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# Chemoinformatic Consideration of Novel Psychoactive Substances: Compilation and Preliminary Analysis of a Categorised Dataset

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- 25 **Conflict of interest declaration:**
- 26 The authors can confirm that no conflicts of interest are present relating the reported work.

### 27 Abstract

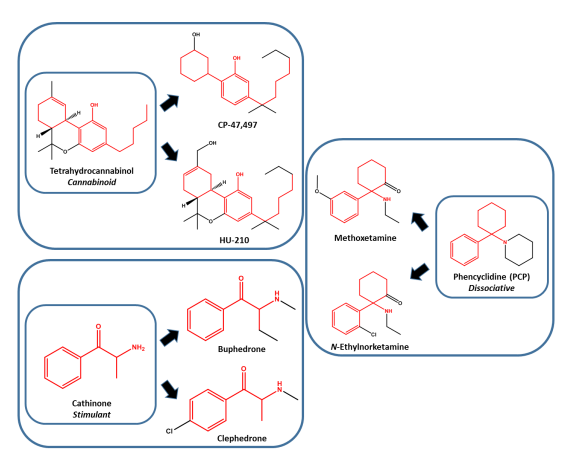
Recent years have seen the emergence into circulation of a growing array of novel psychoactive substances (NPS). Knowledge of the pharmacological profiles and risk liability of these compounds is typically very scarce. Development of chemoinformatic tools enabling prediction of properties within uncharacterised analogues has potential be of particular use. In order to facilitate this, compilation of a chemical inventory comprising known NPS is a necessity.

33 Sourcing a variety of published governmental and analytical reports, a dataset composed of 690 34 distinct acknowledged NPS, complete with defined chemical structures, has been constructed. This is 35 supplemented by a complementary series of 155 established psychoactive drugs of abuse (EPDA). 36 Classification was performed in accordance with their key molecular structural features, subjective 37 effect profiles and pharmacological mechanisms of action. In excess of forty chemical groupings, 38 spanning seven subjective effect categories and six broad mechanisms of pharmacological action, were identified. Co-occurrence of NPS and EPDA within specific classes was common, showcasing 39 40 inherent scope both for chemical read-across and for the derivation of structural alerts.

### 41 **1. Introduction**

42 Over the course of the previous decade, the emergence onto the unregulated market of novel, 43 predominantly synthetic psychoactive compounds - referred to henceforth as "novel psychoactive substances" (NPS) – has grown to constitute an increasing public health concern across much of the 44 developed world.<sup>[1]</sup> Such agents are typically intended to mimic closely the effects associated with 45 established, very often illicit, psychotropic drugs of abuse (examples of which are provided within 46 47 Figure 1.). Their initial presence outside of the boundaries of substance control schedules within many legislative areas has led to their acquisition of the popular descriptor "legal highs".<sup>[2]</sup> Whilst numerous 48 49 nations have since taken action to bring under control the broad chemical classes within which these 50 compounds typically fall, emergence of new analogues is continuous. The yet incomplete knowledge concerning their pharmacological and toxicological profiles ensures therefore that their presence and 51 52 use continues to form an ever-evolving and potentially substantial risk towards consumers.<sup>[3]</sup>





54

55 Figure 1. Scheme outlining identity and chemical structure of a selection of established psychoactive drugs of

abuse, accompanied by relevant novel analogues.

57 NPS may be sourced in practice through an assortment of routes, and in an array of formulations. 58 "Head shops", present both as traditional street-side locations and increasingly online, offer a variety of products either individually or as constituents within mixtures.<sup>[4]</sup> Sold typically under descriptions 59 such as "herbal incense" or "pot pourri", and further commonly referred to as "Spice", cannabimimetic 60 61 blends composed of a variety of synthetic cannabinoid species are acknowledged as constituting a significant proportion of this market.<sup>[5]</sup> Stimulant and empathogenic compounds (distributed 62 63 classically as "bath salts" or "plant food") additionally find wide availability, as do psychedelic tryptamines and lysergamides, opioid agonists and sedatives.<sup>[6]</sup> Commonly sold as "research 64 chemicals", their unregulated sourcing and production allied to the undefined nature of many 65 66 formulations contributes to the uncertainty which surrounds identification of single NPS. The 67 discerning of pharmacological and toxicological properties attributable to them is therefore rendered 68 a demanding and non-trivial task.<sup>[7]</sup> Challenge is additionally posed to the analytical chemist, who must 69 define routes towards the characterisation of an ever-expanding library of structures.<sup>[8]</sup>

70 Attempts to understand in greater depth the impacts upon physical and mental wellbeing associated 71 with the abuse of specific NPS are confounded by a variety of factors. These derive both from the 72 inherent novelty of the compounds, and from the unregulated, often clandestine nature of their 73 production and distribution. Owing to the rapid and continuing emergence of novel substances, there 74 exists in general a paucity of reliable experimental and clinical data concerning their toxicological 75 potential. Case studies acquired from patients who have presented following acute ingestion of a cocktail of NPS - either in the presence or absence of established illicit psychoactive drugs -76 constitute the dominant testimony apparent within the literature.<sup>[9-12]</sup> Such reports display obvious 77 78 limitations with regards to the characterisation of individual compounds, most notably with regards 79 to specific cellular and organ-level toxicities and dependency profiles over extended periods of use.

