To: Drug Policy

By: Representatives Yancey, McLean

HOUSE BILL NO. 4

AN ACT TO AMEND SECTION 41-29-113, MISSISSIPPI CODE OF 1972, TO ADD TIANEPTINE TO SCHEDULE I OF THE UNIFORM CONTROLLED SUBSTANCES ACT; AND FOR RELATED PURPOSES.

- 4 BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MISSISSIPPI:
- 5 **SECTION 1.** Section 41-29-113, Mississippi Code of 1972, is
- 6 amended as follows:
- 7 41-29-113.
- 8 SCHEDULE I
- 9 (a) Schedule I consists of the drugs and other substances,
- 10 by whatever official name, common or usual name, chemical name, or
- 11 brand name designated, that is listed in this section.
- 12 (b) **Opiates.** Unless specifically excepted or unless listed
- 13 in another schedule, any of the following opiates, including their
- 14 isomers, esters, ethers, salts and salts of isomers, esters and
- 15 ethers, whenever the existence of these isomers, esters, ethers
- 16 and salts is possible within the specific chemical designation:
- 17 (1) Acetyl-alpha-methylfentanyl
- 18 (N-[1-(1-methyl-2-phenethyl)-4-piperidinyl]-N-phenylacetamide);

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19
               (2)
                   Acetylmethadol;
20
               (3)
                   Acetyl fentanyl
21
    (N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide);
22
               (4) Acryl fentanyl
23
    (N-(1-phenethylpiperidin-4-yl)-N-phenylacrylamide), also known as
24
    acryloylfentanyl;
25
               (5) AH-7921 (3,4-dichloro-N-[(1-dimethylamino)
26
    cyclohexylmethyl]benzamide);
27
               (6) Allylprodine;
28
               (7)
                   Alphacetylmethadol, except levo-alphacetylmethadol
29
    (levo-alpha-acetylmethadol, levomethadyl acetate, or LAAM);
30
               (8)
                   Alphameprodine;
31
               (9)
                   Alphamethadol;
32
                   Alpha-Methylfentanyl
               (10)
33
    (N-[1-alpha-methyl-beta-phenyl)ethyl-4-piperidyl]propionanilide;
34
    1-(1-methyl-2-phenylethyl)-4-(N-propanilido)piperidine);
35
               (11) Alpha-Methylthiofentanyl
    (N-[1-methyl-2-(2-thienyl)ethyl-4-piperidinyl]-N-phenylpropanamide
36
37
    );
38
               (12) Benzethidine;
39
               (13)
                   Betacetylmethadol;
40
               (14)
                   Beta-Hydroxyfentanyl
41
    (N-[1-(2-hydroxy-2-phenethyl)-4-piperidinyl]-N-phenylpropanamide);
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42
                    Beta-Hydroxy-3-methylfentanyl
43
    (N-[1-(2-hydroxy-2-phenylethyl)-3-methyl-4-piperidinyl]-N-phenylpr
44
    opanamide);
45
               (16)
                   Beta-Hydroxythiofentanyl
46
    (N-[1-[2-hydroxy-2-(thiophen-2-yl)ethyl]piperidin-4-yl]-N-phenylpr
47
    opionamide);
48
               (17)
                    Betameprodine;
49
               (18)
                    Betamethadol;
50
                    Beta-Methyl fentanyl
               (19)
    (N-phenyl-N-(1-(2-phenylpropyl)piperidin-4-yl)propionamide), also
51
52
    known as \beta-methyl fentanyl;
53
               (20)
                    Beta'-Phenyl fentanyl
54
    (N-(1-phenethylpiperidin-4-yl)-N, 3-diphenylpropanamide), also
55
    known as \beta'-phenyl fentanyl or 3-phenylpropanoyl fentanyl;
56
               (21)
                   Betaprodine;
57
               (22)
                    Butyrl fentanyl
58
    (N-(1-phenethylpiperidin-4-yl)-N-phenylbutyramide);
59
               (23) Clonitazene;
60
               (24)
                    Crotonyl fentanyl
61
    ((E)-N-(1-phenethylpiperidin-4-yl)-N-phenylbut-2-enamide);
62
               (25)
                   Cyclopentyl fentanyl
63
    (N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopentanecarboxamide);
64
               (26)
                    Cyclopropyl fentanyl
65
    (N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopropanecarboxamide);
66
               (27)
                     Dextromoramide;
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67
               (28)
                     Diampromide;
68
               (29)
