Novel psychoactive substances (designer drugs): overview and pharmacology of modulators of monoamine signalling

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Summary

Novel psychoactive substances are newly designed drugs (“internet drugs”, “research chemicals”, “legal highs”) potentially posing similar health risks to classic illicit substances. Chemically, many novel psychoactive substances can be classified as phenethylamines, amphetamines, synthetic cathinones, piperazines, pipradrols/aminooindanes, benzofurans, and tryptamines. Pharmacologically, these substances interact with various monoaminergic targets. Typically, stimulants inhibit the transport of dopamine and noradrenaline (pipradrols, pyrovalerone cathinones) or induce the release of these monoamines (amphetamine and methamphetamine-like cathinones), entactogens predominantly enhance serotonin release (phenylpiperazines, aminoindanes, para-substituted amphetamines, and MDMA-like cathinones) similar to MDMA (ecstasy), and hallucinogens (tryptamines, hallucinogenic phenethylamines) are direct agonists at serotonin 5-HT2A receptors. Synthetic cannabinoids are another group of novel substances which all act as agonists at the cannabinoid CB1 receptor similar to THC but are chemically diverse. In particular, the relative serotoninergic vs dopaminergic activity (determined by the dopamine/serotonin transporter inhibition ratio in vitro) can be helpful to predict the desired psychotropic but also the toxic effects of novel substances as well as their potential for addiction. Although the use of novel psychoactive substances mostly produces minor or moderate poisonings, serious complications occur. Serotonergic drugs (entactogens and hallucinogens) are associated with acute serotonin syndrome, hyperthermia, seizures, and hyponatremia. Dopaminergic drugs are highly addictive and acute toxicity includes prolonged stimulation, insomnia, agitation, and psychosis. Agitation, anxiety, paranoia, hypertension, and rarely myocardial infarction and renal failure are seen with synthetic cannabinoids. Treatment is supportive.

Key words: novel psychoactive substances; designer drugs; amphetamines; research chemicals; pharmacology; toxicity; serotonin; dopamine

Introduction

The term “novel psychoactive substances” refers to newly-misused narcotic or psychotropic drugs that may pose a threat to public health comparable to classic previously listed psychotropic substances. Typically, the novel substances or “designer drugs” are analogues or chemical de-
derivatives of controlled substances designed to produce effects similar to the controlled substances they mimic. In contrast to classic illicit drugs ("street drugs"), novel psychoactive substances are typically sold through the Internet ("internet drugs"). The substances are misbranded as "research chemicals", "bath salts", "plant food", and labelled as "not for human consumption". The substances are typically chemically slightly different from already scheduled drugs to circumvent regulations and are therefore also termed "legal highs". The fully synthetic drugs are mostly produced in China and South East Asia and sold worldwide by numerous Internet retail vendors often as less expensive replacements of classic stimulants or narcotics [1]. In the last years we have seen an unprecedented growth in the number of new psychoactive substances on the illicit drug market. In the European Union (EU), 41 novel psychoactive substances were identified for the first time in 2010, 49 in 2011, 73 in 2012, 81 in 2013, and 37 by April 2014 within the European Early Warning System [2]. By 2014, more than 300 novel substances have been detected since 2005. Currently, more than one new substance is identified in one of the EU countries every week. The European Monitoring Centre for Drugs and Drug Addiction (EMCDDA) collects information on these substances from various national sources including police, customs, forensic laboratory networks, health care systems, event organisers, drug checks, and Internet test purchase samples. Internet snapshots are also an important method to describe trends in new substances available for online sale [3]. The EMCDDA also regularly publishes reports on the risk assessments of novel psychoactive substances (http://www.emcdda.europa.eu/activities/action-on-new-drugs). Furthermore, the European Drug Emergency Network (Euro-DEN) collects health emergency data from hospitals to look at trends in the acute harms associated with the use of novel psychoactive drugs [4]. The University Hospital of Basel participates in the Euro-DEN and we characterised several intoxications with novel drugs [5, 6] and in particular their pharmacological properties [7–9]. Data on the clinical characteristics of intoxications with novel psychoactive substances are also typically collected and reported by poison centres [10–13]. In this review I will describe the in vitro pharmacodynamicics of novel psychoactive substances acting on monoamine signalling and how it relates to the clinical effects of these compounds. Several excellent publications have more comprehensively reviewed the pharmacology and toxicology of novel psychoactive substances [14, 15].

