2023 SESSION

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1	HOUSE BILL NO. 2294
2	AMENDMENT IN THE NATURE OF A SUBSTITUTE
3	(Proposed by the House Committee for Courts of Justice
4	on February 3, 2023)
5	(Patron Prior to Substitute—Delegate Kilgore)
6 7	A BILL to amend and reenact §§ 3.2-4112, 3.2-4113, 3.2-4114, 3.2-4114.2, 3.2-4115, 3.2-4116, 3.2-4118, 3.2-4119, 3.2-5145.1, 3.2-5145.2:1, 3.2-5145.4, 3.2-5145.5, 4.1-600, 18.2-247,
8	5.2-4116, 5.2-4119, 5.2-5145.1, 5.2-5145.2, 5.2-5145.4, 5.2-5145.5, 4.1-000, 18.2-247, 18.2-251.1:3, 18.2-371.2, 54.1-3401, 54.1-3408.3, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 18.2-251.1:3, 18.2-371.2, 54.1-3401, 54.1-3408.3, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 18.2-251.1:3, 18.2-371.2, 54.1-3401, 54.1-3408.3, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 18.2-251.1:3, 18.2-371.2, 54.1-3401, 54.1-3408.3, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 18.2-251.1:3, 18.2-371.2, 54.1-3401, 54.1-3408.3, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 18.2-251.1:3, 18.2-371.2, 54.1-3423, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 54.1-3442.7, 54.1-3443, 54.1-3442.7, 54.1-3443, 54.1-3442.7, 54.1-3442.7, 54.1-3443, 54.1-3442.7, 54.1-344444444444444444444444444444444444
9	54.1-3446, 59.1-200, 59.1-203, and 59.1-206 of the Code of Virginia and to amend the Code of
10	Virginia by adding a section numbered 3.2-5145.4:1, relating to tetrahydrocannabinol; industrial
11	hemp; regulated hemp products.
12	Be it enacted by the General Assembly of Virginia:
13	1. That §§ 3.2-4112, 3.2-4113, 3.2-4114, 3.2-4114.2, 3.2-4115, 3.2-4116, 3.2-4118, 3.2-4119,
14	3.2-5145.1, 3.2-5145.2:1, 3.2-5145.4, 3.2-5145.5, 4.1-600, 18.2-247, 18.2-251.1:3, 18.2-371.2, 54.1-3401,
15	54.1-3408.3, 54.1-3423, 54.1-3442.6, 54.1-3442.7, 54.1-3443, 54.1-3446, 59.1-200, 59.1-203, and
16	59.1-206 of the Code of Virginia are amended and reenacted and that the Code of Virginia is
17 18	amended by adding a section numbered 3.2-5145.4:1 as follows: § 3.2-4112. Definitions.
19	As used in this chapter, unless the context requires a different meaning:
20	"Cannabis sativa product" means a product made from any part of the plant Cannabis sativa with a
21	concentration of tetrahydrocannabinol that is greater than that allowed by federal law.
22	"Deal" means to temporarily possess industrial hemp grown in compliance with state or federal law
23	that (i) has not been processed and (ii) was not grown and will not be processed by the person
24	temporarily possessing it.
25 26	"Dealer" means any person who is registered pursuant to subsection A of § 3.2-4115 to deal in industrial hamp. "Dealer" does not include a rate a stablishment that calls or offers for calls a hamp
26 27	industrial hemp. "Dealer" does not include a retail establishment that sells or offers for sale a hemp product.
28	"Dealership" means the location at which a dealer stores or intends to store the industrial hemp in
29	which he deals.
30	"Federally licensed hemp producer" means a person who holds a hemp producer license issued by
31	the U.S. Department of Agriculture pursuant to 7 C.F.R. Part 990.
32	"Grow" means to plant, cultivate, or harvest a plant or crop.
33 34	"Grower" means any person registered pursuant to subsection A of § 3.2-4115 to grow industrial hemp.
35	"Handle" means to temporarily possess industrial hemp grown in compliance with state or federal
36	law that (i) has not been processed and (ii) was not grown by and will not be processed by the person
37	temporarily possessing it.
38	"Handler" means any person who is registered pursuant to subsection A of § 3.2-4115 to handle
39	industrial hemp. "Handler" does not include a retail establishment that sells or offers for sale a hemp
40 41	<i>"Handler's storage site" means the location at which a handler stores or intends to store the</i>
42	industrial hemp he handles.
43	"Hemp product" means a product, including any raw materials from industrial hemp that are used for
44	or added to a food or beverage product, that (i) contains industrial hemp and has completed all stages of
45	processing needed for the product and (ii) when offered for retail sale (a) contains a total
46	tetrahydrocannabinol concentration of no greater than 0.3 percent and (b) contains no more than two
47 48	milligrams of total tetrahydrocannabinol per package.
40 49	"Industrial hemp" means any part of the plant Cannabis sativa, including seeds thereof, whether growing or not, with a concentration of tetrahydrocannabinol that is no greater than that allowed by
50	federal law. "Industrial hemp" includes an industrial hemp extract that has not completed all stages of
51	processing needed to convert the extract into a hemp product.
52	"Process" means to convert industrial hemp into a hemp product.
53	"Processor" means a person registered pursuant to subsection A of § 3.2-4115 to process industrial
54	hemp.
55 56	"Process site" means the location at which a processor processes or intends to process industrial hemp.
50 57	"Production field" means the land or area on which a grower or a federally licensed hemp producer
58	is growing or intends to grow industrial hemp.
59	"Tetrahydrocannabinol" means any naturally occurring or synthetic tetrahydrocannabinol, including

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60 its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of 61 isomers is possible within the specific chemical designation and any preparation, mixture, or substance

containing, or mixed or infused with, any detectable amount of tetrahydrocannabinol. 62

63 "Tetrahydrocannabinol" includes delta-6a(10a), delta-7, delta-8, delta-9, and delta-10

64 tetrahydrocannabinol. For the purposes of this definition, "isomer" means the optical, position, and 65 geometric isomers.

