When good times go bad: managing ‘legal high’ complications in the emergency department

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Abstract: Patients can use numerous drugs that exist outside of existing regulatory statutes in order to get “legal highs.” Legal psychoactive substances represent a challenge to the emergency medicine physician due to the sheer number of available agents, their multiple toxidromes and presentations, their escaping traditional methods of analysis, and the reluctance of patients to divulge their use of these agents. This paper endeavors to cover a wide variety of “legal highs,” or uncontrolled psychoactive substances that may have abuse potential and may result in serious toxicity. These agents include not only some novel psychoactive substances aka “designer drugs,” but also a wide variety of over-the-counter medications, herbal supplements, and even a household culinary spice. The care of patients in the emergency department who have used “legal high” substances is challenging. Patients may misunderstand the substance they have been exposed to, there are rarely any readily available laboratory confirmatory tests for these substances, and the exact substances being abused may change on a near-daily basis. This review will attempt to group legal agents into expected toxidromes and discuss associated common clinical manifestations and management. A focus on aggressive symptom-based supportive care as well as management of end-organ dysfunction is the mainstay of treatment for these patients in the emergency department.

Keywords: legal highs, novel psychoactive substances, toxicology, opioid toxidrome, anticholinergic toxidrome, sympathomimetic toxidrome, hallucinogens, inhalants

Introduction

Much of the recent discussion on “legal highs” has focused on the so-called novel psychoactive substances (NPS), an ever-expanding group of chemicals known as designer drugs that has flooded the drug market in the past decade.¹ These chemicals are creatively marketed and continuously re-designed to stay one step ahead of current regulatory legislation.¹ However, the misuse of legal substances to produce hallucinogenic, stimulant, euphoric, or depressant effects akin to controlled substances has occurred since the earliest efforts at drug regulation. After the 1925 Geneva Opium Convention attempted to establish international control over morphine, drug manufacturers started flooding the market with uncontrolled morphine chemical derivatives.² Furthermore, people seeking psychoactive effects akin to illicit substances have long turned to not only novel designer chemicals but also household products such as propylhexedrine-based inhalers, dextromethorphan (DXM)-based over-the-counter remedies, and nutmeg, to name only a few.

With this expanded definition in mind, this paper endeavors to cover a wide variety of “legal highs,” or uncontrolled psychoactive substances that may have abuse potential
often occur before the onset of their hallucinogenic effects.7 Peripheral effects, including tachycardia and dry mouth, which are rare in the setting of isolated ingestion. An analysis of 1458 cases reported to American poison centers revealed no attributable fatalities.8 Another series of 8 years of poison center data only found 5 reported fatalities attributable to Datura ingestions may present with prolonged toxic effects lasting hours to days, given the half-lives of 2–3 hours for atropine and up to 9–10 hours for scopolamine combined with decreased gastrointestinal (GI) motility.7

In addition to supportive care, management of belladonna alkaloid plant toxicity centers around treating agitation as well as reversing severe anticholinergic effects if needed. Decontamination with activated charcoal is controversial, as most patients will present for medical care when clinically intoxicated and agitated.7 For cases of minor toxicity, benzodiazepines are one mainstay of supportive care.7 For severe agitation, the cholinesterase inhibitor physostigmine is effective in reversing isolated anticholinergic toxicity.7 Due to the risk of triggering a cholinergic crisis, including bradycardia, respiratory distress and seizures, physostigmine should be used with caution, and only with a reversal agent such as atropine on hand.7 The starting dose of physostigmine is 0.5–2 mg in adults infused intravenously no faster than 1 mg/minute, with redosing occasionally necessary because of the drug’s relatively short half-life of 15–40 minutes.7

Diphenhydramine Diphenhydramine (Benadryl18, Johnson & Johnson, New Brunswick, NJ, USA) is a histamine blocker with central effects as well as anticholinergic and sodium channel blocking properties, found in many over-the-counter medications ranging from sleep aids to allergy and cold medications.3 Although thought to have limited abuse potential due to its overly sedating qualities, diphenhydramine is, in fact, commonly abused for its euphoric and hallucinogenic effects at higher dosages.3,9,10 Diphenhydramine is commonly co-formulated with acetaminophen, and in cases of toxic overdoses, the delayed GI motility from diphenhydramine can contribute to delayed and persistent toxic levels of acetaminophen.11

The classic signs of diphenhydramine overdose are an anticholinergic toxidrome combined with profound sedation and at times marked agitation.12 Diphenhydramine has also been known to produce rhabdomyolysis.13 Also, in comparison with other anticholinergics, diphenhydramine is much more likely to cause seizures in acute overdose.14 Diphenhydramine is chemically similar to the tricyclic antidepressants and has sodium channel blockade effects resulting in seizures and wide complex tachyarrhythmias in higher dosages.15 Management of diphenhydramine toxicity centers around control of agitation and hyperthermia, as well as treating rhabdomyolysis, potential tachyarrhythmias, and seizures.16 Although physostigmine has a role in reversing anticholinergic effects, it is contraindicated in cardiotoxic diphenhydramine overdoses due to the risk of asystole, and should not be used if the QRS >100 ms in the setting of a diphenhydramine overdose.17 In case of QRS widening or
tachyarrhythmias, sodium bicarbonate has been used successfully.\textsuperscript{15} Massive overdoses due to diphenhydramine with refractory tachyarrhythmias have been treated successfully with intralipid therapy.\textsuperscript{18}