                     Diethylthiambutene;
69
                     Difenoxin;
               (30)
70
                     Dimenoxadol;
               (31)
71
               (32)
                     Dimepheptanol;
72
               (33)
                     Dimethylthiambutene;
73
                     Dioxaphetyl butyrate;
               (34)
74
               (35)
                     Dipipanone;
75
                     Ethylmethylthiambutene;
               (36)
76
               (37)
                     Etonitazene;
77
               (38)
                     Etoxeridine;
78
               (39)
                     Fentanyl carbamate
79
     (ethyl(1-phenethylpiperidin-4-yl)(phenyl)carbamate);
80
                     Fentanyl-related substances, meaning any substance
    not otherwise listed under another schedule and for which no
81
82
    exemption or approval is in effect under Section 505 of the
83
    Federal Food, Drug, and Cosmetic Act [21 USC 355] that is
84
    structurally related to fentanyl by one or more of the following
85
    modifications:
86
                         Replacement of the phenyl portion of the
                    (A)
87
    phenethyl group by any monocycle, whether or not further
88
    substituted in or on the monocycle;
89
                    (B)
                         Substitution in or on the phenethyl group with
90
    alkyl, alkenyl, alkoxyl, hydroxyl, halo, haloalkyl, amino or nitro
91
    groups;
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92
                     (C)
                          Substitution in or on the piperidine ring with
 93
     alkyl, alkenyl, alkoxyl, ester, ether, hydroxyl, halo, haloalkyl,
     amino or nitro groups;
 94
 95
                          Replacement of the aniline ring with any
                     (D)
96
     aromatic monocycle whether or not further substituted in or on the
97
     aromatic monocycle; and/or
98
                          Replacement of the N-propionyl group by
                     (E)
99
     another acyl group.
100
                     4-Fluoroisobutyryl fentanyl
                (41)
101
     (N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide),
102
     also known as para-fluoroisobutyryl fentanyl);
103
                      2'-Fluoro ortho-fluorofentanyl
                (42)
104
     (N-(1-(2-fluorophenethyl)piperidin-4-yl)-N-(2-fluorophenyl)
105
     propionamide), also known as 2'-fluoro 2-fluorofentanyl;
106
                (43)
                     Furanyl fentanyl
107
     (N-(1-phenethylpiperidin-4-yl)-N-phenylfuran-2-carboxamide);
108
                (44)
                     Furethidine;
109
                    Hydroxypethidine;
                (45)
110
                (46)
                      Isobutyryl fentanyl
111
     (N-(1-phenethylpiperidin-4-yl)-N-phenylisobutyramide);
112
                (47)
                      Isotonitazene (N, N-diethyl-2-(2-(4
113
     isopropoxybenzyl)-5-nitro-1H-benzimidazol-1-yl)ethan-1-amine);
114
                (48)
                     Ketobemidone (including the optical and geometric
115
     isomers);
                     Levomoramide;
116
                (49)
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117
                (50)
                     Levophenacylmorphan;
118
                (51)
                     Methoxyacetyl fentanyl
     (2-methoxy-N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide);
119
120
                (52)
                     4'-Methyl acetyl fentanyl
121
     (N-(1-(4-methyl)piperidin-4-yl)-N-phenylacetamide);
122
                (53)
                     3-Methylfentanyl
123
     (N-[3-methyl-1-(2-phenylethyl)-4-piperidyl]-N-phenylpropanamide);
124
                (54)
                     3-Methylthiofentanyl (N-[3-methyl-1-
125
     (2-thienylethyl)-4-piperidinyl]-N-phenylpropanamide);
126
                (55)
                    Morpheridine;
127
                (56)
                     MPPP (1-methyl-4-phenyl-4-propionoxypiperidine);
128
                    MT-45
                (57)
129
