By: Senator(s) Fillingane

To: Drug Policy; Judiciary, Division B

## SENATE BILL NO. 2119

AN ACT TO AUTHORIZE PHARMACIES TO SELL AND PERSONS TO PURCHASE, WITHOUT A PRESCRIPTION, PRODUCTS THAT CONTAIN CERTAIN 3 QUANTITIES OF PSEUDOEPHEDRINE OR EPHEDRINE; TO REQUIRE PHARMACIES SELLING PRODUCTS AUTHORIZED UNDER THIS ACT TO USE THE NPLEX SYSTEM 5 BEFORE SELLING THOSE PRODUCTS; TO REQUIRE PHARMACIES TO MAINTAIN AN ELECTRONIC LOG OF REQUIRED INFORMATION FOR EACH TRANSACTION; TO 7 REQUIRE THE PURCHASER OF THE PACKAGE TO BE AT LEAST EIGHTEEN YEARS OF AGE, AS SHOWN BY VALID IDENTIFICATION, AND TO SIGN A RECORD OF 8 9 EACH TRANSACTION; TO PROVIDE CRIMINAL PENALTIES FOR VIOLATIONS OF 10 THIS ACT; TO AMEND SECTION 41-29-117, MISSISSIPPI CODE OF 1972, TO 11 CONFORM; AND FOR RELATED PURPOSES. BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MISSISSIPPI:

- 12
- under Section 73-21-105 to sell or distribute to a person, without 14

SECTION 1. (1) (a) It is lawful for a pharmacy registered

- 15 a prescription, products containing not more than three and six
- tenths (3.6) grams per day and not more than seven and two tenths 16
- (7.2) grams per thirty-day period of pseudoephedrine or ephedrine, 17
- 18 and it is lawful for a person to purchase products containing
- 19 those ingredients from a registered pharmacy without a
- 20 prescription.

13

- 21 (b) All products authorized under this subsection (1)
- 22 must be stored by a pharmacy by placing the products behind a

- 23 counter in an area within the pharmacy where the public is not
- 24 permitted.
- 25 (c) Any products authorized under this subsection (1)
- 26 sold by a pharmacy must be sold by an individual licensed as a
- 27 pharmacist or by an employee of the pharmacy under the direct
- 28 supervision and control of a licensed pharmacist.
- 29 (d) No pharmacy may sell or distribute, and no person
- 30 may purchase, more products than allowed under this section unless
- 31 by valid prescription. It is not a defense in a prosecution under
- 32 this section that no money was exchanged during a transaction that
- 33 would otherwise be unlawful under this section.
- 34 (2) A pharmacy selling products in a manner authorized under
- 35 subsection (1) of this section must:
- 36 (a) Use the National Precursor Log Exchange (NPLEX)
- 37 system administered by the National Association of Drug Diversion
- 38 Investigators, provided that the system is available to pharmacies
- 39 or retailers in the state without a charge for accessing the NPLEx
- 40 system, before completing the over-the-counter sale of each
- 41 product authorized under subsection (1) of this section. Before
- 42 completing a sale of an over-the-counter material, compound,
- 43 mixture, or preparation containing any detectable quantity of
- 44 pseudoephedrine or ephedrine, its salts or optical isomers, or
- 45 salts of optical isomers a pharmacy or retailer shall
- 46 electronically submit the information required under subsection
- 47 (b) of this subsection (2) to the NPLEx system. The pharmacy or

- 48 retailer shall not complete the sale if the NPLEx system generates
- 49 a stop-sale alert. The system shall contain an override function
- 50 that may be used by an agent of a retail establishment who is
- 51 dispensing the drug product, and who has a reasonable fear of
- 52 imminent bodily harm if the transaction is not completed. The
- 53 system shall create a record of each use of the override
- 54 mechanism.
- (b) Maintain an electronic log of required information
- 56 for each transaction, and require the purchaser of the package to
- 57 be at least eighteen (18) years of age and provide a valid,
- 58 unsuspended driver's license or nondriver identification card
- 59 issued by this state or another state, a United States Uniformed
- 60 Services Privilege and Identification Card, or a United States or
- 61 foreign passport, and to sign a written or electronic log
- 62 attesting to the validity of the information provided for each
- 63 transaction. The record of each transaction must include the
- 64 information from the identification card as well as the type of
- 65 and government entity issuing the identification card used, the
- 66 name, date of birth, and current address of the purchaser, the
- 67 date and time of the sale, the name of the compound, mixture, or
- 68 preparation being sold, and the total amount, in grams or
- 69 milligrams, of pseudoephedrine or ephedrine being sold.
- 70 (c) Maintain a written log or an alternative electronic
- 71 recordkeeping mechanism if a pharmacy or retailer experiences
- 72 mechanical or electronic failure of the required electronic

