

H. B. 2931

(BY DELEGATE(S) ASHLEY)

[Introduced February 24, 2015; referred to the
Committee on Health and Human Resources; and then to
the Committee on the Judiciary.]

A BILL to amend and reenact §60A-2-204 of the Code of West Virginia, 1931, as amended, relating to adding drugs to the classification of schedule I drugs.

Be it enacted by the Legislature of West Virginia:

That §60A-2-204 of the Code of West Virginia, 1931, as amended, be amended and reenacted to read as follows:

ARTICLE 2. STANDARDS AND SCHEDULES.

§60A-2-204. Schedule I.

- 1 (a) Schedule I shall consist of the drugs and other
- 2 substances, by whatever official name, common or usual name,
- 3 chemical name, or brand name designated, listed in this section.

4 (b) Opiates. Unless specifically excepted or unless listed in
5 another schedule, any of the following opiates, including their
6 isomers, esters, ethers, salts and salts of isomers, esters and
7 ethers, whenever the existence of such isomers, esters, ethers and
8 salts is possible within the specific chemical designation (for
9 purposes of subdivision (34) of this subsection only, the term
10 isomer includes the optical and geometric isomers):

11 (1) Acetyl-alpha-methylfentanyl (N-[1-(1-methyl-2-
12 phenethyl)-4-piperidinyl]—phenylacetamide);

13 (2) Acetylmethadol;

14 (3) Allylprodine;

15 (4) Alphacetylmethadol (except levoalphacetylmethadol also
16 known as levo-alpha-acetylmethadol, levomethadyl acetate, or
17 LAAM);

18 (5) Alphameprodine;

19 (6) Alphamethadol;

20 (7) Alpha-methylfentanyl (N-[1-(alpha-methyl-beta-phenyl)
21 ethyl-4-piperidyl] propionanilide; 1-(1-methyl-2-phenylethyl)-4-
22 (- propanilido) piperidine);

- 23 (8) Alpha-methylthiofentanyl (N-[1-methyl-2-(2-thienyl)
24 ethyl- 4-piperidiny]—phenylpropanamide);
- 25 (9) Benzethidine;
- 26 (10) Betacetylmethadol;
- 27 (11) Beta-hydroxyfentanyl (N-[1-(2-hydroxy-2-phenethyl)
28 -4- piperidiny]-N-phenylpropanamide);
- 29 (12) Beta-hydroxy-3-methylfentanyl (other name: N-[1-(2-
30 hydroxy-2-phenethyl)-3-methyl-4-piperidiny]-N-
31 phenylpropanamide);
- 32 (13) Betameprodine;
- 33 (14) Betamethadol;
- 34 (15) Betaprodine;
- 35 (16) Clonitazene;
- 36 (17) Dextromoramide;
- 37 (18) Diampromide;
- 38 (19) Diethylthiambutene;
- 39 (20) Difenoxin;
- 40 (21) Dimenoxadol;
- 41 (22) Dimepheptanol;
- 42 (23) Dimethylthiambutene;

- 43 (24) Dioxaphetyl butyrate;
- 44 (25) Dipipanone;
- 45 (26) Ethylmethylthiambutene;
- 46 (27) Etonitazene;
- 47 (28) Etoxeridine;
- 48 (29) Furethidine;
- 49 (30) Hydroxypethidine;
- 50 (31) Ketobemidone;
- 51 (32) Levomoramide;
- 52 (33) Levophenacymorphan;
- 53 (34) 3-Methylfentanyl (N-[3-methyl-1-(2-phenylethyl)-4-
- 54 piperidyl]-N-phenylpropanamide);
- 55 (35) 3-methylthiofentanyl (N-[3-methyl-1-(2-thienyl) ethyl-
- 56 4- piperidinyl]—phenylpropanamide);
- 57 (36) Morpheridine;
- 58 (37) MPPP (1-methyl-4-phenyl-4-propionoxypiperidine);
- 59 (38) Noracymethadol;
- 60 (39) Norlevorphanol;
- 61 (40) Normethadone;
- 62 (41) Norpipanone;

