Synthetic Cathinones and Their Rewarding and Reinforcing Effects in Rodents

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Synthetic cathinones, colloquially referred to as “bath salts,” are derivatives of the psychoactive alkaloid cathinone found in Catha edulis (Khat). Since the mid-to-late 2000s, these amphetamine-like psychostimulants have gained popularity amongst drug users due to their potency, low cost, ease of procurement, and constantly evolving chemical structures. Concomitant with their increased use is the emergence of a growing collection of case reports of bizarre and dangerous behaviors, toxicity to numerous organ systems, and death. However, scientific information regarding the abuse liability of these drugs has been relatively slower to materialize. Recently we have published several studies demonstrating that laboratory rodents will readily self-administer the “first generation” synthetic cathinones methylenedioxypyrovalerone (MDPV) and methylone via the intravenous route, in patterns similar to those of methamphetamine. Under progressive ratio schedules of reinforcement, the rank order of reinforcing efficacy of these compounds is MDPV ≥ methamphetamine > methylone. MDPV and methylone, as well as the “second generation” synthetic cathinones α-pyrrolidinovalerophenone (α-PVP) and 4-methylethcathinone (4-MEC), also dose-dependently increase brain reward function. Collectively, these findings indicate that synthetic cathinones have a high abuse and addiction potential and underscore the need for future assessment of the extent and duration of neurotoxicity induced by these emerging drugs of abuse.

1. The Rise of Synthetic Cathinone Use and Abuse

In 2007, a new class of designer drugs known as synthetic cathinones emerged in Europe. Soon afterwards, reports of synthetic cathinone use, abuse, toxicity, and death began to surface in USA [1–12]. The rise of synthetic cathinone use in USA was alarmingly rapid, with poison control centers receiving 0, 304, and 6,156 calls reporting synthetic cathinone toxicity in the years 2009–2011, respectively [13]. Approximately 98% of synthetic cathinones first identified in toxicological investigations were primarily 4-methylmethcathinone (4-MMC, mephedrone), 3,4-methylenedioxypyrovalerone (MDPV), and 3,4-methylenedioxymethcathinone (methylone) [1–12], but as discussed below, many additional synthetic cathinones have since surfaced.

Synthetic cathinones are chemical derivatives of cathinone, a naturally occurring amphetamine-like alkaloid found in the Catha edulis (Khat) shrub. Khat has been utilized for centuries by indigenous peoples of the Horn of Africa and Arabian Peninsula for its stimulant properties [14]. However, due to its high abuse liability, cathinone is classified as a Schedule I controlled substance in USA. In an attempt by manufacturers and distributors to evade the attention of regulatory and law enforcement agencies, synthetic cathinones are falsely marketed and sold as innocuous retail products such as “bath salts,” “plant food,” “research chemicals,” and “glass cleaner,” to name a few. Synthetic cathinones are also often sold under brand names such as “Ivory Wave” and “Vanilla Sky” and are usually labeled “not for human consumption” or “for research purposes only” [15]. These packages typically consist of one or multiple synthetic cathinones and are often mixed with other substances such as caffeine, topical anesthetics, binding and cutting agents, and even other illicit drugs [11, 16]. Regardless of the marketing tactic employed, synthetic cathinones are...
ultimately intended for use as "legal high" alternatives to illicit psychostimulants such as cocaine, methamphetamine, and 3,4-methylenedioxyamphetamine (MDMA, "Ecstasy") [1–12, 17–20].

For synthetic cathinone users, desired psychological effects include increased energy, libido, empathy, euphoria, alertness, and well-being. However, numerous serious adverse effects are associated with synthetic cathinone use [1–12]. Psychological and behavioral complications include confusion, panic, persistent hallucinations and delusions, agitated paranoia, and aggression and violence including suicide, homicide, and infanticide. Synthetic cathinone use is also associated with toxicity of multiple organ systems, sometimes referred to as the "sympathomimetic toxidrome" [7], including chest pain, nausea, vomiting, seizures, hypertension, tachycardia, hyperthermia, cardiac arrest, and death. Despite the high risk of these adverse effects, users of synthetic cathinones frequently report a persistent desire to continue use of these drugs, and prolonged periods of synthetic cathinone use have been reported [2, 3, 6, 8, 12, 21–27], suggesting a high potential for addiction and dependence.