- 73 tracking system until such time as the pharmacy or retailer is
- 74 able to comply with the electronic sales-tracking requirement. No
- 75 person shall purchase, receive or otherwise acquire more than
- 76 three and six-tenths (3.6) grams per day or seven and two-tenths
- 77 (7.2) grams of pseudoephedrine or ephedrine within any thirty-day
- 78 period.
- 79 (3) The National Association of Drug Diversion
- 80 Investigators shall provide real-time access to the NPLEx
- 81 information through the NPLEx online portal to law enforcement in
- 82 the state.
- 83 (4) (a) Pseudoephedrine and ephedrine products dispensed
- 84 pursuant to a legitimate prescription are exempt from this
- 85 section.
- 86 (b) The amounts of pseudoephedrine and ephedrine
- 87 products dispensed to a person pursuant to a legitimate
- 88 prescription shall not be considered under subsection (1)(a) of
- 89 this section.
- 90 (5) A violation of this section is a misdemeanor and is
- 91 punishable as follows:

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- 92 (a) For a first offense, by a fine not to exceed One
- 93 Thousand Dollars (\$1,000.00).
- 94 (b) For a second or subsequent offense, by a fine not
- 95 to exceed Ten Thousand Dollars (\$10,000.00).
- 96 (6) A pharmacist who is the general owner or operator of an
- 97 establishment where pseudoephedrine and ephedrine products are

98	available for sale shall not be penalized under this section for
99	the conduct of an employee if the retailer documents that an
100	employee training program approved by the Mississippi Board of
101	Pharmacy was conducted by the pharmacist. The Mississippi Board
102	of Pharmacy shall develop or approve all training programs for
103	pharmacy employees.
104	(7) A person who resides in a state that requires a
105	prescription for the purchase of pseudoephedrine or ephedrine, or
106	who presents identification from a state that requires a
107	prescription for the purchase of pseudoephedrine or ephedrine, may
108	purchase those products only upon presentation of a valid
109	prescription for the pseudoephedrine or ephedrine.
110	SECTION 2. Section 41-29-117, Mississippi Code of 1972, is
111	amended as follows:
112	41-29-117. (A) The controlled substances listed in this
113	section are included in Schedule III.
114	SCHEDULE III
115	(a) Stimulants. Any material, compound, mixture, or
116	preparation which contains any quantity of the following
117	substances or their salts, isomers, or salts of isomers, of the
118	following substances:
119	(1) Benzphetamine;

(2)

(3)

(4)

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121

122

Chlorphentermine;

Phendimetrazine.

Clortermine;

123	(b) Depressants. Unless listed in another schedule,
124	any material, compound, mixture, or preparation which contains any
125	quantity of the following substances:
126	(1) Any substance which contains any quantity of a
127	derivative of barbituric acid, or any salt of a derivative of
128	barbituric acid, except those substances which are specifically
129	listed in other schedules;
130	(2) Unless specifically excepted or unless listed
131	in another schedule, any compound, mixture or preparation
132	containing any of the following substances or any salt of the
133	substances specifically included in this subsection (2) and one or
134	more other active medicinal ingredients which are not listed in
135	any other schedule:
136	(i) Amobarbital;
137	(ii) Secobarbital;
138	(iii) Pentobarbital;
139	(3) Any suppository dosage form containing any of
140	the following substances or any salt of any of the substances
141	specifically included in this subsection (3) approved by the Food
142	and Drug Administration for marketing only as a suppository:
143	(i) Amobarbital;
144	(ii) Secobarbital;
145	(iii) Pentobarbital;
146	(4) Chlorhexadol;
147	(5) Embutramide:

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                         Any drug product containing
149
     gamma-hydroxybutyric acid, including its salts, isomers and salts
     of isomers, for which an application is approved under Section 505
150
151
     of the Federal Food, Drug and Cosmetic Act;
152
                         Ketamine; its salts, isomers, and salts of
153
     isomers; other names include
154
     (+)-2-(2-chlorophenyl)-2-(methylamino)cyclohexanone;
155
                     (8)
                         Lysergic acid;
156
                         Lysergic acid amide;
                     (9)
157
                     (10) Methyprylon;
158
                     (11)
                          Perampanel; its salts, isomers, and salts of
159
     isomers;
160
                     (12)
                          Sulfondiethylmethane;
161
                          Sulfonethylmethane;
                     (13)
162
                     (14)
                          Sulfonmethane;
163
                     (15)
                          Tiletamine and zolazepam or any salt thereof;
164
     other names for the tiletamine and zolazepam combination product
     include: telazol; other names for tiletamine include:
165
166
     2-(ethylamino)-2-(2-thienyl)-cyclohexanone; other names for
167
     zolazepam include: 4-(2-fluorophenyl)-6,8-dihydro 1,3,
     8-trimethylpyrazolo-[3,4-e](1,4)-diazepin-7(1H)-one, flupyrazapon.
168
169
                (C)
                    Nalorphine.
170
                    Any material, compound, mixture or preparation
                (d)
171
     which contains any quantity of ephedrine or pseudoephedrine,
172
     except for any product that contains any quantity of
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173	pseudoephedr	ine or	ephedrin	ne that	is	sold	d subject	to	the	quantity
174	restrictions	autho	rized in	Section	n 1	of t	this act.			

- 175 (e) **Narcotic drugs.** Any material, compound, mixture,
  176 or preparation containing limited quantities of any of the
  177 following narcotic drugs, or any salts thereof:
- 178 (1) Not more than one and eight-tenths (1.8) grams
  179 of codeine, or any of its salts, per one hundred (100) milliliters
  180 or not more than ninety (90) milligrams per dosage unit, with an
  181 equal or greater quantity of an isoquinoline alkaloid of opium;
- 182 (2) Not more than one and eight-tenths (1.8) grams
  183 of codeine, or any of its salts, per one hundred (100) milliliters
  184 or not more than ninety (90) milligrams per dosage unit, with one
  185 or more active, nonnarcotic ingredients in recognized therapeutic
  186 amounts;
- (3) Not more than one and eight-tenths (1.8) grams
  of dihydrocodeine, or any of its salts, per one hundred (100)
  milliliters or not more than ninety (90) milligrams per dosage
  unit, with one or more active, nonnarcotic ingredients in
  recognized therapeutic amounts;
- (4) Not more than three hundred (300) milligrams
  of ethylmorphine, or any of its salts, per one hundred (100)
  milliliters or not more than fifteen (15) milligrams per dosage
  unit, with one or more active, nonnarcotic ingredients in
  recognized therapeutic amounts;

197	(5) Not more than five hundred (500) milligrams of
198	opium per one hundred (100) milliliters or per one hundred (100)
199	grams, or not more than twenty-five (25) milligrams per dosage
200	unit, with one or more active, nonnarcotic ingredients in
201	recognized therapeutic amounts;
202	(6) Not more than fifty (50) milligrams of
203	morphine, or any of its salts, per one hundred (100) milliliters
204	or per one hundred (100) grams with one or more active,
205	nonnarcotic ingredients in recognized therapeutic amounts.
206	(f) Anabolic steroids. Unless specifically exempted or
207	listed in another schedule, any material, compound, mixture or
208	preparation containing any quantity of any of the following
209	anabolic steroids (any drug or hormonal substance chemically and
210	pharmacologically related to testosterone other than estrogens,
211	progestins, corticosteroids and dehydroepiandrosterone):
212	(1) 3beta,17-dihydroxy-5a-androstane;
213	(2) 3alpha, 17beta-dihydroxy-5a-androstane;
214	(3) 5alpha-androstan-3,17-dione;
215	(4) 1-androstenediol
216	(3beta,17beta-dihydroxy-5alpha-androst-1-ene);
217	(5) 1-androstenediol
218	(3alpha,17beta-dihydroxy-5alpha-androst-1-ene);
219	(6) 4-androstenediol
220	(3beta,17beta-dihydroxy-androst-4-ene);

