

**SENATE FLOOR VERSION**

March 4, 2025

3 COMMITTEE SUBSTITUTE  
FOR  
4 SENATE BILL NO. 860

4 SENATE BILL NO. 860 By: Weaver

An Act relating to the Uniform Controlled Dangerous Substances Act; amending 63 O.S. 2021, Section 2-204, as last amended by Section 3, Chapter 308, O.S.L. 2024 (63 O.S. Supp. 2024, Section 2-204), which relates to Schedule I substances; adding certain substance to Schedule I; and providing an effective date.

13 BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA:

14 SECTION 1. AMENDATORY 63 O.S. 2021, Section 2-204, as  
15 last amended by Section 3, Chapter 308, O.S.L. 2024 (63 O.S. Supp.  
16 2024, Section 2-204), is amended to read as follows:

17       Section 2-204. The controlled substances listed in this section  
18       are included in Schedule I and include any material, compound,  
19       mixture, or preparation that contains any quantity of the following  
20       hallucinogenic substances, their salts, isomers, and salts of  
21       isomers, unless specifically excepted, when the existence of these  
22       salts, isomers, and salts of isomers is possible within the specific  
23       chemical designation.

1       A. Any of the following opiates including their isomers,  
2 esters, ethers, salts, and salts of isomers, esters, and ethers,  
3 unless specifically excepted, when the existence of these isomers,  
4 esters, ethers, and salts is possible within the specific chemical  
5 designation:

- 6       1. Acetylmethadol;
- 7       2. Allylprodine;
- 8       3. Alphacetylmethadol;
- 9       4. Alphameprodine;
- 10      5. Alphamethadol;
- 11      6. Benzethidine;
- 12      7. Betacetylmethadol;
- 13      8. Betameprodine;
- 14      9. Betamethadol;
- 15      10. Betaprodine;
- 16      11. Clonitazene;
- 17      12. Dextromoramide;
- 18      13. Dextrorphan (except its methyl ether);
- 19      14. Diampromide;
- 20      15. Diethylthiambutene;
- 21      16. Dimenoxadol;
- 22      17. Dimepheptanol;
- 23      18. Dimethylthiambutene;
- 24      19. Dioxaphetyl butyrate;

- 1       20. Dipipanone;
- 2       21. Ethylmethylthiambutene;
- 3       22. Etonitazene;
- 4       23. Etoxeridine;
- 5       24. Furethidine;
- 6       25. Hydroxypethidine;
- 7       26. Isotonitazene;
- 8       27. Ketobemidone;
- 9       28. Levomoramide;
- 10      29. Levophenacylmorphan;
- 11      30. Metonitazene;
- 12      31. Morpheridine;
- 13      32. N-desethyl isotonitazene;
- 14      33. N-pyrrolidino protonitazene;
- 15      34. Noracymethadol;
- 16      35. Norlevorphanol;
- 17      36. Normethadone;
- 18      37. Norpipanone;
- 19      38. Phenadoxone;
- 20      39. Phenampromide;
- 21      40. Phenomorphan;
- 22      41. Phenoperidine;
- 23      42. Piritramide;
- 24      43. Proheptazine;

- 1       44. Properidine;  
2       45. Protonitazene;  
3       46. Racemoramide; or  
4       47. Trimeperidine.

5           B. Any of the following opium derivatives, their salts,  
6 isomers, and salts of isomers, unless specifically excepted, when  
7 the existence of these salts, isomers, and salts of isomers is  
8 possible within the specific chemical designation:

- 9       1. Acetorphine;  
10      2. Acetyldihydrocodeine;  
11      3. Benzylmorphine;  
12      4. Codeine methylbromide;  
13      5. Codeine-N-Oxide;  
14      6. Cyprenorphine;  
15      7. Desomorphine;  
16      8. Dihydromorphine;  
17      9. Etorphine;  
18     10. Heroin;  
19     11. Hydromorphenol;  
20     12. Methyldesorphine;  
21     13. Methylhydromorphone;  
22     14. Morphine methylbromide;  
23     15. Morphine methylsulfonate;  
24     16. Morphine-N-Oxide;

1       17. Myrophine;  
2       18. Nicocodeine;  
3       19. Nicomorphine;  
4       20. Normorphine;  
5       21. Phoclodine;  
6       22. Thebacon;  
7       23. N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-acetamide  
8       (Acetyl fentanyl);  
9       24. N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-butenamide  
10      (Crotonyl fentanyl);  
11      25. N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-2-  
12      furancarboxamide (Furanyl fentanyl);  
13      26. N-phenyl-1-(2-phenylethyl)-4-piperidinamine (4-ANPP);  
14      27. N-(1-phenethylpiperidin-4-yl)-N-  
15      phenylcyclopropanecarboxamide (Cyclopropyl fentanyl); or  
16      28. N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-butanamide  
17      (Butyrl fentanyl).