**Classification of novel psychoactive substances**

The new psychoactive substances can be classified based on their psychotrophic effects as stimulants, empathogens/entactogens (3,4-methylenedioxymethamphetamine [MDMA, ecstasy]-like substance) or hallucinogens or according to their chemical family as phenethylamines, amphetamines, cathinones, piperazines, pipradrols/piperidines, aminoindanes, benzofturans, and tryptamines [15]. The synthetic cannabinoids include a large number of chemically diverse substances which act on the cannabinoid CB1 receptor and have mostly hallucinogenic but also some stimulant properties. Currently, the compounds are classified based on their common pharmacological action rather than similar chemical structures. The substances exhibit a rather broad spectrum of dopaminergic, noradrenergic, or serotoninergic pharmacological effects even within their chemical family [7–9, 16]. Accordingly, the psychotropic and clinical toxicological effects are also quite diverse and vary considerably from drug to drug.

**Synthetic cannabinoids**

Herbal mixtures ("spices") have emerged as legal alternatives to cannabis [17]. It was initially assumed that the psychotropic effects were derived from the plants themselves. However, synthetic compounds added to the herbal blends and with action at the CB1 receptor were found to be responsible for the psychopharmacological activity [18]. "Spice" drugs have become popular alternatives to marijuana among teenagers and constitute an exceptionally large class of novel psychoactive substances [2]. Synthetic cannabinoids vary considerably in chemical structure while they uniformly act as agonists at the CB1 receptor [17]. Many of the substance names (abbreviations) typically start with the initials of the chemists (JWH or AM) followed by numbers. Most synthetic cannabinoids produce overall similar effects and toxicity to tetrahydrocannabino1 (THC) [6, 17]. However, synthetic cannabinoids are thought to be associated with more severe psychosis and agitation and more sympathomimetic effects because they are more potent full receptor agonists and also lack cannabinoid, which is contained in natural THC-containing products and has anxiolytic and antipsychotic properties. In fact, agitation, aggression, paranoid thinking and anxiety are common symptoms after consumptions of synthetic cannabinoids [19, 20]. However, more recently a second generation of synthetic cannabinoids has emerged both in the US and in Europe and lead to epidemics associated with more severe toxicity including collapses, seizures, and cardiac toxicity [11]. Cases with acute kidney injury have also been described [21]. The effect of the synthetic cannabinoids is typically short-lasting compared with amphetamine-type substances [5, 20]. Short-time monitoring and supportive care is sufficient to treat most cases with intoxications. Synthetic cannabinoids are not detected by drug screening tests for THC, and highly specialised mass spectrometry techniques and time are required to document the presence of these novel compounds in biological samples [6, 22].

**Phenethylamines and amphetamines**

Phenethylamine is the core structure of many novel psychoactive substances (fig. 1). Amphetamines (alpha-methyl-phenyl-ethyl-amine, fig. 1) are formed by the addition of an alpha methyl group which protects against metabolism by monoamine oxidase [23]. Methylation of the terminal amine results in methamphetamine and greater CNS activity. Amphetamine and methamphetamine (fig. 1) are both classic psychostimulants which have been used clinically, recreationally and by military services since the 1930s [23]. The amphetamine-derivative MDMA (fig. 1) is
not a designer drug or novel psychoactive substance as it has been used for decades. However, MDMA is still one of the most widely used recreational drugs and many novel psychoactive substances were designed to mimic its effects or as substitutes for MDMA in ecstasy pills. MDMA is the prototypical empathogen or entactogen meaning that it produces feelings of empathy or “being touched”. The drug is mostly used to enhance sociability [24]. In controlled studies, MDMA increases emotional empathy, trust, extroversion, and sociality relatively more than stimulants which mainly produce arousal and stimulation [25, 26].