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221
                     (7)
                          5-androstenediol
222
     (3beta, 17beta-dihydroxy-androst-5-ene);
223
                     (8)
                          1-androstenedione ([5alpha]-androst-1-en-3,
224
     17-dione);
225
                     (9)
                         4-androstenedione (androst-4-en-3,17-dione);
226
                     (10) 5-androstenedione (androst-5-en-3,17-dione);
227
                     (11) Bolasterone
228
     (7alpha, 17alpha-dimethyl-17beta-hydroxyandrost-4-en-3-one);
229
                     (12)
                          Boldenone
230
     (17beta-hydroxyandrost-1, 4, -diene-3-one);
231
                     (13)
                           Boldione (androsta-1, 4-diene-3, 17-dione);
232
                     (14) Calusterone
233
     (7beta, 17alpha-dimethyl-17beta-hydroxyandrost-4-en-3-one);
234
                     (15) Clostebol
     (4-chloro-17beta-hydroxyandrost-4-en-3-one);
235
236
                     (16)
                           Dehydrochloromethyltestosterone
237
     (4-chloro-17beta-hydroxy-17alpha-methylandrost-1,4-dien-3-one);
238
                     (17) Desoxymethyltestosterone
239
     (17alpha-methyl-5alpha-androst-2-en-17beta-ol, also known as
240
     madol);
241
                     (18)
                           Delta1-dihydrotestosterone (also known as
242
     1-testosterone) (17beta-hydroxy-5alpha-androst-1-en-3-one);
243
                     (19)
                           4-dihydrotestosterone
244
     (17beta-hydroxy-androstan-3-one);
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245
                     (20)
                          Drostanolone
246
     (17beta-hydroxy-2alpha-methyl-5alpha-androstan-3-one);
247
                     (21) Ethylestrenol
248
     (17alpha-ethyl-17beta-hydroxyestr-4-ene);
249
                     (22)
                         Fluoxymesterone
250
     (9-fluoro-17alpha-methyl-11beta,
251
     17beta-dihydroxyandrost-4-en-3-one);
252
                     (23)
                           Formebolone
253
     (2-formyl-17alpha-methyl-11alpha, 17beta-dihydroxyandrost-1,
254
     4-dien-3-one);
255
                          Furazabol
                     (24)
256
     (17alpha-methyl-17beta-hydroxyandrostano[2,3-c]-furazan);
257
                     (25) 13beta-ethyl-17alpha-hydroxygon-4-en-3-one;
258
                           4-hydroxytestosterone
                     (26)
     (4,17beta-dihydroxyandrost-4-en-3-one);
259
260
                     (27)
                           4-hydroxy-19-nortestosterone
261
     (4,17beta-dihydroxy-estr-4-en-3-one);
262
                     (28) Mestanolone
263
     (17alpha-methyl-17beta-hydroxy-5-androstan-3-one);
264
                     (29) Mesterolone
265
     (1alpha-methyl-17beta-hydroxy-[5alpha]-androstan-3-one);
266
                     (30) Methandienone
267
     (17alpha-methyl-17beta-hydroxyandrost-1,4-dien-3-one);
268
                     (31)
                          Methandriol (17alpha-methyl-3beta,
269
     17beta-dihydroxyandrost-5-ene);
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270
                     (32)
                          Methasterone (2[alpha],
271
     17[alpha]-dimethyl-5[alpha]-androstan-17[beta]-ol-3-one;
272
                     (33)
                         Methenolone
273
     (1-methyl-17beta-hydroxy-5alpha-androst-1-en-3-one);
274
                     (34)
                           17alpha-methyl-3beta,
275
     17beta-dihydroxy-5a-androstane;
276
                           17alpha-methyl-3alpha,
                     (35)
277
     17beta-dihydroxy-5a-androstane;
278
                     (36)
                           17alpha-methyl-3beta,
279
     17beta-dihydroxyandrost-4-ene;
280
                     (37)
                           17alpha-methyl-4-hydroxynandrolone
281
     (17alpha-methyl-4-hydroxy-17beta-hydroxyestr-4-en-3-one);
282
                     (38) Methyldienolone
283
     (17alpha-methyl-17beta-hydroxyestra-4,9(10)-dien-3-one);
284
                     (39)
                           Methyltrienolone
     (17alpha-methyl-17beta-hydroxyestra-4,9-11-trien-3-one);
285
286
                     (40)
                          Methyltestosterone
287
     (17alpha-methyl-17beta-hydroxyandrost-4-en-3-one);
288