     (1-cyclohexyl-4-(1,2-diphenylethyl)piperazine);
130
                (58)
                     Noracymethadol;
131
                (59)
                     Norlevorphanol;
132
                (60)
                     Normethadone;
133
                (61)
                     Norpipanone;
134
                (62)
                     Ocfentanil (N-(2-fluorophenyl)-2-methoxy-N-
135
     (1-phenethylpiperidin-4-yl)acetamide);
136
                (63) Ortho-Fluoroacryl fentanyl
137
     (N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)acrylamide);
138
                (64) Ortho-Fluorobutyryl fentanyl
139
     (N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)butyramide), also
140
     known as 2-fluorobutyryl fentanyl;
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141
                (65) Ortho-Fluorofentanyl
     (N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)propionamide),
142
     also known as 2-fluorofentanyl;
143
144
                (66) Ortho-Fluoroisobutyryl fentanyl
145
     (N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide);
146
                (67)
                    Ortho-Methyl acetylfentanyl
147
     (N-(2-methylphenyl)-N-(1-phenethylpiperidin-4-yl)acetamide), also
148
     known as 2-methyl acetylfentanyl;
149
                (68) Ortho-Methyl methoxyacetyl fentanyl
150
     (2-methoxy-N-(2-methylphenyl)-N-(1-phenethylpiperidin-4-yl)
151
     acetamide), also known as 2-methyl methoxyacetyl fentanyl;
152
                (69)
                     Para-Chloroisobutyryl fentanyl
153
     (N-(4-chlorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide);
154
                     Para-Fluorobutyryl fentanyl
                (70)
155
     (N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)butyramide);
156
                (71)
                     Para-Fluorofentanyl (N-(4-fluorophenyl)
157
     -N-[1-(2-phenylethyl)-4-piperidinyl]propanamide);
158
                     Para-Fluoro furanyl fentanyl N-(4-fluorophenyl)-N-
                (72)
159
     (1-phenethylpiperidin-4-yl)furan-2-carboxamide);
160
                    Para-Methoxybutyryl fentanyl
                (73)
161
     (N-(4-methoxyphenyl)-N-(1-phenethylpiperidin-4-yl)butyramide);
162
                    Para-Methylfentanyl
                (74)
163
     (N-(4-methylphenyl)-N-(1-phenethylpiperidin-4-yl)propionamide),
164
     also known as 4-methylfentanyl);
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165
                (75)
                     PEPAP
166
     (1-(-2-phenethyl)-4-phenyl-4-acetoxypiperidine);
167
                (76)
                     Phenadoxone;
168
                    Phenampromide;
                (77)
169
                (78)
                     Phenomorphan;
170
                (79)
                     Phenoperidine;
171
                     Phenyl fentanyl
                (80)
172
     (N-(1-phenethylpiperidin-4-yl)-N-phenylbenzamide), also known as
173
     benzoyl fentanyl;
174
                (81) Piritramide;
175
                (82) Proheptazine;
176
                    Properidine;
                (83)
177
                (84)
                    Propiram;
178
                (85)
                    Racemoramide;
179
                     Tetrahydrofuranyl fentanyl
                (86)
180
     (N-(1-phenethylpiperidin-4-yl)-N-phenyltetrahydrofuran-2-
181
     carboxamide);
182
                (87)
                     Thiofentanyl
183
     (N-phenyl-N-[1-(2-thienyl)ethyl-4-piperidinyl]propanamide);
184
                (88)
                     Thiofuranyl fentanyl
185
     (N-(1-phenethylpiperidin-4-yl)-N-phenylthiophene-2-carboxamide),
     also known as 2-thiofuranyl fentanyl or thiophene fentanyl;
186
187
                (89) Tilidine;
188
                (90)
                    Trimeperidine;
                     U-47700, (3,4-dichloro-N-
189
                (91)
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190
     [2-(dimethylamino)cyclohexyl]-N-methylbenzamide);
                (92) Valeryl fentanyl
191
192
     (N-(1-phenethylpiperidin-4-yl)-N-phenylpentanamide).
