



An Overview of Recent Developments in the Analytical Detection of New Psychoactive Substances (NPSs)

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5 **of New Psychoactive Substances (NPSs)**
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Abstract

Novel psychoactive substances (NPSs), sometimes referred to as “*legal highs*” in more colloquial environments/the media, are a class of compounds that have been recently made available for abuse (not necessarily recently discovered) which provide similar effects to the traditional well studied illegal drugs but are not always controlled under existing local, regional or international drug legislation.

Following an unprecedented increase in the number of NPSs in the last 5 years (with 101 substances discovered for the first time in 2014 alone) its, occasionally fatal, consequences have been extensively reported in the media. Such NPSs are typically marketed as ‘not for human consumption’ and are instead labelled and sold as plant food, bath salts as well as a whole host of other equally nondescript aliases in order to bypass legislative controls. NPSs are a new multi-disciplinary research field with the main emphasis in terms of forensic identification due to their adverse health effects, which can range from minimal to life threatening and even fatalities. In this mini-review we overview this recent emerging research area of NPSs and the analytical approaches reported to provide detection strategies as well as detailing recent reports towards providing point-of-care/in-the-field NPS (“*legal high*”) sensors.

Keywords: New Psychoactive Substance(s); “*legal highs*”; designer drugs; synthetic cannabinoids; substituted cathinones; Spice; mephedrone; analysis.

Introduction

Novel Psychoactive Substance (NPS) is an umbrella term to refer to substances which mimic the effects of common illicit materials (for example, methamphetamine and cannabis) however they are not controlled by drug legislation such as the Misuse of Drugs Act¹ in the United Kingdom and other similar controls internationally. Designed, in some cases deliberately, to evade international control, NPSs may pose a significant danger to the health and safety of the public. As with controlled substances, NPSs are understood to have potentially negative short-term side effects such as paranoia, psychosis and seizures however these may not always be fully understood on account of the materials often being fairly new and understudied, as such their long term health risks are also not always clearly understood.² The United Nations Office on Drugs and Crime (UNODC) and European Monitoring Centre for Drugs and Drug Addiction (EMCDDA) standardised the term “*New Psychoactive Substance(s)*” and detailed the following sub-categories: Synthetic cannabinoids, Synthetic cathinones, Ketamine, Phenethylamines, Piperazines, Plant-based substances: khat, kratom, salvia divinorum and Miscellaneous: aminoindanes, phencyclidine, tryptamines.

Given the nature of NPSs underhanded production, purposely designed to evade international drug legislation, they are intrinsically marketed and sold as “*legal highs*”. Easily available at ‘head shops’ (a commercial outlet selling cannabis and tobacco paraphernalia), market stalls and the internet; vendors of NPSs are often operating on the edge of legality by being both vague and creative in their description of the products contents and its purported uses. NPSs may be sold as research chemicals, plant food, bath salts, exotic incenses *etc* together with slightly more telling descriptors such as: party pills, herbal highs and smoking blends although these names can often be mercurial, for example, mephedrone (a synthetic cathinone) pre-control was plant food whereas after becoming a controlled substance it was referred to as a ‘research chemical’.

Although given these nondescript aliases, NPSs products often have brand names; examples of “*legal high*” brand names are ‘Benzo Fury’, ‘Afghan Incense’, ‘NRG-1’ and ‘NRG-2’. The name or description given to a NPSs or “*legal high*” product may not always pertain to what is the actual psychoactive substance present, for example mephedrone was detected in products sold as naphyrone or NRG-1 in the UK even after its ban³, another survey found 70% of NRG-1 and NRG-2 products examined contained mixtures of substituted cathinones and not, at the time uncontrolled, naphyrone.⁴ Clearly there are no assurances to the customer of these NPS products that the contents are the same as advertised,

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3 furthermore they may be unwittingly violating drug legislation as the contents within are
4 controlled substances.
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6 Abuse of NPSs has been reported to be increasing since *ca.* 2009 and has continued to
7 be an ever growing market⁵ emerging at an unprecedented rate something also reflected in the
8 online marketplace with the number of online vendors in the UK increasing by more than
9 300% between 2010 and 2011.⁶ New materials made available for abuse appear rapidly and,
10 at times, can gain a ‘foothold’ in the market – such as mephedrone. In 2014, 101 new
11 substances were reported for the first time to the EU early warning system (EWS) run by the
12 EMCDDA up from 81 in 2013 which is also an increase from the 74 substances notified in
13 2012.⁷ Of the findings of the EWS synthetic cannabinoids are the most frequently discovered
14 with 102 detected between 2005 and 2013. A graphical representation of NPSs notified to the
15 EWS between 2005-2014 is shown in Figure 1.⁷
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23 The media has reported on numerous deaths related to “*legal highs*” and given the
24 wide variety of NPS and the ever-changing composition of existing products, a completely
25 new field of research has emerged in the continual development of analytical techniques
26 along with presumptive tests and in-the-field sensors. To date, there are reviews on the
27 chemistry, pharmacology and toxicology of NPSs but no comprehensive review of the
28 current techniques for the analysis of these substances has, to-date, been compiled.
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33 In this mini-review a thorough overview of this new analytical field of NPSs is provided
34 which covers: synthetic cannabinoids (most frequently discovered NPS by the EWS),
35 synthetic cathinones; particularly mephedrone (amidst reports by the Crime Survery for
36 England and Wales [CSEW] detailing mephedrone as the most prevalent of abused NPSs)
37 and in lesser detail pieces of interesting research of the other NPSs notified to the EWS
38 (Visible in Figure 1).⁷
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Synthetic Cathinones

Synthetic cathinones are an amphetamine-like cheap alternative to Ecstasy derived from cathinone; an organic stimulant found in Khat – a plant native to East Africa and the Middle-East and they possess pharmacological similarity to the phenethylamine class of psychoactives (*e.g.* amphetamine and methamphetamine). The effects of synthetic cathinones on the body are reported to have both cardiovascular and neurological side-effects; believed to block the reuptake of norepinephrine, dopamine and serotonin⁸ whilst there are also reports that they also induce the release of more dopamine⁹ suggesting synthetic cathinones act like both methamphetamine and cocaine synchronously.⁸⁻¹²

Internationally there has been a tightening of the legislation regarding synthetic cathinone derivatives, for example, cathinones are illegal in the UK as well as Germany, The United States, Canada and many others.^{13, 14} The European Monitoring Centre for Drugs and Drug Addiction's (EMCDDA) Early Warning System (EWS) has reported 74 new synthetic cathinones between 2005 and 2014, with 30 new substances discovered in the year 2014 alone (Figure 1). Clearly, the epidemic initiated by synthetic cathinones is showing no signs of cessation within the near future hence the development of methods for their detection and quantification is timely and urgently required. Mephedrone in particular, since its availability for abuse, is popular amongst users of “*legal high*” products and despite its classification in 2009 reports from the Crime Survey for England and Wales (CSEW) reveal mephedrone was still being abused in England and Wales in 2014.

Popularly known as ‘bath salts’, ‘research chemicals’ or ‘plant food’, synthetic cathinones are sold under, often mercurial, non-descript brand names such as ‘Energy’ (NRG), Blizzard and Ivory Snow containing warning labels such as ‘not for human consumption’ or ‘not tested for hazards or toxicity’ in an attempt to bypass legislative controls. The active component in a “*legal high*” product can vary wildly, even within the same brand name;^{3, 10, 15} for example mephedrone was detected in products sold as naphyrone or NRG-1 in the UK even after its ban³ while another found 70% of NRG-1 and NRG-2 products examined contained mixtures of substituted cathinones and not, at the time uncontrolled, naphyrone.⁴ Clearly there are no assurances to the customer of these NPS products that the contents are the same as advertised (if at all) and furthermore the customer may be unwittingly violating drug legislation if the products contain controlled substances.

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3 The list of case reports concerning synthetic cathinone-induced intoxication is
4 extensive and ever increasing. In the United States the number of calls to emergency centres,
5 as a result of synthetic cathinone abuse, increased from 303 to 6,100 between 2010 and 2011.
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7 A plethora of case reports are reported in the literature and media spanning a sizeable age
8 range, including both of the sexes and include fatalities and the curious report of the murder
9 of a goat whilst dressed in lingerie.¹⁶ For instance, a female aged 15 had symptoms of nausea,
10 vomiting, altered mental status, euvoaemic hypo-osmotic hyponatremia with encephalopathy
11 and increased intracranial pressure – mephedrone metabolites were found in her urine.¹⁷ A
12 male aged 31 after admitting to taking three 1500 mg packets of “bath salts” and was reported
13 to have hallucinations, paranoia, agitation; elevated serum CPK level, hyperkealemia,
14 dehydration, rhabdomyolysis and acute renal failure.
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21 Considering all the synthetic cathinones discovered, there can be no assertions to
22 which are the being abused but what is evident from the literature is that the most prominent
23 synthetic cathinones found within “legal high” products globally are mephedrone (4’-
24 methylmethcathinone; 4-MMC) and 3’,4’-methylenedioxypropylvalerone (MDPV).¹⁵
25 Mephedrone is more prevalent in Europe and MDPV in the United States;¹⁵ a list of the most
26 prevalent cathinone derivatives¹⁸ abused worldwide can be found in Table 1 although the
27 focus of the review will apply generally towards the detection and quantification of
28 mephedrone.
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35 Studying the patterns of NPS abuse can be difficult as it is frequently based upon self-
36 reported user surveys.¹⁹ This is potentially problematic as in many instances users are, due to
37 poorly labelled products (see earlier), not in fact aware of the substances they are taking. In
38 light of this, numerous groups are making advances towards screening the current NPSs
39 being abused. A number of revered groups using a range of chromatographic techniques
40 including HPLC and GC-MS, with LC-MS methods seemingly the preferred and established
41 technique of choice, have published exhaustively upon the laboratory-based analysis of
42 synthetic cathinones,^{3, 20-40} phase I and II metabolites^{41, 42} and more recently, in light of the
43 often nonenantioselective NPS synthesis, chiral separation of racemic mixtures.⁴³
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50 In 2014 Archer *et al.*¹⁹ analysed urine samples collected from a night club over one
51 weekend. The manuscript with its real and imaginative title, “*Taking the Pissoir – a novel*
52 *and reliable way of knowing what drugs are being used in nightclubs*”, reported the detection
53 of classical recreational drugs and NPSs such as: mephedrone, 3’-
54 trifluoromethylphenylpiperazine and 2-aminoindane using various chromatographic and mass
55 spectrometric methods.¹⁹ Furthermore parent drug/metabolites were also detected for
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3 amphetamine, cocaine, ketamine, 3',4'-methylenedioxyamphetamine (MDMA),
4 mephedrone and 3-trifluoromethylphenylpiperazine (3-TFMPP); this is important as it
5 indicates drugs were being used and not simply discarded into the urinal.¹⁹ In the same year,
6 Leffler *et al.*⁴⁴ (located in the United States) analysed 14 separate street samples wherein 10
7 synthetic cathinones were identified employing a variety of techniques, including gas
8 chromatography with mass spectrometric detection (GC-MS) and flame ionization (GC-
9 FID).⁴⁴ HPLC direct infusion tandem mass spectrometry (MS/MS) was also used to identify
10 compounds which were not available as reference materials. Out of the synthetic cathinones
11 detected:
12 3',4'-methylenedioxypropylvalerone (MDPV), 3',4'-methylenedioxy- α -
13 pyrrolidinobutiophenone (MDPBP), 4'-fluoromethcathinone (4-FMC), butylone,
14 mephedrone, naphyrone, 4'-methylethcathinone (4-MEC), ethcathinone, α -
15 pyrrolidinopentiophenone (α -PVP), and 3'-methyl- α -pyrrolidinopropiophenone (3-MPPP).
16 MDPV was the most prevalent, found in five of the 14 samples and ranging from 11% to
17 73% (w/w) between samples.⁴⁴