Although it is noted that both *in vivo* and *in vitro* experimental data are largely non-existent for the great majority of compounds which have emerged over the preceding 10-15 years, appreciation of

relevant structure-activity relationships may allow for the inference of the capacity of a substance to react towards given adverse outcomes. As such, there exists significant scope for the input of chemoinformatic and predictive toxicological approaches within characterisation of the properties possessed by this diverse range of chemical subtypes. Pooling of related molecules into relevant groups further has the capacity to assist in predicting pharmacology, drawing upon similarity with established drugs whilst simultaneously permitting extrapolation to novel substances as their presence becomes known.

89 The essential first step towards any chemoinformatic consideration of NPS is in the curation of a 90 compound inventory, complete with defined, unambiguous structure relating each constituent 91 molecule. A variety of national and supra-national government and advisory agencies have, over the 92 preceding ten years, issued periodical lists of named compounds considered by their experts to fall 93 within the bracket of NPS. It is from these, complemented by a variety of independent analytical 94 sources, that we have sought to construct an expansive compendium of NPS acknowledged as 95 constituting wider concern. As such, the aim of this study was to compile and categorise known NPS 96 and provide basis for comparison - both structurally and mechanistically - with established 97 psychoactive compounds. Presented is a dataset composed of 690 novel psychoactive substances, 98 classified according to their purported effect profiles, neuropharmacological mode of action and 99 structural composition. Comparison was made with an accessory compilation consisting of 155 100 established psychoactive drugs of abuse, generally possessive of recognised pharmacological and 101 toxicological profiles.

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### 106 **2. Materials and methods**

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### 108 **2.1. Compilation of database**

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Two distinct datasets, one composed solely of recorded NPS and another consisting of established psychoactive drugs of abuse (EPDA), were developed in accordance with protocols described below. In instances whereby compounds were found to occupy both classifications, placement preferentially within the latter grouping was ensured. Each may be found located in its entirety within Supplementary Table 1.

115

### 116 Novel psychoactive substances

117

Information concerning the identities of compounds acknowledged as NPS was accumulated from sources as outlined within Table 1. Amongst the literature drawn upon were reports issued through governmental and supra-governmental entities including the United Nations Office on Drugs and Crime (UNODC) and the European Monitoring Centre for Drugs and Drug Addiction (EMCDDA), alongside a selection of original research publications and reviews developed by independent groups. A comprehensive index of source material, incorporating assignment of origin for each substance, is present within Supplementary Table 1.

125 Established psychoactive drugs of abuse

Substances constituting "illicit" grouping within the DrugBank resource (<u>www.drugbank.ca</u>) were examined for their purported psychoactive properties.<sup>[13]</sup> Those adjudged as possessing no such liability (primarily steroidal compounds utilised for physical effect) were removed from consideration, furnishing a 155-member established psychoactive drug of abuse set.

130

131

Reference	Entries	Reference	Entries
UNODC, 2013 <sup>[14]</sup>	234	Debruyne & Le Boisselier, 2015 <sup>[28]</sup>	122
EMCDDA, 2006 <sup>[15]</sup>	12	Banister <i>et al.</i> , 2015 <sup>[29]</sup>	12
EMCDDA, 2007 <sup>[16]</sup>	6	Banister <i>et al.</i> , 2016 <sup>[30]</sup>	18
EMCDDA, 2008 <sup>[17]</sup>	14	Qian <i>et al.,</i> 2017 <sup>[31]</sup>	9
EMCDDA, 2009 <sup>[18]</sup>	12	Shevyrin <i>et al.,</i> 2014 <sup>[32]</sup>	3
EMCDDA, 2010 <sup>[19]</sup>	24	Shevyrin <i>et al.,</i> 2016 <sup>[33]</sup>	1
EMCDDA, 2011 <sup>[20]</sup>	39	Uchiyama, Matsuda <i>et al.</i> , 2014 <sup>[34]</sup>	13
EMCDDA, 2012 <sup>[21]</sup>	46	Uchiyama, Shimokawa <i>et al.</i> , 2014 <sup>[35]</sup>	8
EMCDDA, 2013 <sup>[22]</sup>	73	Uchiyama <i>et al.,</i> 2015 <sup>[36]</sup>	11
EMCDDA, 2014 <sup>[23]</sup>	74	Nakajima <i>et al.</i> , 2015 <sup>[37]</sup>	4
EMCDDA, 2015 <sup>[24]</sup>	94	Blakey <i>et al.</i> , 2016 <sup>[38]</sup>	8
EMCDDA, 2016 <sup>[25]</sup>	101	Lai <i>et al.,</i> 2015 <sup>[39]</sup>	6
EMCDDA, 2017 <sup>[26]</sup>	60	Coppola & Mondola, 2012 <sup>[40]</sup>	5
NFL Slovenia <sup>[27]</sup>	77		