**Nutmeg**

Widely available, nutmeg is known to provoke hallucinogenic effects in high dosages. According to available case reports, nutmeg reached a peak of abuse in the 1960s and 1970s.\textsuperscript{19} It is well documented as a drug of abuse among prisoners, with Malcolm X famously detailing in his autobiography during his incarceration that a "a penny matchbox full of nutmeg had the kick of 3 or 4 reefer."\textsuperscript{20} Most likely due to unpleasant, dysphoric side effects as well as need to consume a large unpalatable amount, nutmeg abuse is not common among the general population.\textsuperscript{21–23}

Nutmeg’s intoxicating effects are thought to be due to myristicin, the largest chemical portion of the volatile oil of nutmeg.\textsuperscript{24} Myristicin acts as a monoamine oxidase inhibitor (MAOI), increasing the amount of serotonin in the brain.\textsuperscript{25} It has also been found in animal models to be metabolized into 3-methoxy-4,5-methylene dioxyamphetamine, which is a sympathomimetic with hallucinogenic and euphoric effects.\textsuperscript{26}

Nutmeg produces a biological effect similar to an anticholinergic toxidrome, with altered mental status, skin flushing, dry mucus membranes, tachycardia, and hypertension.\textsuperscript{24} A feature to distinguish nutmeg toxicity from an anticholinergic agent is miosis, although a large amount of nutmeg intoxications with mydriatic pupils have been noted in 1 case series.\textsuperscript{27} Treatment for nutmeg intoxications is supportive in nature.\textsuperscript{24} One case series reported a provoked seizure in the setting of nutmeg intoxication in a person with a seizure disorder.\textsuperscript{23} Only 2 fatalities are known in the literature, with both cases being notable for myriad confounding variables.\textsuperscript{22} One regional survey found that nutmeg was commonly associated with concurrent drug exposures.\textsuperscript{23}

**A. muscaria**

\textit{A. muscaria} aka “fly agaric” is a mushroom that grows near pine trees throughout the northern hemisphere, identifiable by its bright red caps with white spots (Figure 1).\textsuperscript{28} Legal to possess and consumed for its hallucinogenic properties, the main psychoactive components of fly agaric are ibotenic acid, an analog of the excitatory neurotransmitter glutamate, as well as muscimol, an analog of the inhibitory neurotransmitter gamma-Aminobutyric acid (GABA).\textsuperscript{29} These contradictory chemicals are thought to result in a subjective experience similar to alcohol intoxication due to GABA-like effects, but with hallucinations due to its concurrent glutamate-like effects.\textsuperscript{30}

The clinical toxidrome of fly agaric has been said to mimic that of an anticholinergic toxidrome, with flushing, fever, and pupillary dilation.\textsuperscript{28,31} Vomiting and diarrhea have also been reported,\textsuperscript{31} although some sources report this to be an infrequent feature of fly agaric intoxication.\textsuperscript{28} Termed “pantherine-muscarine” poisoning, the neurologic effects of fly agaric mimic its concurrent GABA and glutamate analog components, with sometimes rapid alternation between profound sedation and severe agitation.\textsuperscript{31}

Treatment is supportive, with consideration of GI decontamination with charcoal.\textsuperscript{28} Benzodiazepines can be used for adverse excitatory effects such as seizures, but with caution, given potential rapid switching to muscimol-mediated sedation.\textsuperscript{28} Effects can last up to 24 hours, and fatalities are rare.\textsuperscript{28} Also, concurrent ingestions should be considered, including accidental ingestion of the more hepatotoxic cyclopeptide-containing mushrooms such as \textit{Amanita phalloides}.\textsuperscript{30}

**Opioid legal highs**

As the USA grapples with unprecedented levels of opioid abuse, there has been an increased push among prescribers
and regulatory bodies to curb long-term prescribing of these medications.\textsuperscript{32} Opioid-dependent patients seeking to reduce prescription opioid use, or those who have had their prescription sources curtailed by medical providers, have increasingly turned to legally available opioid substitutes to provide euphoric effects as well as to mitigate withdrawal symptoms.\textsuperscript{33} Legal substances that have emerged as opioid substitutes include loperamide, kratom, poppy seed tea (PST), and a number of synthetic designer opioids.

Loperamide
Loperamide (Imodium®, Johnson & Johnson, New Brunswick, NJ, USA) is a peripherally acting opioid derivative used as an over-the-counter antidiarrheal since 1988.\textsuperscript{34} Long considered a drug with low abuse potential, its central penetration is thought to be limited by overall poor bioavailability as well as the action of P-glycoproteins that actively secrete the drug from the central nervous system.\textsuperscript{35} After many years of only scattered reports of toxicity and abuse, loperamide has recently emerged as a drug of abuse in the USA, largely going hand-in-hand with the opioid epidemic.\textsuperscript{35} Known as the “desperate junky’s methadone,” it is mainly abused to alleviate withdrawal symptoms, and by some to provoke an opioid-based euphoria.\textsuperscript{36} Abusers of loperamide typically take 3–4 times the recommended daily dose in 1 sitting.\textsuperscript{35} Some also take advantage of cytochrome inhibitors such as cimetidine and grapefruit juice as well as P-glycoprotein inhibitors such as quinidine and pepper to raise serum levels of the drug.\textsuperscript{35}

Loperamide’s main toxic effects include not only the respiratory and central nervous system depression typical of opioid overdoses, but also cardiotoxicity in the form of life threatening cardiac arrhythmias.\textsuperscript{35} Loperamide in high doses has been found to have sodium and potassium channel blocking effects, resulting in profound cardiotoxic effects.\textsuperscript{37} It has been associated with widening of the QRS complex, prolonged QTc interval, as well as ventricular monomorphic and polymorphic tachycardias resulting in multiple fatalities.\textsuperscript{37,38} Syncope or ventricular dysrhythmias and electrocardiogram abnormalities in a patient suspected of opioid abuse should trigger consideration of loperamide toxicity.\textsuperscript{37} Loperamide will not show up on a standard urine drug screen.\textsuperscript{35}