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27 Earlier reports in Denmark, Pedersen *et al.*³⁵ presented an automated solid-phase
28 extraction (SPE) and ultra-high-performance liquid chromatography (UHPLC) with TOF-MS
29 screening method for 256 illicit compounds in blood and 95 of these compounds were
30 validated with regard to matrix effects, extraction recovery, and process efficiency with the
31 limit of detection (LOD) ranging from 0.001 to 0.1 mg kg⁻¹.³⁵ Application of the technique to
32 the analysis of 1335 forensic traffic cases revealed 992 cases (74%) were positive for one or
33 more traffic-relevant drugs above the Danish legal limits. Commonly abused drugs such as
34 amphetamine, cocaine, and frequent types of benzodiazepines were the major findings.
35 Nineteen less frequently encountered drugs were detected: buprenorphine, butylone, cathine,
36 fentanyl, lysergic acid diethylamide, *m*-chlorophenylpiperazine, MDPV, mephedrone, 4'-
37 methylamphetamine, *p*-fluoroamphetamine, and *p*-methoxy-*N*-methylamphetamine.³⁵

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Even as early as 2011, there have been numerous attempts at constructing screening
methods for substituted cathinones in a number of different matrices, Bell and co-workers²⁹
reported a rapid multi-analyte direct urinalysis LC-MS/MS screening method being able to
detect eight analytes including; 4'-methylmethcathinone (mephedrone), 3',4'-
methylenedioxyamphetamine (bk-MDMA, 'methydone'), 4'-methoxymethcathinone (bk-
PMMA, 'methedrone') and 3', 4'-methylenedioxypropylvalerone (MDPV).²⁹ Using a dilution
of 1 part urine to 4 mobile phase to reduce matrix effects and although not all compounds
were completely chromatographically resolved, there was sufficient specificity to allow target

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3 analyte identification. All the analytes were readily detected at a concentration of 500 ng mL⁻¹
4 offering an attractive method for the routine screen of NPSs.²⁹ The global impact of
5 synthetic cathinones is compounded when substances such as mephedrone and MDPV have
6 been detected following sewage-based epidemiology in Chinese ‘megacities’.⁴⁵
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10 In terms of quantification, Santali *et al.* provided the first fully validated HPLC
11 method for the quantification of mephedrone²² where limits of detection and quantification of
12 0.1 and 0.3 µg mL⁻¹ respectively were reported. Khreit *et al.* further refined this method
13 enabling the detection of both mephedrone and two novel derivatives, 4'-methyl-*N*-
14 ethylcathinone (4-MEC) and 4'-methyl-*N*-benzylcathinone (4-MBC), in seized samples of
15 “NRG-2”. In this case the limits of detection and quantification were reported as 0.03 and
16 0.08 for 4-MEC and 0.05 and 0.14 µg mL⁻¹ for 4-MBC both in their pure form and in the
17 presence of common adulterants such as caffeine and benzocaine.^{3, 23} There has also been
18 work using chromatographic methods on the detection of cathinone based “legal highs” in
19 biological matrices^{24, 37} in which Beyer *et al.* were able to detect and quantify 25 designer
20 cathinones in a validated LC-MS-MS method.³⁷
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28 Other work²⁶ has seen an attempt to screen chronic abuse of mephedrone through GC-
29 MS analysis of hair. The hair was first decontaminated in methylene chloride and incubated
30 overnight in a pH 7 buffer in the presence of deuterated MDMA at 40 degrees Celsius. The
31 work saw 67 hair specimens tested for mephedrone with 13 yielding positive results of
32 concentrations ranging from 0.2 - 313.2 ng mg⁻¹.²⁶ The work showed that like other stimulant
33 drugs, mephedrone is well incorporated into hair and the analytical method reported appears
34 sensitive enough to reveal occasional to regular use of mephedrone.²⁶
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40 Recently direct analysis in real time mass spectrometry (DART-MS) has been utilised
41 to quantify and characterise the multitude of new and emerging NPSs.⁴⁶ Solid synthetic
42 cathinone samples (2-FMC, 2-MEC, 2-FEC and 2-EEC) were sampled directly without pre-
43 treatment and positive ion mass spectra were acquired using a DART-SVP™ ion source
44 interfaced to an AccuTOF mass spectrometer. Further advancements in this methodology by
45 the same authors⁴⁷ has seen the application of a time-of-flight (TOF) mass analyzer along
46 with in-source collision-induced dissociation (CID) spectra to provide data for presumptive
47 analysis of various synthetic cathinones in a similar fashion to GC-MS analysis.⁴⁷ The
48 authors scope for this work is to provide a rapid screening method to quickly respond to the
49 rapid evolution of designer drugs and the consequent testing backlogs that develop.^{46, 47} Ion
50 mobility spectrometry (IMS) has also been applied to the screening of an array of NPSs
51 within the literature with acceptable results.^{48, 49}
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3 Smith *et al.*^{50, 51} provided an alternative to chromatography and proposed a novel
4 sensing protocol based upon the electrochemical methods. Of note is the reduction of the
5 cathinone substitutes; mephedrone and 4-MEC with a scope to provide an on-the-spot
6 analytical screening tool with cyclic voltammetry.⁵⁰ Analysed in pH 4.3 acetate buffer, limits
7 of detection were found to correspond to 11.80 $\mu\text{g mL}^{-1}$ for 4-MMC and 11.60 $\mu\text{g mL}^{-1}$ for 4-
8 MEC.⁵⁰ This work demonstrated for the first time a rapid, accurate, and sensitive method for
9 the quantification of synthetic cathinone components found in seized street “*legal high*”
10 samples (NRG-2) *via* the use of an electrochemical protocol utilizing graphite screen-printed
11 electrodes (GSPEs) which was also independently verified with HPLC.⁵⁰
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18 Interesting developments in the detection of synthetic cathinone derivatives is the use
19 of surface enhanced Raman-spectroscopy (SERS) have also been reported.^{52, 53} In this novel
20 approach, the usually required thin metallic surface (typically gold or silver) was provided by
21 galvanising a British two pence coin with silver. Note that a pre-1992 two pence coin (97%
22 Copper) is required as post-1992 two pence coins are composed of copper-plated steel and
23 have an undefined composition.⁵² Figure 2 shows the concept when dendritic structures are
24 evident on the two pence surface, providing proof of concept for SERS detection of
25 mephedrone, MDMA and aminoindane 5',6'-methylenedioxy-2-aminoindane (MDAI) was
26 demonstrated.⁵² Further developments saw the researchers working towards a new
27 optimization strategy for the SERS detection of mephedrone using a portable Raman system
28 employing a fractional factorial design approach to significantly reduce the statistical
29 experiments whilst maintaining statistical integrity.⁵³ Furthermore, four optimised SERS
30 protocols for which the reproducibility of the SERS signal and the limit of detection of
31 mephedrone were established with an estimated limit of detection of 1.6 $\mu\text{g mL}^{-1}$.⁵³
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42 Another alternative to the well-established chromatographic methods, NPS detection
43 has been reported with the use of immunochemistry, Paillet-Loilier *et al.*¹⁸ noted the use of
44 this technique to test the cross-reactivity of some synthetic cathinones using the semi-
45 quantitative AxSYM amphetamine/methamphetamine II assay in tandem with Fluorescence
46 Polarization Immunoassay (FPIA). Evaluating the responses from aqueous solutions of 14
47 substituted cathinones at 1 mg/L, 10 mg/L and 100 mg/L, the authors observe pentedrone,
48 pentylone, α -pyrrolidinovalerophenone (PVP), and 3',4'-methylenedioxypropylone
49 (MDPV) did not react with the protocol. Some synthetic cathinones, however, reacted in the
50 assay at 10 mg/L: ethylone, mephedrone, methylone, methedrone, and 4'-methylethcathinone
51 (MEC) scrutiny of this reveals that each of these that did react had the least substitutions on
52 the ethylamine chain suggesting the method has limitations to larger molecules.¹⁸
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3 Commercially available enzyme-linked immunosorbent assays have been used to analyse
4 eight synthetic cathinone derivatives amongst 30 designer drugs.⁵⁴ The test demonstrated
5 cross-reactivity at concentrations as low as 0.15 mg L⁻¹ when tested against the Randox
6 Mephedrone/Methcathinone ELISA kit (RANDOX Toxicology, Crumlin, UK), a protocol
7 recently developed for forensic specific cathinone screening in urine and blood specimens.⁵⁴
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11 Presumptive testing of cathinone derivatives was carried out by Nic Daeid and
12 colleagues⁴⁰ as per United Nations recommended guidelines. Various presumptive tests were
13 investigated, however results suggested the Zimmerman test, which relies on the presence of
14 a carbonyl group in close proximity to a methyl group on the same molecule and reaction
15 with 2', 4'-dinitrobenzene to form a Meisenheimer reddish-purple colour, was the most
16 consistently effective test method. A small amount of each test sample was placed into a well
17 of a spotting tile and 2 drops of 1% 2', 4'-dinitrobenzene in methanol followed by 2 drops of
18 15% potassium hydroxide in water were added. Any colour change or other noticeable effect
19 occurring immediately on addition of the reagents was noted and observations were made
20 again after 5 minutes; Specific colour changes were observed in all cases apart from
21 bupropion. Nic Daeid *et al.* have also reported using stable isotopic fractionation/profiling
22 (isotope ratio mass spectrometry; IRMS), to provide a potentially quantifiable link between
23 the precursor (4'-methylpropiofenone) and the illicit drug product (4'-
24 methylmethcathinone) for a particular manufacturer and synthetic route of mephedrone.⁵⁵
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Synthetic Cannabinoids