66 "Total tetrahydrocannabinol" means the sum, after the application of any necessary conversion 67 factor, of the percentage by weight of tetrahydrocannabinol and the percentage by weight of tetrahydrocannabinolic acid. 68

§ 3.2-4113. Production of industrial hemp lawful. 69

A. It is lawful for a grower, his agent, or a federally licensed hemp producer to grow, a dealer 70 handler or his agent to deal in handle, or a processor or his agent to process industrial hemp in the 71 Commonwealth for any lawful purpose. No federally licensed hemp producer or grower or his agent 72 shall be prosecuted under Chapter 11 (§ 4.1-1100 et seq.) of Title 4.1 or § 18.2-247, 18.2-248, 73 74 18.2-248.01, 18.2-248.1, or 18.2-250 for the possession or growing of industrial hemp or any Cannabis 75 sativa with a tetrahydrocannabinol concentration that does not exceed the total delta 9 76 tetrahydrocannabinol concentration percentage established in federal regulations applicable to negligent violations located at 7 C.F.R. § 990.6(b)(3). No dealer handler or his agent or processor or his agent 77 78 shall be prosecuted under Chapter 11 (§ 4.1-1100 et seq.) of Title 4.1 or § 18.2-247, 18.2-248, 79 18.2-248.01, 18.2-248.1, or 18.2-250 or issued a summons or judgment for the possession, dealing 80 handling, or processing of industrial hemp. In any complaint, information, or indictment, and in any action or proceeding brought for the enforcement of any provision of Article 1 (§ 18.2-247 et seq.) of Chapter 7 of Title 18.2 or the Drug Control Act (§ 54.1-3400 et seq.), it shall not be necessary to negate 81 82 83 any exception, excuse, proviso, or exemption contained in this chapter or the Drug Control Act, and the 84 burden of proof of any such exception, excuse, proviso, or exemption shall be on the defendant.

85 B. Nothing in this chapter shall be construed to authorize any person to violate any federal law or 86 regulation.

87 C. No person shall be prosecuted under Chapter 11 (§ 4.1-1100 et seq.) of Title 4.1 or § 18.2-247, 88 18.2-248, 18.2-248.01, 18.2-248.1, or 18.2-250 for the involuntary growth of industrial hemp through the 89 inadvertent natural spread of seeds or pollen as a result of proximity to a production field, dealership 90 handler's storage site, or process site. 91

§ 3.2-4114. Regulations.

92 A. The Board may adopt regulations pursuant to this chapter as necessary to register persons to 93 grow, deal in handle, or process industrial hemp or implement the provisions of this chapter.

94 B. Upon publication by the U.S. Department of Agriculture in the Federal Register of any final rule 95 regarding industrial hemp that materially expands opportunities for growing, producing, or dealing in handling industrial hemp in the Commonwealth, the Board shall immediately adopt amendments 96 97 conforming Department regulations to such federal final rule. Such adoption of regulations by the Board 98 shall be exempt from the provisions of the Administrative Process Act (§ 2.2-4000 et seq.). 99

§ 3.2-4114.2. Authority of Commissioner; notice to law enforcement; report.

100 A. The Commissioner may charge a nonrefundable fee not to exceed \$250 for any application for 101 registration or renewal of registration allowed under this chapter. The Commissioner may charge a 102 nonrefundable fee for the tetrahydrocannabinol testing allowed under this chapter. All fees collected by 103 the Commissioner shall be deposited in the state treasury.

B. The Commissioner shall adopt regulations establishing a fee structure for a registration issued 104 pursuant to § 3.2-4115. With the exception of § 2.2-4031, no provision of the Administrative Process 105 106 Act (§ 2.2-4000 et seq.) or public participation guideline adopted pursuant thereto shall apply to the adoption of any regulation pursuant to this subsection. However, prior to adopting any regulation 107 108 pursuant to this subsection, the Commissioner shall review the recommendation of an advisory panel 109 that shall consider the economic impact of any proposed fee amount on the Commonwealth's industrial 110 hemp industry. The advisory panel shall, at a minimum, include (i) an agribusiness representative or 111 organization, (ii) a farming representative or organization, and (iii) a hemp industry representative or 112 organization. Prior to adopting any regulation pursuant to this subsection, the Commissioner shall publish a notice of opportunity to comment in the Virginia Register of Regulations and post the action 113 114 on the Virginia Regulatory Town Hall. Such notice shall contain (a) a summary of the proposed regulation; (b) the text of the proposed regulation; and (c) the name, address, and telephone number of 115 116 the agency contact person responsible for receiving public comments. Such notice shall be made at least 60 days in advance of the last date prescribed in such notice of submittals of public comment. The 117 legislative review provisions of subsections A and B of § 2.2-4014 shall apply to the promulgation or 118 final adoption process of regulations pursuant to this subsection. The Commissioner shall consider and 119 120 keep on file all public comments received for any regulation adopted pursuant to this subsection.

121 C. The Commissioner may establish an application period for a registration or renewal of registration

122 allowed under this chapter.

123 D. The Commissioner shall notify the Superintendent of State Police of each registration issued by 124 the Commissioner under this chapter and each license submitted to the Commissioner by a federally 125 licensed hemp producer.

126 E. The Commissioner shall forward a copy or appropriate electronic record of each registration 127 issued by the Commissioner under this chapter and each license submitted to the Commissioner by a 128 federally licensed hemp producer to the chief law-enforcement officer of the county or city where 129 industrial hemp will be grown, dealt handled, or processed.

130 F. The Commissioner may monitor the industrial hemp grown, dealt handled, or processed by a 131 person registered pursuant to subsection A of § 3.2-4115 and provide for random sampling and testing 132 of the industrial hemp in accordance with any criteria established by the Commissioner and at the cost 133 of the grower, dealer handler, or processor, for compliance with tetrahydrocannabinol limits and for other appropriate purposes established pursuant to § 3.2-4114. In addition to any routine inspection and 134 sampling, the Commissioner may inspect and sample the industrial hemp at any production field, 135 136 dealership handler's storage site, or process site during normal business hours without advance notice if 137 he has reason to believe a violation of this chapter is occurring or has occurred.

138 G. The Commissioner may require a grower, dealer handler, or processor to destroy, at the cost of 139 the grower, dealer handler, or processor and in a manner approved of and verified by the Commissioner, 140 any Cannabis sativa that the grower grows, in which the dealer deals the handler handles, or that the 141 processor processes that has been tested and is found to have a concentration of tetrahydrocannabinol 142 that is greater than that allowed by federal law, or any Cannabis sativa product that the processor 143 produces.

144 H. Notwithstanding the provisions of subsection G, if the provisions of subdivisions 1 and 2 are 145 included in a plan that (i) is submitted by the Department pursuant to § 10113 of the federal Agriculture 146 Improvement Act of 2018, P.L. 115-334, (ii) requires the Department to monitor and regulate the 147 production of industrial hemp in the Commonwealth, and (iii) is approved by the U.S. Secretary of 148 Agriculture:

149 1. The Commissioner may require a grower, dealer handler, or processor to destroy, at the cost of 150 the grower, dealer handler, or processor and in a manner approved of and verified by the Commissioner, 151 any Cannabis sativa that the grower grows, in which the dealer deals the handler handles, or that the 152 processor processes that has been tested and is found to have a concentration of tetrahydrocannabinol 153 that is greater than 0.6 percent.