Management of loperamide toxicity includes extended consideration of decontamination, treatment of respiratory depression, and monitoring and treatment of potential cardio toxicity.\textsuperscript{37} Charcoal-based GI decontamination has been advised up to 2–4 hours after a large overdose in the absence of contraindications, due to the drug’s GI motility slowing effects.\textsuperscript{37} Naloxone has been used for loperamide-provoked respiratory depression, with one pediatric case needing multiple doses over a 24-hour period, presumably due to the drug’s slow elimination times.\textsuperscript{35} Widened QRS complexes due to loperamide may be reasonably treated with sodium bicarbonate, while long QT intervals should be treated with correction of electrolyte derangements.\textsuperscript{35} It is reasonable to treat loperamide-induced torsades with standard agents, including magnesium, defibrillation, isoproterenol, and electrical pacing.\textsuperscript{37}

Poppy seed tea
Although opium and the poppy plant itself are controlled in the USA, poppy seeds are completely unregulated and can be bought in bulk. Although they do not contain any psychoactive opiates, poppy seeds have been shown to have trace amounts of opium latex containing morphine and codeine on their surface.\textsuperscript{39} This fact has been exploited by those seeking a legal and readily accessible source of opiates by making poppy seed tea (PST).\textsuperscript{40} PST synthesis involves steeping a bulk supply of poppy seeds in water, thus diluting a variable amount of codeine and morphine into a liquid that is then ingested.\textsuperscript{40} PST abuse emerged among opiate-dependent patients in New Zealand after a government crackdown on heroin supply to the country.\textsuperscript{40} The internet provides both a means of purchasing bulk poppy seeds as well as a source of information on PST use and instructions for preparation.\textsuperscript{40} The main limitation on PST use among some users was its foul taste.\textsuperscript{40} Users have reported an onset of 15 minutes and sometimes prolonged effects lasting up to 24 hours.\textsuperscript{40}

Although official case reports of fatalities are sparse, scattered fatalities in the popular media have been reported among young people using PST.\textsuperscript{41} Cases of physiologic dependence on poppy seed concoctions have also been reported.\textsuperscript{42} An issue with the synthesis of PST is the highly variable opiate content found among different seed sources, ranging from 1–2 orders of magnitude for morphine and codeine content based on seed source.\textsuperscript{39} This may make it very difficult for those synthesizing and ingesting opiates by this method to reliably ascertain the morphine and codeine content of their PST, theoretically leading to the potential for overdose.

Kratom
Kratom is the common name for the plant Mitragyna speciosa, a large leafy tree which is indigenous to Southeast Asia, containing psychoactive alkaloids with dose-dependent stimulatory and opioid-like effects.\textsuperscript{23} By way of direct chewing of leaves or through the formation of teas and other mixtures, kratom has been traditionally used throughout Southeast Asia as an antidi-
arrheal, a painkiller, and a cheap opium substitute. Kratom has emerged in the USA as an easily available and legal opioid substitute, available through internet suppliers as well as brick-and-mortar “smoke shops,” with the number of calls to poison centers regarding kratom increasing from 26 in 2010 to 263 in 2015. In late 2016, the Drug Enforcement Administration (DEA) attempted to classify kratom and its main psychoactive alkaloids mitragynine and 7-hydroxymitragynine as Schedule I substances, a move that was met with such fierce public backlash that the agency backed down from their scheduling decision pending further review in October 2016.

Kratom’s main psychoactive alkaloids are mitragynine and 7-hydroxymitragynine, substances that have been shown to have activity at supraspinal μ-opioid receptors, as well as possible stimulant activity through α2 receptors and 5-HT2A receptors. Kratom users commonly report a dose-dependent response with stimulant effects in small doses up to opioid analgesic effects in higher doses. Other subjective effects include euphoria, relaxation, increased energy, with some negative effects, including nausea, vomiting, stomach ache, and constipation.

The safety profile of kratom has not been fully evaluated. Well-controlled data on the toxic effects of kratom on human subjects are lacking, and case reports of toxicity are relatively rare. There have been scattered case reports of intractable seizures that are refractory to benzodiazepines in patients with acute kratom ingestions, although many cases are associated with co-ingestion of other prescription and illicit drugs.

Regarding their tendency to produce opioid toxicity, animal models have generally found less respiratory depression with mitragynine compared with other opioid receptor agonists, and there has not yet been a case report of kratom inducing an opioid-based hypoventilatory state in which naloxone has been used. However, it would be reasonable to give naloxone in the setting of a kratom ingestion with respiratory depression. Kratom has been frequently adulterated with other opioid receptor agonists such as O-desmethyltramadol, resulting in 9 deaths in 1 case series. Long-term use of kratom has been associated with intrahepatic cholestasis and other liver disease, along with hypothyroidism. Patients can also develop physical dependence with withdrawal symptoms that mimic that of opioids, with symptoms such as abdominal pain, sweating, nausea, vomiting, and diarrhea. There have been reports of fatalities with high levels of mitragynine, with kratom presumably as the cause of death, but these reports are also notable for many other co-ingestions such as propylhexedrine, benzodiazepines, and selective serotonin reuptake inhibitors (SSRIs).