Synthetic cannabinoids emerged as a recreational product *ca.* 2008 in the form of aminoalkylindoles such as JWH-018. They were originally investigated by Professor Huffman⁵⁶ as therapeutic compounds, however they were subsequently abandoned due to the unwanted psychoactive side effects. Despite many classes synthetic cannabinoids becoming controlled under drug legislation, there are still many which remain legal whilst still posing threat to the population. As with synthetic cathinone derivatives, there is often limited to no information on the packaging of the products and the active ingredients present can vary greatly between products of the same name.⁵⁷⁻⁶¹ These compounds were first introduced into products known as ‘K2’ and ‘Spice’ with the latter having a market range of: Spice Silver, Spice Gold and Spice Diamond.¹ The products, advertised as incense or smoking mixtures, are typically sold consisting of a few grams of finely cut green/brown plant material as to perhaps replicate the appearance of cannabis whilst being infused with the active synthetic cannabinoid component(s). There are instances of retailers selling the active components as research chemicals (similarly to synthetic cathinones) which arrive as a crystalline powder of high purity.⁶¹

There are various case reports to support the literature and media claims that synthetic cannabinoids have psychoactive effects akin to that of cannabis. Indeed, the components of Spice and related herbal products have been identified as aminoalkylindoles originally synthesised by Huffman and Atwood *et al.* and have demonstrated that JWH-018 is a potent and effective CB₁ receptor agonist.⁶²

Interesting case reports with regards to the effects of the Spice epidemic include a report by Schneir *et al.*,⁶³ who published case studies on two women admitted to a San Diego (USA) emergency department after smoking Spice “Banana Cream Nuke” – disorientated, feeling unusual and “*as if they did not know where they were*”⁶³. Another report describes three cases of the effects of Spice⁶⁴, all having a negative urine drug screen whilst exhibiting agitation, paranoia and tachycardia. Follow up analysis revealed the urine to contain metabolites of JWH-018 and JWH-073⁶⁴. More recent reports also highlight similar observations in adolescents and young adults after intoxication with synthetic cannabinoids.⁶⁵

¹ Ingredients listed on the packaging of products are as follows - Spice Gold: *bay bean, blue lotus, Lion's Tail, Indian Warrior, Dwarf Skullcap, Maconha brava, Pink Lotus, Marshmallow, Red Clover, Rose, Siberian motherwort, Vanilla and honey.* Spice Gold Spirit: *Leonurus, Cardiac, Pedicularis, Canadensis, Scutellaria, Latero flora, Athaea officinalis, Rosa damascene, Vanilla planifolia.* Spice Diamond: *Bay bean, Blue lotus, Lion's tail, Indian Warrior, Dwarf Skullcap, Maconha brava, Pink Lotus, Marshmallow, Red Clover, Rose, Siberian motherwort, vanilla, honey, aroma.* Note the lack of any real ingredients (chemical) and no mention of any aminoalkylindole (JWH compounds) or cyclohexylphenyls (CP compounds).

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3 Vardakou *et al.*⁶⁶ have given an overview of other case reports⁶⁶ and the psychoactive
4 properties of Spice products and “legal highs”.

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6 Laboratory analysis revealed the active components of first generation Spice and
7 related products to be, the previously mentioned, aminoalkylindoles such as JWH-018 and
8 also cyclohexylphenols such as CP-47,497. As their popularity rose through sales in so-called
9 ‘head shops’ as well as on the internet, the substances were legislated as illegal in most
10 countries worldwide;⁶⁷ the range of active synthetic cannabinoid components of first
11 generation Spice products can be observed in Scheme SPICE1. *Note:* the aminoalkylindoles
12 (see Scheme 1) are given the notation of JWH after the academic who first synthesised these
13 compound, Professor J.W. Huffman.
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20 Further confirmation of this came at the end of 2008 when the German company THC
21 Pharma reported JWH-018 was an active ingredient in Spice products.⁶⁸ Following on from
22 this Auwater *et al.*⁶⁹ and Uchiyama *et al.*⁷⁰ identified and characterized the CP 47,497-C8 (see
23 Scheme 1) as its isomer – a synthetic by-product in Spice Silver, Gold and Diamond as well
24 as in products named ‘Yuctan Fire’ and ‘Sence’ which is reported to have 5 to 10 times more
25 analgesic potency that tetrahydrocannabinol.⁷¹
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30 An interesting paper from the point of view of the medical staff that have had to deal
31 with the Spice usage patients has a light-hearted title of: “*Spice*” girls: *Synthetic cannabinoid*
32 *intoxication*.⁶³ The authors noted that a urine drugs-of-abuse immunoassay was negative for
33 amphetamines, barbiturates, benzodiazepines, benzoylecgonine (cocaine metabolite),
34 methadone and opiates, oxycodone, phencyclidine, propoxyphene and
35 tetrahydrocannabinoids. The residue of the patient’s Spice product “*Banana Cream Nuke*”
36 was found to contain the synthetic cannabinoids JWH-018 and JWH-073 (the chemical
37 structure can be seen in Scheme 1) through gas chromatography-mass spectrometry (GC-MS)
38 and high performance liquid chromatography with ultraviolet detection (HPLC-UV). The
39 report highlighted the need for drugs-of-abuse screenings to be able to detect the JWH class
40 of compounds, particularly within a clinical setting.
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48 In Germany Lindigkeit *et al.* analysed Spice Gold with a GC-MS method wherein the
49 herbal mixtures were ground and put through a two hour Soxhlet extraction with petroleum
50 ether.⁵⁸ Analysis revealed the samples contained CP 47,497-C8 and JWH-018 until German
51 health authorities on the 22nd January 2009 prohibited the sale of the active components
52 found in Spice - from this point JWH-018 was absent from Spice, however it wasn’t long
53 until a new analogue, JWH-073, was found to be contained in Spice products.⁵⁸ Because the
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3 manufacturers of such products can readily change the active components in Spice, a rapid
4 method of detecting prohibited compounds in the complex mixtures is highly sought after.

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6 To this end, Emanuel and co-workers⁶⁸ reported for the first time the components of
7 Spice “*Gold Spirit*” using GC-MS (following a simple liquid extraction) alongside the
8 analysis of Spice “*Gold*” and “*Diamond*”; at the time the three most popular Spice products
9 used. Results indicated that Spice “*Gold*” contained CP 47,497-C8 along with ethyl vanillin,
10 α -tocopherol and γ -tocopherol whereas Spice “*Diamond*” contained caffeine, α -tocopherol,
11 γ -tocopherol, palmitic acid along with CP 47,497-C8 and JWH-018. As for Spice “*Gold*
12 *Spirit*”, JWH-018 and α -tocopherol were found to be present.⁶⁸

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14 Other work has of course followed on the analysis of Spice and related herbal
15 products for instance Uchiyama and co-workers⁵⁹ who analysed 46 different herbal products
16 with 44 having synthetic cannabinoids as determined *via* GC-MS and LC-MS. Two major
17 cannabinoids were found; [2-hydroxy-4-(2-methylnonan-2-yl)phenyl]cyclohexan-1-ol
18 (cannabicyclohexanol) and JWH-018 and the analysis of the herbal product (amount of NPS
19 per gram) were found to range from 1.1 to 16.9 mg g⁻¹ and 2.0 to 35.9 mg g⁻¹ respectively.⁵⁹

