WEST VIRGINIA LEGISLATURE 2025 REGULAR SESSION

Introduced

House Bill 3343

By Delegate Worrell

[Introduced March 13, 2025; referred to the

Committee on Health and Human Resources]

Intr HB 2025R3766 1 A BILL to amend and reenact §60A-2-204 of the Code of West Virginia, 1931, as amended, 2 relating to schedule I drugs; and adding a provision relating to the scheduling of crystalline 3 polymorph psilocybin approved by the Food and Drug Administration. Be it enacted by the Legislature of West Virginia: **ARTICLE 2. STANDARDS AND SCHEDULES.** §60A-2-204. Schedule I. 1 (a) Schedule I shall consist of the drugs and other substances, by whatever official name, 2 common or usual name, chemical name, or brand name designated, listed in this section including 3 their isomers, esters, ethers, salts and salts of isomers, esters, and ethers, whenever the 4 existence of such isomers, esters, ethers, and salts is possible within the specific chemical

6 (b) Opiates.

designation.

5

7 Acetyl-alpha-methylfentanyl (N-[1-(1-methyl-2-phenethyl) -4-piperidinyl]—

8 phenylacetamide);

- 9 Acetylmethadol;
- 10 Allylprodine;
- 11 Alphacetylmethadol (except levoalphacetylmethadol also known as levo-alpha-12 acetylmethadol, levomethadyl acetate, or LAAM);
- 13 Alphameprodine;
- 14 Alphamethadol;
- Alpha-methylfentanyl (N-[1-(alpha-methyl-beta-phenyl) ethyl-4-piperidyl] propionanilide;
- 16 1-(1-methyl-2-phenylethyl)-4-((propanilido) piperidine);
- 17 Alpha-methylthiofentanyl (N-[1-methyl-2-(2-thienyl) ethyl-4-piperidinyl]—
- 18 phenylpropanamide);
- 19 Benzethidine;
- 20 Betacetylmethadol;

21	Beta-hydroxyfentanyl (N-[1-(2-hydroxy-2-phenethyl)-4-piperidinyl]-N-
22	phenylpropanamide);
23	Beta-hydroxy-3-methylfentanyl (other name: N-[1-(2- hydroxy-2-phenethyl)-3-methyl-4-
24	piperidinyl]-N-phenylpropanamide);
25	Betameprodine;
26	Betamethadol;
27	Betaprodine;
28	Clonitazene;
29	Dextromoramide;
30	Diampromide;
31	Diethylthiambutene;
32	Difenoxin;
33	Dimenoxadol;
34	Dimepheptanol;
35	Dimethylthiambutene;
36	Dioxaphetyl butyrate;
37	Dipipanone;
38	Ethylmethylthiambutene;
39	Etonitazene;
40	Etoxeridine;
41	Furethidine;
42	Hydroxypethidine;
43	Ketobemidone;
44	Levomoramide;
45	Levophenacylmorphan;
46	3-Methylfentanyl (N-[3-methyl-1-(2-phenylethyl)-4- piperidyl]-N-phenylpropanamide);

47	3-methylthiofentanyl (N-[3-methyl-1-(2-thienyl) ethyl-4- piperidinyl]—phenylpropanamide);
48	Morpheridine;
49	N-Methylnorfentanyl (N-(1-Methyl-4-piperidinyl)-N-phenyl-propanamide,
50	monohydrochloride);
51	Norfentanyl (N-Phenyl-N-4-piperidinyl-propanamide);
52	MPPP (1-methyl-4-phenyl-4-propionoxypiperidine);
53	Noracymethadol;
54	Norlevorphanol;
55	Normethadone;
56	Norpipanone;
57	Para-fluorofentanyl (N-(4-fluorophenyl)-N-[1-(2- phenethyl)-4-piperidinyl] propanamide);
58	PEPAP(1-(-2-phenethyl)-4-phenyl-4-acetoxypiperidine);
59	Phenadoxone;
60	Phenampromide;
61	Phenomorphan;
62	Phenoperidine;
63	Piritramide;
64	Proheptazine;
65	Properidine;
66	Propiram;
67	Racemoramide;
68	Thiofentanyl (N-phenyl-N-[1-(2-thienyl)ethyl-4- piperidinyl]-propanamide);
69	Tilidine;
70	Trimeperidine.
71	(c) Opium derivatives,
72	Acetorphine;

73	Acetyldihydrocodeine;
74	Benzylmorphine;
75	Codeine methylbromide;
76	Codeine-N-Oxide;
77	Cyprenorphine;
78	Desomorphine;
79	Dihydromorphine;
80	Drotebanol;
81	Etorphine (except HCl Salt);
82	Heroin;
83	Hydromorphinol;
84	Methyldesorphine;
85	Methyldihydromorphine;
86	Morphine methylbromide;
87	Morphine methylsulfonate;
88	Morphine-N-Oxide;
89	Myrophine;
90	Nicocodeine;
91	Nicomorphine;
92	Normorphine;
93	Pholcodine;
94	Thebacon.
95	(d) Hallucinogenic substances.
96	Alpha-ethyltryptamine; some trade or other names: etryptamine; Monase; alpha-ethy-1H-
97	indole-3-ethanamine; 3-(2- aminobutyl) indole; alpha-ET; and AET;
98	4-bromo-2, 5-dimethoxy-amphetamine; some trade or other names: 4-bromo-2,5-