Role of dopamine, serotonin and noradrenaline in the pharmacology and toxicology of novel psychoactive substances

Many novel psychoactive substances interact with biogenic amine neurotransmitter transporters. Amphetamines including methamphetamine and MDMA inhibit the dopamine, serotonin and noradrenaline (norepinephrine) transporter (DAT, SERT, NET, respectively) and also release these monoamines through the respective transporter. Methamphetamine predominantly increases dopamine and noradrenaline. MDMA mostly increases serotonin and noradrenaline. The entactogenic effects of MDMA are generally considered to depend on its serotonergic effects [27]. MDMA also increases oxytocin [25, 28] which may mediate its prosocial effects [29]. Consequently, substances which predominantly release serotonin, similar to MDMA, can be expected to produce MDMA-like entactogenic effects. MDMA and similar serotonergic substances are also associated with serotonergic toxicity including serotonin syndrome, hypotension, hyperthermia, and seizures [30–32]. In contrast, psychostimulants such as methamphetamine or methylphenidate are mostly enhancing dopaminergic neurotransmission [7, 33]. Dopamine mediates the reinforcing and addictive properties of drugs of abuse. In contrast, an increase in the serotonergic properties of a substance is associated with a reduced potential for addiction [34]. Consequently the relative dopaminergic to serotonergic properties in vitro (dopamine/serotonin transporter inhibition ratio) of a novel substance can be determined as a useful marker for its potential clinical psychotropic and acute toxic effects [7] (fig. 2). Serotonin release [27] and a DAT/SERT inhibition ratio of typically 0.01–0.1 are expected to result in subjective drug effects similar to those of MDMA or other empathogens [7, 8] (fig. 2). Cocaine has a DAT/SERT inhibition ratio close to unity, and methamphetamine is more selective for the DAT, with a DAT/SERT inhibition ratio >10 and mostly exerting psychostimulant effects in humans [7] (fig. 2). Compounds with a DAT/SERT inhibition ratio >1 are also associated with a high abuse potential [7]. Furthermore, all classic amphetamines and novel psychoactive substances increase the activity of the noradrenergic system [7–9], which results in the sympathomimetic cardiostimulant effects associated with all these substances. The potency of a substance to activate the noradrenergic system inversely correlates with the doses typically used recreationally [7, 35]. The profiles of novel psychoactive substances can be determined in vitro and compared with those of the classic substances where the clinical effects are known. Substances which selectively activate the catecholamine systems such as methamphetamine, or 3,4-methylenedioxyxymethylamphetamine (MDPV) or α-pyrrolidinopentiophenone (α-PVP) [7, 36] are considered highly addictive and their acute toxicity typically includes prolonged insomnia, agitation, and psychotic symptoms [37].

Figure 1

Structures of representative novel psychoactive substances (stimulants and empathogens). Chemically, the substances shown include amphetamines (amphetamine, methamphetamine, MDMA, PMA), cathinones (β-keto-amphetamines such as methedrone, mephedrone, methylene, pentylone, buphedrone, methcathinone, naphlylone, MDPV or α-PVP), an aminoindane (MDAI), piperazines (mCPP and BZP) and pipradrols (2-DMPMP and methylphenidate). Pharmacologically, these substances are psychostimulants or empathogens and primarily interact with monoamine transporters.
Para-(4)-phenyl-substituted (serotonergic) amphetamines

Paramethoxymphetamine (PMA) and paramethoxy-methamphetamine (PMMA) are typically sold as ecstasy [38]. This substitution for MDMA is unwanted because PMA and PMMA are associated with higher morbidity and mortality particularly attributable to hyperthermia [38, 39]. The para-substituted amphetamines are potent noradrenaline and serotonin transporter inhibitors and releasers of these monoamines [8]. Their hyperthermic properties are stronger than those of MDMA [40] and have been associated with serotonergic and adrenergic receptor activation [41]. Therefore, hyperthermic complications are of particular concern when these amphetamines or novel psychoactive substances with a comparable pharmacological profile are used [8]. Novel substances with a predominantly serotonergic action [8] and associated with high serotonergic toxicity are 4-methylthioamphetamine (4-MTA) [42] and methedrone (β-keto-PMA) [43]. 4-MTA is the methyl-thio analog of PMA. 4-MTA produces MDMA-like effects in animals and humans and is typically used by ecstasy users [44]. Methedrone is the β-keto-substituted analog of PMMA. Methedrone is a serotonergic cathinone found in “bath salt” products.