18      C. Any material, compound, mixture, or preparation which  
19      contains any quantity of the following hallucinogenic substances,  
20      their salts, isomers, and salts of isomers, unless specifically  
21      excepted, when the existence of these salts, isomers, and salts of  
22      isomers is possible within the specific chemical designation:

- 23      1. Methcathinone;  
24      2. 3, 4-methylenedioxy amphetamine;

- 1       3. 3, 4-methylenedioxy methamphetamine;
- 2       4. 5-methoxy-3, 4-methylenedioxy amphetamine;
- 3       5. 3, 4, 5-trimethoxy amphetamine;
- 4       6. Bufotenine;
- 5       7. Diethyltryptamine;
- 6       8. Dimethyltryptamine;
- 7       9. 4-methyl-2, 5-dimethoxyamphetamine;
- 8       10. Ibogaine;
- 9       11. Lysergic acid diethylamide;
- 10      12. Marijuana;
- 11      13. Mescaline;
- 12      14. N-benzylpiperazine;
- 13      15. N-ethyl-3-piperidyl benzilate;
- 14      16. N-methyl-3-piperidyl benzilate;
- 15      17. Psilocybin;
- 16      18. Psilocyn;
- 17      19. 2, 5 dimethoxyamphetamine;
- 18      20. 4 Bromo-2, 5-dimethoxyamphetamine;
- 19      21. 4 methoxyamphetamine;
- 20      22. Cyclohexamine;
- 21      23. Salvia Divinorum;
- 22      24. Salvinorin A;

1       25. Thiophene Analog of Phencyclidine. Also known as: 1-(1-(2-  
2 thiienyl) cyclohexyl) piperidine; 2-Thienyl Analog of Phencyclidine;  
3 TPCP, TCP;

4       26. Phencyclidine (PCP);

5       27. Pyrrolidine Analog for Phencyclidine. Also known as 1-(1-  
6 Phenylcyclohexyl) - Pyrrolidine, PCPy, PHP;

7       28. 1-(3-trifluoromethylphenyl) piperazine;

8       29. Flunitrazepam;

9       30. B-hydroxy-amphetamine;

10      31. B-ketoamphetamine;

11      32. 2,5-dimethoxy-4-nitroamphetamine;

12      33. 2,5-dimethoxy-4-bromophenethylamine;

13      34. 2,5-dimethoxy-4-chlorophenethylamine;

14      35. 2,5-dimethoxy-4-iodoamphetamine;

15      36. 2,5-dimethoxy-4-iodophenethylamine;

16      37. 2,5-dimethoxy-4-methylphenethylamine;

17      38. 2,5-dimethoxy-4-ethylphenethylamine;

18      39. 2,5-dimethoxy-4-fluorophenethylamine;

19      40. 2,5-dimethoxy-4-nitrophenethylamine;

20      41. 2,5-dimethoxy-4-ethylthio-phenethylamine;

21      42. 2,5-dimethoxy-4-isopropylthio-phenethylamine;

22      43. 2,5-dimethoxy-4-propylthio-phenethylamine;

23      44. 2,5-dimethoxy-4-cyclopropylmethylthio-phenethylamine;

24      45. 2,5-dimethoxy-4-tert-butylthio-phenethylamine;

- 1       46. 2,5-dimethoxy-4-(2-fluoroethylthio)-phenethylamine;
- 2       47. 5-methoxy-N, N-dimethyltryptamine;
- 3       48. N-methyltryptamine;
- 4       49. A-ethyltryptamine;
- 5       50. A-methyltryptamine;
- 6       51. N, N-diethyltryptamine;
- 7       52. N, N-diisopropyltryptamine;
- 8       53. N, N-dipropyltryptamine;
- 9       54. 5-methoxy-a-methyltryptamine;
- 10      55. 4-hydroxy-N, N-diethyltryptamine;
- 11      56. 4-hydroxy-N, N-diisopropyltryptamine;
- 12      57. 5-methoxy-N, N-diisopropyltryptamine;
- 13      58. 4-hydroxy-N-isopropyl-N-methyltryptamine;
- 14      59. 3,4-Methylenedioxymethcathinone (Methylone);
- 15      60. 3,4-Methylenedioxypyrovalerone (MDPV);
- 16      61. 3-Methylmethcathinone (Metaphedrone);
- 17      62. 4-Methylmethcathinone (Mephedrone);
- 18      63. 4-methoxymethcathinone;
- 19      64. 4-Fluoromethcathinone;
- 20      65. 3-Fluoromethcathinone;
- 21      66. 1-(8-bromobenzo 1,2-b;4,5-b' difuran-4-yl)-2-aminopropane;
- 22      67. 2,5-Dimethoxy-4-chloroamphetamine;
- 23      68. 4-Methylmethcathinone;
- 24      69. Pyrovalerone;