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21 In addition to the identification of the chemical components contained within the
22 Spice product range there is a need to understand the effects of the synthetic cannabinoids on
23 the human metabolism. Sobolevsky⁷² reported for the first the time, urinary metabolites of
24 JWH-018; clearly highly useful for analysis of patients admitted to emergency departments
25 and for the development of point-of-care tests (see the story of the “*Spice girls*” earlier in this
26 mini-review). Using LC-MS and GC-MS, two main monohydroxylated metabolites were
27 identified which are almost completely glucuroconjugated with minor metabolites such as *N*-
28 despentyl hydroxy-, carboxy-, dihydroxy-, and reduced di- and trihydroxy-metabolites.⁷² It
29 should be noted the parent compound (JWH-018) was reported to not be detected in urine.⁷²
30 The authors observed that there are two main metabolites that are valuable for detection of
31 JWH-018 in post-administration urine and LC-MS is a more useful technique as minor
32 metabolites can also be analysed to support analytical findings.⁷² Different analytical
33 approaches on Spice and related products have been reported⁷³⁻⁷⁸ with literature reporting the
34 presence of new cannabimimetic compounds.^{60, 79, 80} Following this pioneering work, there
35 has been a pursuit of studying synthetic cannabinoids in urine.⁸¹⁻⁸⁷ Further work by Moran *et*
36 *al.*⁸⁸ has extended the work of Sobolevsky⁷² and validated an LC-MS/MS method for the
37 quantitation of human urine metabolites of JWH-018 and JWH-073. The work highlighted 6
38 metabolites for each molecule with the primary metabolites being distinguishable between
39 JWH-018 and JWH-073. The authors have also extended this using a solid-phase extraction
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3 approach.⁸⁹ One criticism of the above work exploring the metabolites in urine is the limited
4 population studies – clearly larger studies will be needed to further understand the
5 pharmacology of synthetic cannabinoids. Other research has been devised to quantify
6 cannabinoids in serum and blood.⁹⁰⁻⁹⁴
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10 A different strategy has been to analyse cannabinoids in hair to show long term past
11 consumption.⁹⁵ To this end, Hutter *et al.*⁹⁶ reported the hair testing of 22 synthetic
12 cannabinoids in human hair. The methodology involves a simple ultrasonication of the hair
13 sample in ethanol and has a limit of quantification (LOQ) of 0.5 pg mg⁻¹.⁹⁶ Perhaps more
14 interestingly, synthetic cannabinoids have even been found in the urine of US athletes
15 (although its use to enhance performance is questionable.). Urine samples were collected
16 from 5,956 athletes and analysed via high performance-liquid chromatography-tandem mass
17 spectrometry (HPLC-MS) for the presence of JWH-018, JWH-073 and their metabolites.⁹⁷ In
18 4.5% of the samples, metabolites of both synthetic cannabinoid compounds were detected;
19 metabolites of JWH-018 and JWH-073 (50%), JWH-018 (49%), and only JWH-073 (1%)
20 were detected in positive samples.
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28 The focus of the research above has focussed on laboratory based instrumentation,
29 rightly so in order to unambiguously quantify NPSs but as highlighted in the case of the
30 “*Spice girls*”, synthetic cannabinoids do not react using traditional THC immunoassay tests.
31 To this end Arnston *et al.*⁹⁸ have designed two enzyme linked immunosorbent assays for
32 detection of JWH-018 and JWH-250 in urine. The assay of JWH-018 has significant cross
33 reactivity with several synthetic cannabinoids and their metabolites contrary to the JWH-250
34 assay which exhibits limited cross-reactivity. To start, assays are calibrated at 5 ng mL⁻¹ with
35 the 5-OH metabolite of JWH-018 and the 4-OH metabolite of JWH-250. To validate the
36 method, 114 and 84 samples of urine for JWH-018 and JWH-250 respectively were used and
37 confirmed by using liquid chromatograph tandem mass spectrometry (LC-MS/MS) testing for
38 metabolites of JWH-018, JWH-019, JWH-073, JWH-250 and AM-2201. Accuracy was
39 deemed to be greater than 98% with 95% sensitivity and specificity for both assays.
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48 Another approach of interest is a presumptive test marketed by “Narcotic Testing
49 Supplies & Equipment Store”.⁹⁹ The test works by inserting a small quantity of a suspected
50 sample into a plastic ampoule containing 25 µL reagent and 150 mg of specially treated
51 absorbing crystals (sodium 36%, potassium iodide 98% and 0.2% ethanol) stirring and
52 comparing the colour of the liquid to a pre-determined colour chart clearly visible from
53 Figure 3 however the specificity of such a screening test is questionable.⁹⁹
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3 As components of Spice and related substances become banned, they are replaced
4 with a compound which exhibits similar psychoactive properties yet negating the
5 effectiveness of the newly introduced ban, see the paper: "*Spice: A Never ending story?*" for
6 example.⁵⁸ As such there is an urgent need for a faster laboratory method; for that reason
7 Emanuel *et al.* reported the use of solid probe mass spectrometry alleviating the need for any
8 sample pre-treatment such as liquid-liquid extraction.⁶⁸ Since α -tocopherol is always present
9 in the Spice herbs range, the authors demonstrated that once α -tocopherol was subtracted
10 from the obtained spectra, the fragmentation patterns of CP 47,497-C8 and JWH-018 become
11 'visible'.⁶⁸ This screening methodology is useful for the rapid analysis of the prohibited
12 substances within the Spice product range (as well as related substances) with a positive
13 response nullifying the need for any pre-treatment step (such as liquid-liquid extraction)
14 allowing a full quantification *via* GC-MS or similar approaches *i.e.* LC-MS. Work from
15 Lesiak *et al.*¹⁰⁰ has also attempted to rapidly detect synthetic cannabinoids without the need
16 for sample preparation with the use of direct analysis in real time mass spectrometry (DART-
17 MS)¹⁰⁰ being able to screen for AM-2201, JWH-122, JWH-203, JWH-210 and RCS-4.
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20 To highlight the ever moving field of "*legal highs*" with respect to synthetic
21 cannabinoids, in October 2012 new variants were reportedly found where the structures were
22 a modification of compounds from the 3-naphthoylindole series^{57-60, 69, 70, 80, 101-107} identified
23 from regular seizures made by police in Russia and Belarus.¹⁰¹ Shevyrin *et al.* have reported
24 on the analytical characterisation of these new class of synthetic cannabinoids using GC-
25 HRMS, UHPLC-HRMS, NMR and FT-IR¹⁰¹ providing robust and reliable confirmatory
26 analytical approaches. Reports from South Korea also highlight the ever-changing market
27 detailing the different synthetic cannabinoids which have been identified by their National
28 Forensic Service between 2009 – June 2013.¹⁰⁸ The authors note that whilst initially it was
29 largely naphthoylindoles (*e.g.* JWH-018, JWH-073), phenylacetylindoles (*e.g.* JWH-203,
30 JWH-250), benzoylindols (*e.g.* RCS-2, RCS-4) and CP-47,497 derivatives abused; after
31 legislative bans were introduced, gradually over time, the molecules identified became new,
32 typically halogenated, substances such as cyclopropylindoles (*e.g.* UR-144, XLR-11) and
33 adamantylindoles (*e.g.* APICA, APINACA)¹⁰⁸ which are represented in Scheme 2.
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51 Following the influx of new compounds, groups worldwide moved towards their
52 detection. Scheidweiler *et al.*¹⁰⁹ developed and validated a liquid chromatography–tandem
53 mass spectrometric (LC–MS/MS) method for simultaneously quantifying JWH-018, JWH-
54 019, JWH-073, JWH-081, JWH-122, JWH-200, JWH-210, JWH-250, JWH-398, RCS-4,
55 AM-2201, MAM-2201, UR-144, CP 47,497-C7, CP 47,497-C8 and their metabolites, and
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3 JWH-203, AM-694, RCS-8, XLR-11 and HU-210 parent compounds in urine.¹⁰⁹ Previously
4 there were no extensive synthetic quantitative methods reported in the literature until this
5 work which presented the novel LC-MS/MS protocol quantifying 20 synthetic cannabinoids
6 and 21 metabolites, and semi-quantifying 12 alkyl –hydroxy-metabolites.¹⁰⁹
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10 Continuing from this, another approach towards the detection of the new generation of
11 synthetic cannabinoid agonist, Mohr *et al.*¹¹⁰ applied Enzyme-Linked Immunosorbent Assay
12 (ELISA) towards one of the most prevalent synthetic cannabinoids in urine, UR-144, and
13 XLR-11. Once again testing in urine, the method was validated against liquid
14 chromatography-tandem mass spectrometry with 90 positive and negative control samples
15 for UR-144, XLR-11 and its metabolites.
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Miscellaneous

As reported in the introduction, the novel psychoactive substance epidemic is an ever growing market with a vast array of new materials discovered each year.⁷ To cover every known substance is beyond the scope of this review however; in this section, interesting pieces of research from around the world will be covered.

Piperazines

N-benzylpiperazine (BZP), the structure of which is shown in Scheme 3, is known to be a central nervous system stimulant with its effects reported to be similar to amphetamine in that it also triggers the release of dopamine and norepinephrine whilst inhibiting the uptake of dopamine, norepinephrine and serotonin.¹¹¹ Although BZP is structurally similar to amphetamine it is reported to have only one-tenth the potency.¹¹¹ Marketed as a ‘party pill’ before legal restrictions BZP was viewed as a safe alternative to amphetamines such as MDMA,¹¹² however recently it has varying degrees of legislative control internationally.¹¹³ Its appearance in “*legal high*” samples is still reported^{114, 115} however after being made illegal the prevalence of its use has declined; for example in New Zealand after being made a prohibited substance in 2008, the use of BZP amongst the general population dropped from 15.3% in 2006 to 3.2% in 2009.¹¹⁶

In the UK, the first deaths associated with BZP and 3-TFMPP were three separate fatalities wherein one of both of the drugs were confirmed to be present although not determined to be the direct mechanism of death.¹¹⁷ Dickson *et al.*¹¹⁸ reported that BZP, 3'-TFMPP and MCPP are present in ecstasy tablets since the former, in some nations, is a legal alternative to MDMA. The authors analysed 251 MDMA positive urine samples using GC-MS *via* a liquid-liquid extraction and pentafluoropropionic anhydride (PFPA) derivatisation as sample pre-treatment to screen for 33 drugs potentially present.¹¹⁸ In 36% of the sample, drugs other than MDMA were found to be present; BZP, 3-TFMPP and MCPP were detected in 15%, 7% and 1% of the samples respectively.¹¹⁸

A wide array of analytical approaches have been reported by many different authors such as LC-MS,^{24, 119} capillary electrophoresis,¹²⁰ HPLC-fluorescence,¹²¹ LC with diode array,^{122, 123} GC-MS¹²⁴⁻¹²⁶ and chemiluminescence.¹²⁷ Arbo and co-workers¹²⁸ provided a thorough overview of piperazine compounds as drugs of abuse with the full range of analytical techniques and matrices applied, readers are directed to this paper.¹²⁸

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3 It is clear, something that is generally the case with all “legal highs”, confirmatory
4 laboratory based analysis is well developed. Lesser developed, however, are approaches that
5 could adapted for used in-the-field or within a clinical setting where a near-instantaneous
6 response is required. To this end, currently there are no immunoassays for the detection of
7 piperazines derivatives¹²⁸ and cross-reactivity of these compounds in fluorescence
8 polarization immunoassay using AxSYM[®], amphetamine/methamphetamine assay has been
9 reported.¹²⁹

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11 Recently Philip *et al.*¹³⁰ have reported on the development and validation of a
12 specific colour test using 1', 2'-naphthoquinone-4-sulphonate (NQS) forming an intense
13 bridge orange-red complex with BZP at room temperature. The authors reported that
14 common cutting agents such as glucose and caffeine did not affect the test. 3-TFMPP, MCPP,
15 pCPP, MeOPP and piperazines produced an orange-red colour change where the apparent
16 brilliance of the BZP-NQS complex made it apparently to be distinguishable from the other
17 colour changes with the potential cross-reactants.