154 2. If such a test of Cannabis sativa indicates a concentration of tetrahydrocannabinol that is greater 155 than 0.6 percent but less than one percent, the Commissioner shall allow the grower, dealer handler, or 156 processor to request that the Cannabis sativa be sampled and tested again before he requires its 157 destruction.

158 I. The Commissioner shall advise the Superintendent of State Police or the chief law-enforcement 159 officer of the appropriate county or city when, with a culpable mental state greater than negligence, a 160 grower grows, a dealer deals in a handler handles, or a processor processes any Cannabis sativa with a concentration of tetrahydrocannabinol that is greater than that allowed by federal law or a processor 161 produces a Cannabis sativa product. 162

163 J. The Commissioner may pursue any permits or waivers from the U.S. Drug Enforcement 164 Administration or appropriate federal agency that he determines to be necessary for the advancement of 165 the industrial hemp industry.

166 K. The Commissioner may establish a corrective action plan to address a negligent violation of any 167 provision of this chapter. 168

§ 3.2-4115. Issuance of registrations; exemption.

169 A. The Commissioner shall establish a registration program to allow a person to grow, deal in 170 *handle*, or process industrial hemp in the Commonwealth.

171 B. Any person seeking to grow, deal in *handle*, or process industrial hemp in the Commonwealth 172 shall apply to the Commissioner for a registration on a form provided by the Commissioner. At a 173 minimum, the application shall include:

174 1. The name and mailing address of the applicant;

175 2. The legal description and geographic data sufficient for locating (i) the land on which the 176 applicant intends to grow industrial hemp, (ii) the site at which the applicant intends to deal in handle 177 industrial hemp, or (iii) the site at which the applicant intends to process industrial hemp. A registration 178 shall authorize industrial hemp growth, dealing in handling, or processing only at the location specified 179 in the registration;

180 3. A signed statement indicating whether the applicant has ever been convicted of a felony. A person 181 with a prior felony drug conviction within 10 years of applying for a registration under this section shall 182 not be eligible to be registered;

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183 4. Written consent allowing the sheriff's office, police department, or Department of State Police, if a 184 registration is ultimately issued to the applicant, to enter the premises on which the industrial hemp is 185 grown, dealt in handled, or processed to conduct physical inspections of the industrial hemp and to 186 ensure compliance with the requirements of this chapter. No more than two physical inspections shall be 187 conducted under this subdivision per year, unless a valid search warrant for an inspection has been 188 issued by a court of competent jurisdiction;

189 5. Written consent allowing the Commissioner or his designee to enter the premises on which the 190 industrial hemp is grown, dealt in handled, or processed to conduct inspections and sampling of the 191 industrial hemp to ensure compliance with the requirements of this chapter;

192 6. A statement of the approximate square footage or acreage of the location he intends to use as a 193 production field, dealership handler's storage site, or process site; 194

7. Any other information required by the Commissioner; and

8. The payment of a nonrefundable application fee, in an amount set by the Commissioner.

196 C. Each registration issued pursuant to this section shall be valid for a period of one year from the 197 date of issuance and may be renewed in successive years. Each annual renewal shall require the 198 payment of a registration renewal fee, in an amount set by the Commissioner.

199 D. All records, data, and information filed in support of a registration application submitted pursuant 200 to this section and all information on a hemp producer license issued by the U.S. Department of 201 Agriculture submitted to the Commissioner pursuant to this section shall be considered proprietary and 202 excluded from the provisions of the Virginia Freedom of Information Act (§ 2.2-3700 et seq.).

203 E. Notwithstanding the provisions of subsection B, no federally licensed hemp producer shall be 204 required to apply to the Commissioner for a registration to grow industrial hemp in the Commonwealth. Each federally licensed hemp producer shall submit to the Commissioner a copy of his hemp producer 205 license issued by the U.S. Department of Agriculture pursuant to 7 C.F.R. Part 990. 206

§ 3.2-4116. Registration conditions.

208 A. A person who is not a federally licensed hemp producer shall obtain a registration pursuant to 209 subsection A of § 3.2-4115 prior to growing, dealing in handling, or processing any industrial hemp in 210 the Commonwealth.

B. A person issued a registration pursuant to subsection A of § 3.2-4115 shall:

1. Maintain records that reflect compliance with this chapter;

2. Retain all industrial hemp growing, dealing handling, or processing records for at least three years;

3. Allow his production field, dealership handler's storage site, or process site to be inspected by and 214 215 at the discretion of the Commissioner or his designee, the Department of State Police, or the chief 216 law-enforcement officer of the locality in which the production field, or dealership handler's storage 217 *site*, or process site exists;

218 4. Allow the Commissioner or his designee to monitor and test the grower's, dealer's handler's, or 219 processor's industrial hemp for compliance with tetrahydrocannabinol levels and for other appropriate 220 purposes established pursuant to § 3.2-4114, at the cost of the grower, dealer handler, or processor; and

221 5. If required by the Commissioner, destroy, at the cost of the grower, dealer handler, or processor and in a manner approved of and verified by the Commissioner, any Cannabis sativa that the grower 222 223 grows, the dealer deals in handler handles, or the processor processes that has been tested and, following any re-sampling and retesting as authorized pursuant to the provisions of § 3.2-4114.2, is 224 225 found to have a concentration of tetrahydrocannabinol that is greater than that allowed by federal law, or 226 any Cannabis sativa product that the processor produces. 227

§ 3.2-4118. Forfeiture of industrial hemp grower, handler, or processor registration; violations.

228 A. The Commissioner shall deny the application, or suspend or revoke the registration, of any person who, with a culpable mental state greater than negligence, violates any provision of this chapter. The 229 Commissioner shall provide reasonable notice of an informal fact-finding conference pursuant to 230 231 § 2.2-4019 to any person in connection with the denial, suspension, or revocation of a registration.

232 B. If a registration is revoked as the result of an informal hearing, the decision may be appealed, and 233 upon appeal an administrative hearing shall be conducted in accordance with the Administrative Process 234 Act (§ 2.2-4000 et seq.). The grower, dealer handler, or processor may appeal a final order to the circuit 235 court in accordance with the Administrative Process Act.