- 1       70. N,N-diallyl-5-methoxytryptamine;
- 2       71. 3,4-Methylenedioxy-N-ethylcathinone (Ethylone);
- 3       72. B-keto-N-Methylbenzodioxolylbutanamine (Butylone);
- 4       73. B-keto-Methylbenzodioxolylpentanamine (Pentylone);
- 5       74. Alpha-Pyrrolidinopentiophenone;
- 6       75. 4-Fluoroamphetamine;
- 7       76. Pentedrone;
- 8       77. 4'-Methyl-a-pyrrolidinohexaphenone;
- 9       78. 2,5-dimethoxy-4-(n)-propylphenethylamine;
- 10      79. 2,5-dimethoxyphenethylamine;
- 11      80. 1,4-Dibenzylpiperazine;
- 12      81. N,N-Dimethylamphetamine;
- 13      82. 4-Fluoromethamphetamine;
- 14      83. 4-Chloro-2,5-dimethoxy-N-(2-methoxybenzyl)phenethylamine  
15 (25C-NBOMe);
- 16      84. 4-Iodo-2,5-dimethoxy-N-(2-methoxybenzyl)phenethylamine  
17 (25I-NBOMe);
- 18      85. 4-Bromo-2,5-dimethoxy-N-(2-methoxybenzyl)phenethylamine  
19 (25B-NBOMe);
- 20      86. 1-(4-Fluorophenyl)piperazine;
- 21      87. Methoxetamine;
- 22      88. 3,4-dichloro-N[2-dimethylamino)cyclohexyl]-N-  
23 methylbenzamide;
- 24      89. N-ethyl hexadrone;

- 1       90. Isopropyl-U-47700;
- 2       91. Para-fluorobutyrl fentanyl;
- 3       92. Para-fluorofentanyl (pFF);
- 4       93. Fluoro isobutryrl fentanyl;
- 5       94. 3-Hydroxy Phencyclidine (PCP);
- 6       95. 3-methoxy Phencyclidine (PCP);
- 7       96. Flualprazolam; or
- 8       97. Flubromazolam.

9                   D. Unless specifically excepted or unless listed in a different  
10 schedule, any material, compound, mixture, or preparation which  
11 contains any quantity of the following substances having stimulant  
12 or depressant effect on the central nervous system:

- 13       1. Fenethylline;
- 14       2. Mecloqualone;
- 15       3. N-ethylamphetamine;
- 16       4. Methaqualone;
- 17       5. Gamma-Hydroxybutyric Acid, also known as GHB, gamma-  
18 hydroxybutyrate, 4-hydroxybutyrate, 4-hydroxybutanoic acid, sodium  
19 oxybate, and sodium oxybutyrate;
- 20       6. Gamma-Butyrolactone (GBL) as packaged, marketed,  
21 manufactured, or promoted for human consumption, with the exception  
22 of legitimate food additive and manufacturing purposes;

23  
24

1       7. Gamma Hydroxyvalerate (GHV) as packaged, marketed, or  
2 manufactured for human consumption, with the exception of legitimate  
3 food additive and manufacturing purposes;

4       8. Gamma Valerolactone (GVL) as packaged, marketed, or  
5 manufactured for human consumption, with the exception of legitimate  
6 food additive and manufacturing purposes;

7       9. 1,4 Butanediol (1,4 BD or BDO) as packaged, marketed,  
8 manufactured, or promoted for human consumption with the exception  
9 of legitimate manufacturing purposes; or

10      10. N-ethylpentylone.

11     E. 1. The following industrial uses of Gamma-Butyrolactone,  
12 Gamma Hydroxyvalerate, Gamma Valerolactone, or 1,4 Butanediol are  
13 excluded from all schedules of controlled substances under this  
14 title:

- 15       a. pesticides,
- 16       b. photochemical etching,
- 17       c. electrolytes of small batteries or capacitors,
- 18       d. viscosity modifiers in polyurethane,
- 19       e. surface etching of metal coated plastics,
- 20       f. organic paint disbursements for water soluble inks,
- 21       g. pH regulators in the dyeing of wool and polyamide  
22           fibers,
- 23       h. foundry chemistry as a catalyst during curing,

- 1           i.     curing agents in many coating systems based on  
2                       urethanes and amides,  
3           j.     additives and flavoring agents in food, confectionary,  
4                       and beverage products,  
5           k.     synthetic fiber and clothing production,  
6           l.     tetrahydrofuran production,  
7           m.     gamma butyrolactone production,  
8           n.     polybutylene terephthalate resin production,  
9           o.     polyester raw materials for polyurethane elastomers  
10                  and foams,  
11           p.     coating resin raw material, and  
12           q.     as an intermediate in the manufacture of other  
13                  chemicals and pharmaceuticals.