63 (42) Para-fluorofentanyl (N-(4-fluorophenyl)-N-[1-(2-
64 phenethyl)-4-piperidinyl] propanamide);

65 (43) PEPAP (1-(2-phenethyl)-4-phenyl-4-
66 acetoxypiperidine);

67 (44) Phenadoxone;

68 (45) Phenampromide;

69 (46) Phenomorphan;

70 (47) Phenoperidine;

71 (48) Piritramide;

72 (49) Proheptazine;

73 (50) Properidine;

74 (51) Propiram;

75 (52) Racemoramide;

76 (53) Thiofentanyl (N-phenyl-N-[1-(2-thienyl)ethyl-4-
77 piperidinyl]-propanamide);

78 (54) Tilidine;

79 (55) Trimeperidine.

80 (c) *Opium derivatives*. — Unless specifically excepted or
81 unless listed in another schedule, any of the following opium
82 immediate derivatives, its salts, isomers and salts of isomers

83 whenever the existence of such salts, isomers and salts of
84 isomers is possible within the specific chemical designation:

- 85 (1) Acetorphine;
- 86 (2) Acetyldihydrocodeine;
- 87 (3) Benzylmorphine;
- 88 (4) Codeine methylbromide;
- 89 (5) Codeine-N-Oxide;
- 90 (6) Cyprenorphine;
- 91 (7) Desomorphine;
- 92 (8) Dihydromorphine;
- 93 (9) Drotebanol;
- 94 (10) Etorphine (except HCl Salt);
- 95 (11) Heroin;
- 96 (12) Hydromorphenol;
- 97 (13) Methyldesorphine;
- 98 (14) Methyldihydromorphine;
- 99 (15) Morphine methylbromide;
- 100 (16) Morphine methylsulfonate;
- 101 (17) Morphine-N-Oxide;
- 102 (18) Myrophine;

103 (19) Nicocodeine;

104 (20) Nicomorphine;

105 (21) Normorphine;

106 (22) Pholcodine;

107 (23) Thebacon.

108 (d) *Hallucinogenic substances*. — Unless specifically
109 excepted or unless listed in another schedule, any material,
110 compound, mixture or preparation, which contains any quantity
111 of the following hallucinogenic substances, or which contains
112 any of its salts, isomers and salts of isomers, whenever the
113 existence of such salts, isomers, and salts of isomers is possible
114 within the specific chemical designation (for purposes of this
115 subsection only, the term “isomer” includes the optical, position
116 and geometric isomers):

117 (1) Alpha-ethyltryptamine; some trade or other names:
118 tryptamine; Monase; alpha-ethy-1H-indole-3-ethanamine; 3-(2-
119 aminobutyl) indole; alpha-ET; and AET;

120 (2) 4-bromo-2, 5-dimethoxy-amphetamine; some trade or
121 other names: 4-bromo-2,5-dimethoxy-alpha-
122 methylphenethylamine; 4-bromo-2,5-DMA;

123 (3) 4-Bromo-2,5-dimethoxyphenethylamine; some trade or
124 other names: 2-(4-bromo-2,5-dimethoxyphenyl)-1-aminoethane;
125 alpha- desmethyl DOB; 2C-B, Nexus;

126 (4) (A) N-(2-Methoxybenzyl)-4-bromo-2, 5-
127 dimethoxyphenethylamine. The substance has the acronym 25B-
128 NBOMe.

129 (B) 2-(4-chloro-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)
130 ethanamine (25C-NBOMe).