Synthetic cathinones

Cathinones contain a ketone group at the β-position of the amphetamine (β-keto-amphetamines, fig. 1). The most commonly found synthetic cathinones are mephedrone (4-methylmethcathinone), methylene (β-keto-MDMA), 4-methyllethcathinone (4-MEC), and MDPV [45, 46]. Other abused cathinones include ethylone, methedrone, naphyrone, flephedrone, 3–fluoromethcathinone (3–FMC), pentylole, buphedrone, α-PVP, and others [7, 8, 15, 45]. The pharmacology of the various cathinones has recently been determined [7, 8, 16, 47, 48]. Cathinones can be classified based on their pharmacological profiles (mostly the DAT/SERT inhibition ratio) with relevance to their clinical toxicity [7]. The serotonergic cathinone methedrone predominantly acts on the serotonin system similar to MDMA or PMMA and PMMA as noted above [8]. The cocaine-MDMA-mixed cathinones group includes substances which are roughly equipotent inhibitors of DAT and SERT similar to cocaine and release serotonin similar to MDMA. The group includes mephedrone, 4-MEC, methylene, ethylone, and butylone [7, 8]. For example, the subjective effects of mephedrone have been reported to be similar to those of cocaine [49] but also MDMA [50]. Typically, these drugs produce psychotropic effects similar to MDMA when administered orally but with enhanced psychostimulation similar to cocaine when administered intranasally. All cathinones exhibit higher dopaminergic activity when compared with their non-β-keto amphetamine analogs. This increased dopaminergic property of the cathinones suggests higher stimulant-type effects and greater risk for dependence. For example, mephedrone, which is readily self-administered by rats [51], has been reported to produce strong craving in humans and is said by users to be more addictive than cocaine [49]. In addition, substances with fast brain access have a higher risk of dependence. Naphyrone is very similar to cocaine. Both inhibit all monoamine transporters with equal potency but do not release monoamines [7]. The methamphetamine-like cathinones including cathinone, methcathinone, ethcathinone, flephedrone, and 3–FMC exhibit a DAT/SERT inhibition ratio similar to methamphetamine, with high inhibitory potencies at the DAT and low potencies at the SERT [7, 8]. These cathinones also release dopamine and noradrenaline similar to methamphetamine. Clinically, these cathinones produce similar toxicity to amphetamine, including hypertension, hyperthermia, euphoria, locomotor activation, and hallucinations following higher or repeated doses. The pyrovalerone cathinones include pyrovalerone, MDPV, α-PVP, and others. These substances are very potent and selective inhibitors of the DAT and NET, at least 10–fold more potent than cocaine or methamphetamine. The drugs are no substrates of the transporter and therefore do not release monoamines. The pyrovalerone derivatives also readily cross the blood-brain barrier because of their high lipophilicity [7]. Consistent with its high potency as a dopamine transporter inhibitor, MDPV is a potent reinforcer in rats, similar to methamphetamine [52]. Taken together these pharmacological properties predict high risks of symptomatic intoxication and of addiction in humans. Acute toxic effects of MDPV and α-PVP mainly include acute psychotic symptoms with agitation and hallucinations [12, 37]. Of note, the pharmacology of MDPV is strikingly different from that of MDMA, although MDPV and MDMA both share the characteristic 3,4-methylenedioxy-phenyl group. In fact, caution is needed when making predictions about the pharmacological activity of new phenethylamine-type psychoactive substances based only on the chemical structures. Pharmacological data are needed for each novel designer drug [53].

Ring substituted phenethylamines and amphetamines (2-C and 2-D series)

Addition of methoxy-groups at the 2- and 5-positions with any hydrophobic substitution at the 4-position of the phenethylamine confers hallucinogenic activity (2C-series or
ring substituted phenethylamines). Example drugs of the large 2-C series are 2,5-dimethoxy-4-bromophenethylamine (2C-B) and 2,5-dimethoxy-4-iodophenethylamine (2C-I) (fig. 3). The hallucinogenic properties of these drugs are further enhanced by a methyl-group at the α-carbon (D-Series or ring substituted amphetamine or hallucinogenic amphetamines). Example drugs included in this group are 2,5-dimethoxy-4-methylamphetamine (DOM), 2,5-dimethoxy-4-bromoamphetamine (DOB), and 2,5-dimethoxy-4-iodoamphetamine (DOI) (fig. 3). Nausea and tachycardia but also long-lasting hallucinogenic effects, agitation, and ergotism (vasospasms) have been reported in intoxications with these hallucinogenic amphetamines. The hallucinogenic activity of the 2-C and D series drugs is mediated by an interaction with the serotonergic 5-HT₂A receptor [54]. Various N-methoxybenzyl-substituted phenethylamines have emerged on the EU drug market since 2012 [2]. These drugs such as 4-Iodo-2,5-dimethoxy-N-(2-methoxybenzyl)phenethylamine (25I-NBOMe, fig. 3) exert an even higher potency at the 5-HT₂A receptor and possibly also at other receptors compared with the already very potent classic hallucinogens [55]. Severe and fatal intoxications including agitation, hallucinations, seizures, and hyperthermia have been reported with 25I-NBOMe [56, 57] consistent with serotonergic but also sympathomimetic toxicity. Importantly these novel hallucinogens are extremely potent and psychoactive at microgram doses. This is likely to result in overdosing.