236 C. A person issued a registration pursuant to subsection A of \S 3.2-4115 who negligently (i) fails to 237 provide a description and geographic data sufficient for locating his production field, dealership 238 handler's storage site, or process site; (ii) grows, deals in handles, or processes Cannabis sativa with a tetrahydrocannabinol concentration greater than that allowed by federal law; or (iii) produces a Cannabis 239 240 sativa product shall comply with any corrective action plan established by the Commissioner in 241 accordance with the provisions of subsection E. The Commissioner shall not deem a grower negligent if 242 such grower makes reasonable efforts to grow industrial hemp and grows Cannabis sativa with a tetrahydrocannabinol concentration that does not exceed the total delta-9 tetrahydrocannabinol 243 244 concentration percentage established in federal regulations applicable to negligent violations located at 7

245 C.F.R. § 990.6(b)(3).

246 D. A person who grows, deals in handles, or processes industrial hemp and who negligently fails to 247 register pursuant to subsection A of § 3.2-4115 shall comply with any corrective action plan established 248 by the Commissioner in accordance with the provisions of subsection E.

249 E. A corrective action plan established by the Commissioner in response to a negligent violation of a 250 provision of this chapter shall identify a reasonable date by which the person who is the subject of the 251 plan shall correct the negligent violation and shall require such person to report periodically for not less 252 than two calendar years to the Commissioner on the person's compliance with the provisions of this 253 chapter.

F. No person who negligently violates the provisions of this chapter three times in a five-vear period 254 255 shall be eligible to grow, deal in handle, or process industrial hemp for a period of five years beginning 256 on the date of the third violation. 257

§ 3.2-4119. Eligibility to receive tobacco settlement funds.

258 Industrial hemp growers, dealers handlers, or processors registered under this chapter or federally 259 licensed hemp producers may be eligible to receive funds from the Tobacco Indemnification and 260 Community Revitalization Fund established pursuant to § 3.2-3106.

261 § 3.2-5145.1. Definitions.

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As used in this article, unless the context requires a different meaning:

263 "Food" means any article that is intended for human consumption and introduction into commerce, 264 whether the article is simple, mixed, or compound, and all substances or ingredients used in the 265 preparation thereof. "Food" does not mean drug as defined in § 54.1-3401.

266 "Industrial hemp" means a Cannabis sativa plant that has a concentration of tetrahydrocannabinol 267 that is no greater than that allowed by federal law.

268 "Industrial hemp extract" means an extract (i) of a Cannabis sativa plant that has a concentration of 269 tetrahydrocannabinol that is no greater than that allowed for industrial hemp by federal law and, (ii) that 270 is intended for human consumption, and (iii) when offered for retail sale, that (a) contains a total 271 tetrahydrocannabinol concentration that is no greater than 0.3 percent and (b) contains no more than two milligrams of total tetrahydrocannabinol per package. "Industrial hemp extract" is not a hemp seed-derived ingredient that is approved by the U.S. Food and Drug Administration or is the subject of 272 273 274 a generally recognized as safe notice for which the U.S. Food and Drug Administration had no 275 questions.

276 "Tetrahydrocannabinol" means the same as that term is defined in § 3.2-4112.

"Total tetrahydrocannabinol" means the same as that term is defined in § 3.2-4112.

§ 3.2-5145.2:1. Sellers or manufacturers of industrial hemp extract; penalties.

279 A. Any person who *manufactures*, sells, or offers for sale an industrial hemp extract or food 280 containing an industrial hemp extract shall be subject to the requirements of this chapter and regulations 281 adopted pursuant to this chapter.

282 B. Any person who (i) manufactures, sells, or offers for sale an industrial hemp extract or food 283 containing an industrial hemp extract without first obtaining a permit to do so from the Commissioner 284 pursuant to § 3.2-5100, unless exempt from a permit pursuant to subdivision C 6 of § 3.2-5130; (ii) 285 continues to manufacture, sell, or offer for sale an industrial hemp extract or food containing an 286 industrial hemp extract after revocation or suspension of such permit; (iii) fails to disclose on a form 287 prescribed by the Commissioner that he intends to manufacture, sell, or offer for sale a substance 288 intended to be consumed orally that contains an industrial hemp-derived cannabinoid; (iv) manufactures, 289 sells, or offers for sale a food that (a) has a total tetrahydrocannabinol concentration that is greater 290 than 0.3 percent or (b) contains more than two milligrams of total tetrahydrocannabinol per package; 291 (v) manufactures, offers for sale, or sells in violation of this chapter or a regulation adopted pursuant to 292 this chapter a substance intended to be consumed orally that is advertised or labeled as containing an 293 industrial hemp-derived cannabinoid; or (vi) otherwise violates any provision of this chapter or a 294 regulation adopted pursuant to this chapter, in addition to any other penalties provided, is subject to a 295 civil penalty not to exceed \$10,000 for each day a violation occurs. Such penalty shall be collected by 296 the Commissioner and the proceeds shall be payable to the State Treasurer for remittance to the 297 Department.

298 C. Any person who (i) manufactures, sells, or offers for sale an industrial hemp extract or food 299 containing an industrial hemp extract without first obtaining a permit to do so from the Commissioner 300 pursuant to § 3.2-5100, unless exempt from a permit pursuant to subdivision C 6 of § 3.2-5130; (ii) 301 continues to manufacture, sell, or offer for sale an industrial hemp extract or food containing an 302 industrial hemp extract after revocation or suspension of such permit; (iii) fails to disclose on a form 303 prescribed by the Commissioner that he intends to manufacture, sell, or offer for sale a substance intended to be consumed orally that contains an industrial hemp-derived cannabinoid; (iv) manufactures, 304 305 offers for sale, or sells in violation of this chapter or a regulation adopted pursuant to this chapter a

306 substance intended to be consumed orally that is advertised or labeled as containing an industrial 307 hemp-derived cannabinoid; or (v) otherwise violates any provision of this chapter or a regulation 308 adopted pursuant to this chapter, in addition to any other penalties provided, is guilty of a Class 1 309 misdemeanor. Each day in which a violation occurs shall constitute a separate offense.

310 D. The Commissioner may, in accordance with the Administrative Process Act (§ 2.2-400 et seq.), 311 deny, suspend, or revoke a permit issued pursuant to § 3.2-5100 if the permitted entity is found to have 312 violated subdivision A 69, 70, 71, 72, 73, or 74 of § 59.1-200 by a court of competent jurisdiction.

313 E. This section shall not apply to a person authorized to offer for sale or sell products that are (i) 314 approved for marketing by the U.S. Food and Drug Administration and scheduled in the Drug Control 315 Act (§ 54.1-3400 et seq.) or (ii) dispensed pursuant to Article 4.2 (§ 54.1-3442.5 et seq.) of Chapter 34 316 of Title 54.1. 317

§ 3.2-5145.4. Industrial hemp extract requirements.

A. An industrial hemp extract shall (i) be produced from industrial hemp grown in compliance with 318 319 applicable law and (ii) notwithstanding any authority under federal law to have a greater concentration 320 of tetrahydrocannabinol, when offered for retail sale, (a) have a total tetrahydrocannabinol concentration 321 of no greater than 0.3 percent and (b) contain no more than two milligrams of total 322 tetrahydrocannabinol per package.