193
               (93) Tianeptine and any salt, sulfate, free acid, or
194
     other preparation of Tianeptine, and any salt, sulfate, free acid,
195
     compound, derivative, precursor, or preparation thereof which is
196
     substantially chemically equivalent or identical with Tianeptine.
197
               Opium derivatives. Unless specifically excepted or
198
     unless listed in another schedule, any of the following opium
199
     derivatives, their salts, isomers and salts of isomers, whenever
200
     the existence of these salts, isomers and salts of isomers is
201
     possible within the specific chemical designation:
202
                (1)
                    Acetorphine;
203
                    Acetyldihydrocodeine;
                (2)
204
                (3)
                     Benzylmorphine;
                    Codeine methylbromide;
205
                (4)
206
                (5)
                     Codeine-N-Oxide;
207
                (6)
                     Cyprenorphine;
208
                (7)
                     Desomorphine;
209
                     Dihydromorphine;
                (8)
210
                (9)
                     Drotebanol;
211
                (10)
                     Etorphine (except hydrochloride salt);
212
                (11)
                     Heroin;
213
                     Hydromorphinol;
                (12)
214
                     Methyldesorphine;
                (13)
```

215	(14) Methyldihydromorphine;					
216	(15) Monoacetylmorphine;					
217	(16) Morphine methylbromide;					
218	(17) Morphine methylsulfonate;					
219	(18) Morphine-N-Oxide;					
220	(19) Myrophine;					
221	(20	Nicocodeine;					
222	(21	Nicomorphine;					
223	(22	Normorphine;					
224	(23) Pholcodine;					
225	(24) Thebacon.					
226	(d) Hal	lucinogenic substances. Unless specifically excepted					
227	or unless lis	ted in another schedule, any material, compound,					
228	mixture or preparation which contains any quantity of the						
229	following substances, their salts, isomers (whether optical,						
230	positional, o	r geometric) and salts of isomers, whenever the					
231	existence of	these salts, isomers and salts of isomers is possible					
232	within the sp	ecific chemical designation:					
233	(1)	Alpha-ethyltryptamine;					
234	(2)	4-bromo-2,5-dimethoxy-amphetamine;					
235	(3)	4-bromo-2,5-dimethoxyphenethylamine;					
236	(4)	2,5-dimethoxyamphetamine;					
237	(5)	2,5-dimethoxy-4-ethylamphetamine (DOET);					
238	(6)	2,5-dimethoxy-4-(n)-propylthiophenethylamine					
239	(2C-T-7);						

240	(7) 4-methoxyamphetamine;
241	(8) 5-methoxy-3,4-methylenedioxy-amphetamine;
242	(9) 4-methyl-2,5-dimethoxy-amphetamine;
243	(10) 3,4-methylenedioxy amphetamine;
244	(11) 3,4-methylenedioxymethamphetamine (MDMA);
245	(12) 3,4-methylenedioxy-N-ethylamphetamine (also known
246	as N-ethyl-alpha-methyl-3,4(methylenedioxy)phenethylamine, N-ethyl
247	MDA, MDE, MDEA);
248	(13) N-hydroxy-3,4-methylenedioxyamphetamine (also
249	known as N-hydroxy MDA, N-OHMDA, and
250	N-hydroxy-alpha-methyl-3,4 (methylenedioxy) phenethylamine);
251	(14) 3,4,5-trimethoxy amphetamine;
252	(15) 5-methoxy-N, N-dimethyltryptamine (5-MeO-DMT);
253	(16) Alpha-methyltryptamine (also known as AMT);
254	(17) Bufotenine;
255	(18) Diethyltryptamine;
256	(19) Dimethyltryptamine;
257	(20) 5-methoxy-N, N-diisopropyltryptamine (5-MeO-DIPT);
258	(21) Ibogaine;
259	(22) Lysergic acid diethylamide (LSD);
260	(23) (A) Marijuana (Hemp as defined and regulated