Benzofurans and benzodifurans

Benzofurans and benzodifurans are also groups of ring substituted amphetamine. Benzo furans containing one furan ring are 6-(2-aminomethyl)benzo furan (6-APB, fig. 3) and 5-(2-aminomethyl)benzofuran (5-APB) and several others. These drugs are structurally related to MDMA and similarly inhibit monoamine transporters [16] and release serotonin and noradrenaline (unpublished data by the author). Users report the effects of 5-APB and 6-APB to be comparable to MDMA but more intense [58]. Adverse effects include nausea, sympathomimetic stimulation, and agitation [58]. Benzodifurans are a group also known as the “fly” drugs (bromo-dragon fly, 2C-B-fly [fig. 3], etc.) and are hallucinogen. These drugs may produce paranoia, agitation, tachycardia, and hyperthermia and have been implicated in several fatalities [58].

Piperazines

Piperazines are commonly found in ecstasy pills as substitutes for MDMA [59]. The phenylpiperazines meta-chlorophenylpiperazine (m-CPP, fig. 1) and trifluoromethylphenylpiperazine (TFMPP) are indirect and direct serotonergic agonists without dopaminergic activity. Consistently these drugs are not reinforcing [60]. m-CPP and TFMPP have less desirable psychotropic effects and more adverse effects, including dysphoria, anxiety, and nausea, compared with MDMA [61–63]. Benzylpiperazine (BZP, fig. 1) is an indirect dopamine and noradrenaline agonist without serotonergic properties [9] and exerts stimulant effects in humans [64]. The clinical toxicity of BZP mainly includes hallucinations, agitation, seizures, and hyperthermia [65]. Drug users report more unpleasant effects and hallucinations with BZP than with MDMA [61]. BZP is sometimes combined with m-CPP or TFMPP and sold as ecstasy [59] mimicking the mixed dopaminergic-serotonergic profile of MDMA. However, the BZP-TFMPP combination is not well tolerated at higher doses and produces agitation, anxiety, hallucinations, and vomiting [66].

Aminodindanes

Aminodindanes such as 5,6-methyleneoxy-2-aminodindane (MDAI, fig. 1) and 5-iodoaminodindane (5-IADI) are allegedly less-neurotoxic alternatives to MDMA. MDAI and 5-IADI release serotonin and noradrenaline comparable with MDMA [9]. Consistently the subjective effects of MDAI are also reported to be very similar to those of MDMA [67]. Complications include serotonin syndrome and hyperthermia [67], also similar to MDMA.

Pipradrols/piperidines

Intoxications with “ivory wave”, which contains the pipradrol derivative desoxypripradrol (2-diphenyl-methylpiperidine 2-DPPP, fig. 1) or diphenylprolinol (diphenyl-2-pyrrolidinemethanol [D2PM]) have been reported starting in 2010 [68, 69]. D2PM and 2-DPPM are selective and very potent catecholamine transporter inhibitors without transporter-mediated substrate-releasing properties, similar to methylphenidate [9]. This pharmacological profile is also very similar to MDPV, α-PVP, and other pyrovalerone cathinones [7, 36] and likely associated with high abuse liability and an increased risk of psychiatric complications. The clinical toxicity of 2-DPPM and D2PM is long-lasting (24-72 h) and involves sympathomimetic stimulation with hypertension, agitation, hallucinations, and insomnia [45, 68, 69].