28 *Aminoindanes*

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30 Aminoindanes are a group of synthetic compounds characterised by the presence of
31 a phenethylamine skeleton, they are currently not controlled globally¹³¹ and have more
32 recently been found to be contained in “legal high” products sold as powders akin to
33 synthetic cathinones.^{132, 133} 2-Aminoindane has a basic ring structure that is similar to
34 amphetamine (and therefore by proxy, substituted cathinones also) that can be chemically
35 modified and the following derivatives (Scheme 4); 5', 6'-methylenedioxy-2-aminoindane
36 (MDAI), 5', 6'-methylenedioxy-*N*-methyl-2-aminoindane (MDMAI), 5'-iodo-2-aminoindane
37 (5-IAI), and 5'-methoxy-6'-methyl-2-aminoindane (MMAI) have all reportedly been found in
38 “legal highs”.¹³²

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40 A number of aminoindane compounds have been thoroughly characterized by Casale
41 and Hays¹³⁴ who provided analytical protocols in the form of NMR, MS and IR for 5-IAI, 4-
42 IAI, their synthetic intermediates and impurities in order to assist forensic analysts.¹³⁴ There
43 is other work that reports a LC-MS/MS screening method for 26 analytes,³⁴ including MDAI,
44 and such an approach is designed to provide screening, within a clinical toxicology setting,
45 for the potential misuse of “legal highs” via analysis of urine.³⁴

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47 Partiuclarly of note, work by Elie and co-workers reports that microcrystalline
48 identification of MDAI, mephedrone and *N*-benzylpiperazine (BZP) is possible.¹¹⁴ In this
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3 protocol the illicit compound is dissolved into methanol and diluted with water to produce a
4 content of 10% (v/v) with mercury chloride (10 gL^{-1} + 10% methanol) used as the
5 microcrystalline agent.¹¹⁴ This approach involves dropping 10 μL of the drug solution with
6 10 μL of the reagent solution onto a glass slide; the resulting structures were optically imaged
7 following assisted nucleation (gently swirling a plastic pipette tip in the freshly mixed
8 drop).¹¹⁴ Figure 4 shows the observed crystal structure which is compared to the crystal
9 structure of illicit drugs. The MDAI free base (Figure 4bi) was found to form flat serrated
10 blades of various dimensions which become irregular with increasing sizes. Smaller crystals
11 are observed to be single blades whereas larger crystals develop two dimensional bunch
12 structures - after drying larger blade crystals are evident. It was noted that crystals grew
13 within 60s following assisted nucleation indicating the potential for a fast presumptive test
14 strategy.¹¹⁴ The uniqueness of these tests were determined through comparisons of MDAI
15 structure with a range of illicit drugs, indicating that potentially this approach is feasible to
16 identify the MDAI structure in a real sample containing other illicit drugs. To this end the
17 authors¹¹⁴ purchased “legal high” samples and utilised their microcrystalline presumptive test
18 approach which when collaborated with FTIR/GC-MS.
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32 *Salvinorin A (Salvia divinorum)*

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35 *Salvia divinorum* is a hallucinogenic psychoactive herb local to Oaxaca in Central
36 Mexico and for centuries has been used by cultures indigenous to the region.^{135, 136} This rare
37 member of the mint family is also known as ‘magic mint’ and more colloquially: ‘ska Maria’,
38 ‘ska Pastora’, ‘hierba de Maria’, ‘hojas de la Pastora’ all names which pertain to the belief
39 that *S.divinorum* is the reincarnation of the Virgin Mary.¹³⁷ The use of this plant as a
40 psychoactive substance has spread globally, its major constituent – salvinorin A (SA) is a
41 known selective opioid antagonist and to this end emphasis in the literature has been put on
42 detecting SA.¹³⁵ A dosage between 200–500 μg of SA has been found to induce profound
43 hallucinations with feelings of physical or mental displacement as well as experiencing
44 extraordinary illusions.¹³⁸ Recently studies have postured SAs effects involve the
45 endocannabinoid system.¹³⁹
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53 To analyse intact *S. divinorum* leaves for the presence of SA there has been the
54 employing of both thin layer chromatography using desorption electrospray ionization mass
55 spectrometry (TLC-DESI-MS)¹⁴⁰ and thin layer chromatography teamed with gas
56 chromatography/mass spectrometry (TLC-GS/MS).¹⁴¹ By utilizing these techniques, the
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3 authors of both techniques were able to confirm the presence of salvinorin A in a submitted
4 plant material suspected to be *Salvia divinorum*.^{140, 141}
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6 Pichini and co-workers¹⁴² attempted the detection of Salvinorin A in different
7 biological matrices opposed to the solid leaf matter. Utilising a gas chromatography mass
8 spectrometric protocol, it was applied to detecting SA in plasma, urine, saliva and sweat.¹⁴²
9 Following validation with 17-alpha-methyltestosterone as an internal standard the method
10 was applied to the analysis of urine, saliva and sweat from two consumers after smoking 75
11 mg plant leaves. Salvinorin A was detected in urine (2.4 and 10.9 ng/mL) and saliva (11.1
12 and 25.0 ng/mL), but not in sweat patches from consumers.¹⁴² The quantification of SA in
13 plasma and cerebrospinal fluid (from a rhesus monkey) has also been attempted and
14 sucesfully completed using a negative ion liquid chromatography-mass spectrometry
15 atmospheric pressure chemical ionization (LC-MS/APCI).¹⁴³ Using the United States Food
16 and Drug Administration (FDA) guidelines the authors of the method concluded the
17 technique had a lower limit of quantification (LLOQ) of 2 ng/mL for 0.5 mL of plasma
18 samples over the linear range 2-1000 ng/mL.¹⁴³
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31 *Mitragynine (Kratom)*

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34 Mitragynine is an indole alkaloid derived from the plant *Mitragyna speciosa* which is
35 indigenous to Thailand and other Southeast Asian countries. This is a common “legal high”
36 and is known commonly as *Kratom* which is also the chemical’s Thai name. The leaves of
37 the *M. Speciosa* was historically used as an opium substitute as well as being used
38 traditionally by villages in southern Thailand as a medicine for diarrhoea, muscle pain and
39 hypertension in addition to also being used by agricultural workers and labourers to relieve
40 tiredness and improve efficiency.¹⁴⁴ Its study remains pertinent as reports of a fatality
41 associated with *Kratom* are as recent as 2013.¹⁴⁵
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48 Interestingly, mitragynine is the major constituent of *Kratom* reported to be 66.2%
49 based on the crude base from the young leaves.^{146, 147} Levels of mitragynine in adults plants
50 from Thailand have been reported to be approximately over 60% whereas in Malaysia only
51 over 10%. Payanmtheine and the mitragynine diastereomer speciogynine were the second
52 most abundant alkaloids and the mitragynine diastereomer speciogynine was the third
53 abundant alkaloid in both plants.¹⁴⁸
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3 The pharmacology of mitragynine has been extensively studied and has been reported
4 to have analgesic activity on the opioid system.^{144, 149-151} Unlike the case of other NPSs
5 reported in this review where they have emerged and analytical techniques have had to be
6 developed/invented for their quantification, mitragynine, due to its historical use analytical
7 methods already exist and are generally applied to facilitate pharmacological studies. To this
8 end, Janchawee¹⁴⁴ reported the first analytical methodology utilising HPLC-UV. A linear
9 range of 0.1 – 10 µg mL⁻¹ was reported with a LOD of 0.03 µg mL⁻¹ and LOQ of 0.1 µg mL⁻¹.
10 Their protocol was applied to determine the pharmacokinetic characteristic of mitragynine in
11 the serum of rats following oral administration.
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18 As the leaves of *Kratom* became sold as “legal highs” in many other countries
19 Kikura-Hanajiri and colleagues¹⁴⁶ reported the detection of mitrogynine and 7-OH-
20 mitragynine (oxidative derivatives of mitragynine)¹⁵² in 13 “legal high” products using LC-
21 ESI-MS. The authors found that 11 of the 13 products were found to contain mitragynine and
22 7-OH-mitragynine with their content found to range from 1 to 6% and in the latter 0.01 to
23 0.04%.¹⁴⁶ Other researchers have directed research to study the methods of mitragynine in
24 biological matrices using LC-MS¹⁵³⁻¹⁵⁵ and UHPLC-UV.^{156, 157}
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30 From inspection of the literature, it is evident that there are multiple ways for the
31 detection and quantification of *Kratom* ingestion/consumption with detection levels as low as
32 0.02 µg mL⁻¹.¹⁵⁸ For example Arndt and co-workers reported a upon a case of a drug and
33 rehabilitation centre reporting an analysis for *Krypton* (another name for *Kratom*) in the urine
34 of a former opiate addicted woman.¹⁵⁹ The immunological drug screenings were performed
35 with test strips and a cloned enzyme donor immunoassay wherein alkaloids and tramadol
36 metabolites were analysed by LC-MS/MS. The immunoassays yielded negative responses for
37 amphetamines, barbituates, benzodiazepines, benzoylecgonine, buprenorphine,
38 ethylgluconoride, methadone, opiates, oxycodone and THC-COOH just as the test strips were
39 negative from tramadol and its metabolites. The LC-MS/MS detected the alkaloids typically
40 found in *Kratom* (mitragynine, speciociliatine, speciogynine, mitraciliatine and paynantheine
41 – detection of these alkaloids served sufficient proof of *Kratom* abuse and after confrontation
42 with data the patient admitted to several infusions of the plant.¹⁵⁹
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Conclusion and future challenges

The work described in this review demonstrates the range of new analytical methods and techniques applied to the detection and quantification of NPSs, which have recently emerged on the recreational drugs market. Given the rapidly evolving nature of the recreational drugs market, in terms of the number of new substances being identified (101 new substances, in Europe, in 2014); the ease at which these substances are available through on-line vendors or “*head shops*”; the freely-available information regarding NPS production and/or pharmacology and the lack of globalised drug/precursor control legislation - makes the current analytical, forensic and legal challenges clearly apparent. These issues coupled with the limited availability and range of certified primary reference standards; fully validated, simple and cheap laboratory-based analytical methods and selective and sensitive in-field testing technology highlights the growing gap in knowledge and necessitates economic investment and focused research in this underfunded area.