**Table 1.** Summary of literature sources from which NPS identities were drawn.

### **2.2. Acquisition and visualisation of chemical structures**

In instances where not provided explicitly within source publications, molecular structures
 corresponding to listed compounds were obtained through online resources including PubChem
 (www.pubchem.gov), ChemSpider (www.chemspider.com) and the New Synthetic Drugs Database
 (http://www.nsddb.eu/).<sup>[41, 42]</sup> Details concerning structural composition were coded for each entry as
 SMILES strings.<sup>[43]</sup> Visualisation was achieved subsequently through use of ChemAxon MarvinView
 software (version 1.6).<sup>[44]</sup>

### **2.3.** Grouping and classification of compounds

146 Grouping with respect to psychoactive effect

147 Classification as regards psychotropic influence was performed with reference to descriptions present

148 within source literature. Ancillary information, as required, was obtained through use of the Erowid

149 online resource (<u>www.erowid.org</u>).<sup>[45]</sup>

151 Grouping with respect to pharmacological mechanism of action

152 Assorted literature sources, referenced in the text, were employed in order to attribute the dominant

- 153 neuropharmacological mechanism to constituent compounds.
- 154
- 155 Grouping with respect to molecular structural features

156 Molecules were visualised in accordance with protocols described above. Chemical and 157 pharmacological knowledge was employed in order constitute groups related by shared, biologically-158 relevant structural motifs. Those falling outside of such categories were termed "unclassified".

159

### 160 **2.4.** Principal component analysis of chemical space

161 Descriptors relating to the physicochemical and structural properties of compounds contained within 162 NPS and EPDA sets were determined through use of CORINA Symphony Descriptors Community 163 Edition (v. 2, MN-AM, Nuremberg, Germany: www.mn-am.com/services/corinasymphonydescriptors). 164 Further series of parameters, centred upon the presence within structures of definitive chemical 165 fingerprints, were developed through assistance of the ChemoTyper application (v. 1.1, MN-AM, Nuremberg, Germany) with reference to established ToxPrint chemotypes.<sup>[46]</sup> Physicochemical and 166 167 structural descriptors (in total 31, refer to Supplementary Table 3 for their identity) were integrated into combined arrays, from which principal components were extracted using Principal Component 168 169 Analysis within the Minitab Statistical software (v. 18.1, State College PA, USA). Visualisation, in the 170 form of scatter plots, was achieved through use of this same program.

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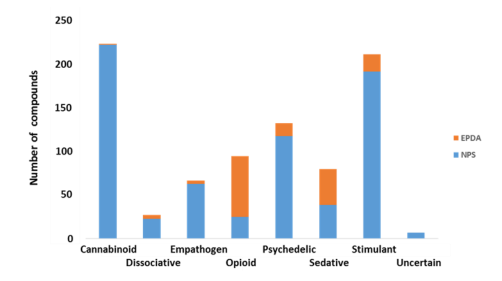
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### 175 **3. Results**

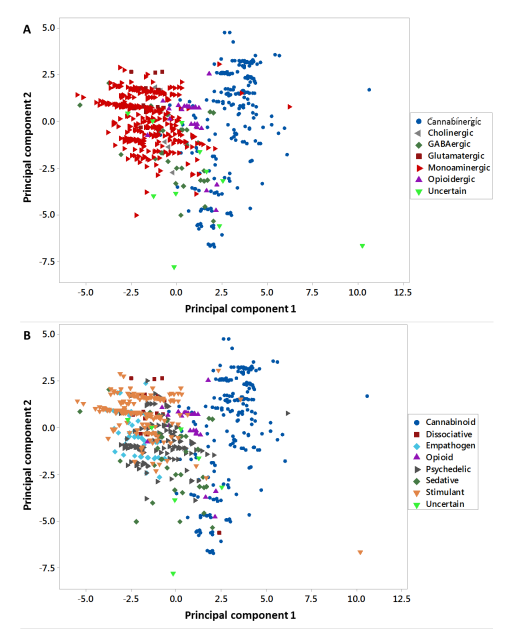
### 176 **3.1. Overview and analysis of dataset**

177 A total of 690 compounds characterised as NPS were identified from within the aforementioned 178 sources. With regards to purported psychoactive properties (as displayed visually within Figure 2), 223 179 were distinguished as cannabinoids, 192 as stimulants, 118 as psychedelics, 63 as empathogens, 39 as 180 sedatives, 25 as opioids, and 20 as dissociatives. Owing to insufficient attestation coupled with 181 structural obscurity, 10 compounds, labelled "uncertain", had no definitive effect or effects attributed. 182 367 of these compounds influenced monoaminergic transmission, 223 cannabinergic, 36 GABAergic, 183 25 opioidergic, 19 glutamatergic and 6 cholinergic (with 14 uncertain). Substances were further 184 partitioned, where appropriate, into one of 35 distinct chemical groupings. A selection of 43 isolated 185 compounds defied such categorisation, and were in turn listed "unclassified". From the Drugbank 186 "Illicit" dataset, a sum of 155 psychoactive compounds was gathered. In all, 70 could be identified as 187 opioids, 40 as sedatives, 21 as stimulants, 15 as psychedelics, 4 as dissociatives, 4 as empathogens and 188 1 as cannabinoid. These entries spanned 23 distinct chemical classifications, incorporating six absent 189 amongst NPS.