Tryptamines

Tryptamines contain an indole structure joined to an ethylamine group in simple tryptamines (for a classification see [15]). Natural tryptamines include serotonin and hallucinogens such as psilocybin in magic mushrooms and dimethyltryptamine (DMT, fig. 3) in Ayahuasca. Ergolines are more complex tryptamines and include the prototypic hallucinogen LSD. Commonly misused novel synthetic tryptamines (designer hallucinogens) are alpha-methyltryptamine (AMT) [13] and 4-hydroxy-N-methyl-N-ethyltryptamine (4-HO-MET) [45] (fig. 3). Tryptamine-induced hallucinations are mostly mediated by direct interaction with serotonergic 5-HT₂A receptors [54]. However, many tryptamines also release serotonin [70] and in some cases dopamine and noradrenaline [71]. As a result serotonin syndrome and sympathomimetic toxicity may occur and tryptamines have stimulant as well as hallucinogenic properties [13]. Typically visual hallucinations predominate. Consistent with their action as serotonergic agonists tryptamines are not addictive [71].
Prevalence of novel psychoactive substance use

In a German survey, 9% of 15-18 year-old school students reported having smoked spice products at least once in their lives and 2% also within the last 30 days. Other “legal highs” (research chemicals, bath salts) had been used by 3% at least once and in 1% within the last 30 days [72]. In contrast, in the UK, life-time prevalence rates of mephedrone use of 20.3% have been reported from some areas in 2010 [73].

In Switzerland, the prevalence of novel psychoactive substances use is yet unknown. In 2009–2010, 37% of people using classic drugs such as amphetamines or MDMA tested positive for novel psychoactive substances in hair [74]. For example, m-CPP was found in 10.5%, mephedrone in 3%, 4-fluoromethamphetamine in 4% [74]. However, many substances were possibly not yet used at the time of analysis or were not yet detectable. Considering our ongoing surveillance data collections from emergency department visits, presentations of intoxications with novel psychoactive substances are infrequent compared with those with alcohol, cannabis or cocaine [4, 75].

Treatment of intoxications

Table 2 lists typically reported acute medical problems associated with novel substances from each class. The clinical toxicity of novel psychoactive substances is generally similar to that of other amphetamines including MDMA. The majority of patients (85-95%) presenting at emergency departments with acute medical problems associated with the use of novel psychoactive substances are minor or moderate poisonings [12, 45, 76]. Common clinical features are hypertension, tachycardia, chest pain, agitation, and hallucinations [12, 76, 77]. Severe and fatal poisonings manifest as serotonin syndrome [67], hyperthermia [38, 78], seizures, and brain oedema due to hyponatraemia [79]. These are also the severe complications of MDMA use [31, 80]. Treatment is mainly supportive. Heart rate, blood pressure, and body temperature should be monitored. Depending on clinical features laboratory tests including electrolytes, creatine kinase, liver enzymes, and cardiac enzymes may be indicated. Acute treatment of a sympathomimetic toxidrome primarily includes benzodiazepines and fluid replacement to control agitation, cardiovascular stimulation and hyperthermia. The use of haloperidol without benzodiazepines is generally not recommended because seizure and dysrhythmia thresholds are lowered and negative drug-induced psychological effects including anxiety may increase [81]. Hypertension should be treated primarily with nitrates. β-blockers should be avoided because of unopposed α-adrenergic stimulation resulting in further increases in blood pressure [82]. Phentolamine may be useful but α-blockade increases stimulant-induced tachycardia [83]. Combined α-β-blockade with carvedilol reduced MDMA-induced elevations in blood pressure, heart rate and also body temperature [84], but it is not a routine treatment for stimulant-induced toxicity [85]. Physical cooling and relaxation may be needed in cases of severe hyperthermia, but antipyretics are of no use.

<table>
<thead>
<tr>
<th>Table 1: Monoamine transporter inhibition.</th>
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<tbody>
<tr>
<td>Substance</td>
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<tr>
<td>---------------------------</td>
</tr>
<tr>
<td>PMA</td>
</tr>
<tr>
<td>m-CPP</td>
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<tr>
<td>MDMA</td>
</tr>
<tr>
<td>Methedrone</td>
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<tr>
<td>MDAI</td>
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<tr>
<td>Mepedrone</td>
</tr>
<tr>
<td>Naphryrone</td>
</tr>
<tr>
<td>Cocaine</td>
</tr>
<tr>
<td>Methylene</td>
</tr>
<tr>
<td>BZP</td>
</tr>
<tr>
<td>Pentylene</td>
</tr>
<tr>
<td>Methamphetamine</td>
</tr>
<tr>
<td>Buphedrone</td>
</tr>
<tr>
<td>Methcathinone</td>
</tr>
<tr>
<td>Amphetamine</td>
</tr>
<tr>
<td>MDPV</td>
</tr>
<tr>
<td>2-DPMP</td>
</tr>
<tr>
<td>PV</td>
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<tr>
<td>Methylphenidate</td>
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</tbody>
</table>

NET = noradrenaline transporter; DAT = dopamine transporter; SERT = serotonin transporter. Values are concentrations at which the transporter is inhibited by 50%. A low value indicates greater potency.