131 (C) 2-(4-iodo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)
132 ethanamine (25I-NBOMe)

133 (5) 2,5-dimethoxyamphetamine; some trade or other names:
134 2,5-dimethoxy-alpha-methylphenethylamine; 2,5-DMA;

135 ~~(5)~~ (6) 2,5-dimethoxy-4-ethylamphet-amine; some trade or
136 other names: DOET;

137 ~~(6)~~ (7) 2,5-dimethoxy-4-(n)-propylthiophenethylamine (other
138 name: 2C-T-7);

139 ~~(7)~~ (8) 4-methoxyamphetamine; some trade or other names:
140 4 - m e t h o x y - a l p h a - m e t h y l p h e n e t h y l a m i n e ;
141 paramethoxyamphetamine; PMA;

142 ~~(8)~~ (9) 5-methoxy-3, 4-methylenedioxy-amphetamine;

- 143 ~~(9)~~ (10) 4-methyl-2,5-dimethoxy-amphetamine; some trade
144 and other names: 4-methyl-2,5-dimethoxy-alpha-
145 methylphenethylamine; “DOM”; and “STP”;
- 146 ~~(10)~~ (11) 3,4-methylenedioxy amphetamine;
- 147 ~~(11)~~ (12) 3,4-methylenedioxymethamphetamine (MDMA);
- 148 ~~(12)~~ (13) 3,4-methylenedioxy-N-ethylamphetamine (also
149 known as – ethyl-alpha-methyl-3,4 (methylenedioxy)
150 phenethylamine, N-ethyl MDA, MDE, MDEA);
- 151 ~~(13)~~ (14) N-hydroxy-3,4-methylenedioxyamphetamine (also
152 known as – hydroxy-alpha-methyl-3,4 (methylenedioxy)
153 phenethylamine, and – hydroxy MDA);
- 154 ~~(14)~~ (15) 3,4,5-trimethoxy amphetamine;
- 155 ~~(15)~~ (16) 5-methoxy-N,N-dimethyltryptamine (5-MeO-
156 DMT);
- 157 ~~(16)~~ (17) Alpha-methyltryptamine (other name: AMT);
- 158 ~~(17)~~ (18) Bufotenine; some trade and other names:
159 3-(beta-Dimethylaminoethyl)-5-hydroxyindole; 3-(2-
160 dimethylaminoethyl) -5-indolol; N, N-dimethylserotonin; 5-
161 hydroxy-N,N- dimethyltryptamine; mappine;

- 162 ~~(18)~~ (19) Diethyltryptamine; sometrade and other names: N,
163 N-Diethyltryptamine; DET;
- 164 ~~(19)~~ (20) Dimethyltryptamine; some trade or other names:
165 DMT;
- 166 ~~(20)~~ (21) 5-Methoxy-N,N-diisopropyltryptamine (5-MeO-
167 DIPT);
- 168 ~~(21)~~ (22) Ibogaine; some trade and other names: 7-Ethyl-6,
169 6 Beta, 7, 8, 9, 10, 12, 13-octahydro-2-methoxy-6, 9-methano-
170 5H- pyrido [1', 2': 1, 2] azepino [5,4-b] indole; Tabernanthe
171 iboga;
- 172 ~~(22)~~ (23) Lysergic acid diethylamide;
- 173 ~~(23)~~ (24) Marihuana;
- 174 ~~(24)~~ (25) Mescaline;
- 175 ~~(25)~~ (26) Parahexyl-7374; some trade or other names: 3-
176 Hexyl -1-hydroxy-7, 8, 9, 10-tetrahydro-6, 6, 9-trimethyl-6H-
177 dibenzo [b,d] pyran; Synhexyl;
- 178 ~~(26)~~ (27) Peyote; meaning all parts of the plant presently
179 classified botanically as *Lophophora williamsii* Lemaire,
180 whether growing or not, the seeds thereof, any extract from any
181 part of such plant, and every compound, manufacture, salts,

182 immediate derivative, mixture or preparation of such plant, its
183 seeds or extracts;

184 ~~(27)~~ (28) N-ethyl-3-piperidyl benzilate;

185 ~~(28)~~ (29) N-methyl-3-piperidyl benzilate;

186 ~~(29)~~ (30) Psilocybin;

187 ~~(30)~~ (31) Psilocyn;

188 ~~(31)~~ (32) Tetrahydrocannabinols; synthetic equivalents of
189 the substances contained in the plant, or in the resinous
190 extractives of Cannabis, sp. and/or synthetic substances,
191 immediate derivatives and their isomers with similar chemical
192 structure and pharmacological activity such as the following:

193 delta-1 Cis or trans tetrahydrocannabinol, and their optical
194 isomers;

195 delta-6 Cis or trans tetrahydrocannabinol, and their optical
196 isomers;

197 delta-3,4 Cis or trans tetrahydrocannabinol, and its optical
198 isomers;

199 (Since nomenclature of these substances is not
200 internationally standardized, compounds of these structures,

201 regardless of numerical designation of atomic positions
202 covered.)