99 dimethoxy-alpha-methylphenethylamine; 4-bromo- 2,5-DMA; 4-Bromo-2,5-dimethoxyphenethylamine; some trade or other names: 2-(4-bromo-2,5-100 101 dimethoxyphenyl)-1-aminoethane; alpha- desmethyl DOB; 2C-B, Nexus; 102 N-(2-Methoxybenzyl)-4-bromo-2, 5-dimethoxyphenethylamine. The substance has the 103 acronym 25B-NBOMe; 104 2-(4-chloro-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl) ethanamine (25C-NBOMe); 105 2-(4-iodo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl) ethanamine (25I-NBOMe); 106 2.5-dimethoxyamphetamine; some trade or other names: 2,5-dimethoxy-alpha-107 methylphenethylamine; 2,5-DMA; 108 2.5-dimethoxy-4-ethylamphet-amine; some trade or other names; DOET; 109 2,5-dimethoxy-4-(n)-propylthiophenethylamine (other name: 2C-T-7); 110 4-methoxyamphetamine; some trade other names: 4-methoxy-alphaor 111 methylphenethylamine; paramethoxyamphetamine; PMA; 112 3-Hydoxy-phencyclidine (other name hydroxy PCP); 113 5-methoxy-3, 4-methylenedioxy-amphetamine; 114 4-methyl-2,5-dimethoxy-amphetamine; some trade and other names: 4-methyl-2,5dimethoxy-alpha-methylphenethylamine; "DOM"; and "STP"; 115 116 3,4-methylenedioxy amphetamine; 117 3,4-methylenedioxymethamphetamine (MDMA); 118 3,4-methylenedioxy-N-ethylamphetamine (also known as (ethyl-alpha-methyl-3,4 119 (methylenedioxy) phenethylamine, N-ethyl MDA, MDE, MDEA); 120 N-hydroxy-3,4-methylenedioxyamphetamine (also known as (hydroxy-alpha-methyl-3,4 121 (methylenedioxy) phenethylamine, and (hydroxy MDA); 122 3,4,5-trimethoxy amphetamine; 123 5-methoxy-N,N-dimethyltryptamine (5-MeO-DMT); 124 Alpha-methyltryptamine (other name: AMT);

125	Bufotenine; some trade and other names: 3-(beta-Dimethylaminoethyl)-5-
126	hydroxyindole;3-(2-dimethylaminoethyl) -5-indolol; N, N-dimethylserotonin; 5-hydroxy-N,N-
127	dimethyltryptamine; mappine;
128	Diethyltryptamine; sometrade and other names: N, N-Diethyltryptamine; DET;
129	Dimethyltryptamine; some trade or other names: DMT;
130	5-Methoxy-N,N-disopropyltryptamine (5-MeO-DIPT);
131	Ibogaine; some trade and other names: 7-Ethyl-6, 6 Beta, 7, 8, 9, 10, 12, 13-octahydro-2-
132	methoxy-6, 9-methano-5H- pyrido [1', 2': 1, 2] azepino [5,4-b] indole; Tabernanthe iboga;
133	Lysergic acid diethylamide;
134	Marihuana; Marijuana (Cannabis, sp.);
135	Mescaline;
136	Parahexyl-7374; some trade or other names: 3-Hexyl -1-hydroxy-7, 8, 9, 10-tetrahydro-6,
137	6, 9-trimethyl-6H-dibenzo [b,d] pyran; Synhexyl;
138	Peyote; meaning all parts of the plant presently classified botanically as Lophophora
139	williamsii Lemaire, whether growing or not, the seeds thereof, any extract from any part of such
140	plant, and every compound, manufacture, salts, immediate derivative, mixture, or preparation of
141	such plant, its seeds or extracts;
142	N-ethyl-3-piperidyl benzilate;
143	N-methyl-3-piperidyl benzilate;
144	Psilocybin;
145	Psilocyn;
146	Tetrahydrocannabinols; synthetic equivalents of the substances contained in the plant, or
147	in the resinous extractives of Cannabis, sp. and/or synthetic substances, immediate derivatives
148	and their isomers with similar chemical structure and pharmacological activity including, but not
149	limited to the following:
150	delta-1 Cis or trans tetrahydrocannabinol, and their optical isomers;

