoint, and the value of any treatment for addiction will depend on how it alters what people do. Environmental determinants such as schedules of reinforcement and exposure to drug-related cues are critical to the etiology and progression of addiction, and manipulating these environmental variables can have robust therapeutic effects. For example, one of the most effective clinical means of maintaining abstinence in addicted individuals is through contingency management, a treatment strategy in which nondrug reinforcers are used to reinforce behavior incompatible with drug use (76). Thus, in developing clinical interventions, conducting laboratory research with animal models, and generally working toward a comprehensive theoretical account of addiction, it is important to appreciate that addiction is a behavioral phenomenon that involves a complex interaction between the environment, drugs, and the brain.
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