2. Chemistry and Pharmacology of Synthetic Cathinones

The chemical structure of cathinone is strikingly similar to that of D-amphetamine (Figure 1), with cathinone differing only in the presence of a ketone oxygen atom (C=O) on the β position of the side chain. However, as recently pointed out by Glennon [28], some synthetic cathinones have chemical structures that are entirely novel such that the amphetamine analogs upon which they are based have received little if any scientific study. As mentioned above, during the initial detection of these drugs in USA, approximately 98% of all synthetic cathinones encountered by law enforcement were either mephedrone, MDPV, or methylone (see Figure 1 for chemical structures). Citing an imminent threat to public health, the U.S. Department of Justice temporarily placed mephedrone, MDPV, and methylone into Schedule I status in October, 2011, which was followed by permanent classification of mephedrone and MDPV as such in August 2012 [29]. Methylone remained under temporary scheduling status pending further collection of data and information and was permanently placed into Schedule I status in April 2013 [30].

Synthetic cathinones are considered "designer drugs" since their chemical structure can be easily altered by as few as one or two atoms to create new chemical entities. Due to the ease with which synthetic cathinones can be chemically modified to create unique chemical entities, over 40 other synthetic cathinones have been identified in clandestine drug markets [28], including the "second generation" synthetic cathinones naphyrone (naphthylpyrvalerone), 3-fluoromethcathinone (3-FMC), methedrone (4-methoxymethcathinone), β-keto-N-methylbenzodioxolylbutanamine (bk-MBDB, butylone), β-keto-methylbenzo-dioxolypentanamine (bk-MBDB, butylone), 4-methyl-N-ethylcathinone (4-MEC), 4-methyl-pyrrolidinopropiophenone (4-MePPP), α-pyrrolidinopentaphenone (α-PVP), 2-methylamino-l-phenylpentan-l-one (pentedrone), and 4-fluoro-N-methylcathinone (4-FMC, flephedrone). Many of these were placed into temporary Schedule I status in February 2014 while additional data on abuse liability are collected [31].

Synthetic cathinones exert their psychostimulant and sympathomimetic effects either by promoting the release of monoamine neurotransmitters (dopamine, DA; norepinephrine, NE; and serotonin, 5-HT) via reversal of plasma membrane monoamine transporters or by inhibiting the reuptake of monoamines from the synaptic cleft [32–42]. The resulting excessive monoamine levels in the synaptic cleft lead to overstimulation of postsynaptic DA, NE, and 5-HT receptors in the brain and periphery, which results in their psychological, behavioral, and toxic effects. These neuropharmacological actions of synthetic cathinones are remarkably similar to those of more traditional illicit psychostimulants, such as cocaine (a monoamine reuptake inhibitor with high affinity for DA versus other plasma membrane monoamine transporters), D-amphetamine and methamphetadine (monoamine releasers with higher affinity for DA and NE versus 5-HT), and MDMA (a monoamine releaser with a higher affinity for 5-HT versus other plasma membranemonoaminetransporters). It is becoming increasingly apparent that, like traditional psychostimulants, different synthetic cathinones have diverse mechanisms of action [28]. For example, mephedrone is a broad spectrum monoamine releasing agent [32, 33, 35, 38, 39, 41], MDPV is a long-acting inhibitor of plasma membrane DA and NE transporters [33, 41, 43], and methylone is a monoamine releasing agent with a higher affinity for plasma membrane 5-HT transporters [32, 38, 41]. Although studies on "second generation" synthetic cathinones are few, thus far it appears that these newer analogues have similar neurochemical actions [38, 41, 44, 45].

3. Laboratory Rodents Readily Self-Administer MDPV and Methylene

The intravenous self-administration (IVSA) model is generally considered to be the "gold standard" of preclinical paradigms for assessing the abuse liability of psychoactive compounds. Our laboratory routinely uses this paradigm in rats to assess the potential therapeutic efficacy of pharmacological compounds as potential antiaddiction medications, and we have recently utilized this paradigm to assess the potential abuse liability of synthetic cathinones [46, 47].

In this procedure, first an indwelling intravenous catheter is surgically implanted into the jugular vein while the other end is tunneled under the skin and connected to a vascular access port implanted under the skin and exiting between the scapulae. Following recovery from surgery, the animal is placed in a drug self-administration chamber equipped with two levers that are interfaced with a computer and a syringe pump (see Figure 2). A sterile drug solution (such as MDPV or methylone dissolved in physiological saline) is
Figure 1: Similar chemical structures of cathinone, D-amphetamine, and methamphetamine (top row), the "first generation" synthetic cathinones mephedrone, MDPV, and methylone (middle row), and the "second generation" synthetic cathinones α-PVP and 4-MEC.

