House Bill 231 (COMMITTEE SUBSTITUTE)

By: Representatives Broadrick of the 4th, Hawkins of the 27th, and Gravley of the 67th

:

A BILL TO BE ENTITLED AN ACT

- 1 To amend Chapter 13 of Title 16 of the Official Code of Georgia Annotated, relating to
- 2 controlled substances, so as to change certain provisions relating to Schedules I, II, IV, and
- 3 V controlled substances; to change certain provisions relating to the definition of dangerous
- 4 drug; to provide for related matters; to provide for an effective date; to repeal conflicting
- 5 laws; and for other purposes.

6 BE IT ENACTED BY THE GENERAL ASSEMBLY OF GEORGIA:

7 SECTION 1.

- 8 Chapter 13 of Title 16 of the Official Code of Georgia Annotated, relating to controlled
- 9 substances, is amended in Code Section 16-13-25, relating to Schedule I controlled
- substances, by adding two new subparagraphs to paragraph (1) to read as follows:
- 11 "(RR) 3,4-dichloro-N-[(1-dimethylamino)cyclohexylmethyl]benzamide (AH-7921);
- 12 (SS) 3,4-dichloro-N-(2-(dimethylamino)cyclohexyl)-N-methylbenzamide (U-47700);"
- SECTION 2.
- 14 Said chapter is further amended in Code Section 16-13-25, relating to Schedule I controlled
- substances, by revising subparagraphs (CC), (EE), (JJ), (KK), (LL), (MM), (NN), (RR), and
- 16 (FFF) of and by adding new subparagraphs to paragraph (3) as follows:
- 17 "(CC) 3-methylfentanyl Reserved;"
- 18 "(EE) Para-flurofentanyl Reserved;"
- 19 "(JJ) Alpha-Methylthiofentanyl Reserved;
- 20 (KK) Acetyl-Alpha-Methylfentanyl Reserved;
- 21 (LL) 3-Methylthiofentanyl Reserved;
- 22 (MM) Beta-Hydroxyfentanyl Reserved;
- 23 (NN) Thiofentanyl Reserved;"
- 24 "(RR) Beta-Hydroxy-3-Methylfentanyl Reserved;"
- 25 "(FFF) 4-Fluoromethcathinone Fluoromethcathinone;"

26	"(EEEE) 1-(1-benzofuran-6-yl)propan-2-amine (6-APB);
27	(FFFF) 1-(1-benzofuran-5-yl)-N-ethylpropan-2-amine (5-EAPB);"
28	SECTION 3.
29	Said chapter is further amended in Code Section 16-13-25, relating to Schedule I controlled
30	substances, by revising subparagraphs (B) and (C) of paragraph (4) as follows:
31	"(B) N-(1-benzyl-4-piperidyl)-N-phenylpropanamide (benzyl-fentanyl) Reserved;
32	(C) N-(1-(2-thienyl)methyl-4-piperidyl)-N-phenylpropanamide (thenylfentanyl)
33	Reserved;"
34	SECTION 4.
35	Said chapter is further amended in Code Section 16-13-25, relating to Schedule I controlled
36	substances, by substituting the "." at the end of subparagraph (V) of paragraph (12) with a
37	";" and by adding new paragraphs to read as follows:
38	"(13) The fentanyl analog structural class, including any of the following derivatives.
39	their salts, isomers, or salts of isomers, unless specifically utilized as part of the
40	manufacturing process by a commercial industry of a substance or material not intended
41	for human ingestion or consumption, as a prescription administered under medical
42	supervision, or for research at a recognized institution, whenever the existence of these
43	salts, isomers, or salts of isomers is possible within the specific chemical designation or
44	unless specifically excepted or listed in this or another schedule, structurally derived from
45	fentanyl, and whether or not further modified in any of the following ways:
46	(A) Substitution anywhere on the phenethyl group with:
47	(i) Alkyl group;
48	(ii) Hydroxyl group;
49	(iii) Halide group;
50	(B) Replacement of the phenethyl group with:
51	(i) Thienyl ethyl group, which can be further substituted with:
52	(I) Alkyl group;
53	(II) Hydroxyl group;
54	(III) Halide group;
55	(ii) Oxotetrazol ethyl group, which can be further substituted with:
56	(I) Alkyl group;
57	(II) Hydroxyl group;
58	(III) Halide group;
59	(iii) Alkyl group;
60	(iv) Thienyl methyl group, which can be further substituted with:

61 (I) Alkyl group; 62 (II) Hydroxyl group; 63 (III) Halide group; 64 (v) Benzyl group, which can be further substituted with: 65 (I) Alkyl group; 66 (II) Hydroxyl group; 67 (III) Halide group; (vi) Furanyl ethyl group, which can be further substituted with: 68 69 (I) Alkyl group; 70 (II) Hydroxyl group; 71 (III) Halide group; 72 (vii) Phenyl alkyl group, which can be further substituted with: 73 (I) Alkyl group; 74 (II) Hydroxyl group; 75 (III) Halide group; (viii) Pyridinyl ethyl group, which can be further substituted with: 76 77 (I) Alkyl group; 78 (II) Hydroxyl group; 79 (III) Halide group; 80 (ix) Diazole ethyl group, which can be further substituted with: 81 (I) Alkyl group; 82 (II) Hydroxyl group; 83 (III) Halide group; 84 (IV) Nitro group; 85 (x) Thiazole ethyl group, which can be further substituted with: 86 (I) Alkyl group; 87 (II) Hydroxyl group; 88 (III) Halide group; 89 (xi) Benzoxazolinone ethyl group, which can be further substituted with: 90 (I) Alkyl group; 91 (II) Hydroxyl group; 92 (III) Halide group; 93 (C) Substitution anywhere on the piperidine ring with: 94 (i) Alkyl group; 95 (ii) Allyl group; 96 (iii) Phenyl group; (iv) Ester group; 97

98 (v) Ether group; 99 (vi) Pyridine group, which can be further substituted with: 100 (I) Alkyl group; 101 (II) Hydroxyl group; 102 (III) Halide group; 103 (vii) Thiazole group, which can be further substituted with: 104 (I) Alkyl group; 105 (II) Hydroxyl group; 106 (III) Halide group; 107 (viii) Oxadiazole group, which can be further substituted with: 108 (I) Alkyl group; 109 (II) Hydroxyl group; 110 (III) Halide group; 111 (IV) Ether group; 112 (D) Substitution anywhere on the propanamide group with: 113 (i) Cyclic alkyl group; 114 (ii) Acyclic alkyl group: 115 (iii) Methoxy group; 116 (E) Replacement of the propanamide group with: 117 (i) Acryloyl amino group; 118 (ii) Acetamide group, which itself can be further substituted with a cyclic alkyl 119 group; 120 (iii) Methoxy acetamide group; 121 (iv) Furanyl amide group; 122 (F) Substitution anywhere on the phenyl ring with: 123 (i) Halide group; 124 (ii) Methoxy group; 125 (iii) Alkyl group; (G) Replacement of the phenyl ring with the pyrazine ring; 126 (14) The piperidinyl-sulfonamide structural class, including any of the following 127 compounds, derivatives, their salts, isomers, or salts of isomers, halogen analogues, or 128 homologues, unless specifically utilized as part of the manufacturing process by a 129 130 commercial industry of a substance or material not intended for human ingestion or consumption, as a prescription administered under medical supervision, or for research 131 at a recognized institution, whenever the existence of these salts, isomers, or salts of 132 isomers, halogen analogues, or homologues is possible within the specific chemical 133 134 designation or unless specifically excepted or listed in this or another schedule,

135 structurally derived from piperidinyl-sulfonamide, and whether or not further modified 136 in any of the following ways: 137 (A) By substitution at the 1-position of the piperidinyl ring with any of the following: (i) Alkyl group; 138 139 (ii) Phenyl alkyl group; 140 (iii) Amino substituted phenyl alkyl group; 141 (iv) Nitro substituted phenyl alkyl group; 142 (v) Cycloalkyl group; 143 (vi) Alkenyl substituent group; (B) By substitution at the 3-position or 4-position of the piperidinyl ring with any of 144 145 the following: 146 (i) Halide group; 147 (ii) Alkyl group; 148 (iii) Alkoxy substituent; 149 (C) By substitution on the sulfonamide with any of the following: 150 (i) Pyridyl group; 151 (ii) Alkyl group; 152 (iii) Phenyl group; 153 (iv) Phenyl alkyl group; (v) Alkoxy substituted phenyl group; 154 155 (vi) Halogen substituted phenyl group; 156 (vii) Nitro substituted phenyl group; 157 (viii) Amino substituted phenyl group; 158 (ix) Alkanoylamino substituted phenyl group; 159 (x) Amido substituted phenyl group; 160 (15) The 1-cyclohexyl-4-(1,2-diphenylethy)-piperazine (MT-45) structural class, including any of the following derivatives, their salts, isomers, or salts of isomers, unless 161 162 specifically utilized as part of the manufacturing process by a commercial industry of a 163 substance or material not intended for human ingestion or consumption, as a prescription administered under medical supervision, or for research at a recognized institution, 164 whenever the existence of these salts, isomers, or salts of isomers is possible within the 165 specific chemical designation or unless specifically excepted or listed in this or another 166 schedule, structurally derived from 1-cyclohexyl-4-(1,2-diphenylethy)-piperazine 167 (MT-45), and whether or not further modified in any of the following ways: 168 (A) Replacement of the cyclohexyl group with any of the following: 169 170 (i) Cycloheptyl group;