261	under Sections 69-25-201 through 69-25-221 and Cannabidiol
262	contained in a legend drug product approved by the Federal Food
263	and Drug Administration or obtained under Section 41-29-136 are
264	<pre>exempt under Schedule I);</pre>

265	(B) Hashish;					
266	(24) Mescaline;					
267	(25) Parahexyl;					
268	(26) Peyote;					
269	(27) N-ethyl-3-piperidyl benzilate;					
270	(28) N-methyl-3-piperidyl benzilate;					
271	(29) Psilocybin;					
272	(30) Psilocyn;					
273	(31) Tetrahydrocannabinols, meaning					
274	tetrahydrocannabinols contained in a plant of the genus Cannabis					
275	(cannabis plant), as well as the synthetic equivalents of the					
276	substances contained in the cannabis plant, or in the resinous					
277	extractives of such plant, and/or synthetic substances,					
278	derivatives, and their isomers with similar chemical structure and					
279	pharmacological activity to those substances contained in the					
280	plant such as the following:					
281	(A) 1 cis or trans tetrahydrocannabinol;					
282	(B) 6 cis or trans tetrahydrocannabinol;					
283	(C) 3,4 cis or trans tetrahydrocannabinol.					
284	(Since nomenclature of these substances is not					
285	internationally standardized, compounds of these structures,					
286	regardless of atomic positions, are covered.)					
287	("Tetrahydrocannabinols" excludes dronabinol and nabilone.)					
288	For purposes of this paragraph, tetrahydrocannabinols do not					

289	include hemp or hemp products regulated under Sections 69-25-201
290	through 69-25-221.
291	However, the following products are exempted from control:
292	(i) THC-containing industrial products made
293	from cannabis stalks (e.g., paper, rope and clothing);
294	(ii) Processed cannabis plant materials used
295	for industrial purposes, such as fiber retted from cannabis stalks
296	for use in manufacturing textiles or rope;
297	(iii) Animal feed mixtures that contain
298	sterilized cannabis seeds and other ingredients (not derived from
299	the cannabis plant) in a formula designed, marketed and
300	distributed for nonhuman consumption;
301	(iv) Personal care products that contain oil
302	from sterilized cannabis seeds, such as shampoos, soaps, and body
303	lotions (if the products do not cause THC to enter the human
304	body);
305	(v) Hemp as regulated under Sections
306	69-25-201 through 69-25-221; and
307	(vi) Any product derived from the hemp plant
308	designed for human ingestion and/or consumption that is approved
309	by the United States Food and Drug Administration;
310	(32) Phencyclidine;
311	(33) Ethylamine analog of phencyclidine (PCE);
312	(34) Pyrrolidine analog of phencyclidine (PHP, PCPy);

(35) Thiophene analog of phencyclidine;

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314
                (36)
                      1-[1-(2-thienyl)cyclohexyl] pyrrolidine (TCPy);
315
                      4-methylmethcathinone (mephedrone);
                (37)
                      3,4-methylenedioxypyrovalerone (MDPV);
316
                (38)
317
                      2-(2,5-dimethoxy-4-ethylphenyl)ethanamine (2C-E);
                (39)
318
                (40)
                      2-(2,5-dimethoxy-4-methylphenyl)ethanamine (2C-D);
319
                (41)
                     2-(4-chloro-2,5-dimethoxyphenyl)ethanamine (2C-C);
320
                     2-(4-iodo-2,5-dimethoxyphenyl)ethanamine (2C-I);
                (42)
321
     or 2,5-dimethoxy-4-iodophenethylamine;
322
                (43) 2-[4-(ethylthio)-2,5-dimethoxyphenyl]ethanamine