203 ~~(32)~~ (33) Ethylamine analog of phencyclidine; some trade or
204 other names: N-ethyl-1-phenylcyclohexylamine, (1-
205 phenylcyclohexyl) ethylamine, N-(1-phenylcyclohexyl)
206 ethylamine, cyclohexamine, PCE;

207 ~~(33)~~ (34) Pyrrolidine analog of phencyclidine; some trade or
208 other names: 1-(1-phenylcyclohexyl)-pyrrolidine, PCPy, PHP;

209 ~~(34)~~ (35) Thiophene analog of phencyclidine; some trade or
210 other names: 1-[1-(2-thienyl)-cyclohexyl]-piperidine, 2-
211 thienyl analog of phencyclidine; TPCP, TCP;

212 ~~(35)~~ (36) 1[1-(2-thienyl)cyclohexyl]pyrrolidine; some other
213 names: TCPy.

214 ~~(36)~~ (37) 4-methylmethcathinone (Mephedrone);

215 ~~(37)~~ (38) 3,4-methylenedioxypropylamphetamine (MDPV);

216 ~~(38)~~ (39) 2-(2,5-Dimethoxy-4-ethylphenyl)ethanamine (2C-
217 E);

218 ~~(39)~~ (40) 2-(2,5-Dimethoxy-4-methylphenyl)ethanamine
219 (2C-D)

- 220 ~~(40)~~ (41) 2-(4-Chloro-2,5-dimethoxyphenyl) ethanamine
221 (2C-C)
- 222 ~~(41)~~ (42) 2-(4-Iodo-2,5-dimethoxyphenyl)ethanamine (2C-I)
- 223 ~~(42)~~ (43) 2-[4-(Ethylthio)-2,5-dimethoxyphenyl]ethanamine
224 (2C-T-2)
- 225 ~~(43)~~ (44) 2-[4-(Isopropylthio)-2,5-dimethoxyphenyl]
226 ethanamine (2C-T-4)
- 227 ~~(44)~~ (45) 2-(2,5-Dimethoxyphenyl)ethanamine (2C-H)
- 228 ~~(45)~~ (46) 2-(2,5-Dimethoxy-4-nitro-phenyl)ethanamine
229 (2C-N)
- 230 ~~(46)~~ (47) 2-(2,5-Dimethoxy-4-(n)-propylphenyl)ethanamine
231 (2C-P)
- 232 ~~(47)~~ (48) 3,4-Methylenedioxy-N-methylcathinone
233 (Methylone)
- 234 ~~(48)~~ (49) 2,5-dimethoxy-4-(n)-propyltghiophenethylamine
235 (2C-T-7, itsoptical isomers, salts and salts of isomers)
- 236 ~~(49)~~ (50) 5-methoxy-N,N-dimethyltryptamine some trade or
237 other names: 5-methoxy-3-[2-(dimethylamino)ethyl]indole; 5-
238 MeO-DMT(5-MeO-DMT)
- 239 ~~(50)~~ (51) Alpha-methyltryptamine (other name: AMT)

240 ~~(51)~~ (52) 5-methoxy-N,N-diisopropyltryptamine (other
241 name: 5-MeO-DIPT)

242 ~~(52)~~ (53) Synthetic Cannabinoids as follows:

243 (A) 2-[(1R,3S)-3-hydroxycyclohexyl]-5-(2-methyloctan-2-
244 yl)phenol) {also known as CP 47,497 and homologues};

245 (B) rel-2-[(1S,3R)-3-hydroxycyclohexyl] -5-(2-
246 methylnonan-2-yl)phenol {also known as CP 47,497-C8
247 homolog};