323 B. In addition to the requirements of this chapter, an industrial hemp extract or food containing an 324 industrial hemp extract shall comply with regulations adopted by the Board pursuant to § 3.2-5145.5. 325

§ 3.2-5145.4:1. Labeling and packaging requirements.

326 A. An industrial hemp extract or food containing an industrial hemp extract shall be contained in 327 child-resistant packaging, as defined in § 4.1-600.

B. An industrial hemp extract or food containing an industrial hemp extract shall be packaged and 328 equipped with a label that states, in English and in a font no less than 1/16 of an inch, (i) all 329 330 ingredients contained in the industrial hemp extract or food containing an industrial hemp extract, (ii) the amount of such industrial hemp extract or food containing an industrial hemp extract that 331 332 constitutes a single serving, and (iii) the number of milligrams and percent of total tetrahydrocannabinol 333 per serving and number of milligrams and percent of total tetrahydrocannabinol per package.

334 C. Any industrial hemp extract or food containing an industrial hemp extract that contains 335 tetrahydrocannabinol (i) shall be equipped with a label that states that the industrial hemp extract or 336 food containing an industrial hemp extract contains tetrahydrocannabinol and (ii) may not be sold to 337 persons younger than 21 years of age.

338 D. An industrial hemp extract or food containing an industrial hemp extract, when offered for retail 339 sale, shall be accompanied by a certificate of analysis, produced by an independent laboratory that is 340 registered with the U.S. Drug Enforcement Administration and is accredited pursuant to standard 341 ISO/IEC 17025 of the International Organization for Standardization by a third-party accrediting body, 342 that states the total tetrahydrocannabinol concentration of the substance or the total 343 tetrahydrocannabinol concentration of the batch from which the substance originates. The certificate of 344 accreditation pursuant to standard ISO/IEC 17025 issued by the third-party accrediting body to the 345 independent laboratory shall be available for review at the location at which the industrial hemp extract 346 or food containing an industrial hemp extract is offered for sale or sold.

E. A manufacturer shall identify each batch of an industrial hemp extract or a food containing an 347 348 industrial hemp extract with a unique code for traceability. Julian date coding or any other system developed and documented by the manufacturer for assigning a unique code to a batch may be used. 349 350 The batch identification shall appear and be legible on the label of an industrial hemp extract or food 351 containing an industrial hemp extract.

352 F. The label of an industrial hemp extract or food containing an industrial hemp extract shall not 353 contain a claim indicating the product is intended for diagnosis, cure, mitigation, treatment, or 354 prevention of disease, which shall render the product a drug, as that term is defined in 21 U.S.C. 355 § 321(g)(1). An industrial hemp extract or food containing an industrial hemp extract with a label that 356 contains a claim indicating the product is intended for diagnosis, cure, mitigation, treatment, or 357 prevention of disease shall be considered misbranded. 358

§ 3.2-5145.5. Regulations.

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A. The Board is authorized to adopt regulations for the efficient enforcement of this article.

360 B. The Board shall adopt regulations identifying contaminants of an industrial hemp extract or a food containing an industrial hemp extract and establishing tolerances for such identified contaminants. 361

362 C. The Board shall adopt regulations establishing labeling requirements for an industrial hemp extract 363 or a food containing an industrial hemp extract. Such regulations shall require that any industrial hemp 364 extract or food containing an industrial hemp extract that contains tetrahydrocannabinol be equipped with a label that states (i) that the industrial hemp extract or food containing an industrial hemp extract 365 366 contains tetrahydrocannabinol and may not be sold to persons younger than 21 years of age, (ii) all ingredients contained in the industrial hemp extract or food containing an industrial hemp extract, (iii) 367

368 the amount of such industrial hemp extract or food containing an industrial hemp extract that constitutes 369 a single serving, and (iv) the total percentage and milligrams of tetrahydrocannabinol included in the 370 industrial hemp extract or food containing an industrial hemp extract and the number of milligrams of 371 tetrahydrocannabinol that are contained in each serving.

372 D. The Board shall adopt regulations establishing batch testing requirements for industrial hemp
373 extracts. The Board shall require that batch testing of industrial hemp extracts be conducted by an
374 independent testing laboratory that meets criteria established by the Board.

375 E. D. With the exception of § 2.2-4031, neither the provisions of the Administrative Process Act 376 (§ 2.2-4000 et seq.) nor public participation guidelines adopted pursuant thereto shall apply to the 377 adoption of any regulation pursuant to this section. Prior to adopting any regulation pursuant to this 378 section, the Board shall publish a notice of opportunity to comment in the Virginia Register of 379 Regulations and post the action on the Virginia Regulatory Town Hall. Such notice of opportunity to 380 comment shall contain (i) a summary of the proposed regulation; (ii) the text of the proposed regulation; and (iii) the name, address, and telephone number of the agency contact person responsible for receiving 381 382 public comments. Such notice shall be made at least 60 days in advance of the last date prescribed in 383 such notice for submittals of public comment. The legislative review provisions of subsections A and B 384 of § 2.2-4014 shall apply to the promulgation or final adoption process for regulations pursuant to this 385 section. The Board shall consider and keep on file all public comments received for any regulation 386 adopted pursuant to this section.

387 § 4.1-600. Definitions.

393

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388 As used in this subtitle, unless the context requires a different meaning:

389 "Advertisement" or " advertising" means any written or verbal statement, illustration, or depiction
390 that is calculated to induce sales of retail marijuana, retail marijuana products, marijuana plants, or
391 marijuana seeds, including any written, printed, graphic, digital, electronic, or other material, billboard,
392 sign, or other outdoor display, publication, or radio or television broadcast.

"Authority" means the Virginia Cannabis Control Authority created pursuant to this subtitle.

"Board" means the Board of Directors of the Virginia Cannabis Control Authority.

395 "Cannabis Control Act" means Subtitle II (§ 4.1-600 et seq.).

396 "Child-resistant" means, with respect to packaging or a container, (i) specially designed or 397 constructed to be significantly difficult for a typical child under five years of age to open and not to be 398 significantly difficult for a typical adult to open and reseal and (ii) for any product intended for more 399 than a single use or that contains multiple servings, resealable.

400 "Cultivation" or "cultivate" means the planting, propagation, growing, harvesting, drying, curing,
 401 grading, trimming, or other similar processing of marijuana for use or sale. "Cultivation" or "cultivate"
 402 does not include manufacturing or testing.