323
     (2C-T-2);
324
                (44)
325
     2-[4-(isopropylthio)-2,5-dimethoxyphenyl]ethanamine (2C-T-4);
326
                     2-(2,5-dimethoxyphenyl)ethanamine (2C-H);
                (45)
327
                     2-(2,5-dimethoxy-4-nitro-phenyl)ethanamine (2C-N);
                (46)
328
                (47) 2-(2,5-dimethoxy-4-(n)-propylphenyl)ethanamine
329
     (2C-P);
330
                     3,4-methylenedioxy-N-methylcathinone (methylone);
                (48)
                (49)
331
332
     2-(4-bromo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)ethanamine
333
     (25B-NBOMe; 2C-B-NBOMe; 25B; Cimbi-36);
334
                (50)
335
     2-(4-chloro-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)ethanamine
336
     (25C-NBOMe; 2C-C-NBOMe; 25C; Cimbi-82);
337
                (51)
     2-(4-iodo-2,5-dimethoxyphenyl)-N-(2-methoxybenzyl)ethanamine or
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339
     N-[(2-methoxyphenyl)methyl]ethanamine (25I-NBOMe; 2C-I-NBOMe; 25I;
340
     Cimbi-5);
341
                     7-bromo-5-(2-chlorophenyl)-1,3-dihydro-2H-1,
342
     4-benzodiazepin-2-one (also known as Phenazepam);
343
                (53)
                     7-(2-chlorophenyl)-4-ethyl-13-methyl-3-thia-1,8,
344
     11,12-tetraazatricyclo[8.3.0.0]trideca-2(6),4,7,10,12-pentaene
345
     (also known as Etizolam);
346
                (54)
                     Salvia divinorum;
347
                     Synthetic cannabinoids. Unless specifically
                (55)
     excepted or unless listed in another schedule, any material,
348
349
     compound, mixture, or preparation which contains any quantity of a
350
     synthetic cannabinoid found in any of the following chemical
351
     groups, whether or not substituted to any extent, or any of those
352
     groups which contain any synthetic cannabinoid salts, isomers, or
353
     salts of isomers, whenever the existence of such salts, isomers,
     or salts of isomers is possible within the specific chemical
354
355
     designation, including all synthetic cannabinoid chemical
356
     analogues in such groups:
357
                     (A)
                          (6aR, 10aR) - 9 - (hydroxymethyl) - 6,
358
     6-dimethyl-3-(2-methyloctan-2-yl)-6a,7,10,10a-tetrahydrobenzo[c]
359
     chromen-1-ol (also known as HU-210 or
360
     1,1-dimethylheptyl-11-hydroxy-delta8-tetrahydrocannabinol);
361
                          Naphthoylindoles and naphthylmethylindoles,
                     (B)
362
     being any compound structurally derived from 3-(1-naphthoyl)indole
     or 1H-indol-3-yl-(1-naphthyl) methane, whether or not substituted
363
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364	in the	indole	ring	to	any	extent,	or	in	the	naphthyl	ring	to	any
365	extent	;											

- 366 (C) Naphthoylpyrroles, being any compound
 367 structurally derived from 3-(1-naphthoyl)pyrrole, whether or not
 368 substituted in the pyrrole ring to any extent, or in the naphthyl
 369 ring to any extent;
- 370 (D) Naphthylmethylindenes, being any compound 371 structurally derived from 1-(1-naphthylmethyl)indene, whether or 372 not substituted in the indene ring to any extent or in the 373 naphthyl ring to any extent;
- 374 (E) Phenylacetylindoles, being any compound 375 structurally derived from 3-phenylacetylindole, whether or not 376 substituted in the indole ring to any extent or in the phenyl ring 377 to any extent;