**Synthetic opioids**

In tandem with the opioid epidemic, synthetic designer opioids such as AH-7921, U-47700, MT-45, and numerous fentanyl (Sublimaze®, Janssen Pharmaceutica, Beerse, Belgium) analogs have emerged onto the drug market in the past few years. In an effort to stem the flood of these agents, the DEA has continuously added synthetic opioids to Schedule I, including U-47700, AH-7921, and many fentanyl analogs. Still unscheduled, an opioid known as MT-45, a piperazine derivative, has found abuse as a legally available opioid. MT-45 overdose produces an opioid toxidrome, and acute ingestions resulting in respiratory depression have been successfully reversed with naloxone. Users of MT-45 find it to have less opioid-euphoric effects at a relatively higher financial cost and with more adverse effects, likely putting it to have less opioid-euphoric effects at a relatively higher financial cost and with more adverse effects, likely putting it to habitual MT-45 abuse include hair loss, hearing loss, dermatitis, and cataracts.

A more substantial public health threat than MT-45 is the re-emergence of fentanyl and a wide range of fentanyl analogs to the illicit drug market. The abuse of fentanyl and novel fentanyl analogs dates to the 1970s, marketed in pure form such as 3-methylfentanyl being sold as “China White,” but also as surreptitious additions to traditional illicit drugs such as heroin and cocaine. Periodic epidemics of fentanyl abuse has resulted in sporadic outbursts of morbidity and mortality over the past few decades. In 2016 alone, the DEA saw the use of 13 different fentanyl analogs on the illicit drug market, some of which are unscheduled. Fentanyl analogs vary widely in potency compared with morphine, ranging from acetylfentanyl, which is 15.7 times as potent, up to 3-methylfentanyl, which may be 569 times as potent. A major consideration for the emergency medicine provider is that overdoses of fentanyl and the more potent fentanyl analogs may require substantially more naloxone than that of heroin for their reversal, up to 12 mg in a case series from Illinois.

**Sympathomimetic legal highs**

There are many “legal highs” that result in a sympathomimetic toxidrome. Many of them are abused by those seeking similar effects as the classic illicit psychostimulants such as cocaine, methamphetamine, and 3,4-methylenedioxy-N-methylamphetamine (MDMA). Others are abused for their serotonin-mediated hallucinogenic effects, but are notable for producing sympathomimetic toxicity. Many of these agents will share similarities in their treatment approach, which entails benzodiazepines, active cooling for hyperthermia, and...
specific management of end-organ complications related to hypertension and vasoconstrictive effects.

**Synthetic cathinones**

A Schedule I substance, cathinone is a monoamine amphetamine analog with stimulatory properties found in the shrub *Catha edulis*, commonly known as khat.56 Khat is endemic to the Horn of Africa as well as the Southeast Arabian peninsula, where it is only chewed fresh for its stimulatory effects due to cathinone’s quick degradation.56 Since cathinone’s discovery, many chemical derivatives have been synthesized and researched for possible therapeutic effects, the most successful of which is the antidepressant bupropion (Wellbutrin®, GlaxoSmithKline, Brentford, UK) (m-chloro-N-tert-butyl-cathinone).56

In the past decade, many synthetic cathinones have been developed and exploited for their strong stimulant effects as “not for human consumption” legal highs, marketed as “bath salts” or “plant food.”56 When consumed, these substances have variable stimulatory effects comparable with methamphetamine or MDMA.57 Cathinone and mephedrone do share chemical structural similarities with methamphetamine (Figure 2). Synthetic cathinones reached a peak of 6137 reports in 2012, declining to 382 in 2016,58 likely due to federal and state efforts at scheduling and regulating these psychoactive substances.59 Following a pattern of other NPS, new agents continue to be synthesized and marketed outside federal control according to the DEA’s Emerging Threat Report.54

The toxicologic effects of synthetic cathinones mimic those of the amphetamines as well as MDMA, and treatment is primarily supportive around managing the agitation, fever, hyponatremia, and end-organ complications that can sometimes occur with these agents.57 Full evaluation of the toxicologic effects of cathinones is complicated by the breadth and emergence of new agents, as well as the common presence of co-ingestions.57 Notably, the main adverse effects include both cardiovascular (hypertension, tachycardia, angina, myocarditis) and neuropsychiatric (agitation, aggression, dystonia, and hyperreflexia) complications.57 There have been a number of case reports of hyponatremia in the context of synthetic cathinone use should be managed much like MDMA-related hyponatremia, with fluid restriction for mild cases up to the use of hyperosmotic agents for cerebral edema.57 Other noted laboratory abnormalities include metabolic acidosis, elevated creatinine kinase, elevated liver enzymes, and acute renal failure.57

**Benzofurans**

Commonly known as “Benzo Fury,” the benzofurans refer to the phenethylamine MDA (3,4-methylenedioxymethamphetamine) derivatives 5-(2-aminopropyl)benzofuran (5-APB) and 6-(2-am-inopropyl)benzofuran (6-APB), currently unscheduled chemicals abused for their stimulant as well as entactogenic effects similar to the Schedule I substance MDMA.60 Benzofurans have been found to inhibit reuptake of serotonin, dopamine, andnorepinephrine, and they also have affinity for adrenergic and 5-HT receptors.60

The subjective effects of benzofurans have been described as much like MDMA, but with more adverse effects in the form of nausea, dry mouth, dry eyes, insomnia, diarrhea, palpitations, headache, and various adverse psychological symptoms, including visual and auditory hallucinations.60 The benzofurans have also been shown to have extended persistence of clinical effects, lasting for more than 48 hours in over half of reported cases.60

Like many of the NPS, detailed toxicological information on the benzofurans are lacking, and acute ingestions are often concurrent with other intoxicants.61 Case reports point to dramatic sympathomimetic symptoms, including tachycardia, hypertension, mydriasis, hyperthermia, agitation, and tremors after ingestion of benzofurans.60 There have also been reports of myocardial ischemia, indicating perhaps more profound vasoconstrictive effects than MDMA.60 These substances have been implicated in more than 10 deaths.50 Management involves supportive care for sympathomimetic toxicity with benzodiazepines, as well as assessment for hypertensive emergency and rhabdomyolysis.