Figure 2: The rat intravenous self-administration paradigm for assessment of the abuse liability of synthetic cathinones. Upon pressing of one of two levers (designated the active lever) in an operant self-administration chamber (center), a computer-controlled syringe pump (left) delivers a solution containing MDPV or methylone to an indwelling venous catheter via infusion tubing connected to a liquid swivel. Each drug infusion is accompanied by simultaneous presentation of tone and illumination of a stimulus light located above the lever. Presses on the opposite lever (designated the inactive lever) have no consequences but are recorded to ensure establishment of correct response contingency of drug delivery. The graph on the right represents typical response patterns across daily experimental sessions during the acquisition of drug self-administration.

placed in a syringe and delivered by a computer-controlled syringe pump located outside the apparatus. The syringe is connected to a single-channel liquid swivel, which allows free rotation of the animal while maintaining a continuous flow of the drug solution. In addition to the two response levers, stimulus lights and a speaker provide visual and auditory cues during drug infusions. Presses on one of the levers designated the “active” lever result in a computer-controlled drug infusion and simultaneous brief presentation of auditory and visual cues. Presses on the other “inactive” lever have no programmed consequences at any time during the experiment (as a control for nonspecific behavior). To avoid overdose or toxicity due to multiple drug infusions in close temporal proximity, immediately following each drug infusion a “timeout” period is introduced (~20 sec), whereby additional active lever presses do not result in additional drug infusions. Drug self-administration sessions are typically 2–6 hr in length and are conducted daily, 7 days per week.

There are many advantages of intravenous self-administration procedures as a model for human drug-taking behavior, including the following: (1) the drug is administered voluntarily by the animal (as opposed to passive administration by the experimenter); (2) it is administered directly into the bloodstream mimicking intravenous drug use in humans and results in rapid brain penetrance; (3) drug-taking behavior can be temporally examined within and between self-administration sessions; (4) the effects of candidate therapeutic pharmacological compounds or other experimental manipulations on drug self-administration can be determined; and (5) the number of responses that must
be exerted by the animal in order to receive a drug infusion (called the “ratio”) can be increased exponentially (called a “progressive ratio” procedure) until the animal “gives up” and no longer performs the operant task (called the “breakpoint”). This latter method is used to measure the level of motivation to self-administer the drug as well as the efficacy of the reinforcer. In addition, when given longer daily access (i.e., 6 hr or more per session) to psychostimulants with high addiction potential, rats will gradually and significantly escalate their daily drug intake. This escalated drug intake is a cardinal feature of the transition from drug abuse to addiction, and a sustained enhancement in drug intake produces dysregulated brain stress and executive control systems that parallel clinical observations of drug addicts [48–50]. Moreover, extended access to drugs with lower abuse liability generally do not result in escalation of intake, and drug intake patterns usually remain episodic [51].

Recently, our laboratory has demonstrated robust intravenous self-administration of MDPV and methylone in laboratory rats [46, 47]. When tested for self-administration of MDPV under limited access conditions (daily sessions that were 2 hr in length), rats readily acquired self-administration of MDPV at all doses tested (0.05, 0.1, and 0.2 mg/kg/infusion). Under a progressive ratio schedule of reinforcement, which requires an exponentially increasing number of lever presses to obtain each subsequent drug infusion within a single test session (i.e., 1 lever press for the first infusion, 2 for the second, 4 for the third, and so on), a positive relationship between MDPV dose and breakpoints for drug reinforcement was observed, suggesting that higher doses of MDPV are associated with increasing reinforcer efficacy and motivation to obtain the drug. By comparison, responding for a low dose of methamphetamine (0.05 mg/kg/infusion) produced breakpoints that were similar in magnitude to the same dose of MDPV. When the length of the self-administration session was increased to 6 hr/day, we observed an escalation of drug intake over time for the 0.1 and 0.2 mg/kg/infusion doses of MDPV, with the highest dose producing the most robust escalation of intake that was similar in magnitude to the 0.05 mg/kg/infusion dose of methamphetamine. When rats were tested for self-administration of methylene under limited access conditions [46], rats did not display robust self-administration for the lowest dose of methylene tested (0.05 mg/kg/infusion) but readily self-administered higher doses of methylene (0.1, 0.2, and 0.5 mg/kg/infusion). However, unlike MDPV, methylene did not lead to escalation of intake under extended access conditions for any dose tested. This is particularly intriguing since methylene is a β-ketone derivative of MDMA, and self-administration of this parent drug also does not reliably produce escalation of intake under extended access conditions [52–54], which is perhaps also reflective of its preferential affinity for plasma membrane serotonin versus dopamine transporters. Despite the lack of escalation, and underscoring the dangers of synthetic cathinone use, 2 rats in the highest methylene dose group (0.5 mg/kg/infusion) self-administered the drug to the point of seizure and death during a 6 hr session, and similar case reports of methylene-induced deaths in humans have been reported [55–58]. Finally, as with MDPV, we observed a positive relationship between methylene dose and breakpoints for self-administration under progressive ratio conditions. These findings indicate that MDPV and methylene are readily self-administered intravenously by laboratory rats and are in agreement with studies from other laboratories showing that MDPV, as well as the synthetic cathinone mephedrone, is self-administered intravenously by rodents [35, 59–61]. In addition, our studies indicate that escalated intake of synthetic cathinones, a cardinal feature of addiction, can also be observed in laboratory rodents.