248 (C) [(6aR)-9-(hydroxymethyl)-6, 6-dimethyl-3-(2-
249 methyloctan-2-yl)-6a, 7,10,10a-tetrahydrobenzo[c]chromen-1-
250 ol)] {also known as HU-210};

251 (D) (dexanabinol); (6aS,10aS)-9-(hydroxymethyl)-6,6-
252 dimethyl-3-(2-methyloctan-2-yl)-6a,7,10,10a-tetrahydrobenzol
253 [c]chromen-1-ol) {also known as HU-211};

254 (E) 1-Pentyl-3-(1-naphthoyl)indole {also known as JWH-
255 018};

256 (F) 1-Butyl-3-(1-naphthoyl)indole {also known as JWH-
257 073};

258 (G) (2-methyl-1-propyl-1H-indol-3-yl)-1-naphthalenyl-
259 methanone {also known as JWH-015};

260 (H) (1-hexyl-1H-indol-3-yl)-1-naphthalenyl-methanone

261 {also known as JWH-019};

262 (I) [1-[2-(4-morpholinyl) ethyl] -1H-indol-3-yl]-1-

263 naphthalenyl-methanone {also known as JWH-200};

264 (J) 1-(1-pentyl-1H-indol-3-yl)-2-(3-hydroxyphenyl)-

265 ethanone {also known as JWH-250};

266 (K) 2 - ((1 S , 2 S , 5 S) - 5 - h y d r o x y - 2 - (3 -

267 hydroxypropyl)cyclohexyl)-5-(2-methyloctan-2-yl)phenol {also

268 known as CP 55,940};

269 (L) (4-methyl-1-naphthalenyl) (1-pentyl-1H-indol-3-yl) -

270 methanone {also known as JWH-122};

271 (M) (4-methyl-1-naphthalenyl) (1-pentyl-1H-indol-3-yl) -

272 methanone {also known as JWH-398};

273 (N) (4-methoxyphenyl)(1-pentyl-1H-indol-3-yl)methanone

274 {also known as RCS-4};

275 (O) 1-(1-(2-cyclohexylethyl) -1H-indol-3-yl) -2-(2-

276 methoxyphenyl) ethanone {also known as RCS-8};

277 (P) 1-pentyl-3-[1-(4-methoxynaphthoyl)]indole (JWH-081);

278 (Q) 1-(5-fluoropentyl)-3-(1-naphthoyl)indole (AM2201);

279 and

- 280 (R) 1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole (AM694).
- 281 ~~(53)~~(54) Synthetic cannabinoids or any material, compound,
282 mixture or preparation which contains any quantity of the
283 following substances, including their analogues, congeners,
284 homologues, isomers, salts and salts of analogues, congeners,
285 homologues and isomers, as follows:
- 286 (A) CP 47,497 AND homologues, 2-[(1R,3S)-3-
287 Hydroxycyclohexyl]-5-(2-methyloctan-2-YL)phenol);
- 288 (B) HU-210, [(6AR,10AR)-9-(hydroxymethyl)-6,
289 6-dimethyl-3-(2-Methyloctan-2-YL)-6A,7,10,
290 10A-tetrahydrobenzo[C]chromen-1-OL)];
- 291 (C)HU-211, (dexanabinol, (6AS,10AS)-9-(hydroxymethyl)-
292 6,6-Dimethyl-3-(2-methyloctan-2-YL)-
293 6A,7,10,10tetrahydrobenzo[C]chromen-1-OL);
- 294 (D) JWH-018, 1-pentyl-3-(1-naphthoyl)indole;
- 295 (E) JWH-019, 1-hexyl-3-(1-naphthoyl)indole;
- 296 (F) JWH-073, 1-butyl-3-(1-naphthoyl)indole;
- 297 (G) JWH-200, (1-(2-morpholin-4-ylethyl)indol-3-yl)-
298 Naphthalen-1-ylmethanone;
- 299 (H) JWH-250, 1-pentyl-3-(2-methoxyphenylacetyl)indole.]