Future advances can be expected in the following areas: (i) Design and development of miniaturised in-field detection systems for NPSs in bulk samples or adulterated products (such as alcoholic drinks); (ii) Rapid, non-evasive bioanalytical methods for detection of the principle metabolites of common NPSs; (iii) simple, selective and validated laboratory-based chromatographic methods for the discrimination of new psychoactive substances, their isomers and their principle metabolites in biological matrices and; (iv) impurity profiling and/or source identification of common NPSs.

Clearly, the “*war on drugs*” is showing no sign of relenting in the near future and the principle challenge facing law enforcement agencies is to be ‘one-step-ahead’ of the clandestine drug manufacturers. By working collectively, analytical chemists, policy makers, law enforcement and forensic practitioners can suitably identify potential classes of molecules that may become the next generation of NPSs and develop advanced methods/technologies for the simultaneous detection/quantification of these substances thereby legislating against potentially dangerous compounds before they pose a serious threat to human health.

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Table 1 List of the most common synthetic cathinones, recreated from reference ¹⁸.

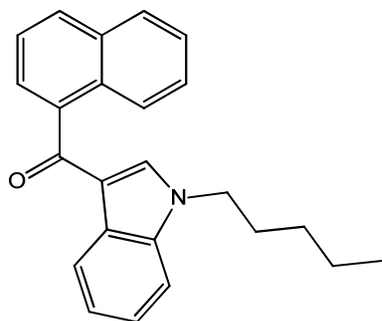
Usual names	Chemical name
Amfepramone or diethylpropion	2-diethylamino-1-phenyl-1-propanone
Benzedrone or methylbenzylcathinone or 4-MBC	1-(4-methylphenyl)-2-benzylamino-1-propanone
BMDB	2-benzylamino-1-(3,4-methylenedioxyphenyl)-1-butanone
BMDP or 3,4-MDBC	2-benzylamino-1-(3,4-methylenedioxyphenyl)-1-propanone
Brephedrone or 4-bromomethcathinone or 4-BMC	1-(4-bromophenyl)-2-(methylamino)-1-propanone
Buphedrone	2-(methylamino)-1-phenyl-1-butanone
Bupropion	1-(3-chlorophenyl)-2-(tertbutylamino)-1-propanone
Butylone or bk-MBDB	2-(methylamino)-1-(3,4-methylenedioxyphenyl)-1-butanone
Cathinone	2-amino-1-phenyl-1-propanone
Dibutylone or methylbutylone or bk-DMBDB	2-(dimethylamino)-1-(3,4-methylenedioxyphenyl)-1-butanone
Dimethylone or bk-MDDMA	1-(1,3-benzodioxol-5-yl)-2-(dimethylamino)-1-butanone
Dimethylmethcathinone or 3,4-DMMC	1-(3,4-dimethylphenyl)-2-(methylamino)-1-propanone
Ephedrone or methcathinone	2-(methylamino)-1-(4ethylphenyl)-1-propanone
Ethylbuphedrone or NEB	2-(ethylamino)-1-phenyl-1-butanone
Ethylcathinone or ethcathinone or ethylpropion	2-(ethylamino)-1-phenyl-1-propanone
Ethylmethcathinone or 4-EMC	2-(methylamino)-1-phenyl-1-propanone
Ethylone or bk-MDEA	2-(ethylamino)-1-(3,4-methylenedioxyphenyl)-1-propanone
Eutylone ou bk-EBDB	1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-1-butanone
Flephedrone or 4-fluoromethcathinone or 4-FMC	2-(methylamino)-1-(4-fluorophenyl)-1-propanone
Fluorocathinone or 'FC	2-amino-1-(4-fluorophenyl)-1-propanone
Fluoromethcathinone or 3-FMC	2-(methylamino)-1-(3-fluorophenyl)-1-propanone
Isoethcathinone	2-(ethylamino)-1-phenyl-2-propanone
Isopentadrone	2-(methylamino)-1-phenyl-2-pentanone
MDMPP	1-(3,4-methylenedioxyphenyl)-2-methyl-2-pyrrolidinyl-1-propanone
MDPBP	1-(3,4-methylenedioxyphenyl)-2-(1-pyrrolidinyl)-1-butanone
MDPPP	1-(3,4-methylenedioxyphenyl)-2-(1-pyrrolidinyl)-1-propanone
MDPV or MDPK	1-(3,4-methylenedioxyphenol)-2-pyrrolidinyl-1-pentanone
Mephedrone or 4-methylmethcathinone or 4-MMC	2-(methylamino)-1-(4-methylphenyl)-1-propanone
Metamfepramone or dimethylcathinone or dimethylpropion	2-dimethylamino-1-phenyl-1-propanone
Methedrone or 4-methoxymethcathinone or bk-PMMA	1-(4-methoxyphenyl)-2-(methylamino)-1-propanone
Methylbuphedrone or 4Me-MABP or bk-N-methyl-4-MAB	2-(methylamino)-1-(4-methylphenyl)-1-butanone
Methylethcathinone or 4-MEC	2-(ethylamino)-1-(4-methylphenyl)-1-propanone
Methylmethcathinone or 3-MMC	2-(methylamino)-1-(3-methylphenyl)-1-propanone
Methylone or MDMC or bk-MDMA	2-methylamino-1-[3,4-methylenedioxyphenyl]-1-propanone
MOPPP	4'-methoxy- α -pyrrolidinovalerophenone
MPBP	1-(4-methylphenyl)-2-(1-pyrrolidinyl)-1-butanone
MPHP	4'-methyl- α -pyrrolidinovalerophenone

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3	MPPP	4'-methyl- α -pyrrolidinovalerophenone
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5	Naphyrone	1-naphthalen-2-yl-2-pyrrolidin-1-yl-1-pentanone
6	Propylbutylone or bk-PBDB	2-(propylamino)-1-(3,4-methylenedioxyphenyl)-1-butanone
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8	Pentedrone or ethyl-methcathinone	2-(methylamino)-1-phenyl-1-pentanone
9	Pentylone	2-(methylamino)-1-(3,4-methylenedioxyphenyl)-1-pentanone
10	PBP	1-phenyl-2-(1-pyrrolidinyl)-1-butanone
11	PEP	1-phenyl-2-(1-pyrrolidinyl)-1-heptanone
12	PPP	1-phenyl-2-(1-pyrrolidinyl)-1-propanone
13	PVP	1-phenyl-2-(1-pyrrolidinyl)-1-pentanone
14		
15	Pyrovalerone	11-(4-methylphenyl)-2-(1-pyrrolidinyl)-1-pentanone
16	Trimethylmethcathinone or 2,4,5-TMMC	2-(methylamino)-1-(2,4,5-trimethylphenyl)-1-propanone
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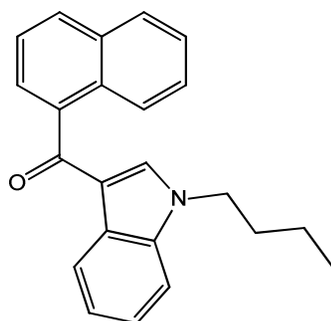
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6 **Scheme 1.** Chemical structures of synthetic cannabinoids found in herbal products such as the Spice range,⁶⁷ scheme reproduced from reference
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8 ⁶⁷ with permission from UNODC.
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11 **1) Aminoalkylindoles**

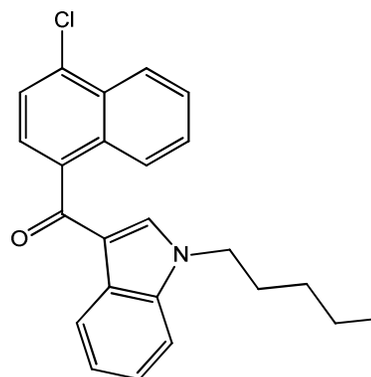
12 a) Naphthoylindoles



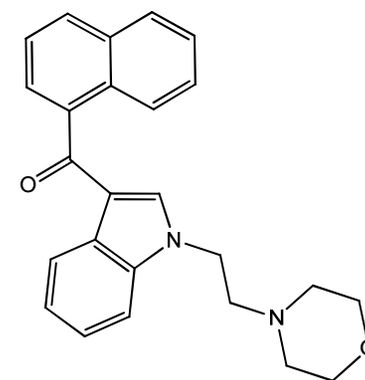
JWH-018



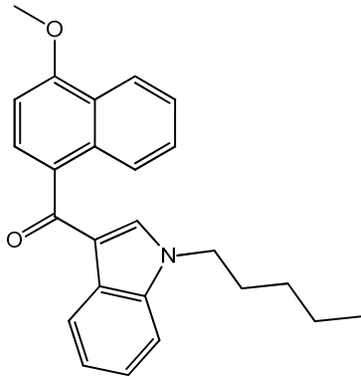
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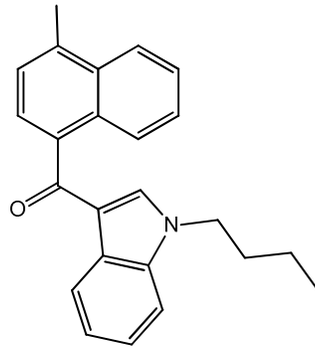
JWH-398