4. Synthetic Cathinones Increase Brain Reward Circuit Function

It is well accepted that drugs of abuse exert their rewarding (euphorogenic) and reinforcing effects via interactions with brain reward circuitry [62–65]. This circuitry, known as the mesocorticolimbic pathway, is primarily comprised of dopaminergic neurons in the ventral tegmental area (VTA) that project rostrally to form the medial forebrain bundle (MFB) in the lateral hypothalamus and form dopaminergic synapses in forebrain regions including the nucleus accumbens (NAC) and prefrontal cortex (PFC). However, confinement of the neural basis of drug reward and reinforcement to this singular dopaminergic pathway is overly simplistic, as many studies have indicated that other ascending monoaminergic fibers that form the MFB, such as noradrenergic neurons from the locus coeruleus and serotoninergic neurons from the raphe nuclei, also contribute to brain reward function [64, 66–69].

A widely used method for assessing functional activity of the brain reward circuitry in behaving animals is the intracranial self-stimulation (ICSS) paradigm [68, 70–73]. In this procedure, which is depicted in Figure 3, a laboratory animal (typically a rat or mouse) performs an operant response such as nose poke, lever press, or rotation of a wheel manipulandum in order to receive a short pulse of electrical current via a chronically implanted electrode into a specific brain region, typically the MFB. The implanted electrode is connected to a computer-controlled current generator via an electrical commutator. The animal quickly learns that the operant response reliably leads to electrical stimulation of the reward circuitry that presumably produces subjective pleasurable effects, and in our experience rats will exert up to several thousand ICSS responses in a 30 min period [46, 47]. Different laboratories use a variety of stimulation parameters such as varying the intensity of the current delivered (μA), duration of pulses (msec), frequency of electrical pulse as a function of pulse/interpulse interval, and the waveform of pulses [74]. However, the two most extensively used ICSS paradigms in evaluating abuse liability of psychoactive compounds are the rate-frequency curve-shift procedure [75] and the discrete-trial current threshold intensity procedure [74, 76, 77], the latter of which is utilized by our laboratory [46, 47, 51].

In the discrete-trials current threshold procedure, following acquisition of operant responding, discrete-trial training
begins where each trial is initiated with a “free” (nonresponse contingent) stimulation, followed by an intertrial interval (ITI, average 7.5 sec) where responses yield no programmed consequences. After the ITI, an operant response yields stimulation identical to the free stimulation. Once animals learn to inhibit responding during the ITI, baseline current intensity threshold training begins. These procedures begin at a predetermined level of current intensity (i.e., 120 μA). Trials are conducted in blocks (3 to 5 trials each), consisting of a free stimulation, ITI, and response-contingent stimulation and current intensity that remain the same for the entire block. Correct responding on the majority of trials results in a lowering of the current intensity by a fixed increment (e.g., 5 μA) for the next block of trials, whereas failure to correctly respond to most trials results in an increase in the current intensity by the same increment for the next trial block. This stimulation titration procedure progresses through 2 ascending and 2 descending series of current intensities. Across these 2 ascending and descending series, ICSS thresholds are calculated as the mean (in μA) across blocks that do or do not support responding. Thus, ICSS thresholds represent the minimum current intensity at which the animal will reliably respond for electrical stimulation of the MFB. Following stabilization of ICSS thresholds across several days of testing, a dose of a candidate drug of abuse (such as a synthetic cathinone) is administered prior to a test session. In our laboratory, we assess changes in current intensity thresholds across a range of doses of each drug in a semirandom counterbalanced design, with multiple determinations at each dose to provide increased reliability of changes in current intensity thresholds (see [46, 47, 51] for details). Resulting decreases in ICSS thresholds (relative to baseline or saline administration) are indicative of hedonic rewarding effects of the drug (i.e., less current is required to activate the reward circuitry due to prior activation by the drug administered), whereas elevations in ICSS thresholds are indicative of aversive or dysphoric effects (i.e., more current is required to activate the reward circuitry due to the aversive nature of the drug administered). ICSS procedures have been used for over 50 years and have consistently revealed that nearly all drugs that are abused by humans, including psychostimulants, lower ICSS thresholds [51, 71, 74–76].