300 ~~(54)~~ (55) Synthetic cannabinoids including any material,
301 compound, mixture or preparation that is not listed as a
302 controlled substance in Schedule I through V, is not a federal
303 Food and Drug Administration approved drug or used within
304 legitimate and approved medical research and which contains
305 any quantity of the following substances, their salts, isomers,
306 whether optical positional or geometric, analogues, homologues
307 and salts of isomers, analogues and homologues, unless
308 specifically exempted, whenever the existence of these salts,
309 isomers, analogues, homologues and salts of isomers, analogues
310 and homologues if possible within the specific chemical
311 designation:

312 (A) Tetrahydrocannabinols meaning tetrahydrocannabinols
313 which are naturally contained in a plant of the genus cannabis as
314 well as synthetic equivalents of the substances contained in the
315 plant or in the resinous extractives of cannabis or synthetic
316 substances, derivatives and their isomers with analogous
317 chemical structure and or pharmacological activity such as the
318 following:

319 (i) DELTA-1 CIS OR trans tetrahydrocannabinol and their
320 Optical isomers.

321 (ii) DELTA-6 CIS OR trans tetrahydrocannabinol and their
322 optical isomers.

323 (iii) DELTA-3,4 CIS or their trans tetrahydrocannabinol and
324 their optical isomers.

325 (B) Naphthoylindoles or any compound containing a 3-(1-
326 Naphthoyl) indole structure with substitution at the nitrogen atom
327 of the indole ring whether or not further substituted in the indole
328 ring to any extent and whether or not substituted in the naphthyl
329 ring to any extent. This shall include the following:

330 (i) JWH 015;

331 (ii) JWH 018;

332 (iii) JWH 019;

333 (iv) JWH 073;

334 (v) JWH 081;

335 (vi) JWH 122;

336 (vii) JWH 200;

337 (viii) JWH 210;

338 (ix) JWH 398;

339 (x) AM 2201;

340 (xi) WIN 55,212.

341 ~~(55)~~ (56) Naphylmethyloindoles or any compound containing
342 a 1indol-3-yl-(1-naphthyl) methane structure with a substitution
343 at the nitrogen atom of the indole ring whether or not further
344 substituted in the indole ring to any extent and whether or not
345 substituted in the naphthyl ring to any extent. This shall include,
346 but not be limited to, JWH 175 and JWH 184.

347 ~~(56)~~ (57) Naphthoypyrroles or any compound containing a
348 3-(1- Naphthoyl) pyrrole structure with substitution at the
349 nitrogen atom of the pyrrole ring whether or not further
350 substituted in the pyrrole ring to any extent and whether or not
351 substituted in the naphthyl ring to any extent. This shall include,
352 but not be limited to, JWH 147 and JWH 307.

353 ~~(57)~~ (58) Naphthylmethyloindenes or any compound
354 containing a Naphthylideneindene structure with substitution at
355 the 3- Position of the indene ring whether or not further
356 substituted in the indene ring to any extent and whether or not
357 substituted in the naphthyl ring to any extent. This shall include,
358 but not be limited to, JWH 176.

359 ~~(58)~~ (59) Phenylacetylindoles or any compound containing
360 a 3- Phenylacetylindole structure with substitution at the
361 nitrogen atom of the indole ring whether or not further
362 substituted in the indole ring to any extent and whether or not
363 substituted in the phenyl ring to any extent. This shall include
364 the following:

365 (A) RCS-8, SR-18 OR BTM-8;

366 (B) JWH 250;

367 (C) JWH 203;

368 (D) JWH 251;

369 (E) JWH 302.

370 ~~(59)~~ (60) Cyclohexylphenols or any compound containing a
371 2-(3- hydroxycyclohexyl) phenol structure with a substitution at
372 the 5-position of the phenolic ring whether or not substituted in
373 the cyclohexyl ring to any extent. This shall include the
374 following:

375 (A) CP 47,497 and its homologues and analogs;

376 (B) Cannabicyclohexanol;

377 (C) CP 55,940.