151	delta-6 Cis or trans tetrahydrocannabinol, and their optical isomers;
152	delta-3,4 Cis or trans tetrahydrocannabinol, and its optical isomers;
153	delta-8 Cis or trans tetrahydrocannabinol and its optical isomers; and
154	delta-10 Cis or trans tetrahydrocannabinol and its optical isomers;
155	(Since nomenclature of these substances is not internationally standardized, compounds
156	of these structures, regardless of numerical designation of atomic positions covered.)
157	Delta-8-tetrahydrocannabinol-O (delta-8-THC-0), Delta-9-tetrahydrocannabinol (delta-9-
158	THC-0) and Synthetic and non-naturally occurring cannabinoids.
159	The provisions of this section related to tetrahydrocannabinols are inapplicable to products
160	or substances lawfully manufactured, distributed, or possessed under the provisions of §19-12E-
161	1 et seq. and Chapter 16H of this code.
162	Ethylamine analog of phencyclidine; some trade or other names: N-ethyl-1-
163	phenylcyclohexylamine, (1-phenylcyclohexyl) ethylamine, N-(1-phenylcyclohexyl) ethylamine,
164	cyclohexamine, PCE;
165	Pyrrolidine analog of phencyclidine; some trade or other names: 1-(1-phenylcyclohexyl)-
166	pyrrolidine, PCPy, PHP;
167	Thiophene analog of phencyclidine; some trade or other names: 1-[1-(2-thienyl)-
168	cyclohexyl]-piperidine, 2-thienylanalog of phencyclidine; TPCP, TCP;
169	1[1-(2-thienyl)cyclohexyl]pyrroldine; some other names: TCPy;
170	4-methylmethcathinone (Mephedrone);
171	3,4-methylenedioxypyrovalerone (MDPV);
172	2-(2,5-Dimethoxy-4-ethylphenyl)ethanamine (2C-E);
173	2-(2,5-Dimethoxy-4-methylphenyl)ethanamine (2C-D);
174	2-(4-Chloro-2,5-dimethoxyphenyl)ethanamine (2C-C);
175	2-(4-lodo-2,5-dimethoxyphenyl)ethanamine (2C-I);
176	2-[4-(Ethylthio)-2,5-dimethoxyphenyl]ethanamine (2C-T-2);

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177
              2-[4-(Isopropylthio)-2.5-dimethoxyphenyl]ethanamine (2C-T-4);
178
              2-(2,5-Dimethoxyphenyl)ethanamine (2C-H);
179
              2-(2,5-Dimethoxy-4-nitro-phenyl)ethanamine (2C-N);
180
              2-(2,5-Dimethoxy-4-(n)-propylphenyl)ethanamine (2C-P);
181
              3.4-Methylenedioxy-N-methylcathinone (Methylone):
182
              2,5-dimethoxy-4-(n)-propyltghiophenethylamine (2C-T-7, itsoptical isomers, salts and
183
       salts of isomers;
184
              5-methoxy-N,N-dimethyltryptamine some trade or other names: 5-methoxy-3-[2-
185
       (dimethylamino)ethyl]indole; 5-MeO-DMT(5-MeO-DMT);
186
              Alpha-methyltryptamine (other name: AMT);
187
              5-methoxy-N,N-diisopropyltryptamine (other name: 5-MeO-DIPT);
188
              Synthetic Cannabinoids as follows:
189
              2-[(1R,3S)-3-hydroxycyclohexyl]-5- (2-methyloctan-2-yl)phenol) {also known as CP
190
       47,497 and homologues};
191
              rel-2-[(1S,3R)-3-hydroxycyclohexyl] -5-(2-methylnonan-2-yl)phenol {also known as CP
192
       47,497-C8 homolog};
                                                                                        7,10,10a-
193
              [(6aR)-9-(hydroxymethyl)-6,
                                               6-dimethyl-3-(2-methyloctan-2-yl)-6a,
194
       tetrahydrobenzo[c]chromen-1-ol)] {also known as HU-210};
195
              (dexanabinol);
196
              (6aS,10aS)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl)-6a,7,10,10a-
197
       tetrahydrobenzol[c]chromen-1-ol) {also known as HU-211};
198
              1-Pentyl-3-(1-naphthoyl)indole {also known as JWH-018};
199
              1-Butyl-3-(1-naphthoyl)indole {also known as JWH-073};
200
              (2-methyl-1-propyl-1H-indol-3-yl)-1-napthalenyl-methanone {also known as JWH-015};
201
              (1-hexyl-1H-indol-3-yl)-1-naphthalenyl-methanone {also known as JWH-019};
202
              [1-[2-(4-morpholinyl) ethyl] -1H-indol-3-yl]-1-naphthalenyl-methanone {also known as
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203
       JWH-200):
204
              1-(1-pentyl-1H-indol-3-yl)-2-(3-hydroxyphenyl)-ethanone {also known as JWH-250};
205
              2-((1S,2S,5S)-5-hydroxy-2- (3-hydroxtpropyl)cyclohexyl) -5-(2-methyloctan-2-yl)phenol
206
       {also known as CP 55,940};
207
              (4-methyl-1-naphthalenyl) (1-pentyl-1H-indol-3-yl) -methanone {also known as JWH-122};
208
              (4-methyl-1-naphthalenyl) (1-pentyl-1H-indol-3-yl) -methanone {also known as JWH-398;
209
              (4-methoxyphenyl)(1-pentyl-1H-indol-3-yl)methanone {also known as RCS-4};
210
              1-(1-(2-cyclohexylethyl) -1H-indol-3-yl) -2-(2-methoxyphenyl) ethanone {also known as
211
       RCS-8};
212
              1-pentyl-3-[1-(4-methoxynaphthoyl)]indole (JWH-081);
213
              1-(5-fluoropentyl)-3-(1-naphthoyl)indole (AM2201); and
214
              1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole (AM694).
215
              Synthetic cannabinoids:
216
              CP 47,497 AND homologues, 2-[(1R,3S)-3-Hydroxycyclohexyl]-5-(2-methyloctan-2-
217
              YL)phenol);
218
              HU-210, [(6AR,10AR)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-Methyloctan-2-YL)-6A,7,10,
219
       10A-tetrahydrobenzo[C] chromen-1-OL)];
220
              HU-211, (dexanabinol, (6AS,10AS)-9-(hydroxymethyl)-6,6-Dimethyl-3-(2-methyloctan-2-
221
       YL)-6A,7,10,10atetrahydrobenzo[C]chromen-1-OL);
222
              JWH-018, 1-pentyl-3-(1-naphthoyl)indole;
223
              JWH-019, 1-hexyl-3-(1-naphthoyl)indole;
224
              JWH-073, 1-butyl-3-(1-naphthoyl)indole;
225
              JWH-200, (1-(2-morpholin-4-ylethyl)indol-3-yl)- Naphthalen-1-ylmethanone;
226
              JWH-250, 1-pentyl-3-(2-methoxyphenylacetyl)indole.]
                       2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3,3-dimethylbutanoate
227
              Methyl
                                                                                              (5F-
228
       ADB);
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229	Methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3-methylbutanoate (5F-AMB);
230	Methyl 2-(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3-methylbutanoate (FUB-
231	AMB);
232	N-(adamantan-1-yl)-1-(5-fluoropentyl)-1H-indazole-3-carboxamide (5F-APINACA);
233	N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(4-fluorobenzyl)-1H-indazole-3-carboxamide
234	(ADB-FUBINACA);
235	Methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-3,3-dimethylbutanoate
236	(MDMB-CHMICA);
237	Methyl 2-(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3,3-dimethylbutanoate (MDMB-
238	FUBINACA);
239	Tetrahydrocannabinols:
240	DELTA-1 CIS OR trans tetrahydrocannabinol and their Optical isomers.
241	DELTA-6 CIS OR trans tetrahydrocannabinol and their optical isomers.
242	DELTA-3,4 CIS or their trans tetrahydrocannabinol and their optical isomers.
243	Synthetic Phenethylamines
244	2-(4-iodo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)ethanamine (25I-NBOMe/ 2C-I-
245	NBOMe);
246	2-(4-chloro-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)ethanamine (25C-NBOMe/2C-C-
247	NBOMe);
248	2-(4-bromo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)ethanamine (25B-NBOMe/ 2C-B-
249	NBOMe);
250	Synthetic Opioids (including their isomers, esters, ethers, salts and salts of isomers, esters
251	and ethers):
252	N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide (acetyl fentanyl);
253	furanyl fentanyl;
254	3,4-dichloro-N-[2-(dimethylamino)cyclohexyl]-N-methylbenzamide (also known as U-