14       2. At the request of any person, the Director of the Oklahoma  
15       State Bureau of Narcotics and Dangerous Drugs Control may exempt any  
16       other product containing Gamma-Butyrolactone, Gamma Hydroxyvalerate,  
17       Gamma Valerolactone, or 1,4 Butanediol from being included as a  
18       Schedule I controlled substance if such product is labeled,  
19       marketed, manufactured, and distributed for legitimate industrial  
20       use in a manner that reduces or eliminates the likelihood of abuse.

21       3. In making a determination regarding an industrial product,  
22       the Director, after notice and hearing, shall consider the  
23       following:

- 24           a.     the history and current pattern of abuse,

- 1           b. the name and labeling of the product,  
2           c. the intended manner of distribution, advertising, and  
3           promotion of the product, and  
4           d. other factors as may be relevant to and consistent  
5           with the public health and safety.

6       4. The hearing shall be held in accordance with the procedures  
7       of the Administrative Procedures Act.

8           F. Any material, compound, mixture, or preparation, whether  
9       produced directly or indirectly from a substance of vegetable origin  
10      or independently by means of chemical synthesis, or by a combination  
11      of extraction and chemical synthesis, that contains any quantity of  
12      the following substances, or that contains any of their salts,  
13      isomers, and salts of isomers when the existence of these salts,  
14      isomers, and salts of isomers is possible within the specific  
15      chemical designation:

16       1. JWH-004;

17       2. JWH-007;

18       3. JWH-009;

19       4. JWH-015;

20       5. JWH-016;

21       6. JWH-018;

22       7. JWH-019;

23       8. JWH-020;

24       9. JWH-030;

1       10. JWH-046;

2       11. JWH-047;

3       12. JWH-048;

4       13. JWH-049;

5       14. JWH-050;

6       15. JWH-070;

7       16. JWH-071;

8       17. JWH-072;

9       18. JWH-073;

10      19. JWH-076;

11      20. JWH-079;

12      21. JWH-080;

13      22. JWH-081;

14      23. JWH-082;

15      24. JWH-094;

16      25. JWH-096;

17      26. JWH-098;

18      27. JWH-116;

19      28. JWH-120;

20      29. JWH-122;

21      30. JWH-145;

22      31. JWH-146;

23      32. JWH-147;

24      33. JWH-148;

1       34. JWH-149;

2       35. JWH-150;

3       36. JWH-156;

4       37. JWH-167;

5       38. JWH-175;

6       39. JWH-180;

7       40. JWH-181;

8       41. JWH-182;

9       42. JWH-184;

10      43. JWH-185;

11      44. JWH-189;

12      45. JWH-192;

13      46. JWH-193;

14      47. JWH-194;

15      48. JWH-195;

16      49. JWH-196;

17      50. JWH-197;

18      51. JWH-198;

19      52. JWH-199;

20      53. JWH-200;

21      54. JWH-201;

22      55. JWH-202;

23      56. JWH-203;

24      57. JWH-204;

1           58. JWH-205;  
2           59. JWH-206;  
3           60. JWH-207;  
4           61. JWH-208;  
5           62. JWH-209;  
6           63. JWH-210;  
7           64. JWH-211;  
8           65. JWH-212;  
9           66. JWH-213;  
10          67. JWH-234;  
11          68. JWH-235;  
12          69. JWH-236;  
13          70. JWH-237;  
14          71. JWH-239;  
15          72. JWH-240;  
16          73. JWH-241;  
17          74. JWH-242;  
18          75. JWH-243;  
19          76. JWH-244;  
20          77. JWH-245;  
21          78. JWH-246;  
22          79. JWH-248;  
23          80. JWH-249;  
24          81. JWH-250;

1           82. JWH-251;  
2           83. JWH-252;  
3           84. JWH-253;  
4           85. JWH-262;  
5           86. JWH-292;  
6           87. JWH-293;  
7           88. JWH-302;  
8           89. JWH-303;  
9           90. JWH-304;  
10          91. JWH-305;  
11          92. JWH-306;  
12          93. JWH-307;  
13          94. JWH-308;  
14          95. JWH-311;  
15          96. JWH-312;  
16          97. JWH-313;  
17          98. JWH-314;  
18          99. JWH-315;  
19          100. JWH-316;  
20          101. JWH-346;  
21          102. JWH-348;  
22          103. JWH-363;  
23          104. JWH-364;  
24          105. JWH-365;

1       106. JWH-367;  
2       107. JWH-368;  
3       108. JWH-369;  
4       109. JWH-370;  
5       110. JWH-371;  
6       111. JWH-373;  
7       112. JWH-386;  
8       113. JWH-387;  
9       114. JWH-392;  
10      115. JWH-394;  
11      116. JWH-395;  
12      117. JWH-397;  
13      118. JWH-398;  
14      119. JWH-399;  
15      120. JWH-400;  
16      121. JWH-412;  
17      122. JWH-413;  
18      123. JWH-414;  
19      124. JWH-415;  
20      125. CP-55, 940;  
21      126. CP-47, 497;  
22      127. HU-210;  
23      128. HU-211;  
24      129. WIN-55, 212-2;