DAT/SERT ratio = IC50 DAT / IC50 SERT. A low DAT/SERT ratio (<0.1) indicates tenfold greater relative serotonin vs dopaminergic activity similar to MDMA. A high DAT/SERT ratio (>10) indicates greater relative dopaminergic vs serotonin activity similar to methamphetamine.

Data are from Simmler et al. 2013 and 2014 and unpublished by the author (PVP).
be sampled and analysed by liquid chromatography-mass spectrometry by a specialised laboratory. Typically the identity of the novel psychoactive substances will not be readily available for the management of the acute intoxication. Nevertheless, the novel psychoactive substances should be identified to better document the substances and their associated toxicity.

**Conclusion**

Stimulants inhibit the transport of dopamine and noradrenaline (pipradrols, pyrovalerone cathinones) or induce the release of these monoamines (amphetamine and methamphetamine-like cathinones), entactogens predominantly enhance serotonin release (phenylpiperazines, aminoindanes, para-substituted amphetamines, and MDMA-like cathinones) similar to MDMA, and hallucinogens (tryptamines, hallucinogenic phenylethamines) are direct agonists at serotonergic 5-HT2A receptors. Synthetic cannabinoids act as agonists at the cannabinoid CB1 receptor similar to THC. In particular, the dopamine-serotonin transporter inhibition ratio *in vitro* can be helpful to predict the expected psychotropic but also the toxic effects of novel substances.

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**Table 2:** Acute clinical toxicity associated with novel psychoactive substances.

<table>
<thead>
<tr>
<th>Substances</th>
<th>Classification</th>
<th>Leading acute toxicity</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMA</td>
<td>(Para-substituted) amphetamine, empathogen</td>
<td>Serotonergic toxidrome, hyperthermia, nausea, seizures, fatalities</td>
</tr>
<tr>
<td>MDMA</td>
<td>Amphetamine, empathogen</td>
<td>Serotonergic and sympathomimetic toxidrome, hyperthermia, hyponatraemia, renal and liver failure</td>
</tr>
<tr>
<td>MDAI</td>
<td>Aminoindane, empathogen</td>
<td>Serotonergic and sympathomimetic toxidrome, hyperthermia</td>
</tr>
<tr>
<td>Mephedrone, methylene</td>
<td>(Cocaine-MDMA-mixed) cathinone, empathogen/stimulant</td>
<td>Sympathomimetic toxidrome, agitation, vomiting, psychosis, chest pain, seizures, insomnia</td>
</tr>
<tr>
<td>3-FMC, flephedrone, methcathinone</td>
<td>(Methamphetamine-like) cathinone, stimulant</td>
<td>Sympathomimetic toxidrome, psychosis, agitation, chest pain, insomnia</td>
</tr>
<tr>
<td>MDPV, o-PVP</td>
<td>(Pyrovalerone)-cathinone, stimulant, high potency drug</td>
<td>Psychosis, agitation, convulsive behaviour, sympathomimetic toxidrome, chest pain, prolonged insomnia</td>
</tr>
<tr>
<td>m-CPP, TFMPP</td>
<td>(Phenyl)piperazine, entactogen</td>
<td>Serotonergic toxicity, nausea, vomiting, anxiety, headache, dizziness, dysphoria, confusion, hallucinations, tachycardia</td>
</tr>
<tr>
<td>BZP</td>
<td>(Benzy)l)piperazine, stimulant</td>
<td>Mostly sympathomimetic toxicity, agitation, anxiety</td>
</tr>
<tr>
<td>2-DPMP, methylphenidate</td>
<td>Pipradrol, stimulant</td>
<td>Sympathomimetic toxidrome, insomnia, agitation, hallucination, anxiety, Insomnia</td>
</tr>
<tr>
<td>6-APB</td>
<td>Benzofuran, empathogen</td>
<td>Serotonergic and sympathomimetic toxidrome, nausea, agitation, dizziness, hyperthermia</td>
</tr>
<tr>
<td>&quot;fly&quot; drugs</td>
<td>Benzodifurans, hallucinogens</td>
<td>Psychois, agitation, hyperthermia, sympathomimetic toxicity, vasospasms, limb pain/Ischaemia, seizures, fatalities</td>
</tr>
<tr>
<td>2C-B</td>
<td>(Ring-substituted) phenethylamine, hallucinogen</td>
<td>Psychosis, agitation, vomiting, vasospasms</td>
</tr>
<tr>
<td>25-I-NBOMe</td>
<td>&quot;NBOM&quot;-phenethylamine, hallucinogen, very high potency drug</td>
<td>Serotonergic and sympathomimetic toxidrome, psychosis, agitation, seizures, hyperthermia</td>
</tr>
<tr>
<td>AMT</td>
<td>Tryptamine, hallucinogen</td>
<td>Serotonergic and sympathomimetic toxidrome, psychosis, agitation, hyperthermia, nausea</td>
</tr>
<tr>
<td>Synthetic cannabinoids</td>
<td>Cannabinoid, hallucinogen</td>
<td>Psychois, agitation, anxiety, sympathomimetic toxidrome, chest pain, myocardial infarction, renal injury, seizure, vomiting</td>
</tr>
</tbody>
</table>