191 **Figure 2.** Numerical composition of psychoactive effect groups.

Principal component analysis of physicochemical and structural properties was performed upon the NPS dataset. Outcomes are expressed visually within plots (Figure 3), detailing comparison of scores obtained between principal components 1 and 2. Evident within Figure 3A, the dominant groupings of cannabinergic and monoaminergic agents are seen to occupy areas of chemical space largely distinct from one-another. Grouping according to psychoactive effect (Figure 3B) illustrates extent of overlap between monoamine-like stimulant, empathogen and psychedelic agents.



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Figure 3. Principal component analyses of NPS dataset. Scores relating first principal components, with
 compounds grouped In accordance with their pharmacological mechanism of action (A) and psychoactive effect
 profile (B).

Substance inventories may be viewed in their entirety through accessing of Supplementary Data. Supplementary Table 1 incorporates the sum of relevant data concerning compound nomenclature, structure and classification. For summary of chemical and psychoactive effect classification overlap, Supplementary Table 2 should be consulted.

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## 210211 **3.2.1. Monoaminergic**

3.2. Consideration of psychoactive categories

211 212

213 Pathways of dopaminergic, adrenergic and serotonergic transmission hold integral roles within 214 regulation of cognition, perception and emotion. Perturbation in the functioning of these systems 215 relates closely, dependent upon mechanistic specificity, to a range of psychoactive influences 216 extending from therapeutic alleviation of depression to induction of intense psychedelic and 217 hallucinogenic experience. There are in practice numerous physiological processes associated with 218 neurotransmitter regulation as modulated through the actions of neuroactive substances, and as such 219 the pharmacology of such compounds is varied. Whilst receptor agonism and antagonism is a feature 220 within selected classes, enhancement of synaptic neurotransmitter concentration through induction of release or inhibition of reuptake forms a generally dominant mode of action.<sup>[47, 48]</sup> 221

In the overwhelming majority of instances, a close chemical similarity to endogenous neurotransmitters is apparent (as highlighted within Figure 4). Functionalisation of the phenylethylamine unit central within catecholamines dopamine (DA) and NA permits rational design of compounds possessive of a spectrum of stimulant, empathogenic and psychedelic effects. Tryptamine-derived serotonin (5-HT) mimics, as direct 5-HT receptor agonists, are further notable for their hallucinogenic influence.

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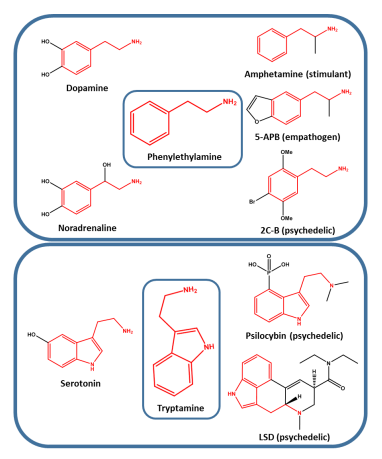
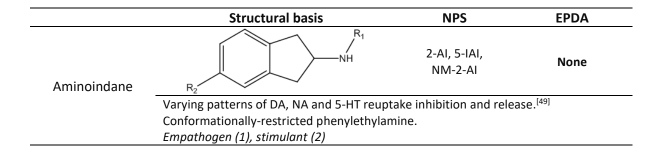


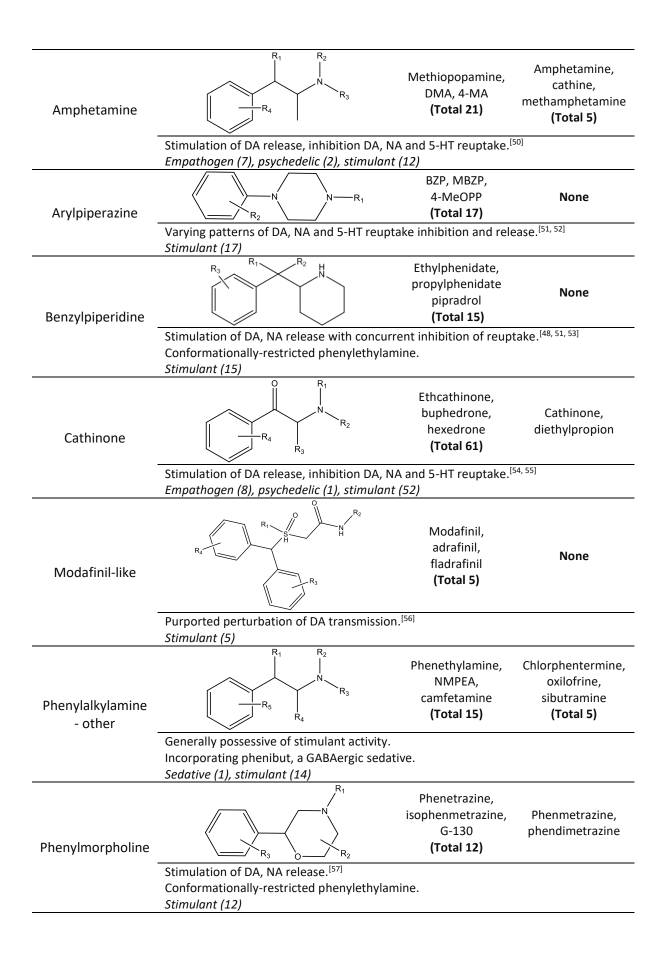
Figure 4. Overview of shared structural motifs common to endogenous neurotransmitters and monoaminergicNPS.