- 378 (F) Cyclohexylphenols, being any compound 379 structurally derived from 2-(3-hydroxycyclohexyl)phenol, whether 380 or not substituted in the cyclohexyl ring to any extent or in the 381 phenolic ring to any extent;
- 382 (G) Benzoylindoles, whether or not substituted in the indole ring to any extent or in the phenyl ring to any extent;
- 384 (H) Adamantoylindoles, whether or not substituted 385 in the indole ring to any extent or in the adamantoyl ring system 386 to any extent;

387	(I) Tetrahydro derivatives of cannabinol and
388	3-alkyl homologues of cannabinol or of its tetrahydro derivatives,
389	except where contained in cannabis or cannabis resin;
390	(J) 3-Cyclopropylmethanone indole or
391	3-Cyclobutylmethanone indole or 3-Cyclopentylmethanone indole by
392	substitution at the nitrogen atom of the indole ring, whether or
393	not further substituted in the indole ring to any extent, whether
394	or not substituted on the cyclopropyl, cyclobutyl or cyclopentyl
395	rings to any extent;
396	(K) Quinolinyl ester indoles, being any compound
397	structurally derived from 1H-indole-3carboxylic acid-8-quinolinyl
398	ester, whether or not substituted in the indole ring to any extent
399	or the quinolone ring to any extent;
400	(L) 3-carboxamide-1H-indazoles, whether or not
401	substituted in the indazole ring to any extent and substituted to
402	any degree on the carboxamide nitrogen and
403	3-carboxamide-1H-indoles, whether or not substituted in the indole
404	ring to any extent and substituted to any degree on the
405	carboxamide nitrogen;
406	(M) Cycloalkanemethanone Indoles, whether or not
407	substituted at the nitrogen atom on the indole ring, whether or
408	not further substituted in the indole ring to any extent, whether

or not substituted on the cycloalkane ring to any extent;

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410
                (56)
                      Naphthalen-1-yl
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- 1-(5-fluoropentyl)-1H-indole-3-carboxylate, also known as NM2201 411
- 412 or CBL2201;
- 413 1-(5-fluoropentyl)-N-(2-phenylpropan-2-yl)-1H-
- 414 pyrrolo[2,3-b]pyridine-3-carboxamide, also known as
- 415 5F-CUMYL-P7AICA;
- 416 (58) 1-(4-methoxyphenyl)-N-methylpropan-2-amine, also
- 417 known as para-methoxymethamphetamine or PMMA.
- 418 Depressants. Unless specifically excepted or unless
- listed in another schedule, any material, compound, mixture, or 419
- 420 preparation which contains any quantity of the following
- 421 substances having a depressant effect on the central nervous
- 422 system, including their salts, isomers, and salts of isomers,
- 423 whenever the existence of such salts, isomers, and salts of
- 424 isomers is possible within the specific chemical designation:
- 425 (1)Clonazolam,
- 426 6-(2-chlorophenyl)-1-methyl-8-nitro-4H-[1,2,4] triazolo[4,3-a][1,4]
- 427 benzodiazepine;
- 428 (2) Flualprazolam,
- 429 8-chloro-6-(2-fluorophenyl)-1-methyl-4H-[1,2,4]triazolo[4,3-a][1,4]
- 430]benzodiazepine;

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- 431 (3) Flubromazepam,
- 432 7-bromo-5-(2-fluorophenyl)-1,3-dihydro-2H-1,4-benzodiazepin-2-one;
- 433 Flubromazolam, (4)

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435
     benzodiazepin;
436
                    Gamma-hydroxybutyric acid (other names include:
437
     GHB, gamma-hydroxybutyrate; 4-hydroxybutyrate; 4-hydroxybutanoic
438
     acid; sodium oxybate; sodium oxybutyrate);
439
                (6)
                    Mecloqualone;
440
                (7)
                    Methaqualone.