**Benzodifurans: Bromo-DragonFLY, 2C-B-FLY, 3C-B-FLY**

The benzodifurans, commonly known as the Bromo-dragonFLY compounds, refer to a relatively novel group of hallucinogenic drugs that saw escalating amounts of abuse as legally available research chemicals over the past decade.53 They exert their potent hallucinogenic effects through 5-HT2A agonism and serotonergic release.64 Its first member and archetype is Bromo-dragonFLY, an amphetamine derivative in which the phenyl group is buttressed by 2 dihydrofuran

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**Figure 2** Chemical similarity of amphetamine, cathinone, and mephedrone; note the addition of the ketone functional group to the latter two compounds.
rings that give the compound its characteristic “dragonfly” shape as well as purported greater potency and duration of action than many other hallucinogenic phenethylamines (Figure 3). Its subjective effects have been described as akin to lysergic acid diethylamide (LSD), although onset may be delayed up to 6 hours, and effects may last up to 3 days.

Case reports indicate that Bromo-dragonFL Y compounds have potent sympathomimetic and vasoconstrictive toxicity. Clinical presentations of Bromo-dragonFL Y ingestions seeking medical care are notable for sympathomimetic toxidromes in the form of tachycardia, hypertension, mydriasis, diaphoresis, agitation, and seizures. Seizure onset can be delayed, up to 8 hours after ingestion in one case. There have also been cases of extended and refractory vasoconstriction lasting days after overdoses, resulting in auto-amputation of digits, renal failure, and rhabdomyolysis. Deaths have been attributed to the drug, including fatalities caused by massive overdoses in Oklahoma that occurred after Bromo-dragonFL Y was mistaken for the less potent derivative compound 2C-B-FL Y.

Bromo-dragonFL Y overdose treatment is supportive. Sympathomimetic effects and seizures have been treated successfully with large and repeated doses of benzodiazepines. The drug’s purported severe vasoconstrictive effects have proven refractory to a wide variety of vasodilatory therapies in available case reports.

Propylhexedrine

A longstanding “legal high,” abused due to its relatively low cost and widespread availability, propylhexedrine (Benzedrex®, B.F. Ascher & Co., Inc., Lenexa, KS, USA) is a cyclohexylamine nasal decongestant that is chemically similar to methamphetamine, lacking the aromatic ring (Figure 4). Delivered via an over-the-counter nasal inhaler, propylhexedrine’s mechanism as a nasal decongestant is through alpha adrenergic stimulation, resulting in localized vasoconstriction of the nasal mucosa. In higher doses, it also causes release of dopamine, norepinephrine, and serotonin from presynaptic vesicles much like the amphetamines. Since its introduction as a safer replacement of previous amphetamine-based inhalers, propylhexedrine has emerged as a legal substitute for methamphetamine. It is commonly abused by dismantling the inhaler and either ingesting the drug-soaked cotton rods whole, or by soaking the rod in lemon water and injecting or consuming the liquid. Its toxic effects reflect the fact that it is, compared with amphetamine, relatively more effective at promoting peripheral adrenergic stimulation rather than dopaminergic and serotonergic release.

Toxicity of propylhexedrine mimics that of the amphetamines, but with seemingly more profound end-organ complications from its relatively higher hypertensive and vasoconstrictive effects, especially with intravenous injection. Clinical toxicity presents with a sympathomimetic-like toxidrome but with profound tachycardia and hypertension. The case literature is littered with examples of significant end-organ damage from propylhexedrine’s peripheral effects, including cardiogenic shock and pulmonary edema in a 22-year-old who suffered a large anterior myocardial infarction, as well as cases of brainstem dysfunction characterized by ophthalmoplegia. One case series from Texas details an association between propylhexedrine injection and 15 cases of sudden death. Management of propylhexedrine toxicity centers around management of sympathomimetic toxicity and careful attention to and treatment of possible end-organ damage from its marked hypertensive effects.

Mescaline

Mescaline is a serotonergic phenethylamine hallucinogen most famously found in peyote, a small spineless cactus found in the deserts between the USA and Mexico that has been traditionally consumed by Native American tribes. Although mescaline and peyote itself are Schedule I substances, other mescaline-producing cacti such as San Pedro cactus and Peruvian Torch cactus are not controlled, and can be legally
obtained through horticultural suppliers. Mescaline is a relatively rare drug of abuse, accounting for only 63 total exposures reported to USA poison control centers in 2015.

Mescaline is chemically related to amphetamines, and cases of toxicity produce a sympathomimetic-like toxidrome. Sympathetic symptoms such as mydriasis, tachycardia, and agitation are commonly seen. Despite its sympathomimetic effects in overdoses, life threatening toxic effects are rare. In 1 case series, GI symptoms were found to predominate, with marked abdominal pains, diarrhea, and vomiting that led medical providers to misdiagnose 1 ingestion as a possibly surgical abdomen. Management of mescaline intoxication is like that for many other hallucinogens: low stimulation environment and benzodiazepines with supportive care as needed.