403 "Edible marijuana product" means a marijuana product intended to be consumed orally, including
 404 marijuana intended to be consumed orally or marijuana concentrate intended to be consumed orally.

405 "Immature plant" means a nonflowering marijuana plant that is no taller than eight inches and no
406 wider than eight inches, is produced from a cutting, clipping, or seedling, and is growing in a container.
407 "Licensed" means the holding of a valid license granted by the Authority.

408 "Licensee" means any person to whom a license has been granted by the Authority.

"Manufacturing" or "manufacture" means the production of marijuana products or the blending,
infusing, compounding, or other preparation of marijuana and marijuana products, including marijuana
extraction or preparation by means of chemical synthesis. "Manufacturing" or "manufacture" does not
include cultivation or testing.

413 "Marijuana" means any part of a plant of the genus Cannabis, whether growing or not, its seeds or 414 resin; and every compound, manufacture, salt, derivative, mixture, or preparation of such plant, its seeds, its resin, or any extract containing one or more cannabinoids. "Marijuana" does not include (i) the 415 416 mature stalks of such plant, fiber produced from such stalk, or oil or cake made from the seed of such 417 plant, unless such stalks, fiber, oil, or cake is combined with other parts of plants of the genus Cannabis-418 "Marijuana" does not include (i); (ii) industrial hemp, as defined in § 3.2-4112, that is possessed by a 419 person registered pursuant to subsection A of § 3.2-4115 or his agent or (ii); (iii) industrial hemp, as 420 defined in § 3.2-4112, that is possessed by a person who holds a hemp producer license issued by the 421 U.S. Department of Agriculture pursuant to 7 C.F.R. Part 990; (iv) a hemp product, as defined in 422 § 3.2-4112, containing a tetrahydrocannabinol concentration of no greater than 0.3 percent that is derived 423 from industrial hemp, as defined in § 3.2-4112, that is grown, dealt, or processed in compliance with 424 state or federal law; (v) an industrial hemp extract, as defined in § 3.2-5145.1; or (vi) any substance 425 containing a tetrahydrocannabinol isomer, ester, ether, salt, or salts of such isomer, ester, or ether that 426 has been placed by the Board of Pharmacy into one of the schedules set forth in the Drug Control Act 427 (§ 54.1-3400 et seq.) pursuant to § 54.1-3443.

428 "Marijuana concentrate" means marijuana that has undergone a process to concentrate one or more

429 active cannabinoids, thereby increasing the product's potency. Resin from granular trichomes from a430 marijuana plant is a concentrate for purposes of this subtitle.

"Marijuana cultivation facility" means a facility licensed under this subtitle to cultivate, label, and 431 432 package retail marijuana; to purchase or take possession of marijuana plants and seeds from other 433 marijuana cultivation facilities; to transfer possession of and sell retail marijuana, immature marijuana 434 plants, and marijuana seeds to marijuana wholesalers and retail marijuana stores; to transfer possession 435 of and sell retail marijuana, marijuana plants, and marijuana seeds to other marijuana cultivation 436 facilities; to transfer possession of and sell retail marijuana to marijuana manufacturing facilities; and to sell immature marijuana plants and marijuana seeds to consumers for the purpose of cultivating 437 marijuana at home for personal use. 438

439 "Marijuana establishment" means a marijuana cultivation facility, a marijuana testing facility, a
 440 marijuana manufacturing facility, a marijuana wholesaler, or a retail marijuana store.

441 "Marijuana manufacturing facility" means a facility licensed under this subtitle to manufacture, label,
442 and package retail marijuana and retail marijuana products; to purchase or take possession of retail
443 marijuana from a marijuana cultivation facility or another marijuana manufacturing facility; and to
444 transfer possession of and sell retail marijuana and retail marijuana products to marijuana wholesalers,
445 retail marijuana stores, or other marijuana manufacturing facilities.

"Marijuana paraphernalia" means all equipment, products, and materials of any kind that are either
designed for use or are intended for use in planting, propagating, cultivating, growing, harvesting,
manufacturing, compounding, converting, producing, processing, preparing, strength testing, analyzing,
packaging, repackaging, storing, containing, concealing, ingesting, inhaling, or otherwise introducing into
the human body marijuana.

451 "Marijuana products" means (i) products that are composed of marijuana and other ingredients and452 are intended for use or consumption, ointments, and tinctures or (ii) marijuana concentrate.

"Marijuana testing facility" means a facility licensed under this subtitle to develop, research, or test
marijuana, marijuana products, and other substances.

"Marijuana wholesaler" means a facility licensed under this subtitle to purchase or take possession of
retail marijuana, retail marijuana products, immature marijuana plants, and marijuana seeds from a
marijuana cultivation facility, a marijuana manufacturing facility, or another marijuana wholesaler and to
transfer possession and sell or resell retail marijuana, retail marijuana products, immature marijuana
plants, and marijuana seeds to a marijuana cultivation facility, marijuana manufacturing facility, retail
marijuana store, or another marijuana wholesaler.

461 "Non-retail marijuana" means marijuana that is not cultivated, manufactured, or sold by a licensed462 marijuana establishment.

463 "Non-retail marijuana products" means marijuana products that are not manufactured and sold by a464 licensed marijuana establishment.

"Place or premises" means the real estate, together with any buildings or other improvements thereon,
designated in the application for a license as the place at which the cultivation, manufacture, sale, or
testing of retail marijuana or retail marijuana products shall be performed, except that portion of any
such building or other improvement actually and exclusively used as a private residence.

"Public place" means any place, building, or conveyance to which the public has, or is permitted to
have, access, including restaurants, soda fountains, hotel dining areas, lobbies and corridors of hotels,
and any park, place of public resort or amusement, highway, street, lane, or sidewalk adjoining any
highway, street, or lane.

473 "Residence" means any building or part of a building or structure where a person resides, but does
474 not include any part of a building that is not actually and exclusively used as a private residence, nor
475 any part of a hotel or club other than a private guest room thereof.
476 "Retail marijuane" means any first in the structure of the structure where a person resides, but does

476 "Retail marijuana" means marijuana that is cultivated, manufactured, or sold by a licensed marijuana477 establishment.

478 "Retail marijuana products" means marijuana products that are manufactured and sold by a licensed479 marijuana establishment.

480 "Retail marijuana store" means a facility licensed under this subtitle to purchase or take possession of
481 retail marijuana, retail marijuana products, immature marijuana plants, or marijuana seeds from a
482 marijuana cultivation facility, marijuana manufacturing facility, or marijuana wholesaler and to sell retail
483 marijuana, retail marijuana products, immature marijuana plants, or marijuana seeds to consumers.