378 ~~(60)~~ (61) Benzoylindoles or any compound containing a 3-
379 (benzoyl) indole structure with substitution at the nitrogen atom
380 of the indole ring whether or not further substituted in the indole
381 ring to any extent and whether or not substituted in the phenyl
382 ring to any extent. This shall include the following:

383 (A) AM 694;

384 (B) Pravadoline WIN 48,098;

385 (C) RCS 4;

386 (D) AM 679.

387 ~~(61)~~ (62) [2,3-dihydro-5 methyl-3-(4-
388 morpholinylmethyl)pyrrolo [1,2,3-DE]-1,4-benzoxazin-6-YL]-
389 1-naphthalenymethanone. This shall include WIN 55,212-2.

390 ~~(62)~~ (63) Dibenzopyrans or any compound containing a 11-
391 hydroxydelta 8-tetrahydrocannabinol structure with substitution
392 on the 3-pentyl group. This shall include HU-210, HU-211, JWH
393 051 and JWH 133.

394 ~~(63)~~ (64) Adamantoylindoles or any compound containing
395 a 3-(1- Adamantoyl) indole structure with substitution at the
396 nitrogen atom of the indole ring whether or not further

397 substituted in the adamantoyl ring system to any extent. This
398 shall include AM1248.

399 ~~(64)~~ (65) Tetramethylcyclopropylindoles or any compound
400 containing A 3-tetramethylcyclopropylindole structure with
401 substitution at the nitrogen atom of the indole ring whether or
402 not further substituted in the indole ring to any extent and
403 whether or not substituted in the tetramethylcyclopropyl ring to
404 any extent. This shall include UR-144 and XLR-11.

405 ~~(65)~~(66) N-(1-Adamantyl)-1-pentyl-1h-indazole-3-
406 carboxamide. This shall include AKB48.

407 ~~(66)~~ (67) Any other synthetic chemical compound that is a
408 Cannabinoid receptor type 1 agonist as demonstrated by binding
409 studies and functional assays that is not listed in Schedules II,
410 III, IV and V, not federal Food and Drug Administration
411 approved drug or used within legitimate, approved medical
412 research. Since nomenclature of these substances is not
413 internationally standardized, any immediate precursor or
414 immediate derivative of these substances shall be covered.

415 (68) Tryptamines:

416 (A) 5-methoxy-N-methyl-N-isopropyltryptamine (5-MeO-

417 MiPT)

418 (B) 4-hydroxy-N,N-diisopropyltryptamine (4-HO-DiPT)

419 (C) 4-hydroxy-N-methyl-N-isopropyltryptamine (4-HO-

420 MiPT)

421 (D) 4-hydroxy-N-methyl-N-ethyltryptamine (4-HO-MET)

422 (E) 4-acetoxy-N,N-diisopropyltryptamine (4-AcO-DiPT)

423 (F) 5-methoxy- α -methyltryptamine (5-MeO-AMT)

424 (G) 4-methoxy-N,N-Dimethyltryptamine (4-MeO-DMT)

425 (H) 4-hydroxy Diethyltryptamine (4-HO-DET)

426 (I) 5-methoxy-N,N-diallyltryptamine (5-MeO-DALT)

427 (J) 4-acetoxy-N,N-Dimethyltryptamine (4-AcO DMT)

428 (K) 4-hydroxy Diethyltryptamine (4-HO-DET)

429 (e) *Depressants*. — Unless specifically excepted or unless

430 listed in another schedule, any material, compound, mixture, or

431 preparation which contains any quantity of the following

432 substances having a depressant effect on the central nervous

433 system, including its salts, isomers and salts of isomers

434 whenever the existence of such salts, isomers and salts of

435 isomers is possible within the specific chemical designation:

436 (1) Mecloqualone;

437 (2) Methaqualone.