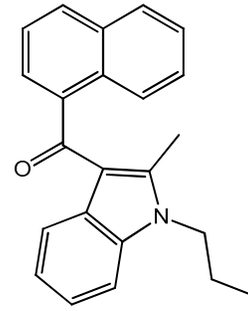
JWH-200



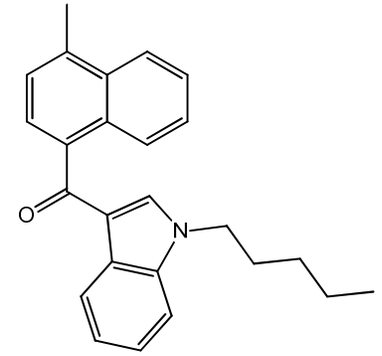
JWH-081



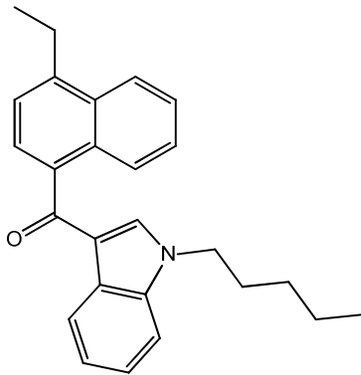
4-Methyl-JWH-073



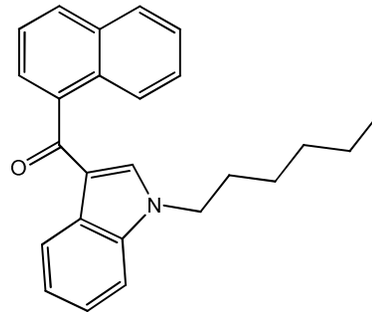
JWH-015



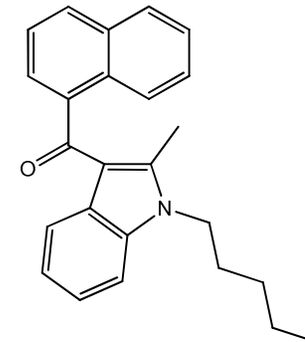
JWH-122



JWH-210



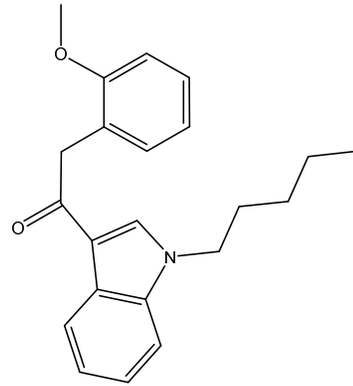
JWH-019



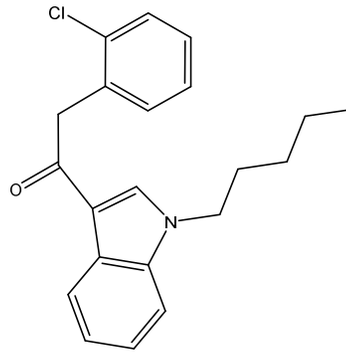
JWH-007

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b) Phenylacetylindoles

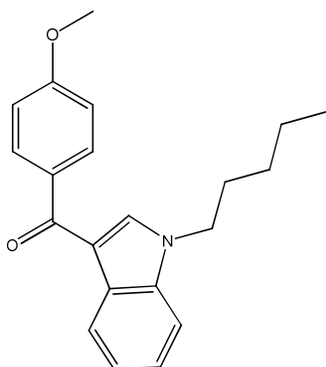


JWH-250

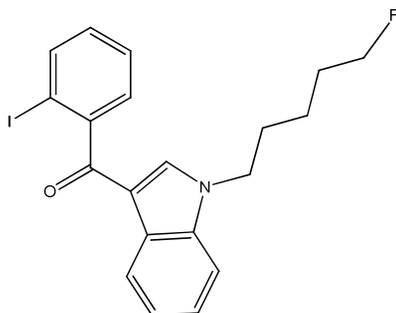


JWH-203

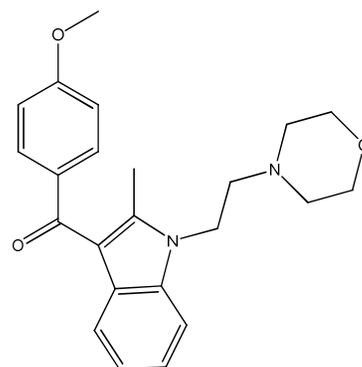
c) Benzoylindoles



RCS-4

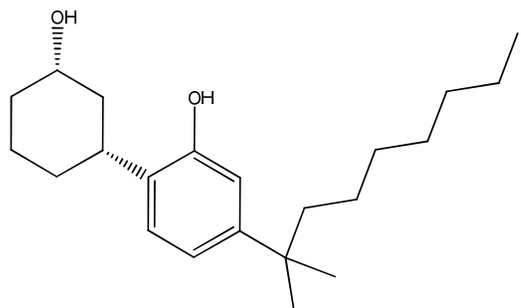


AM-694

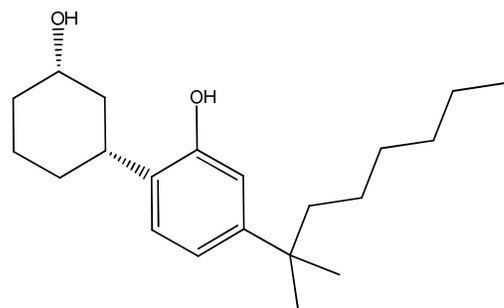


WIN-48,098

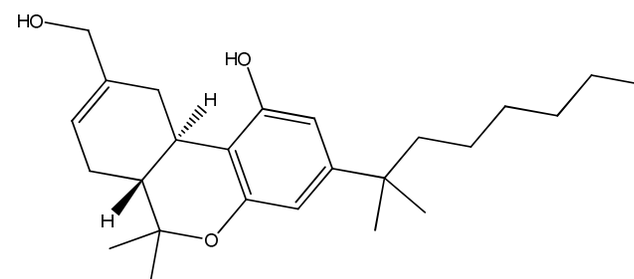
2) Cyclohexylphenoles



CP-47,497-C8

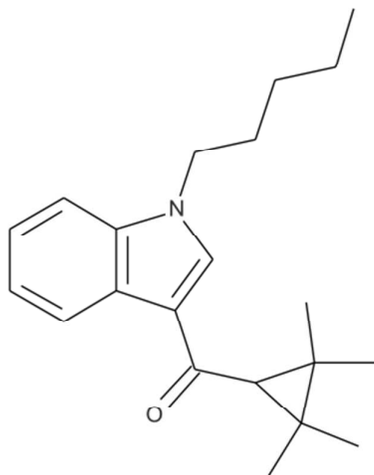


CP-47,497

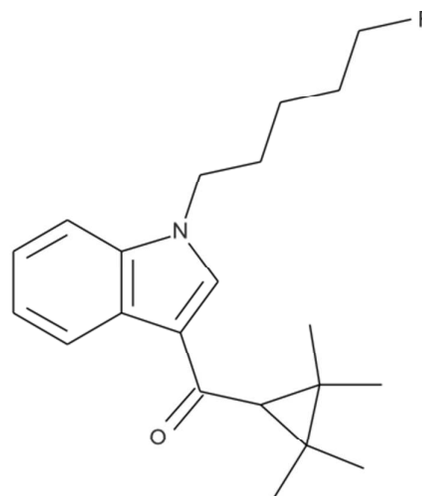


HU-210

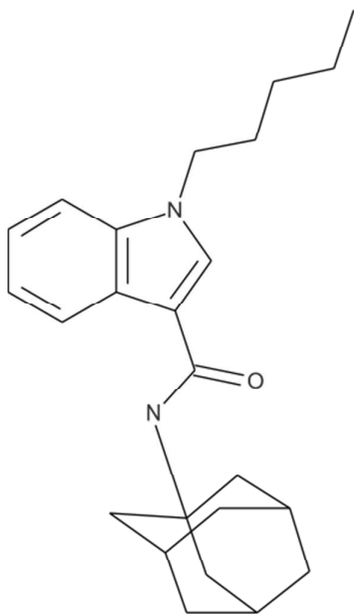
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4 **Scheme 2** Chemical structures of synthetic cathinones discovered after legislative bans were
5 introduced: cyclopropylindoles *e.g.* UR-144, XLR-11 and adamantylindoles (APICA and
6 APINACA)
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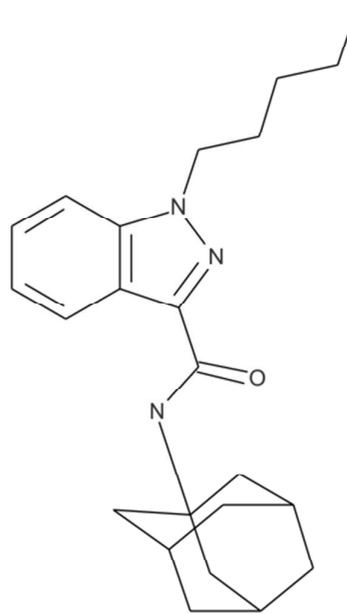
UR-144



XLR-11

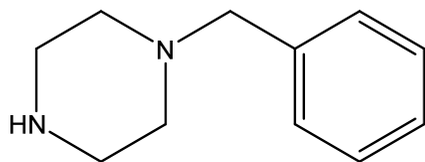


APICA

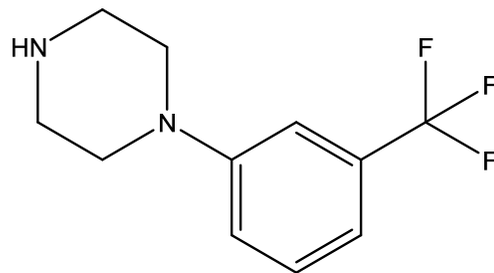


APINACA

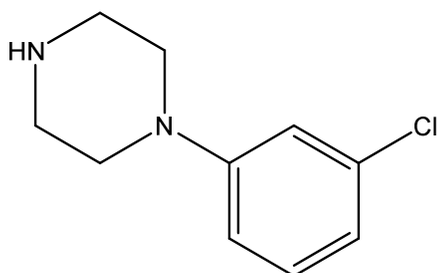
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3 **Scheme 3.** Benzylpiperazine and other piperazines derivatives which have been historically
4 abused.
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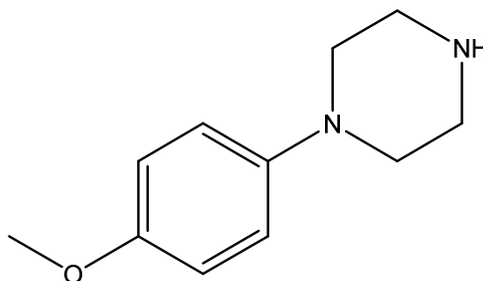
Benzylpiperazine