255	47700);			
256	N-(1-phenethylpiperidin-4-yl)-N-phenylbutyramide, also known as N-(1-			
257	phenethylpiperidin-4-yl)-N-phenylbutanamide, (butyryl fentanyl);			
258	N-[1-[2-hydroxy-2-(thiophen-2-yl)ethylpiperidin-4-yl]-N-phenylpropionamide, also known			
259	as N-[1-[2-hydroxy-2-(2-thienyl)ethyl]-4-piperidinyl]-N-phenylpropanamide, (beta-			
260	hydroxythiofentanyl);			
261	N-(1-phenethylpiperidin-4-yl)-N-phenylacrylamide (acryl fentanyl);			
262	N-(1-phenethylpiperidin-4-yl)-N-phenylisobutyramide (isobutyryl fentanyl);			
263	N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopentanecarboxamide (cyclopropyl fentanyl);			
264	2-(2,4-dichlorophenyl)-N-((1S,2S)-2-(dimethylamino)cyclohexyl)-N-methylacetamide			
265	(also known as U-48800);			
266	Trans-3,4-dichloro-N-[2-(diethylamino)cyclohexyl]-N-methyl-benzamide (also known as			
267	U-49900);			
268	Trans-3,4-dichloro-N-[2-(dimethylamino)cyclohexyl]-N-methyl-benzeneacetamide (also			
269	known as U-51754);			
270	2-(2-(4-butoxybenzyl)-5-nitro-1H-benzimidazol-1-yl)-N,N-diethylethan-1-amine			
271	(butonitazene);			
272	2-(2-(4-ethoxybenzyl)-1H-benzimidazol-1-yl)-N,N-diethylethan-1-amine			
273	(etodesnitazene);			
274	N,N-diethyl-2-(2-(4-fluorobenzyl)-5-nitro-1H-benzimidazol-1-yl)ethan-1-amine			
275	(flunitazene);			
276	N,N-diethyl-2-(2-(4-methoxybenzyl)-1H-benzimidazol-1-yl)ethan-1-amine			
277	(metodesnitazene);			
278	N,N-diethyl-2-(2-(4-methoxybenzyl)-5-nitro-1H-benzimidazol-1-yl)ethan-1-amine			
279	(metonitaze);			
280	2-(4-ethoxybenzyl)5-nitro-1-(2-(pyrrolidin-1-yl)ethyl)-1 H-benzimidazole (N-			