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### 233 3.2.1.1 Stimulant

Characterised by a capacity to invoke senses of wakefulness and heightened energy, the typical stimulant belongs to the broad family of substituted phenylethylamines (as outlined within Table 2). Cathinone and pyrrolidinophenone derivatives are notably numerous, forming as they do common constituents within "bath salt" blends.<sup>[6]</sup> Tropane cocaine analogues and modafinil mimics form notable categories based upon alternative structural motifs.





Pyrrolidinophenone		α-PVP, 4'-Fluoro-α-PVP, α-PNP <b>(Total 44)</b>	None
	Stimulation of DA release, inhibition DA, N	IA and 5-HT reuptake. <sup>I</sup>	[58]
	Stimulant (44)		
Tropane and analogues		Dichloropane, nitracaine, dimethocaine <b>(Total 8)</b>	Cocaine, ecgonine, benzoylecgonine
-	Inhibition of DA, NA and 5-HT reuptake. <sup>[48,</sup>	,59]	
	Hyoscine and hysocamine alternatively fur		eliriants.
	Stimulant (6), psychedelic (2)	-	
	Methoxyphenylalkylamine – other (4),	4-EA NBOMe,	Amineptine,
Minor	non-specified alkaloid (4),	2-MA, vanoxerine	aminorex, pemoline
	unclassified (5)	(Total 13)	(Total 4)

242

Table 2. Overview of key structural features, prominent category entries and recovered EPDA analogues relatedto chemical groupings prevalent amongst stimulant NPS.

### 243 3.2.1.2. Empathogen

Such compounds are characterised by their broad similarity in psychoactive effect to methylenedioxymethamphetamine (MDMA) – described commonly as the induction of stimulation and euphoria accompanied by heightened feelings of social connectivity.<sup>[59, 60]</sup> Their distinctive properties are associated with increased serotonergic potency, likely a function of the fused tryptamine-like heterocyclic units apparent in benzofurans and methylenedioxyphenylalkylamines (detailed in Table 3).

	Structural basis	NPS	EPDA
Benzofuran	R <sub>4</sub> N R <sub>2</sub> R <sub>3</sub>	5-APB, 6-APB, 5-EAPB <b>(Total 15)</b>	None
	Inhibition of reuptake and stimulation of HT <sub>2</sub> receptor. <sup>[60]</sup>	release of DA, NA and	d 5-HT. Agonism at 5-
	Empathogen (14), psychedelic (1)		
Methylenedioxy- phenylalkylamine	$R_{5}$ $R_{7}$ $R_{4}$ $R_{3}$	Ethylone, butylone, EDMA <b>(Total 29)</b>	MDMA, MMDA, tenamfetamine <b>(Total 4)</b>
	Inhibition of reuptake and stimulation of at $5-HT_2$ receptor. <sup>[54, 61]</sup>	release of DA, NA and	d 5-HT. Weak agonism
	Empathogen (27), psychedelic (2)		

Oxazoline		3,4-DMAR, 4,4'-DMAR, N-Methyl aminorex derivative	Aminorex, 4-methylaminorex, pemoline
	Inhibition of DA, NA and 5-HT reuptake, a Bears conformationally-restricted phenyle		
	Empathogen (3)		
Minor	Aminoindane (1), amphetamine (7), cathinone (8), methoxyphenylalkylamine – other (2), tryptamine (1)	Mephedrone, 4-FA, 5-API <b>(Total 19)</b>	None

Table 3. Overview of key structural features, prominent category entries and recovered EPDA analogues related
 to chemical groupings prevalent amongst empathogen NPS.