3-Trifluoromethylphenylpiperazine (TFMPP)

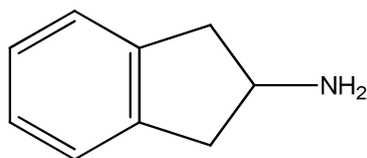


meta-Chlorophenylpiperazine (mCPP)

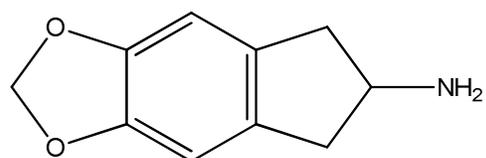


para-Methoxyphenylpiperazine (MeOPP)

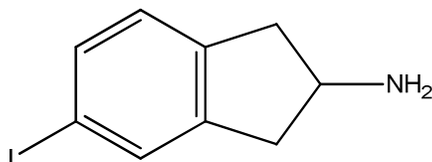
Scheme 4 2-Aminoindane and its derivatives, all of which have been found in “legal high” samples.



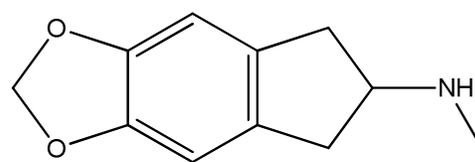
2-Aminoindane



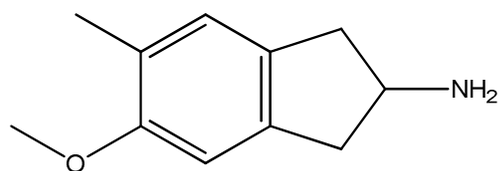
MDAI



5-IAI

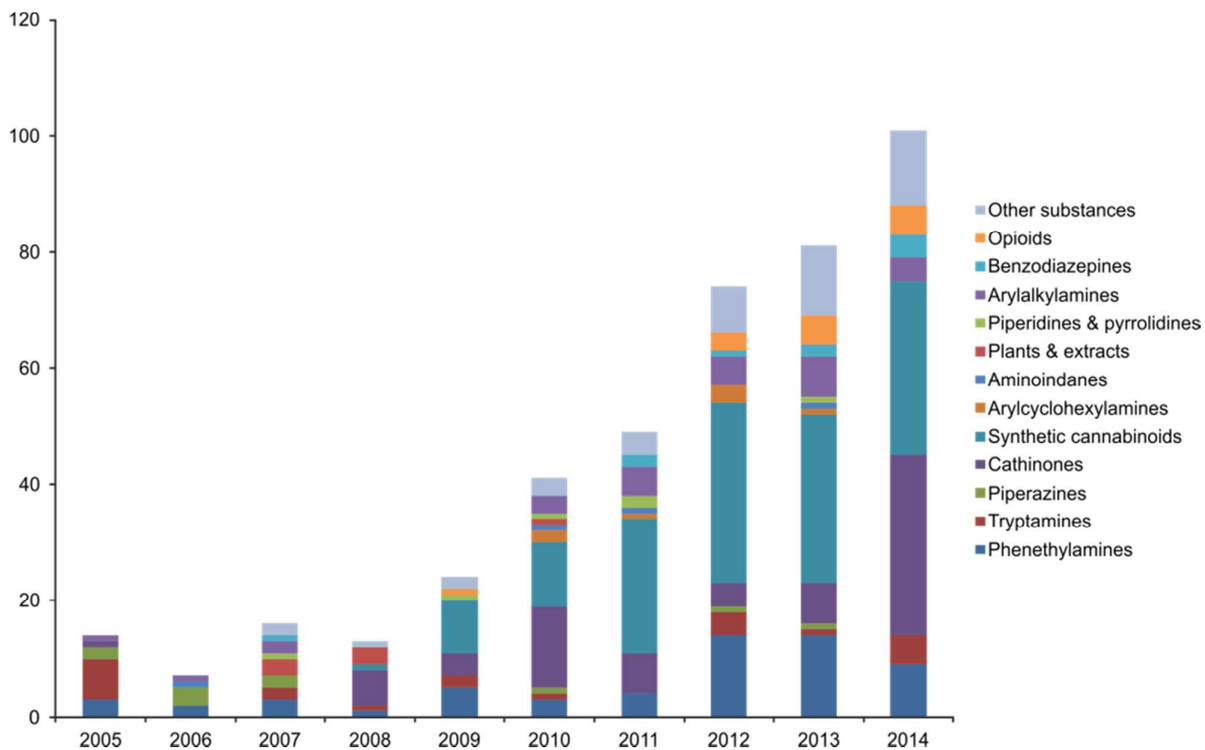


MDMAI

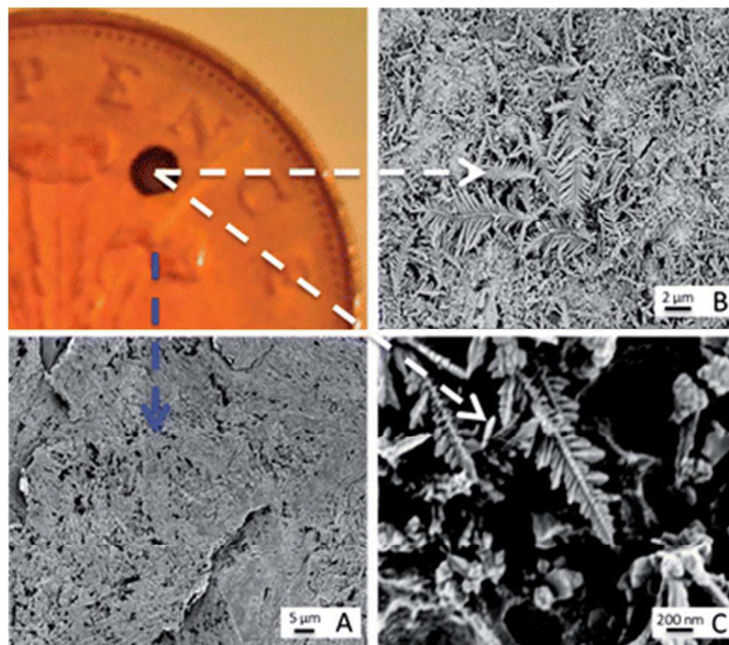


MMAI

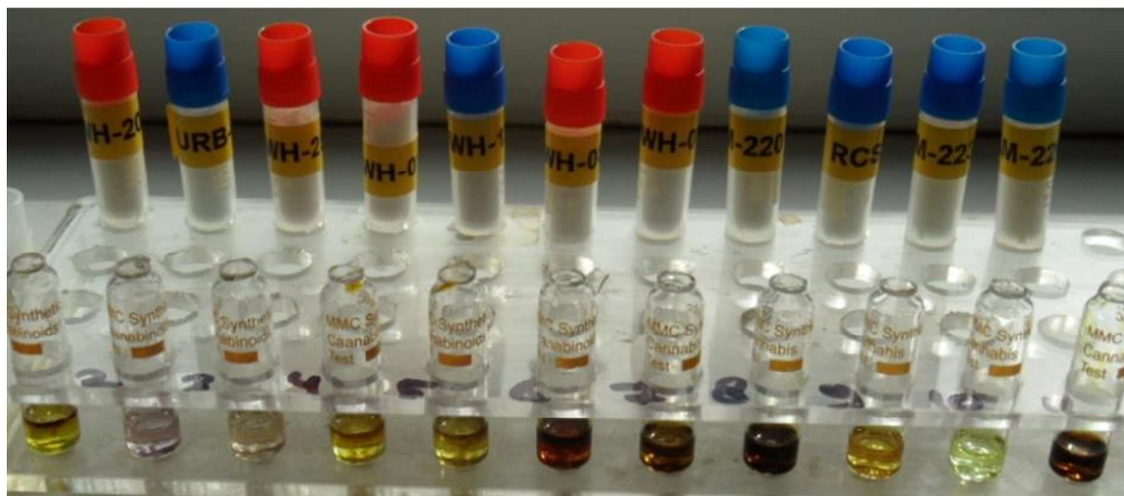
Figure 1 A graphical representation of novel psychoactive substances notified to the EWS between 2005-2014. Reproduced from reference 7 with permission of the EMCDDA.



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3 **Figure 2.** Characterisation of galvanic displacement. The optical image (top left) shows a
4 clean British 2p coin, with silver deposited onto its surface. (A) shows an SEM of the rough
5 surface of the two pence after cleaning. The SEM in (B) shows the silver dendritic structures
6 that are formed on the coins surface once 10 μL of AgNO_3 was left to mature for 20 s at room
7 temperature (23 $^\circ\text{C}$). The fern like structures are magnified in (C) and show that secondary
8 crystalline domains grow perpendicular from a primary silver backbone.⁵² – Reproduced
9 from reference ⁵² with permission of The Royal Society of Chemistry.
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3 **Figure 3.** Visual representation of synthetic cannabinoid presumptive test, reproduced from
4 reference ⁹⁹ with permission of Narcotic Testing Supplies & Equipment Store.
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3 **Figure 4** Microcrystals formed with mercury chloride and (a) mephedrone ($c = 10 \text{ g L}^{-1}$), (b_i)
4 MDAI freebase ($c = 1 \text{ g L}^{-1}$), (b_{ii}) MDAI hydrochloride ($c = 1 \text{ g L}^{-1}$) and (c) BZP ($c = 1 \text{ g L}^{-1}$).¹¹⁴ Reproduced from reference ¹¹⁴ with permission of Elsevier.
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