281	pyrrolidino etoitazene, etonitazepyne);
282	N,N-diethyl-2-(5-nitro-2-(4- propoxybenzyl)-1H-benzimidazol-1- yl)ethan-1-amine
283	(protonitazene);
284	N-pyrrolidino etonitazene;
285	Etodesnitazene;
286	Isotonitazene;
287	Protonitazene;
288	Metonitazene;
289	Butonitazene;
290	Metodesnitazene;
291	Flunitazene;
292	Opioid Receptor Agonist
293	AH-7921 (3,4-dichloro-N- (1dimethylamino)cyclohexylmethyl]benzamide).
294	Naphthoylindoles or any compound containing a 3-(-1- Napthoyl) indole structure with
295	substitution at the nitrogen atom of the indole ring whether or not further substituted in the indole
296	ring to any extent and whether or not substituted in the naphthyl ring to any extent. This shall
297	include the following:
298	JWH 015;
299	JWH 018;
300	JWH 019;
301	JWH 073;
302	JWH 081;
303	JWH 122;
304	JWH 200;
305	JWH 210;
306	JWH 398;

307	AM 2201; and
308	WIN 55,212.

Naphylmethylindoles or any compound containing a 1hindol-3-yl-(1-naphthyl) methane structure with a substitution at the nitrogen atom of the indole ring whether or not further substituted in the indole ring to any extent and whether or not substituted in the naphthyl ring to any extent. This shall include, but not be limited to, JWH 175 and JWH 184.

Naphthoylpyrroles or any compound containing a 3-(1- Naphthoyl) pyrrole structure with substitution at the nitrogen atom of the pyrrole ring whether or not further substituted in the pyrrole ring to any extent and whether or not substituted in the naphthyl ring to any extent. This shall include, but not be limited to, JWH 147 and JWH 307.

Naphthylmethylindenes or any compound containing a Naphthylideneindene structure with substitution at the 3- Position of the indene ring whether or not further substituted in the indene ring to any extent and whether or not substituted in the naphthyl ring to any extent. This shall include, but not be limited to, JWH 176.

Phenylacetylindoles or any compound containing a 3- Phenylacetylindole structure with substitution at the nitrogen atom of the indole ring whether or not further substituted in the indole ring to any extent and whether or not substituted in the phenyl ring to any extent. This shall include the following:

- 325 RCS-8, SR-18 OR BTM-8;
- 326 JWH 250;
- 327 JWH 203;
- 328 JWH 251; and
- 329 JWH 302.

Cyclohexylphenols or any compound containing a 2-(3- hydroxycyclohexyl) phenol structure with a substitution at the 5-position of the phenolic ring whether or not substituted in the cyclohexyl ring to any extent. This shall include the following:

333	CP 47,497 and its homologues and analogs;
334	Cannabicyclohexanol; and
335	CP 55,940.
336	Benzoylindoles or any compound containing a 3-(benzoyl) indole structure with
337	substitution at the nitrogren atom of the indole ring whether or not further substituted in the indole
338	ring to any extent and whether or not substituted in the phenyl ring to any extent. This shall include
339	the following:
340	AM 694;
341	Pravadoline WIN 48,098;
342	RCS 4; and
343	AM 679.
344	[2,3-dihydro-5 methyl-3-(4-morpholinylmethyl)pyrrolo [1,2,3-DE]-1, 4-benzoxazin-6-YL]-1-
345	napthalenymethanone. This shall include WIN 55,212-2.
346	Dibenzopyrans or any compound containing a 11-hydroxydelta 8-tetrahydrocannabinol
347	structure with substitution on the 3-pentyl group. This shall include HU-210, HU-211, JWH 051,
348	and JWH 133.
349	Adamantoylindoles or any compound containing a 3-(-1- Adamantoyl) indole structure with
350	substitution at the nitrogen atom of the indole ring whether or not further substituted in the
351	adamantoyl ring system to any extent. This shall include AM1248.
352	Tetramethylcyclopropylindoles or any compound containing A 3-
353	tetramethylcyclopropylindole structure with substitution at the nitrogen atom of the indole ring
354	whether or not further substituted in the indole ring to any extent and whether or not substituted in
355	the tetramethylcyclopropyl ring to any extent. This shall include UR-144 and XLR-11.
356	N-(1-Adamantyl)-1-pentyl-1h-indazole-3-carboxamide. This shall include AKB48.
357	Any other synthetic chemical compound that is a Cannabinoid receptor type 1 agonist as
358	demonstrated by binding studies and functional assays that is not listed in Schedules II, III, IV, and