               Stimulants. Any material, compound, mixture or
441
           (f)
442
     preparation which contains any quantity of the following central
443
     nervous system stimulants including optical salts, isomers and
444
     salts of isomers unless specifically excepted or unless listed in
445
     another schedule:
446
                (1)
                    Aminorex;
447
                    N-benzylpiperazine (also known as BZP and
                (2)
448
     1-benzylpiperazine);
449
                (3)
                    Cathinone;
450
                    Fenethylline;
                (4)
451
                    Methcathinone;
                (5)
452
                (6)
                    4-methylaminorex (also known as
453
     2-amino-4-methyl-5-phenyl-2-oxazoline);
454
                (7)
                    N-ethylamphetamine;
455
                    Any material, compound, mixture or preparation
                (8)
456
     which contains any quantity of N, N-dimethylamphetamine. (Other
457
     names include: N,N,-alpha-trimethyl-benzeneethanamine and
458
     N, N-alpha-trimethylphenethylamine);
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8-bromo-6-(2-fluorophenyl)-1-methyl-4H-[1,2,4]triazolo[4,3-a][1,4]

459	(9) Synthetic cathinones. (A) Unless listed in
460	another schedule, any compound other than bupropion that is
461	structurally derived from 2-Amino-1-phenyl-1-propanone by
462	modification in any of the following ways:
463	(i) By substitution in the phenyl ring to any
464	extent with alkyl, alkoxy, alkylenedioxy, haloalkyl or halide
465	substituents, whether or not further substituted in the phenyl
466	ring by one or more other univalent substituents;
467	(ii) By substitution at the 3-position with
468	an alkyl substituent;
469	(iii) By substitution at the nitrogen atom
470	with alkyl or dialkyl groups, or by inclusion of the nitrogen atom
471	in a cyclic structure.
472	(B) The compounds covered in this paragraph (9)
473	include, but are not limited to, any material, compound, mixture
474	or preparation which contains any quantity of a synthetic
475	cathinone found in any of the following compounds, whether or not
476	substituted to any extent, or any of these compounds which contain
477	any synthetic cathinone, or salts, isomers, or salts of isomers,
478	whenever the existence of such salts, isomers or salts of isomers
479	is possible, unless specifically excepted or listed in another
480	schedule:
481	(i) 4-methyl-N-ethylcathinone ("4-MEC");
482	(ii) 4-methyl-alpha-pyrrolidinopropiophenone
483	("4-MePPP");

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484
                          (iii)
                                Alpha-pyrrolidinopentiophenone
485
     ("\alpha-PVP");
486
                          (iv)
487
     1-(1,3-benzodioxol-5-yl)-2-(methylamino)butan-1-one ("butylone");
488
                          (v) 2-(methylamino)-1-phenylpentan-1-one
489
     ("pentedrone");
490
                          (vi)
491
     1-(1,3-benzodioxol-5-yl)-2-(methylamino)pentan-1-one
492
     ("pentylone");
493
                                 4-fluoro-N-methylcathinone ("4-FMC");
                          (vii)
494
                          (viii) 3-fluoro-N-methylcathinone ("3-FMC");
495
                          (ix)
     1-(naphthalen-2-yl)-2-(pyrrolidin-1-yl)pentan-1-one ("naphyrone");
496
497
                          (x) Alpha-pyrrolidinobutiophenone ("\alpha-PBP");
498
     and
499
                          (xi)
500
     1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-pentan-1-one
501
     (N-ethylpentylone, ephylone).
502
          SECTION 2. This act shall take effect and be in force from
503
     and after July 1, 2023.
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