484 "Sale" and "sell" includes soliciting or receiving an order for; keeping, offering, or exposing for sale;
485 peddling, exchanging, or bartering; or delivering otherwise than gratuitously, by any means, retail
486 marijuana or retail marijuana products.

487 "Special agent" means an employee of the Virginia Cannabis Control Authority whom the Board has designated as a law-enforcement officer pursuant to this subtitle.

489 "Testing" or "test" means the research and analysis of marijuana, marijuana products, or other
490 substances for contaminants, safety, or potency. "Testing" or "test" does not include cultivation or

491 manufacturing.

492 "Tetrahydrocannabinol" means the same as that term is defined in § 3.2-4112.

493 "Total tetrahydrocannabinol" means the same as that term is defined in § 3.2-4112.

494 § 18.2-247. Use of terms "controlled substances," "marijuana," "Schedules I, II, III, IV, V, and 495 VI," "imitation controlled substance," and "counterfeit controlled substance" in Title 18.2.

496 A. Wherever the terms "controlled substances" and "Schedules I, II, III, IV, V, and VI" are used in
497 Title 18.2, such terms refer to those terms as they are used or defined in the Drug Control Act
498 (§ 54.1-3400 et seq.).

B. The term "imitation controlled substance" when used in this article means (i) a counterfeit
controlled substance or (ii) a pill, capsule, tablet, or substance in any form whatsoever which is not a
controlled substance subject to abuse, and:

502 1. Which by overall dosage unit appearance, including color, shape, size, marking and packaging or
503 by representations made, would cause the likelihood that such a pill, capsule, tablet, or substance in any
504 other form whatsoever will be mistaken for a controlled substance unless such substance was introduced
505 into commerce prior to the initial introduction into commerce of the controlled substance which it is
506 alleged to imitate; or

507 2. Which by express or implied representations purports to act like a controlled substance as a
508 stimulant or depressant of the central nervous system and which is not commonly used or recognized for
509 use in that particular formulation for any purpose other than for such stimulant or depressant effect,
510 unless marketed, promoted, or sold as permitted by the U.S. Food and Drug Administration.

511 C. In determining whether a pill, capsule, tablet, or substance in any other form whatsoever, is an 512 "imitation controlled substance," there shall be considered, in addition to all other relevant factors, 513 comparisons with accepted methods of marketing for legitimate nonprescription drugs for medicinal 514 purposes rather than for drug abuse or any similar nonmedicinal use, including consideration of the 515 packaging of the drug and its appearance in overall finished dosage form, promotional materials or 516 representations, oral or written, concerning the drug, and the methods of distribution of the drug and 517 where and how it is sold to the public.

518 D. The term "marijuana" when used in this article means any part of a plant of the genus Cannabis, 519 whether growing or not, its seeds or resin; and every compound, manufacture, salt, derivative, mixture, 520 or preparation of such plant, its seeds, its resin, or any extract containing one or more cannabinoids. 521 "Marijuana" does not include (i) the mature stalks of such plant, fiber produced from such stalk, oil or 522 cake made from the seed of such plant, unless such stalks, fiber, oil or cake is combined with other 523 parts of plants of the genus Cannabis- Marijuana does not include (i); (ii) industrial hemp, as defined in 524 § 3.2-4112, that is possessed by a person registered pursuant to subsection A of § 3.2-4115 or his agent; 525 (iii) industrial hemp, as defined in § 3.2-4112, that is possessed by a person who holds a hemp 526 producer license issued by the U.S. Department of Agriculture pursuant to 7 C.F.R. Part 990; or (iii) (iv) 527 a hemp product, as defined in § 3.2-4112, containing a tetrahydrocannabinol concentration of no greater 528 than 0.3 percent that is derived from industrial hemp, as defined in § 3.2-4112, that is grown, dealt, or 529 processed in compliance with state or federal law; (v) an industrial hemp extract, as defined in 530 § 3.2-5145.1; or (vi) any substance containing a tetrahydrocannabinol isomer, ester, ether, salt or salts 531 of such isomer, ester, or ether that has been placed by the Board of Pharmacy into one of the schedules set forth in the Drug Control Act (§ 54.1-3400 et seq.) pursuant to § 54.1-3443. 532

E. The term "counterfeit controlled substance" means a controlled substance that, without
authorization, bears, is packaged in a container or wrapper that bears, or is otherwise labeled to bear, the
trademark, trade name, or other identifying mark, imprint or device or any likeness thereof, of a drug
manufacturer, processor, packer, or distributor other than the manufacturer, processor, packer, or
distributor who did in fact so manufacture, process, pack or distribute such drug.

F. The term "tetrahydrocannabinol" means any naturally occurring or synthetic
tetrahydrocannabinol, including its salts, isomers, and salts of isomers whenever the existence of such
salts, isomers, and salts of isomers is possible within the specific chemical designation and any
preparation, mixture, or substance containing, or mixed or infused with, any detectable amount of
tetrahydrocannabinol. "Tetrahydrocannabinol" includes delta-6a(10a), delta-7, delta-8, delta-9, and
delta-10-tetrahydrocannibinol. For the purposes of this definition, "isomer" means the optical, position,
and geometric isomers.

545 G. The term "total tetrahydrocannabinol" means the sum, after the application of any necessary
 546 conversion factor, of the percentage by weight of tetrahydrocannabinol and the percentage by weight of tetrahydrocannabinolic acid.

548 *H*. The Department of Forensic Science shall determine the proper methods for detecting the concentration of delta-9-tetrahydrocannabinol (THC) tetrahydrocannabinol in substances for the purposes
550 of this title, *Chapter 11 (§ 4.1-1100 et seq.) of Title 4.1,* and §§ § 54.1-3401 and 54.1-3446. The testing methodology shall use post-decarboxylation testing or other equivalent method and shall consider the

552 potential conversion of delta-9-tetrahydrocannibinol tetrahydrocannabinolic acid (THC-A) into THC

553 *tetrahydrocannabinol*. The test result shall include the total available THC derived from the sum of the 554 THC and THC-A content.

555 § 18.2-251.1:3. Possession or distribution of cannabis oil, or industrial hemp; laboratories; 556 Department of Agriculture and Consumer Services, Department of Law employees.

557 A. No person employed by an analytical laboratory to retrieve, deliver, or possess cannabis oil or 558 industrial hemp samples from a permitted pharmaceutical processor, a registered industrial hemp grower, 559 a federally licensed hemp producer, or a registered industrial hemp processor for the purpose of performing required testing shall be prosecuted under Chapter 11 (§ 4.1-1100 et seq.) of Title 4.1 or 560 § 18.2-248, 18.2-248.1, 18.2-250, or 18.2-255 for the possession or distribution of cannabis oil or 561 industrial hemp or for storing cannabis oil or industrial hemp for testing purposes in accordance with 562 regulations promulgated by the Board of Pharmacy and the Board of Agriculture and Consumer 563 564 Services.