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### 255 **3.2.1.3.** Psychedelic

- 256 NPS appearing under the description "psychedelic" are noted for their induction of altered states of
- 257 perception characterised by visual hallucination and profound changes in cognition. As direct agonists
- 258 at selected 5-HT receptors (refer to Table 4), tryptamine serotonin analogues and dimethoxy-
- 259 substituted phenylalkylamines constitute the bulk of this class.<sup>[63, 64]</sup>
- 260

	Structural basis	NPS	EPDA
xC- Phenylalkylamine		2C-C, 2C-I, 2C-N <b>(Total 23)</b>	2С-В, 2С-Т-7
	Agonism and antagonism across 5-HT <sub>2</sub> rece	ptors. <sup>[65]</sup>	
	Dimethoxy substituent essential in induction	on of hallucinogenic ef	fect. <sup>[66]</sup>
	Psychedelic (23)		
xC-NBx- Phenylalkylamine	$R_4$ $R_1$ $R_2$ $R_3$ $R_2$	25B-NBOMe 25C-NBOMe 25N-NBOMe <b>(Total 21)</b>	None
	Agonism at 5-HT <sub>2</sub> receptors. <sup>[67]</sup> Dimethoxy substituent essential in induction	on of ballucinogenic of	fect
	Psychedelic (21)	in of Handelilogenic er	

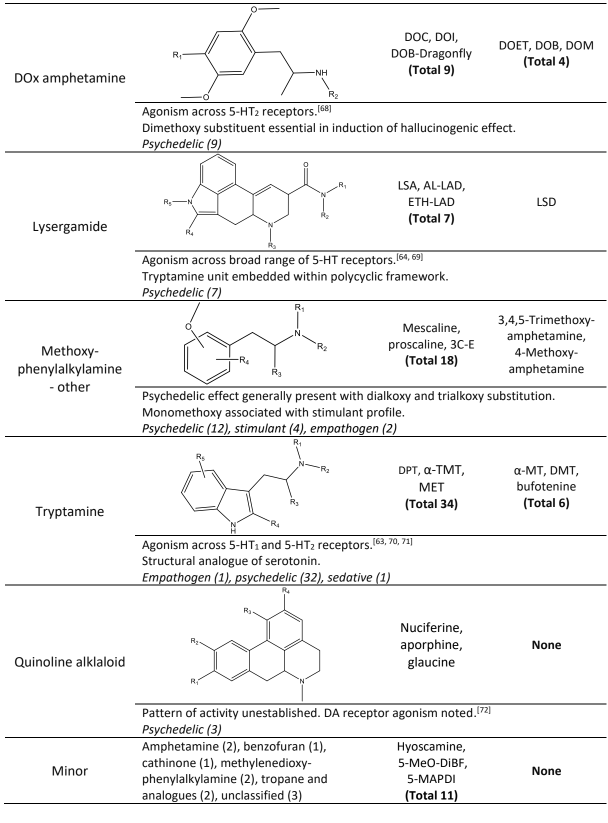


Table 4. Overview of key structural features, prominent category entries and recovered EPDA analogues related
 to chemical groupings prevalent amongst psychedelic NPS.

### 266 3.2.2. Cannabinergic

### 267 Cannabinoid

A variety of synthetic agonists active at cannabinoid CB<sub>1</sub> and CB<sub>2</sub> receptors have, through consequence of the popularity of "Spice"-style blends, entered into circulation.<sup>[73]</sup> With exception of the notable class of THC-like cyclohexylphenols, the great majority of developed compounds display structures – typically carbonyl-substituted indole and indazole derivatives – distinct from natural endogenous or phytochemical activators (listed in full within Table 5).

	Structural basis	NPS	EPDA
Cyclohexylphenol	R <sub>3</sub> OH R <sub>2</sub> R <sub>1</sub>	HU-210, HU-308 CP-47,497 <b>(Total 10)</b>	тнс
	Agonist at CB receptors. <sup>[73]</sup> Structural analogues of THC.		
Indole-alkyl carboxamide		ADBICA, STS-135, MN-25 <b>(Total 57)</b>	None
	Agonist at CB receptors.		
ndole-alkyl ketone		UR-144, AB-001, AM-1248 <b>(Total 14)</b>	None
	Agonist at CB receptors.		
Indole-aryl carboxamide		SGT-25, MN-24, PX-1 <b>(Total 28)</b>	None
	Agonist at CB receptors.		
Indole-naphthyl ketone	R <sub>5</sub> R <sub>4</sub> R <sub>1</sub> R <sub>2</sub> R <sub>3</sub>	JWH-018, JWH-200, AM-2201 <b>(Total 55)</b>	None

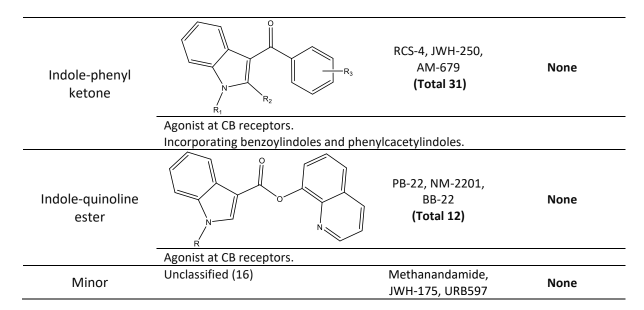


Table 5. Overview of key structural features, prominent category entries and recovered EPDA analogues related
 to selected chemical groupings prevalent amongst cannabinoid NPS.