We have recently assessed the ability of acute systemic administration of the “first generation” synthetic cathinones MDPV [47] and methylone [46] to alter ICSS thresholds in laboratory rats. For MDPV, we found that all doses tested (0.1, 0.5, 1, and 2 mg/kg) produced significant reductions in ICSS thresholds (~20–40%) compared with those following saline administration, with the most robust effects observed after the 1 mg/kg dose. We observed far less robust effects following administration of methylone, with doses of 0.1 and 0.5 mg/kg producing no effects on ICSS thresholds, whereas higher doses (1, 3, 5, and 10 mg/kg) produced reductions in ICSS thresholds ranging from ~5 to 15% in parallel with increasing doses. These less robust effects of methylone versus MDPV on ICSS thresholds indicate that methylone may be less potent in activating brain reward circuit function, which could be attributable to its higher affinity for serotonin versus dopamine transporters [32, 38] and is likely related to the lack of escalated intake in extended access during self-administration. This reduced activation of brain reward circuitry by methylone, as compared to classical psychostimulants such as methamphetamine, has also been observed following administration of MDMA [78], the amphetamine analog of methylone, which also possesses a similar 5-HT/DA transporter affinity ratio [32].

We have also recently conducted a study on the potential ICSS threshold-lowering effects of the “second generation” synthetic cathinones α-PVP and 4-MEC [79]. When rats were administered α-PVP (0.1, 0.3, 1, or 5 mg/kg), significant
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reductions (∼14–20%) in ICSS thresholds were observed following the 0.3 and 1 mg/kg doses. When rats were administered 4-MEC (0.3, 1, 3, 10, 30, or 100 mg/kg), significant reductions (∼10–15%) in ICSS thresholds were observed following the 10 and 30 mg/kg doses. However, for both drugs, the highest doses tested produced increases (∼19% for α-PVP at 5 mg/kg and ∼28% for 4-MEC at 100 mg/kg) in ICSS thresholds, suggesting aversive effects of these high doses. Our ICSS studies confirm reports by other laboratories that synthetic cathinones activate brain reward circuitry [80, 81].

5. Conclusions and Future Directions

We have demonstrated that laboratory rats will voluntarily self-administer the “first generation” synthetic cathinones MDPV and methylone via the intravenous route. Furthermore, MDPV, but not methylene, leads to escalation of intake following extended access to the drug. We also have demonstrated that MDPV and methylene, as well as the “second generation” synthetic cathinones α-PVP and 4-MEC, lower current intensity thresholds for ICSS, indicating activation of brain reward circuitry. Along with existing case reports and preliminary epidemiological studies in humans [1–12], these findings clearly indicate that synthetic cathinones possess a significant abuse liability and potential for addiction and should be considered an emerging class of abused drugs that warrant appropriate legislative control and the development of interventions for detoxification and treatment.

However, numerous questions regarding the effects of synthetic cathinones on the brain still remain to be answered. For example, do synthetic cathinones possess affinity for any molecular entities other than monoamine transporters which might contribute to their neurobiological or behavioral effects? What are the mechanisms underlying the persistent psychotomimetic effects of synthetic cathinones? What is the feasibility of utilizing cognitive-behavioral, pharmacological, or other approaches for treating dependence on synthetic cathinones? What are the lasting effects of synthetic cathinones on neuronal plasticity and function, gliotransmission, cerebrovascular function, cell viability, gene expression, and epigenetic processes? Answers to this latter question are being actively pursued by our laboratory. Specifically, we are currently conducting studies on the effects of synthetic cathinone self-administration on macrostructural changes and neurotoxicity in the brain, with the ultimate goal of identifying potential mechanisms and treatment avenues for reversing lasting deleterious effects of synthetic cathinones in the brain. The field of synthetic cathinone research is only in its infancy, and while initial legislative efforts have attempted to curb the availability and use of these drugs of abuse, it is clear that many more “designer drugs” of this drug class (and others) will continue to evolve numerous steps ahead of policymakers, scientists, educators, and treatment professionals alike.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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References

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(4-MEC), two synthetic cathinones commonly found in "second-generation" bath salts,” Submitted to International Journal of Neuropsychopharmacology.