Sympathomimetic toxidrome typically includes hypertension, tachycardia, mydriasis, agitation, and sweating. Serotonergic toxidrome typically includes tremor, clonus, hyperreflexia, sweating, hyperthermia, mydriasis, agitation, and confusion. Psychosis includes hallucinations and paranoia.

AMT = alpha-methyltryptamine; o-PVP = alpha-pyrolidinopropiophenone; 6-APB = 6-(2-amino-propyl)benzofuran; BZP = benzylpiperazine; 2C-B = 2,5-dimethoxy-4-bromophenethylamine; m-CPP = meta-chlorophenylpiperazine; 2-DPMP = 2-diphenylmethlpiperidine; 3-FMC = 3-fluoromethcathinone; MDAI = 5,6-methylenedioxy-2-aminoindane; MDMA = 3,4-methylenedioxyamphetamine; MDPV = 3,4-methylenedioxypyrovalerone; 25I-NBOMe = 4-ido-2,5-dimethoxy-N-(2-methoxybenzyl)phenethylamine; PMA = paramethoxyamphetamine; TFMPP = trifluoromethylphenylpiperazine.
References


41 Carmo H, Remiao F, Carvalho F, Fernandes E, de Boer D, dos Reys LA, et al. 4–Methylthioamphetamine–induced hyperthermia in mice:


Figure 1

Structures of representative novel psychoactive substances (stimulants and empathogens). Chemically, the substances shown include amphetamines (amphetamine, methamphetamine, MDMA, PMA), cathinones (β-keto-amphetamines such as methedrone, mephedrone, methylone, pentylone, buphedrone, methcathinone, α-PVP), an aminoindane (MDAI), piperazines (mCPP and BZP) and pipradrols (2–DPMP and methylphenidate). Pharmacologically, these substances are psychostimulants or empathogens and primarily interact with monoamine transporters.
Figure 2
Relative dopamine/serotonin inhibition potencies of selected novel psychoactive substances. Dopamine to serotonin transporter (DAT/SERT) inhibition ratios (mean ± 95% confidence intervals) for novel substances are shown in comparison with those of classic empathogens/entactogens (MDMA, ecstasy) and stimulants (cocaine, amphetamine, and methamphetamine). The ratios derived from in vitro studies help to predict the typically unknown clinical toxicity of novel substances. A low DAT/SERT inhibition ratio (<0.1) indicates tenfold greater relative serotonergic vs dopaminergic activity similar to MDMA. A high DAT/SERT inhibition ratio (>10) indicates greater relative dopaminergic vs serotonergic activity similar to methamphetamine. A high DAT/SERT inhibition ratio is a pharmacological characteristic associated with more stimulant effects and with higher potential for addiction.

Figure 3
Structures of representative novel psychoactive substances (hallucinogenic phenethylamines, furans and difurans, and hallucinogenic tryptamines). Chemically, the shown substances include phenethylamines of the 2–C series (2C-B, 2C-I, and 25I-NBOMe) and of the 2–D series (DOM and DOI), a bezofuran (6–APB), a benzodifuran (2C-B-fly), and tryptamines (DMT, AMT, and 4–OH-MET). Pharmacologically, all presented substances are hallucinogens and potent serotonin 5–HT2A receptor agonists except for 6–APB which is an empathogen and indirect serotonin agonist similar to MDMA.