438 (f) *Stimulants*. — Unless specifically excepted or unless
439 listed in another schedule, any material, compound, mixture, or
440 preparation which contains any quantity of the following
441 substances having a stimulant effect on the central nervous
442 system, including its salts, isomers and salts of isomers:

443 (1) Aminorex; some other names: aminoxaphen; 2-amino-5-
444 phenyl-2-oxazoline; or 4,5-dihydro-5-phenyl-2-oxazolamine;

445 (2) Cathinone; some trade or other names: 2-amino-1-
446 phenyl-1- propanone, alpha-aminopropiophenone, 2-
447 aminopropiophenone and norephedrone;

448 (3) Fenethylamine;

449 (4) Methcathinone, its immediate precursors and immediate
450 derivatives, its salts, optical isomers and salts of optical isomers;
451 some other names: (2-(methylamino)-propionophenone; alpha-
452 (methylamino)propionophenone; 2-(methylamino)-1-
453 phenylpropan-1- one; alpha—methylaminopropiophenone;
454 monomethylpropion; 3,4-methylenedioxypropionophenone and/or
455 mephedrone; 3,4-methylenedioxypropionophenone (MPVD);

456 ephedrone; N-methylcathinone; methylcathinone; AL-464; AL-
457 422; AL- 463 and UR1432;

458 (5) (+-) cis-4-methylaminorex; ((+)-cis-4,5-dihydro-4-
459 methyl- 5-phenyl-2-oxazolamine);

460 (6) N-ethylamphetamine;

461 (7) N,N-dimethylamphetemine; also known as N,N-alpha-
462 trimethyl-benzeneethanamine; N,N-alpha-
463 trimethylphenethylamine.

464 (8) Alpha-pyrrolidinopentiophenone, also known as alpha-
465 PVP, optical isomers, salts and salts of isomers.

466 (9) Substituted amphetamines:

467 (A) 2-Fluoroamphetamine

468 (B) 3-Fluoroamphetamine

469 (C) 4-Fluoroamphetamine

470 (D) 2-chloroamphetamine

471 (E) 3-chloroamphetamine

472 (F) 4-chloroamphetamine

473 (G) 2-Fluoromethamphetamine

474 (H) 3-Fluoromethamphetamine

475 (I) 4-Fluoromethamphetamine

476 (J) 4-chloromethamphetamine

477 (g) Temporary listing of substances subject to emergency
478 scheduling. Any material, compound, mixture or preparation
479 which contains any quantity of the following substances:

480 (1) N-[1-benzyl-4-piperidyl]-N-phenylpropanamide
481 (benzylfentanyl), its optical isomers, salts, and salts of isomers.

482 (2) N-[1-(2-thienyl)methyl-4-piperidyl]-N-
483 phenylpropanamide (thenylfentanyl), its optical isomers, salts
484 and salts of isomers.

485 (3) N-benzylpiperazine, also known as BZP.

486 (h) The following controlled substances are included in
487 Schedule I:

488 (1) Synthetic Cathinones or any compound, except
489 bupropion or compounds listed under a different schedule, or
490 compounds used within legitimate and approved medical
491 research, structurally derived from 2- Aminopropan-1-one by
492 substitution at the 1-position with Monocyclic or fused
493 polycyclic ring systems, whether or not the compound is further
494 modified in any of the following ways:

495 (A) By substitution in the ring system to any extent with
496 Alkyl, alkylenedioxy, alkoxy, haloalkyl, hydroxyl or halide
497 Substituents whether or not further substituted in the ring system
498 by one or more other univalent substituents.

499 (B) By substitution at the 3-position with an acyclic alkyl
500 substituent.

501 (C) By substitution at the 2-amino nitrogen atom with alkyl,
502 dialkyl, benzyl or methoxybenzyl groups.

503 (D) By inclusion of the 2-amino nitrogen atom in a cyclic
504 structure.

505 (2) Any other synthetic chemical compound that is a
506 Cannabinoid receptor type 1 agonist as demonstrated by binding
507 studies and functional assays that is not listed in Schedules II,
508 III, IV and V, not federal Food and Drug Administration
509 approved drug or used within legitimate, approved medical
510 research.

NOTE: The purpose of this bill is to expand the state's Schedule 1 Drug list.

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