                     (41)
                          Mibolerone
289
     (7alpha, 17alpha-dimethyl-17beta-hydroxyestr-4-en-3-one);
290
                     (42) 17alpha-methyl-Deltal-dihydrotestosterone (17b
291
     beta-hydroxy-17alpha-methyl-5alpha-androst-1-en-3-one) (also known
     as 17-alpha-methyl-1-testosterone);
292
293
                     (43) Nandrolone (17beta-hydroxyestr-4-en-3-one);
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294
                     (44)
                          19-nor-4-androstenediol
295
     (3beta, 17beta-dihydroxyestr-4-ene);
296
                     (45) 19-nor-4-androstenediol
297
     (3a, 17beta-dihydroxyestr-4-ene);
298
                     (46) 19-nor-5-androstenediol
299
     (3beta, 17beta-dihydroxyestr-5-ene);
300
                     (47) 19-nor-5-androstenediol
301
     (3alpha, 17beta-dihydroxyestr-5-ene);
302
                     (48)
                         19-nor-4,9(10)-androstadienedione
303
     (estra-4, 9(10) -diene3, 17-dione,
304
     19-norandrosta-4,9(10)-diene-3,17-dione);
305
                     (49) 19-nor-4-androstenedione
306
     (estr-4-en-3,17-dione);
307
                     (50) 19-nor-5-androstenedione
308
     (estr-5-en-3,17-dione);
309
                     (51) Norbolethone
310
     (13beta, 17alpha-diethyl-17beta-hydroxygon-4-en-3-one);
311
                     (52) Norclostebol
312
     (4-chloro-17beta-hydroxyestr-4-en-3-one);
313
                     (53) Norethandrolone
314
     (17alpha-ethyl-17beta-hydroxyestr-4-en-3-one);
315
                     (54) Normethandrolone
     (17alpha-methyl-17beta-hydroxyestr-4-en-3-one);
316
317
                     (55) Oxandrolone
     (17alpha-methyl-17beta-hydroxy-2-oxa-[5alpha]-androstan-3-one);
318
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319
                     (56)
                           Oxymesterone
320
     (17alpha-methyl-4,17beta-dihydroxyandrost-4-en-3-one);
321
                     (57) Oxymetholone
322
     (17alpha-methyl-2-hydroxymethylene-17beta-hydroxy-[5alpha]-
323
     androstan-3-one);
324
                     (58) Prostanozol
325
     (17[beta]-hydroxy-5[alpha]-androstano[3,2-c]pyrazole)
326
                     (59)
                          Stanozolol
327
     (17alpha-methyl-17beta-hydroxy-[5alpha]-androst-2-eno[3,2-c]-
     pyrazole);
328
329
                           Stenbolone
                     (60)
330
     (17beta-hydroxy-2-methyl-[5alpha]-androst-1-en-3-one);
331
                     (61) Testolactone
332
     (13-hydroxy-3-oxo-13,17-secoandrosta-1,4-dien-17-oic acid
333
     lactone);
334
                     (62)
                           Testosterone
335
     (17beta-hydroxyandrost-4-en-3-one);
336
                          Tetrahydrogestrinone
                     (63)
337
     (13beta, 17alpha-diethyl-17beta-hydroxygon-4, 9, 11-trien-3-one);
338
                     (64) Trenbolone
339
     (17beta-hydroxyestr-4,9,11-trien-3-one);
340
                     (65) Any salt, ester, or ether of a drug or
341
     substance described in this paragraph. Except such term does not
342
     include an anabolic steroid that is expressly intended for
     administration through implants to cattle or other nonhuman
343
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344	species and that has been approved by the Secretary of Health and
345	Human Services for such administration. If any person prescribes,
346	dispenses, or distributes such steroid for human use, the person
347	shall be considered to have prescribed, dispensed or distributed
348	an anabolic steroid within the meaning of this paragraph.

- 349 (g) Any material, compound, mixture or preparation 350 which contains any quantity of buprenorphine or its salts.
- 351 (h) Dronabinol (synthetic) in sesame oil and
  352 encapsulated in a soft gelatin capsule in a United States Food and
  353 Drug Administration approved drug product.
- (B) Any material, compound, mixture or preparation which contains any quantity of a Schedule III controlled substance other than butalbital, and is listed as an exempt substance in 21 CFR, Section 1308.22, 1308.24, 1308.26, 1308.32 or 1308.34, shall be exempted from the provisions of the Uniform Controlled Substances Law.
- 360 **SECTION 3.** This act shall take effect and be in force from and after January 1, 2022.