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(ii) Cyclooctyl group;

172	(B) Substitution on the diphenyl groups with any of the following:
173	(i) Hydroxyl group;
174	(ii) Halide;
175	(iii) Alkoxy group;
176	(iv) Alkyl group;
177	(v) Ester group;
178	(vi) Phenyl ether group."
179	SECTION 5.
180	Said chapter is further amended in Code Section 16-13-26, relating to Schedule II controlled
181	substances, by adding new subparagraphs to paragraph (2) to read as follows:
182	"(C.5) Carfentanil;"
183	"(V.2) Thiafentanil;"
184	SECTION 6.
185	Said chapter is further amended in Code Section 16-13-26, relating to Schedule II controlled
186	substances, by revising subparagraph (E) of paragraph (3) as follows:
187	"(E) Carfentanil Reserved;"
188	SECTION 7.
189	Said chapter is further amended in Code Section 16-13-28, relating to Schedule IV controlled
190	substances, by revising paragraph (1) of subsection (b) as follows:
191	"(1) By substitution at the 2-position with a ketone <u>or a thione;</u> "
192	SECTION 8.
193	Said chapter is further amended in Code Section 16-13-29, relating to Schedule V controlled
194	substances, by deleting "or" at the end of paragraph (5), by substituting the "." at the end of
195	paragraph (6) with a ";", and by adding a new paragraph to read as follows:
196	"(7) Brivaracetam."
197	SECTION 9.
198	Said chapter is further amended in Code Section 16-13-71, relating to the definition of a
199	dangerous drug, by adding new paragraphs to subsection (b) to read as follows:
200	"(13.531) Adalimumab-atto;"
201	"(68.13) Atezolizumab;"
202	"(97.4) Bezlotoxumab;"
203	"(217.4) Crisaborole;"

204 "(244.2) Defibrotide;" "(331.053) Elbasvir;" 205 206 "(355.6) Etanercept-szzs;" 207 "(355.8) Eteplirsen;" "(430.7) Grazoprevir;" 208 209 "(472.51) Infliximab-dyyb;" "(506.97) Ixekizumab;" 210 "(520.2) Lifitegrast;" 211 212 "(528.1) Lixisenatide;" "(658.7) Nusinersen;" 213 214 "(661.03) Obeticholic acid;" "(661.05) Obiltoxaximab;" 215 "(661.96) Olaratumab;" 216 217 "(663.36) Omalizumab;" "(663.6) OnabotulinumtoxinA;" 218 "(769.37) Prasterone;" 219 220 "(835.5) Reslizumab;" 221 "(848.2) Rucaparib;" 222 "(1027.53) Velpatasvir;" "(1027.57) Venetoclax;" 223 224 **SECTION 10.** 225 Said chapter is further amended in Code Section 16-13-71, relating to the definition of a 226 dangerous drug, by revising paragraphs (13.55), (198.05), and (673) of subsection (b) as 227 follows: "(13.55) Adapalene — See exceptions;" 228 229 "(198.05) Clobazam;" "(673) Reserved Oxymetazoline;" 230 231 **SECTION 11.** Said chapter is further amended in Code Section 16-13-71, relating to the definition of a 232 233 dangerous drug, by adding a new paragraph to subsection (c) to read as follows:

"(0.5) Adapalene — when used with a strength up to 0.1 percent in a topical skin

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product;"

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236	SECTION 12.
237	This Act shall become effective upon its approval by the Governor or upon its becoming law

without such approval.

239 **SECTION 13.**

240 All laws and parts of laws in conflict with this Act are repealed.