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### 278 3.2.3. GABAergic

### 279 Sedative

280 Exclusively inhibitory in effect, potentiation of signalling through GABA receptors imparts sedative

and depressant outcome. GABAergic drug classes, including benzodiazepines and quinazolines,

282 function in general as allosteric receptor agonists, occupying distinct binding sites.<sup>[74]</sup> Kavalactones –

a selection of natural products isolated from the roots of kava (*Piper methysticum*) – exert effects

through an apparently distinct mechanism.<sup>[75]</sup> Further covered, exclusively under the heading of

285 EPDA (and hence omitted from inclusion within Table 6), is the barbiturate class.

	Structural basis	NPS	EPDA
Benzodiazepine	R <sub>5</sub> R <sub>5</sub> R <sub>1</sub> R <sub>2</sub>	Etizolam, nitrazolam, phenazepam <b>(Total 21)</b>	Diazepam, midazolam, prazepam <b>(Total 20)</b>
	Allosteric agonism of GABAA recept	or. <sup>[76]</sup>	
	Sedative (21)		
Kavalactone		Kavain, methysticin, yangonin <b>(Total 6)</b>	None
	Potentiation of GABA signalling three	ough undefined mechanism	[77, 78].
	Sedative (6)		

Quinazoline	R <sub>3</sub> N R <sub>1</sub> R <sub>2</sub>	Etaqualone, afloqualone, mebroqualone <b>(Total 4)</b>	Methaqualone
	Allosteric agonism of GABA <sub>A</sub> receptor. <sup>[1]</sup> Sedative (4)	79]	
Minor	Non-specified alkaloid (1), phenylalkylamine – other (1), tryptamine (1), unclassified (5)	5-HTP, 1,4-butanediol, zopiclone (Total 8)	Pregabalin, fospropofol, GHE <b>(Total 9)</b>

Table 6. Overview of key structural features, prominent category entries and recovered EPDA analogues related
 to chemical groupings prevalent amongst sedative NPS.

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291 3.2.4. Glutamatergic

- 292
- 293 Dissociative

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295 Whilst three primary classes of excitatory ionotropic glutamate receptor are characterised, it is those

296 of the NMDA variety which are considered of greatest pharmacological relevance. Antagonists,

297 notably analogues of ketamine and phencyclidine (PCP), are associated with unique forms of

298 dissociative anaesthesia – incorporating states typically characterised by hallucination, "out-of-body"

299 experience and sedation.<sup>[80, 81]</sup> Table 7 details the prominent chemical groupings.

	Structural basis	NPS	EPDA
Aryl- cyclohexylamine	R <sub>1</sub> R <sub>2</sub> R <sub>4</sub> R <sub>3</sub>	Methoxetamine, deschloroketamine, 4-MeO-PCP <b>(Total 14)</b>	PCP, PCPy, tenocyclidine <b>(Total 4)</b>
	Non-competitive antagonism at NN	IDA receptor. <sup>[82]</sup>	
	Dissociative (14)		
Diarylethylamine	R <sub>1</sub> R <sub>2</sub> R <sub>3</sub>	Ephenidine, diphenidine, NPDPA <b>(Total 5)</b>	None
	Non-competitive antagonism at NN	IDA receptor. <sup>[83]</sup>	
	Incorporates opioidergic MT-45.		
	Dissociative (4), opioid (1)		
Minor	Unclassified (2)	Salvinorin A, memantine	None

300

301 **Table 7.** Overview of key structural features, prominent category entries and recovered EPDA analogues related

302 to chemical groupings prevalent amongst dissociative NPS.

- 303 **3.2.5. Opioidergic**
- 304 Opioid
- 305

Agonists at the major subclasses of opioid receptor ( $\delta$ ,  $\kappa$ ,  $\mu$  and nociceptin) are capable of inducing potent analgaesic effect, coupled commonly with mild euphoria.<sup>[84]</sup> Dependence liability is notably high.<sup>[85]</sup> A variety of categories, including the numerous analogues of morphine, methadone and pethidine (excluded from Table 8) occur exclusively as EPDA.

- **Structural basis** NPS **EPDA** СІ AH-7921, None U-47700, U-49900 Dichlorobenzamide k۹ Agonism across range of opioid receptor subtypes.[86] Opioid (3) Acetylfentanyl, Fentanyl, valerylfentanyl, carfentanil, furanylfentanyl lofentanil Fentanyl derivative (Total 16) (Total 22) Agonism across range of opioid receptor subtypes.<sup>[87]</sup> Opioid (16) Diarylethylamine (1), W-15, non-specified alklaloid (3), unclassified mitragynine, Minor None (2) akuammine (Total 6)
- 310

Table 8. Overview of key structural features, prominent category entries and recovered EPDA analogues related
 to chemical groupings prevalent amongst opioid NPS.

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322 4. Discussion

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324 The recent emergence into circulation of an expanding library of novel psychoactive substances (NPS) 325 constitutes an evolving risk to public health. Efforts to define the landscape of identified compounds 326 with respect to their effect profiles and structural features have proved challenging on account both 327 of the novelty and obscurity of many, and further of the generally narrow scope of reports attesting 328 their detection and characterisation. As such, the intentions of this study have been to collate from 329 accessible source material an expansive inventory of definitively-acknowledged NPS. Like entries were 330 classified with respect to chemical, pharmacological and psychoactive similarity and, where 331 appropriate, related to analogous established drugs of abuse.