1       130. AM-2201;  
2       131. AM-2233;  
3       132. JWH-018 adamantyl-carboxamide;  
4       133. AKB48;  
5       134. JWH-122 N-(4-pentenyl) analog;  
6       135. MAM2201;  
7       136. URB597;  
8       137. URB602;  
9       138. URB754;  
10      139. UR144;  
11      140. XLR11;  
12      141. A-796,260;  
13      142. STS-135;  
14      143. AB-FUBINACA;  
15      144. AB-PINACA;  
16      145. PB-22;  
17      146. AKB48 N-5-Fluorpentyl;  
18      147. AM1248;  
19      148. FUB-PB-22;  
20      149. ADB-FUBINACA;  
21      150. BB-22;  
22      151. 5-Fluoro PB-22; or  
23      152. 5-Fluoro AKB-48.  
24

1       G. In addition to those substances listed in subsection F of  
2 this section, unless specifically excepted or unless listed in  
3 another schedule, any material, compound, mixture, or preparation  
4 which contains any quantity of a synthetic cannabinoid found to be  
5 in any of the following chemical groups:

6       1. Naphthoylindoles: any compound containing a 3-(1-  
7 naphthoyl)indole structure with or without substitution at the  
8 nitrogen atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl,  
9 alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-  
10 (N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-  
11 2-pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,  
12 (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
13 halophenyl group, whether or not further substituted on the indole  
14 ring to any extent, and whether or not substituted on the naphthyl  
15 ring to any extent. Naphthoylindoles include, but are not limited  
16 to:

- 17       a. 1-[2-(4-morpholinyl)ethyl]-3-(1-naphthoyl)indole (JWH-  
18                   200),  
19       b. 1-(5-fluoropentyl)-3-(1-naphthoyl)indole (AM2201),  
20       c. 1-pentyl-3-(1-naphthoyl)indole (JWH-018),  
21       d. 1-butyl-3-(1-naphthoyl)indole (JWH-073),  
22       e. 1-pentyl-3-(4-methoxy-1-naphthoyl)indole (JWH-081),  
23       f. 1-propyl-2-methyl-3-(1-naphthoyl)indole (JWH-015),  
24       g. 1-hexyl-3-(1-naphthoyl)indole (JWH-019),

- h. 1-pentyl-3-(4-methyl-1-naphthoyl)indole (JWH-122),
  - i. 1-pentyl-3-(4-ethyl-1-naphthoyl)indole (JWH-210),
  - j. 1-pentyl-3-(4-chloro-1-naphthoyl)indole (JWH-398),
  - k. 1-pentyl-2-methyl-3-(1-naphthoyl)indole (JWH-007),
  - l. 1-pentyl-3-(7-methoxy-1-naphthoyl)indole (JWH-164),
  - m. 1-pentyl-2-methyl-3-(4-methoxy-1-naphthoyl)indole (JWH-098),
  - n. 1-pentyl-3-(4-fluoro-1-naphthoyl)indole (JWH-412),
  - o. 1-[1-(N-methyl-2-piperidinyl)methyl]-3-(1-naphthoyl)indole (AM-1220),
  - p. 1-(5-fluoropentyl)-3-(4-methyl-1-naphthoyl)indole (MAM-2201), or
  - q. 1-(4-cyanobutyl)-3-(1-naphthoyl)indole (AM-2232);

2. Naphthylmethylindoles: any compound containing a 1H-indol-

15 3-yl-(1-naphthyl)methane structure with or without substitution at

16 the nitrogen atom of the indole ring by an alkyl, haloalkyl,

17 cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl,

18 | halobenzyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-

19 morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-

20 morpholinyl)methyl, (tetrahydropyran-4-yl)methyl, 1-methylazepanyl,

21 phenyl, or halophenyl group, whether or not further substituted on

the indole ring to any extent, and whether or not substituted on the

23 naphthyl ring to any extent. Naphthylmethylindoles include, but are

24 not limited to, (1-pentylinol-3-yl) (1-naphthyl) methane (JWH-175);

1       3. Naphthoylpyrroles: any compound containing a 3-(1-  
2 naphthoyl)pyrrole structure with or without substitution at the  
3 nitrogen atom of the pyrrole ring by an alkyl, haloalkyl,  
4 cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl,  
5 halobenzyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-  
6 morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-  
7 morpholinyl)methyl, (tetrahydropyran-4-yl)methyl, 1-methylazepanyl,  
8 phenyl, or halophenyl group, whether or not further substituted on  
9 the pyrrole ring to any extent, and whether or not substituted on  
10 the naphthyl group to any extent. Naphthoylpyrroles include, but  
11 are not limited to:

- a. 1-hexyl-2-phenyl-4-(1-naphthoyl)pyrrole (JWH-147),
  - b. 1-pentyl-5-(2-methylphenyl)-3-(1-naphthoyl)pyrrole (JWH-370),
  - c. 1-pentyl-3-(1-naphthoyl)pyrrole (JWH-030), or
  - d. 1-hexyl-5-phenyl-3-(1-naphthoyl)pyrrole (JWH-147);

17       4. Naphthylideneindenes: any compound containing a 1-(1-  
18 naphthylmethylene)indene structure with or without substitution at  
19 the 3-position of the indene ring by an alkyl, haloalkyl,  
20 cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl,  
21 halobenzyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-  
22 morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-  
23 morpholinyl)methyl, (tetrahydropyran-4-yl)methyl, 1-methylazepanyl,  
24 phenyl, or halophenyl group, whether or not further substituted on

the indene group to any extent, and whether or not substituted on the naphthyl group to any extent. Naphthylmethylindenes include, but are not limited to, (1-[(3-pentyl)-1H-inden-1-ylidene)methyl]naphthalene (JWH-176);

5       5. Phenylacetylindoles: any compound containing a 3-  
6 phenylacetylindole structure with or without substitution at the  
7 nitrogen atom of the indole ring by alkyl, haloalkyl, cyanoalkyl,  
8 alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-  
9 (N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-  
10 2-pyrrolidinyl)methyl, 1-(N-methyl-3- morpholinyl)methyl,  
11 (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
12 halophenyl group, whether or not further substituted on the indole  
13 ring to any extent, and whether or not substituted on the phenyl  
14 ring to any extent. Phenylacetylindoles include, but are not  
15 limited to:

- a. 1-pentyl-3-(2-methoxyphenylacetyl)indole (JWH-250),
  - b. 1-(2-cyclohexylethyl)-3-(2-methoxyphenylacetyl)indole (RCS-8),
  - c. 1-pentyl-3-(2-chlorophenylacetyl)indole (JWH-203),
  - d. 1-pentyl-3-(2-methylphenylacetyl)indole (JWH-251),
  - e. 1-pentyl-3-(4-methoxyphenylacetyl)indole (JWH-201), or
  - f. 1-pentyl-3-(3-methoxyphenylacetyl)indole (JWH-302);

23       6. Cyclohexylphenols: any compound containing a 2-(3-  
24 hydroxycyclohexyl)phenol structure with or without substitution at

1 the 5-position of the phenolic ring by an alkyl, haloalkyl,  
2 cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl,  
3 halobenzyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-  
4 morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-  
5 morpholinyl)methyl, (tetrahydropyran-4-yl)methyl, 1-methylazepanyl,  
6 phenyl, or halophenyl group, and whether or not further substituted  
7 on the cyclohexyl ring to any extent. Cyclohexylphenols include,  
8 but are not limited to:

- 9 a. 5-(1,1-dimethylheptyl)-2-[(1R,3S)-3-  
10 hydroxycyclohexyl]-phenol (CP-47,497),
- 11 b. 5-(1,1-dimethyloctyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-  
12 phenol (cannabicyclohexanol; CP-47,497 C8 homologue),  
13 or
- 14 c. 5-(1,1-dimethylheptyl)-2-[(1R,2R)-5-hydroxy-2-(3-  
15 hydroxypropyl)cyclohexyl]-phenol (CP 55, 940);

16 7. Benzoylindoles: any compound containing a 3-(benzoyl)indole

17 structure with or without substitution at the nitrogen atom of the  
18 indole ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl,  
19 cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-(N-methyl-  
20 2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-  
21 pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,  
22 (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
23 halophenyl group, whether or not further substituted on the indole  
24 ring to any extent, and whether or not substituted on the phenyl

1 group to any extent. Benzoylindoles include, but are not limited  
2 to:

- 3 a. 1-pentyl-3-(4-methoxybenzoyl)indole (RCS-4),
- 4 b. 1-[2-(4-morpholinyl)ethyl]-2-methyl-3-(4-  
methoxybenzoyl)indole (Pravadoline or WIN 48, 098),
- 5 c. 1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole (AM-694),
- 6 d. 1-pentyl-3-(2-iodobenzoyl)indole (AM-679), or
- 7 e. 1-[1-(N-methyl-2-piperidinyl)methyl]-3-(2-  
iodobenzoyl)indole (AM-2233);

10 8. Cyclopropoylindoles: Any compound containing a 3-

11 (cyclopropoyl)indole structure with substitution at the nitrogen  
12 atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl,  
13 cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-(N-methyl-  
14 2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-  
15 pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,  
16 (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
17 halophenyl group, whether or not further substituted in the indole  
18 ring to any extent, and whether or not substituted in the  
19 cyclopropoyl ring to any extent. Cyclopropoylindoles include, but  
20 are not limited to:

- 21 a. 1-pentyl-3-(2,2,3,3-tetramethylcyclopropoyl)indole  
22 (UR-144),
- 23 b. 1-(5-chloropentyl)-3-(2,2,3,3-  
24 tetramethylcyclopropoyl)indole (5Cl-UR-144), or

1           c. 1-(5-fluoropentyl)-3-(2,2,3,3-

2                 tetramethylcyclopropoyl)indole (XLR11);

3         9. Indole Amides: Any compound containing a 1H-Indole-3-

4 carboxamide structure with or without substitution at the nitrogen

5 atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl,

6 cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-(N-methyl-

7 2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-

8 pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,

9 (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or

10 halophenyl group, whether or not substituted at the carboxamide

11 group by an adamantyl, naphthyl, phenyl, benzyl, quinolinyl,

12 cycloalkyl, 1-amino-3-methyl-1-oxobutan-2-yl, 1-amino-3,3-dimethyl-

13 1-oxobutan-2-yl, 1-methoxy-3-methyl-1-oxobutan-2-yl, 1-methoxy-3,3-

14 dimethyl-1-oxobutan-2-yl or pyrrole group, and whether or not

15 further substituted in the indole, adamantyl, naphthyl, phenyl,

16 pyrrole, quinolinyl, or cycloalkyl rings to any extent. Indole

17 Amides include, but are not limited to:

18           a. N-(1-adamantyl)-1-pentyl-1H-indole-3-carboxamide

19                     (2NE1),

20           b. N-(1-adamantyl)-1-(5-fluoropentyl)-1H-indole-3-

21                 carboxamide (STS-135),

22           c. N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentyl-1H-

23                 indole-3-carboxamide (ADBICA),

24

- 1           d. N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(5-  
2                 fluoropentyl)-1H-indole-3-carboxamide (5F-ADBICA),  
3  
4           e. N-(naphthalen-1-yl)-1-pentyl-1H-indole-3-carboxamide  
5                 (NNE1),  
6  
7           f. 1-(5-fluoropentyl)-N-(naphthalene-1-yl)-1H-indole-3-  
8                 carboxamide (5F-NNE1),  
9  
10          g. N-benzyl-1-pentyl-1H-indole-3-carboxamide (SDB-006),  
11                 or  
12          h. N-benzyl-1-(5-fluoropentyl)-1H-indole-3-carboxamide  
13                 (5F-SDB-006);

14       10. Indole Esters: Any compound containing a 1H-Indole-3-  
15       carboxylate structure with or without substitution at the nitrogen  
16       atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl,  
17       cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-(N-methyl-  
18       2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-  
19       pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,  
20       (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
21       halophenyl group, whether or not substituted at the carboxylate  
22       group by an adamantyl, naphthyl, phenyl, benzyl, quinolinyl,  
23       cycloalkyl, 1-amino-3-methyl-1-oxobutan-2-yl, 1-amino-3,3-dimethyl-  
24       1-oxobutan-2-yl, 1-methoxy-3-methyl-1-oxobutan-2-yl, 1-methoxy-3,3-  
     dimethyl-1-oxobutan-2-yl or pyrrole group, and whether or not  
     further substituted in the indole, adamantyl, naphthyl, phenyl,

1 pyrrole, quinolinyl, or cycloalkyl rings to any extent. Indole  
2 Esters include, but are not limited to:

- 3 a. quinolin-8-yl 1-pentyl-1H-indole-3-carboxylate (PB-  
4 22),
- 5 b. quinolin-8-yl 1-(5-fluoropentyl)-1H-indole-3-  
6 carboxylate (5F-PB-22),
- 7 c. quinolin-8-yl 1-(cyclohexylmethyl)-1H-indole-3-  
8 carboxylate (BB-22),
- 9 d. naphthalen-1-yl 1-(4-fluorobenzyl)-1H-indole-3-  
10 carboxylate (FDU-PB-22), or
- 11 e. naphthalen-1-yl 1-(5-fluoropentyl)-1H-indole-3-  
12 carboxylate (NM2201);

13 11. Adamantanoylindoles: Any compound containing an

14 adamanyl-(1H-indol-3-yl)methanone structure with or without  
15 substitution at the nitrogen atom of the indole ring by an alkyl,  
16 haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl,  
17 benzyl, halobenzyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-  
18 morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-  
19 morpholinyl)methyl, (tetrahydropyran-4-yl)methyl, 1-methylazepanyl,  
20 phenyl, or halophenyl group, whether or not further substituted in  
21 the indole ring to any extent, and whether or not substituted in the  
22 adamanyl ring to any extent. Adamantanoylindoles include, but are  
23 not limited to:

- 1           a. adamantan-1-yl[1-[ (1-methyl-2-piperidinyl)methyl]-1H-  
2                 indol-3-yl]methanone (AM1248), or  
3           b. adamantan-1-yl-(1-pentyl-1H-indol-3-yl)methanone (AB-  
4                 001);

5           12. Carbazole Ketone: Any compound containing (9H-carbazole-3-  
6                 yl) methanone structure with or without substitution at the nitrogen  
7                 atom of the carbazole ring by an alkyl, haloalkyl, cyanoalkyl,  
8                 alkenyl, cycloalkylmethyl, cycloalkylethyl, benzyl, halobenzyl, 1-  
9                 (N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-  
10                 2-pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,  
11                 (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
12                 halophenyl group, with substitution at the carbon of the methanone  
13                 group by an adamantyl, naphthyl, phenyl, benzyl, quinolinyl,  
14                 cycloalkyl, 1-amino-3-methyl-1-oxobutan-2-yl, 1-amino-3,3-dimethyl-  
15                 1-oxobutan-2-yl, 1-methoxy-3-methyl-1-oxobutan-2-yl, 1-methoxy-3,3-  
16                 dimethyl-1-oxobutan-2-yl or pyrrole group, and whether or not  
17                 further substituted at the carbazole, adamantyl, naphthyl, phenyl,  
18                 pyrrole, quinolinyl, or cycloalkyl rings to any extent. Carbazole  
19                 Ketones include, but are not limited to, naphthalen-1-yl(9-pentyl-  
20                 9H-carbazol-3-yl)methanone (EG-018);

21           13. Benzimidazole Ketone: Any compound containing  
22                 (benzimidazole-2-yl) methanone structure with or without  
23                 substitution at either nitrogen atom of the benzimidazole ring by an  
24                 alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl,

1    cycloalkylethyl, benzyl, halobenzyl, 1-(N-methyl-2-  
2    piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-  
3    pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl,  
4    (tetrahydropyran-4-yl)methyl, 1-methylazepanyl, phenyl, or  
5    halophenyl group, with substitution at the carbon of the methanone  
6    group by an adamantyl, naphthyl, phenyl, benzyl, quinolinyl,  
7    cycloalkyl, 1-amino-3-methyl-1-oxobutan-2-yl, 1-amino-3,3-dimethyl-  
8    1-oxobutan-2-yl, 1-methoxy-3-methyl-1-oxobutan-2-yl, 1-methoxy-3,3-  
9    dimethyl-1-oxobutan-2-yl or pyrrole group, and whether or not  
10   further substituted in the benzimidazole, adamantyl, naphthyl,  
11   phenyl, pyrrole, quinolinyl, or cycloalkyl rings to any extent.

12   Benzimidazole Ketones include, but are not limited to:

- 13        a.    naphthalen-1-yl(1-pentyl-1H-benzo[d]imidazol-2-  
14                      1)methanone (JWH-018 benzimidazole analog), or  
15        b.    (1-(5-fluoropentyl)-1H-benzo[d]imidazol-2-  
16                      yl)(naphthalen-1-yl)methanone (FUBIMINA); and

17        14.   Modified by Replacement: any compound defined in this  
18   subsection that is modified by replacement of a carbon with nitrogen  
19   in the indole, naphthyl, indene, benzimidazole, or carbazole ring.

20        H.   Any material, compound, mixture, extract, or preparation  
21   that contains a prohibited kratom product as provided in paragraphs  
22   3 and 4 of subsection A of Section 1-1432.4 of this title.

23        I.   Any prescription drug approved by the federal Food and Drug  
24   Administration under the provisions of Section 505 of the Federal

1 Food, Drug, and Cosmetic Act, Title 21 of the United States Code,  
2 Section 355, that is designated, rescheduled, or deleted as a  
3 controlled substance under federal law by the United States Drug  
4 Enforcement Administration shall be excluded from Schedule I and  
5 shall be prescribed, distributed, dispensed, or used in accordance  
6 with federal law upon the issuance of a notice, final rule, or  
7 interim final rule by the United States Drug Enforcement  
8 Administration designating, rescheduling, or deleting as a  
9 controlled substance such a drug product under federal law, unless  
10 and until the State Board of Pharmacy takes action pursuant to  
11 Section 2-201 of this title. If the Board of Pharmacy does not take  
12 action pursuant to Section 2-201 of this title, the drug product  
13 shall be deemed to be designated, rescheduled, or deleted as a  
14 controlled substance in accordance with federal law and in  
15 compliance with the Uniform Controlled Dangerous Substances Act.

16 SECTION 2. This act shall become effective November 1, 2025.

17 COMMITTEE REPORT BY: COMMITTEE ON PUBLIC SAFETY  
18 March 4, 2025 - DO PASS AS AMENDED BY CS

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