**Dissociative legal highs**

N-methyl-D-aspartate (NMDA) receptor antagonists such as phencyclidine (PCP) and ketamine are abused in recreational settings for their dissociative effects. Ketamine was officially scheduled in 1999 and in the decade to follow the recreational drug market saw the synthesis and emergence of legal ketamine analogs such as methoxetamine and others. Furthermore, the over-the-counter cough remedy DXM has long been abused for its dissociative effects.

**Methoxetamine, Diphenidine, 2-Methoxydiphenidine**

Methoxetamine, an analog of ketamine that has NMDA antagonist activity, has been marketed via the internet as a legal high, and oftentimes as a safer alternative to ketamine abuse. After many European countries moved to ban the drug in the last 5 years in response to reports of fatalities, methoxetamine was then quickly joined by a flood of other “research chemicals” with dissociative properties, such as diphenidine, 2-methoxydiphenidine, as well as PCP analogs such as 3-methoxy-PCP and 4-methoxy-PCP. Many of these agents have not been fully described in terms of their exact pharmacologic and toxic properties, but given their subjective dissociative effects, they are presumed to be potent NMDA antagonists. Methoxetamine is ingested orally or absorbed sublingually, but can also be insufflated or even injected intramuscularly and intravenously. Methoxetamine presents clinically with dissociative effects as well as sympathomimetic effects, including tachycardia and hypertension.

Much of the morbidity of methoxetamine and other novel dissociative psychoactive substances seem to be centered around their neurocognitive effects as well as sympathomimetic qualities and propensity to be taken with other stimulatory psychoactive substances. Management is supportive, with benzodiazepines and intravenous (IV) fluids. Reports of diphenidine and methoxydiphenidine intoxications in Europe were noted to have a high percentage of co-ingestion with cannabis as well as other substances, resulting in severe intoxications. Patients commonly presented with hypertension, tachycardia, anxiety, and altered mental status requiring supportive care with benzodiazepines and sometimes propofol. A high proportion of the 20 reported deaths due to methoxetamine intoxication were also notable for numerous co-ingestions of illicit as well as other legal NPS.

**Dextromethorphan**

DXM is a selective α receptor agonist that has been used for many decades in the USA as an over-the-counter cough remedy. Since its introduction in the 1950s, DXM preparations have been abused for hallucinogenic and dissociative effects at high doses, a fact that led to its brief removal from the market in the 1970s and reformulation into worse-tasting compounds thought to have less abuse potential. Akin to many other legal highs, teenagers and young adults comprise the highest proportion of DXM abusers, among whom DXM abuse is known as “robotripping.” Those abusing DXM-containing over-the-counter formulations can also exhibit signs of toxicity from coformulatory compounds, including hepatotoxic effects from acetaminophen, anticholinergic effects from diphenhydramine, depressant effects from ethanol, and sympathomimetic effects from pseudoephedrine. Those who wish to abuse DXM in purer forms can chemically extract it from over-the-counter preparations, as well as buy pure powder legally from online chemical distributors.

DXM ingestion produces dissociative effects through its metabolism by cytochrome CYP2D6 to dextrorphan, an NMDA antagonist similar to PCP and ketamine. Like other dissociative agents, dextrorphan is also thought to exert adrenergic effects by inhibiting peripheral and central catecholamine reuptake. Further, as an isomer of the opioid levorphanol, DXM at high doses can affect opioid receptors, resulting in signs of opioid toxicity, including respiratory depression and miosis.

The subjective effects of DXM abuse have been noted to entail 3 dose-dependent plateaus, with low doses producing mild stimulant and euphoric effects, middle doses producing an effect like concurrent alcohol and marijuana intoxication, and large doses producing a dissociative effect like ketamine or PCP intoxication. Clinically, presentations of DXM ingestions largely depend on the dose ingested, producing mainly effects on the central nervous system, including...
restlessness, lethargy, ataxia, slurred speech, as well as hyporeflexia.97 Patients can also present with sympathomimetic effects, including diaphoresis and hypertension.97 Effects on the ophthalmologic exam produce varying results, including everything from rotary nystagmus and mydriasis to the miosis seen in opioid ingestions.97 Onset of symptoms usually occurs 30 to 60 minutes after ingestion, and will persist for an average of 6 hours, making any asymptomatic presentation 4 hours post-ingestion at very low risk of later developing toxicity.89

The major toxic effects of DXM are managed with supportive care, and are centered around benzodiazepines for acute agitation, management of toxicity from coformulatory agents, as well as possible serotonin syndrome.89 Guidelines also promote the use of activated charcoal up to 1 hour after the reported time of ingestion.89 It is reasonable to give naloxone to comatose or bradypneic patients in whom DXM ingestion is suspected, as opioid effects or opioid co-ingestants may be present.89 Long-term DXM abuse has produced cases of physical dependence in which abrupt cessation of the drug results in physical withdrawal symptoms of vomiting, myalgias, diarrhea, and night sweats.89

A major concern with DXM ingestion, even at therapeutic dosages in concert with certain other serotonin-affective drugs, is the risk of serotonin syndrome. There have been cases of serotonin syndrome when DXM was taken at antitussive therapeutic dosages in patients on therapeutic MAOIs such as phenelzine, as well as selective serotonin reuptake inhibitors such as fluoxetine.89 Such presentations should be managed aggressively with IV fluids, benzodiazepines, and active cooling measures.89

As DXM is commonly found in bromide salt, there have been theoretical concerns that massive ingestions will result in bromide toxicity as well as false elevations of chloride and even negative anion gaps due to misidentification of bromide for chloride by laboratory autoanalyzers. However, the only available evidence of bromism in the form of elevated chloride and a low (or negative) anion gap in the context of DXM use comes from a single case report.98 A more recent case report of DXM-related clinical bromism occurred when DXM was used with an older bromide-containing pharmaceutical.99

**Hallucinogenic and other compounds**

There are many legal hallucinogenic compounds that lack a discrete toxidrome. Many do increase serotonergic transmission resulting in stimulatory effects, including tachycardia and hypertension. Agitation and neuropsychiatric symptoms predominate with these agents.