332 In total, a sum of 690 distinct novel substances were identified, supplemented by 155 established 333 drugs of abuse. It is apparent that, considered broadly, composition in terms of psychoactive profile 334 amongst the NPS set exhibits significant variation from that noted across EPDA (refer to Section 3.1.). 335 This is illustrated starkly in the preponderance of synthetic cannabinoids present within the former 336 (matching solely in effect against THC), and additionally by the comparative dominance of opioids – 337 notably the exclusive classes of morphine, methadone and pethidine analogues – amongst the latter. 338 Whilst the development and spread of cannabimimetics represents a recent phenomenon, the 339 establishment over many decades of opiate-like substances within clinical practice has contributed 340 towards the characterisation of their liability towards abuse and in turn to their scheduling.<sup>[88]</sup>

There remains a substantial number of chemical categories co-occurring within both novel and established sets. Contributing substantially towards impetus behind the development of NPS has been the desire to circumvent existing legislation concerning control of well-characterised recreational or abuse-liable drugs.<sup>[9]</sup> As such, the synthesis of structural analogues through minor modification of known compounds with an intention of retaining or even potentiating desired psychoactive outcome has assisted greatly in spurring the upturn in emergence of new substances (notably amongst the readily-adapted monoaminergic phenylalkylamines). Analogues of amphetamine and cathinone are

348 accordingly plentiful, whilst similarly well-represented are methylenedioxy entries mimicking configuration of MDMA and hallucinogenic methoxy-substituted 2C- and DOx equivalents.<sup>[63, 89]</sup> 349 350 Despite the general obscurity of a great number of these newer molecules, aspects of their 351 psychoactive and toxicological profiles be inferred with confidence through application of the more 352 extensive knowledge accrued within their established relatives - methodology akin to that of "readacross".<sup>[90-92]</sup> Such a principle which can similarly be extended to function within all chemically-related 353 354 categories incorporating at least a single EPDA analogue and across which pharmacological 355 mechanism of action can be reliably postulated as shared. This is a list which may include, but would 356 not be limited to, the serotonergic tryptamines and lysergamides, glutaminergic arylcyclohexylamines, GABAergic benzodiazepines and opioid fentanyl analogues. 357

358 In contrast to the aforementioned structural mimics, which correspond closely to recognised 359 psychoactive substances, a variety of classes exhibit novelty and distinctness in molecular composition. 360 In such instances the breadth and quality of study data relating the properties of member compounds is typically inferior, and cross-group extrapolation of effects a more substantial challenge. 361 362 Consideration of attributed pharmacological mechanism of action, alongside governing structure-363 activity relationships, adopts greater importance. A variety of notable categories fall under this broad 364 description, including benzofuran phenylalkylamines, diaryethylamines and the great majority of 365 synthetic cannabinoids. Uncharacterised benzofurans might reliably be inferred to possess 366 empathogenic qualities as a function of their structural similarity to the methylenedioxy MDMA derivatives, implying a monoaminergic mode of action (common to phenylalkylamines) distinguished 367 by further weak serotonin receptor agonism.<sup>[61]</sup> Diarylethylamines likewise share great 368 369 correspondence with NMDA antagonist arylcyclohexylamines – a class of dissociatives including 370 amongst its number the extensively-studied ketamine and PCP.

The single largest effect category present within NPS, definitive characterisation of synthetic
 cannabinoid action presents unique challenges. Of the 223 compounds identified, a mere ten (each of

the cyclohexylphenol class) bear structural relation to THC. Composing the remainder are an array of functionalised nitrogen heterocycle derivatives, distinct in composition from established psychoactives. It therefore follows that whilst the shared mechanism of cannabinergic receptor agonism ensures predictability in short-term subjective effects, inference of the physical and psychological consequences of continued use constitutes a greater trial. Ease of functionalisation ensures that the development of novel analogues remains ongoing, with the composition of cannabimimetic blends showing great variety.<sup>[5]</sup>

380 Examples considered across the above text provide broad overviews of how predictive approaches, 381 based upon consideration of molecular similarity, might be employed in order to credibly infer the 382 properties of the multitude of uncharacterised NPS. Drawing and collating from a variety of 383 authoritative sources, an extensive survey of the chemical landscape is presented. A total of 647 of 384 the 690 identified substances (94%) may be placed into one of the 35 defined structural groupings – 385 a practice which greatly orders and simplifies understanding of the set. Of these classes, 17 are seen 386 to co-occur amongst EPDA – thus granting scope for direct comparison of effect profile. 387 Pharmacological system of action is attributable within 676 members (98%) – furnishing mechanistic 388 rationale which will enhance confidence in proposed structure-activity relationships. To the 389 knowledge of the authors, this represents the most thorough unified structural repository of NPS – in 390 terms both of numerical composition and of pharmacological consideration – present within the 391 literature at this time. Provision of unambiguous structural identifiers for each entry, in the form of 392 SMILES strings, allows further for ready research use.

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