359 V, not federal Food and Drug Administration approved drug or used within legitimate, approved medical research. Since nomenclature of these substances is not internationally standardized, 360 361 any immediate precursor or immediate derivative of these substances shall be covered. 362 Tryptamines: 363 5- methoxy- N- methyl-N-isopropyltryptamine (5-MeO-MiPT); 364 4-hydroxy-N,N-diisopropyltryptamine (4-HO-DiPT); 365 4-hydroxy-N-methyl-N-isopropyltryptamine (4-HO-MiPT); 366 4-hydroxy-N-methyl-N-ethyltryptamine (4-HO-MET): 367 4-acetoxy-N,N-diisopropyltryptamine (4-AcO-DiPT); 368 5-methoxy-α-methyltryptamine (5-MeO-AMT): 369 4-methoxy-N,N-Dimethyltryptamine (4-MeO-DMT); 370 4-hydroxy Diethyltryptamine (4-HO-DET); 371 5- methoxy- N,N- diallyltryptamine (5-MeO-DALT); 372 4-acetoxy-N,N-Dimethyltryptamine (4-AcO DMT); 373 4-hydroxy Diethyltryptamine (4-HO-DET); FDU-PB-22 (1-Naphthyl 1-(4-fluorobenzyl)-1H-indole-3-carboxylate); 374 375 FUB-PB-22 (Quinolin-8-yl 1-(4-fluorobenzyl)-1H-indole-3-carboxylate); 376 5-Fluoro-MN-24 (1-(5-Fluoropentyl)-N-(naphthalen-1-yl)-1H-indole-3-carboxamide); 377 MN-24 (N-(naphthalen-1-yl)-1-pentyl-1H-indole-3-carboxamide); 378 SDB-005 (Naphthalen-1-yl 1-pentyl-1H-indazole-3-carboxylate); 379 SDB-006 (1-Pentyl-N-(phenylmethyl)-1H-indole-3-carboxamide); 380 Methyl-Ethylaminopentiophenone; 381 FUB-AMB (Methyl(1-(4-fluorobenzyl)-1H-indazole-3-carbonyl)-L-valinate); 382 5-Fluoro-SDB-005 Indole (Naphthalen-1-yl 1-(5-fluoropentyl)-1H-indole-3-carboxylate); 383 5F-AB-PINACA (N-(1-Amino-3-methyl-1-oxobutan-2-yl)-1-(5-fluoropentyl)-1H-indazole-3-384 carboxamide);

385	MMB-CHMICA	(Methyl	2-(1-(cyclohexylmethyl)-1H-indo	ole-3-carboxamido)-3-
386	methylbutanoat);			
387	MN-24 (N-(naphthalen-1-yl)-1-pentyl-1H-indole-3-carboxamide);			
388	SDB-005 (Naphthalen-1-yl 1-pentyl-1H-indazole-3-carboxylate);			
389	SDB-006 (1-Pentyl-N-(phenylmethyl)-1H-indole-3-carboxamide);			
390	Ethcathinone (2-(ethylamino)-1-phenyl-1-propanone, monohydrochloride);			
391	Methyl-Ethylaminopentiophenone;			
392	FUB-AMB (Methyl(1-(4-fluorobenzyl)-1H-indazole-3-carbonyl)-L-valinate);			linate);
393	5-Fluoro-SDB-005 Inc	dole (Naphth	alen-1-yl 1-(5-fluoropentyl)-1H-ind	dole-3-carboxylate);
394	5F-AB-PINACA (N-	(1-Amino-3-r	nethyl-1-oxobutan-2-yl)-1-(5-fluor	opentyl)-1H-indazole-
395	3- carboxamide);			
396	MMB-CHMICA	(Methyl	2-(1-(cyclohexylmethyl)-1H-indo	ole-3-carboxamido)-3-
397	methylbutanoat);			
398	Bromazolam (8-brom	o-1-methyl-6	-phenyl-4H-[1,2,4]triazolo[4,3-a][1	1,4]benzodiazepine);
399	Clonazolam	(6-(2-chlorop	henyl)-1-methyl-8-nitro-4	H-[1,2,4]triazolo[4,3-
400	a][1,4]benzodiazepine);			
401	Cloniprazepam (5	-(2-chloroph	enyl)-1-(cyclopropylmethyl)-1,3-di	ihydro-7-nitro-2H-1,4-
402	benzodiazepin-2-one);			
403	Etizolam (4-(2-chl	orophenyl)-2	e-ethyl-9-methyl-6H-thieno[3,2-f]	[1,2,4]triazolo[4,3-
404	a][1,4]diazepine);			
405	Flualprazolam	(8-ch	ıloro-6-(2-fluorophenyl)-1-methyl-4	4H-[1,2,4]triazolo[4,3-
406	a][1,4]benzodiazepine);			
407	Flubromazepam (7-br	omo-5-(2-flu	orophenyl)-1,3-dihydro-2H-1,4-be	nzodiazepin-2-one);
408	Flubromazolam	(8-bro	omo-6-(2-fluorophenyl)-1-methyl-4	4H-[1,2,4]triazolo[4,3-
409	a][1,4]benzodiazepine);			
410	Flunitrazolam	(6-(2-fluoro	phenyl)-1-methyl-8-nitro-4H-benz	to[f][1,2,4]triazolo[4,3-