**Salvia**

Native to Oaxaca, Mexico, *Salvia divinorum* is a leafy plant of the mint family that has been long recognized for its psychoactive properties when inhaled or sublingually absorbed.100 Salvia is not currently controlled at a federal level in the USA, although it is banned by several individual states.100,101 The active component in salvia is the chemical salvinorin A, a strong hallucinogen with potency similar to LSD but a mechanism of action novel among the hallucinogens, acting at κ-opioid receptors.102 The subjective effects of salvia have been variably described, with some likening it to LSD but others to a very strong cannabis intoxication.103,104 Salvia intoxication has a quick onset, within 30 seconds, and a total duration of hallucinogenic effects lasting usually <10 minutes.104

There is a lack of reports of salvia intoxication resulting in any serious physiologic toxicity.107 A 10-year survey of a major poison control system’s experience with salvia intoxications showed that most patients reporting sole salvia ingestions required only simple monitoring.105 Patients in this survey who presented with clinical instability or persistently altered mental status were usually under the influence of other concurrent intoxicants, in one case a suspected belladonna alkaloid ingestion.105 If a patient reports salvia intoxication and becomes unstable or dramatically altered over time, one should suspect coingestion of other toxic substances or possible concurrent medical or psychiatric illness.

**Lysergic acid hydroxyethylamide (LSA) plants**

A 5-HT<sub>1A</sub> agonist, LSA is a naturally occurring hallucinogen that is chemically related to LSD and about one-tenth as potent.106 LSA can be found in the seeds of the ornamental plants *Ipomoea violacea* (morning glory) and *Argyreia nervosa* (Hawaiian baby woodrose). Although seeds are often treated with an emetic by seed producers in order to discourage abuse, morning glory and woodrose seeds have been long abused as a legally available LSD substitute.106 After ingestion of crushed seeds, onset of hallucinogenic effects usually occurs within 1 hour and can last up to 10 hours.106

Major side effects include nausea, vomiting, and abdominal discomfort, which are thought to limit the abuse potential of LSA-containing seeds.107 Fatalities are rare, and those reported are usually secondary to self-harm due to acute agitation.108 In addition to producing hallucinogenic effects and agitation, the LSA-containing plants have been known to produce tachycardia, hypertension, pupillary dilation, flushing and polyuria.109 In the ED, suspected ingestion can
be managed through sensory isolation and benzodiazepines as needed.109

Synthetic cannabinoids

With street names such as K2 and “Spice,” the synthetic cannabinoids (SCs) are a heterogeneous and continually growing group of chemicals designed to produce the effects of the prototypical chemical tetrahydrocannabinol (THC) by acting on cannabinoid receptors.110 Given they usually share little to no similarity with THC, almost all will escape detection through standard urine toxicology screens.110 Many have been added to Schedule I.111 However, illicit drug manufacturers continue to design new analogs to escape existing regulations.110 The chemicals in liquid form are typically spread onto plant material that is then smoked or ingested, consumed outright, or vaporized and inhaled through electronic cigarettes.110

SCs are chemically heterogeneous but what they have in common is their affinity for cannabinoid receptors.112 A major difference between the SCs and the psychoactive compounds found in cannabis is in their relative affinity for cannabinoid receptors. Unlike marijuana, which contains over 50 known psychoactive chemicals, many of which have attenuating effects on THC, which is itself a partial agonist of cannabinoid receptors, many of the SCs are full agonists that demonstrate a much stronger affinity for these target receptors.113 It has been hypothesized that this is why, while marijuana results in little overt toxicity, many of the SCs have been found to have marked toxic effects, including excited delirium, seizures, psychosis, cardiotoxic effects, kidney injury, coma, and death.110 According to 1 survey of cannabis and synthetic cannabinoid users, users of SCs were much more likely to seek medical care for their symptoms.113 The risk of agitation and cardiotoxicity exceeds cannabis by 3.8 and 9.2 times, respectively.114

Physical examination findings of those using SCs have been found to be similar to those with cannabis intoxication, with delayed pupillary light reactions, slurred speech, and retardation of movements.115 More severe clinical presentations may present with nausea and vomiting, abdominal pain, and excited delirium. Cases of rhabdomyolysis, acute kidney injury, and even myocardial infarction have been reported.115 SCs have also been associated with new onset refractory status epilepticus.116 There have also been emerging cases of arterial thromboemboli in otherwise healthy young people using SCs.117 Management centers around control of the excited delirium state as well as management of end-organ damage.115 Effects usually last around 8 hours for most of the chemicals in question.115

Ayahuasca

Ayahuasca, also known as hoasca, yage, and natema, is a hallucinogenic tea brewed traditionally among Amazonian tribes for religious purposes from the jungle vine Banisteriopsis caapi together with the leaves of the Psychotria viridis plant.118 P. viridis contains most of the hallucinogenic activity through the compound N,N-dimethyltryptamine (DMT), a potent 5-HT agonist similar to LSD.119 By itself, DMT by oral ingestion is inactivated due to first pass metabolism by GI MAO.119 However, when combined with MAO-inhibiting beta-carboline alkaloids found in B. caapi, enough DMT is absorbed to produce psychedelic effects.119

The number of ayahuasca exposures has risen in the last 10 years in the USA, with a peak in 2012 at around 100 reported exposures.120 Although DMT is a Schedule I drug, and people have been prosecuted for possessing ayahuasca preparations, the plants themselves remain uncontrolled and legal to possess.120

The most common clinical manifestations of ayahuasca intoxication as reported to poison centers were hallucinations, tachycardia, agitation, hypertension, mydriasis, and vomiting.120 Altered mental status requiring intubation, seizures, and a few fatalities have been reported.120 Case reports of fatalities are limited by lack of forensic and toxicologic information, but some experts have suggested possible dangerous effects, including serotonin syndrome between ayahuasca and other serotonergic drugs such as prescription MAOIs and SSRIs.121 Treatment of ayahuasca intoxications consists of symptomatic supportive care and benzodiazepines for agitation or anxiety.120

Kava kava

Kava refers to a traditional beverage prepared from the tropical shrub plant Piper methysticum, native to the South Pacific. Used for centuries for its euphoric and anxiolytic effects, kava has seen increased use in the USA, where it is considered a health supplement.122 Kava is prepared typically by mixing crushed root or prepared extracts or powders with water or coconut milk, and it is also taken in pill form.123 The psychoactive components of kava are thought to be a group of lipophilic substances known as the kavalactones that exert their effects through numerous possible mechanisms ranging from enhanced GABA receptor binding to reversible inhibition of MAO B.124
Reports of acute kava toxicity resulting in adverse events requiring medical care are rare, accounting for 20 cases reported nationally in 2015.76,125,126 Reported adverse events include headaches, dyspnea, nausea, as well as dizziness, for which treatment is supportive.127,128 There have been reports of neurologic toxicity in the form of acute ataxia as well as dystonic reactions responsive to anticholinergics.129,130 Kava has been shown to have more profound acute mental status depressive effects when combined with benzodiazepines and alcohol.126,131,132 Excessive use of kava over a long term has been associated with cases of hepatotoxicity resulting in fulminant liver failure.133 Laboratory findings in chronic kava users have shown elevated liver enzymes, as well as lower blood lymphocyte counts.134 Physical exam findings in chronic kava users include skin photosensitivity, which usually resolves after cessation of use.135

**Inhalants**

Long abused as readily available “legal highs,” inhalants refer to a heterogeneous group of usually lipophilic chemicals that are inhaled for their immediate consciousness-altering effects.136 Inhalants are most commonly abused among adolescents, and the most common types of abused chemicals include the volatile hydrocarbons found in widely available products such as air fresheners, paint, gasoline, adhesives, and propellants.137 The major non-hydrocarbon inhalant of abuse is nitrous oxide, which is found in whipped cream recharging canisters.136 According to available surveys, reports of abuse of volatile hydrocarbons seem to be decreasing,137 while reports of abuse of nitrous oxide seem to be increasing.138 In 1 survey, the inhalants with the highest reported mortality included nitrous oxide, butane, propane, and air fresheners.137

Patients abusing inhalants may present with a wide variety of subjective complaints, including dyspnea, palpitations, lightheadedness, sneezing and coughing, as well as GI complaints, including nausea and vomiting.136 Odors may persist in the breath for several hours following inhalation.139 Patients may also present with acute airway compromise due to chemical injury from aerosolized substances.140 Other clinical presentations are myriad, and may include altered mental status, headaches, and GI disturbances.136

The inhalational hydrocarbons of abuse have numerous toxic effects on the body, with cardiotoxic, neurotoxic, hepatotoxic, and nephrotoxic effects having been reported.136 The most profound cardiotoxic effect of hydrocarbon inhalation includes what is known as sudden sniffing death syndrome, which is thought to be due to cardiac arrhythmias caused by a sudden catecholamine surge in the setting of volatile hydrocarbon-sensitized myocardium.139 Some patients with cardiac arrhythmias due to inhalants survive long enough to present to the ED.141 In this setting, propranolol and other beta blockers have been used as agents to decrease myocardial catecholamine sensitivity, which is thought to be the mechanism of inhalant cardiotoxicity.141 Toluene and other hydrocarbon inhalants are also thought to sometimes cause QTc prolongation and torsades de pointes, as well as myocarditis.142,144 Acute toluene inhalation has also been associated with marked metabolic derangements such as rhabdomyolysis and type 1 renal tubular acidosis, with hyperchloremia, hypokalemia, and a normal anion gap.143

Besides possible acute fatal asphyxiation and even baro-trauma from inhalation, the major toxic effects of nitrous oxide, aka “whippets” as they are known, are neurologic. Nitrous oxide inactivates vitamin B12, decreasing conversion of homocysteine to methionine, ultimately resulting in demyelination within the central and peripheral nervous systems.145 This classically results in subacute combined degeneration, especially around the dorsal columns of the spinal cord.145 Clinically, chronic nitrous oxide abuse may present with peripheral neuropathic effects, including tingling and numbness all the way up to loss of dexterity, poor balance, and leg weakness.146 Although the neurologic changes may be irreversible, some function may be restored through cessation and B12 supplementation.136

**Conclusion**

Existing outside of regulatory drug statutes, the drugs that can be used by patients to get “legal highs” truly represent an astounding array of chemicals, some of which are quite easily obtainable. “Legal highs” can represent a challenge to the emergency medicine physician due to the sheer number of available agents, their multiple toxidromes and presentations, their escaping traditional methods of analysis, and the reluctance of patients to divulge their use of these agents. However, a focus on aggressive symptom-based supportive care is the mainstay of treatment for these patients in the ED.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