411 a][1,4]diazepine); 412 Nifoxipam (5-(2-fluorophenyl)-1,3-dihydro-3-hydroxy-7-nitro-2H-1,4-benzodiazepin-2-413 one); 414 Nitrazolam (1-methyl-8-nitro-6-phenyl-4H-[1,2,4]triazolo[4,3-a][1,4]benzodiazepine); and 415 Pyrazolam (8-bromo-1-methyl-6-(2-pyridinyl)-4H-[1,2,4]triazolo[4,3-416 a][1,4]benzodiazepine). 417 (e) Depressants. 418 4-CN-CUMYL-BUTINACA (1-(4-Cyanobutyl)-N-(2-phenylpropan-2- yl)-1H-indazole-3-419 carboxamide); 420 Alpha-Phenylacetoacetonitrile (3-Oxo-2-phenylbutanenitrile); 421 2-Fluoro Deschloroketamine (2-(2-Fluorophenyl)-2-(methylamino)-cyclohexanone, 422 monohydrochloride); 423 4-MEAP (2-(Ethylamino)-1-(4-methylphenyl)pentan-1-one); 424 Mecloqualone; 425 Methaqualone; 426 Bromazolam (8-bromo-1-methyl-6-phenyl-4H-[1,2,4]triazolo[4,3-a][1,4]benzodiazepine); (6-(2-chlorophenyl)-1-methyl-8-nitro-4 427 Clonazolam H-[1,2,4]triazolo[4,3 428 a][1,4]benzodiazepine); 429 Cloniprazepam (5-(2-chlorophenyl)-1-(cyclopropylmethyl)-1,3-dihydro-7-nitro-2H-1,4-430 benzodiazepin-2-one); 431 (4-(2-chlorophenyl)-2-ethyl-9-methyl-6H-thieno[3,2-f] [1,2,4]triazolo[4,3-Etizolam 432 a][1,4]diazepine); 433 Flualprazolam (8-chloro-6-(2-fluorophenyl)-1-methyl-4H-[1,2,4]triazolo[4,3-434 a][1,4]benzodiazepine); 435 Flubromazepam (7-bromo-5-(2-fluorophenyl)-1,3-dihydro-2H-1,4-benzodiazepin-2-one); 436 Flubromazolam (8-bromo-6-(2-fluorophenyl)-1-methyl-4H-[1,2,4]triazolo[4,3-

437	a][1,4]benzodiazepine);
438	Flunitrazolam (6-(2-fluorophenyl)-1-methyl-8-nitro-4H-benzo[f][1,2,4]triazolo[4,3-
439	a][1,4]diazepine);
440	Nifoxipam (5-(2-fluorophenyl)-1,3-dihydro-3-hydroxy-7-nitro-2H-1,4-benzodiazepin-2-
441	one);
442	Nitrazolam (1-methyl-8-nitro-6-phenyl-4H-[1,2,4]triazolo[4,3-a][1,4]benzodiazepine);
443	Pyrazolam (8-bromo-1-methyl-6-(2-pyridinyl)-4H-[1,2,4]triazolo[4,3-
444	a][1,4]benzodiazepine);
445	Declazepam (7-Chloro-5-(2-chlorophenyl)-1-methyl-1,3-dihydro-2H-1,4-benzodiazepin-2-
446	one); and
447	Deschloroetizolam (2-Ethyl-9-methyl-4-phenyl-6H-thieno[3,2- f][1,2,4]triazolo[4,3-
448	a][1,4]diazepine);
449	(f) Stimulants.
450	Aminorex; some other names: aminoxaphen; 2-amino-5- phenyl-2-oxazoline; or 4,5-
451	dihydro-5-phenyl-2-oxazolamine;
452	Cathinone; some trade or other names: 2-amino-1-phenyl-1- propanone, alpha-
453	aminopropiophenone, 2-aminopropiophenone and norephedrone;
454	Fenethylline;
455	Methcathinone, its immediate precursors and immediate derivatives, its salts, optical
456	isomers and salts of optical isomers; some other names: (2-(methylamino)-propiophenone; alpha-
457	(methylamino)propiophenone; 2-(methylamino)-1-phenylpropan-1- one; alpha—-
458	methylaminopropiophenone; monomethylpropion; 3,4-methylenedioxypyrovalerone and/or
459	mephedrone;3,4-methylenedioxypyrovalerone (MPVD); ephedrone; N-methylcathinone;
460	methylcathinone; AL-464; AL-422; AL- 463 and UR1432;
461	(+-) cis-4-methylaminorex; ((+-)cis-4,5-dihydro-4-methyl- 5-phenyl-2-oxazolamine);
462	N-ethylamphetamine:

463	N,N-dimethylamphetemine; also known as N,N-alpha- trimethyl-benzeneethanamine;
464	N,N-alpha-trimethylphenethylamine;
465	Alpha-pyrrolidinopentiophenone, also known as alpha-PVP, optical isomers, salts and
466	salts of isomers;
467	Substituted amphetamines:
468	2-Fluoroamphetamine;
469	3-Fluoroamphetamine;
470	4-Fluoroamphetamine;
471	2-chloroamphetamine;
472	3-chloroamphetamine;
473	4-chloroamphetamine;
474	2-Fluoromethamphetamine;
475	3-Fluoromethamphetamine;
476	4-Fluoromethamphetamine;
477	4-chloromethamphetamine;
478	Ethcathinone (2-(ethylamino)-1-phenyl-1-propanone, monohydrochloride);
479	Alpha-PHP (1-Phenyl-2-(pyrrolidin-1-yl)hexan-1-one);
480	MPHP (1-(4-Methylphenyl)-2-(pyrrolidin-1-yl)hexan-1-one);
481	PV8 (1-Phenyl-2-(pyrrolidin-1-yl)heptan-1-one);
482	4-Chloro-Alpha-PVP (1-(4-chlorophenyl)-2-(pyrrolidin-1-yl)pentan-1-one);
483	N-Ethylhexedrone (2-(Ethylamino)-1-phenylhexan-1-one);
484	Methoxetamine (2-(Ethylamino)-2-(3-methoxyphenyl)-cyclohexanone); and
485	3-Fluorophenmetrazine (2-(3-Fluorophenyl)-3-methylmorpholine);
486	(g) Temporary listing of substances subject to emergency scheduling. Any material,
487	compound, mixture, or preparation which contains any quantity of the following substances:
488	N-[1-benzyl-4-piperidyl]-N-phenylpropanamide (benzylfentanyl), its optical isomers, salts,

489	and salts of isomers;
490	N-[1-(2-thienyl)methyl-4-piperidyl]-N-phenylpropanamide (thenylfentanyl), its optical
491	isomers, salts, and salts of isomers.
492	N-benzylpiperazine, also known as BZP;
493	Cyclopentyl fentanyl (N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopentanecarboxamide);
494	4-fluorobutyryl fentanyl (N-(4-fluorophenyl)-N-[1-(2-phenylethyl)piperidin-4-yl]-
495	butyramide);
496	Isobutyryl fentanyl (2-methyl-N-phenyl-N-[1-(2-phenylethyl)piperidin-4-yl]-propanamide);
497	Methoxyacetyl fentanyl (2-methoxy-N-phenyl-N-[1-(2-phenylethyl)piperidin-4-yl]-
498	acetamide);
499	3-methylbutyryl fentanyl (N-[3-methyl-1-(2-phenylethyl)piperidin-4-yl]-N-
500	phenylbutyramide);
501	4-methoxybutyryl fentanyl (N-(4-methoxyphenyl)-N-(1-phenethylpiperidin-4-
502	yl)butyramide);
503	Ocfentanil (N-(2-fluorophenyl)-2-methoxy-N-[1-(2-phenylethyl)piperidin-4-yl]-acetamide);
504	Tetrahydrofuran fentanyl (N-(1-phenethylpiperidin-4-yl)-N-phenyltetrahydrofuran-2-
505	carboxamide); and
506	Valeryl fentanyl (N-phenyl-N-[1-(2-phenylethyl)piperidin-4-yl]pentanamide).
507	(h) The following controlled substances are included in Schedule I:
508	Synthetic Cathinones or any compound, except bupropion or compounds listed under a
509	different schedule, or compounds used within legitimate and approved medical research,
510	structurally derived from 2- Aminopropan-1-one by substitution at the 1-position with Monocyclic
511	or fused polycyclic ring systems, whether or not the compound is further modified in any of the
512	following ways:
513	By substitution in the ring system to any extent with Alkyl, alkylenedioxy, alkoxy, haloalkyl,
514	hydroxyl, or halide Substituents whether or not further substituted in the ring system by one or

515	more other univalent substituents;
516	By substitution at the 3-position with an acyclic alkyl substituent;
517	By substitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl or methoxybenzy
518	groups;
519	By inclusion of the 2-amino nitrogen atom in a cyclic structure; or
520	Any other synthetic chemical compound that is a Cannabinoid receptor type 1 agonist as
521	demonstrated by binding studies and functional assays that is not listed in Schedules II, III, IV, and
522	V, not federal Food and Drug Administration approved drug or used within legitimate, approved
523	medical research.
524	(i) Notwithstanding §60A-2-204(d) of this code, if a drug that contains the pharmaceutical
525	composition of crystalline polymorph psilocybin, known as COMP360 or any such trade name
526	approved by the United States Food and Drug Administration, is approved by the United States
527	Food and Drug Administration, it shall be lawful to prescribe, distribute, and market based upon
528	the recommendations of the United States Food and Drug Administration

NOTE: The purpose of this bill is to add crystalline polymorph psilocybin in the permitted list of distributed and prescribed drugs, if scheduled or descheduled by the Food and Drug Administration.

Strike-throughs indicate language that would be stricken from a heading or the present law and underscoring indicates new language that would be added.