B. No employee of the Department of Agriculture and Consumer Services or of the Department of *Law* shall be prosecuted under § 18.2-247, 18.2-248, 18.2-248.01, 18.2-248.1, or 18.2-250 for the
possession or distribution of industrial hemp or any substance containing tetrahydrocannabinol when
possession of industrial hemp or any substance containing tetrahydrocannabinol is necessary in the
performance of his duties.

§ 18.2-371.2. Prohibiting purchase or possession of tobacco products, nicotine vapor products,
alternative nicotine products, and hemp products intended for smoking by a person under 21
years of age or sale of tobacco products, nicotine vapor products, alternative nicotine products,
and hemp products intended for smoking to persons under 21 years of age; civil penalties.

A. No person shall sell to, distribute to, purchase for, or knowingly permit the purchase by any
person less than 21 years of age, knowing or having reason to believe that such person is less than 21
years of age, any tobacco product, nicotine vapor product, alternative nicotine product, or hemp product
intended for smoking.

578 Tobacco products, nicotine vapor products, alternative nicotine products, and hemp products intended 579 for smoking may be sold from a vending machine only if the machine is (i) posted with a notice, in a 580 conspicuous manner and place, indicating that the purchase or possession of such products by persons 581 under 21 years of age is unlawful and (ii) located in a place that is not open to the general public and is 582 not generally accessible to persons under 21 years of age. An establishment that prohibits the presence 583 of persons under 21 years of age unless accompanied by a person 21 years of age or older is not open 584 to the general public.

585 B. No person less than 21 years of age shall attempt to purchase, purchase, or possess any tobacco 586 product, nicotine vapor product, alternative nicotine product, or hemp product intended for smoking. The 587 provisions of this subsection shall not be applicable to the possession of tobacco products, nicotine 588 vapor products, alternative nicotine products, or hemp products intended for smoking by a person less than 21 years of age (i) making a delivery of tobacco products, nicotine vapor products, alternative 589 590 nicotine products, or hemp products intended for smoking in pursuance of his employment or (ii) as part 591 of a scientific study being conducted by an organization for the purpose of medical research to further 592 efforts in cigarette and tobacco use prevention and cessation and tobacco product regulation, provided 593 that such medical research has been approved by an institutional review board pursuant to applicable 594 federal regulations or by a research review committee pursuant to Chapter 5.1 (§ 32.1-162.16 et seq.) of 595 Title 32.1. This subsection shall not apply to purchase, attempt to purchase, or possession by a law-enforcement officer or his agent when the same is necessary in the performance of his duties. 596

597 C. No person shall sell a tobacco product, nicotine vapor product, alternative nicotine product, or 598 hemp product intended for smoking to any individual who does not demonstrate, by producing a driver's 599 license or similar photo identification issued by a government agency, that the individual is at least 21 600 years of age. Such identification is not required from an individual whom the person has reason to 601 believe is at least 21 years of age or who the person knows is at least 21 years of age. Proof that the 602 person demanded, was shown, and reasonably relied upon a photo identification stating that the 603 individual was at least 21 years of age shall be a defense to any action brought under this subsection. In **604** determining whether a person had reason to believe an individual is at least 21 years of age, the trier of 605 fact may consider, but is not limited to, proof of the general appearance, facial characteristics, behavior, 606 and manner of the individual.

607 This subsection shall not apply to mail order or Internet sales, provided that the person offering the tobacco product, nicotine vapor product, alternative nicotine product, or hemp product intended for smoking for sale through mail order or the Internet (i) prior to the sale of the tobacco product, nicotine vapor product, alternative nicotine product intended for smoking verifies that the purchaser is at least 21 years of age through a commercially available database that is regularly used by businesses or governmental entities for the purpose of age and identity verification and (ii) uses a method of mailing, shipping, or delivery that requires the signature of a person at least 21 years of age

614 before the tobacco product, nicotine vapor product, alternative nicotine product, or hemp product 615 intended for smoking will be released to the purchaser.

616 D. The provisions of subsections B and C shall not apply to the sale, giving, or furnishing of any
617 tobacco product, nicotine vapor product, alternative nicotine product, or hemp product intended for
618 smoking to any active duty military personnel who are 18 years of age or older. An identification card
619 issued by the Armed Forces of the United States shall be accepted as proof of age for this purpose.

E. A violation of subsection A or C by an individual or by a separate retail establishment that
involves a nicotine vapor product, alternative nicotine product, hemp product intended for smoking, or
tobacco product other than a bidi is punishable by a civil penalty not to exceed \$100 for a first
violation, a civil penalty not to exceed \$200 for a second violation, and a civil penalty not to exceed
\$500 for a third or subsequent violation.

625 A violation of subsection A or C by an individual or by a separate retail establishment that involves 626 the sale, distribution, or purchase of a bidi is punishable by a civil penalty in the amount of \$500 for a first violation, a civil penalty in the amount of \$1,000 for a second violation, and a civil penalty in the 627 628 amount of \$2,500 for a third or subsequent violation. Where a defendant retail establishment offers 629 proof that it has trained its employees concerning the requirements of this section, the court shall 630 suspend all of the penalties imposed hereunder. However, where the court finds that a retail 631 establishment has failed to so train its employees, the court may impose a civil penalty not to exceed 632 \$1,000 in lieu of any penalties imposed hereunder for a violation of subsection A or C involving a 633 nicotine vapor product, alternative nicotine product, hemp product intended for smoking, or tobacco 634 product other than a bidi.

A violation of subsection B is punishable by a civil penalty not to exceed \$100 for a first violation and a civil penalty not to exceed \$250 for a second or subsequent violation. A court may, as an alternative to the civil penalty, and upon motion of the defendant, prescribe the performance of up to 20 hours of community service for a first violation of subsection B and up to 40 hours of community service for a second or subsequent violation. If the defendant fails or refuses to complete the community service as prescribed, the court may impose the civil penalty. Upon a violation of subsection B, the judge may enter an order pursuant to subdivision A 9 of § 16.1-278.8.

642 Any attorney for the Commonwealth of the county or city in which an alleged violation occurred
643 may bring an action to recover the civil penalty, which shall be paid into the state treasury. Any
644 law-enforcement officer may issue a summons for a violation of subsection A, B, or C.

645 F. 1. Cigarettes and hemp products intended for smoking shall be sold only in sealed packages 646 provided by the manufacturer, with the required health warning. The proprietor of every retail 647 establishment that offers for sale any tobacco product, nicotine vapor product, alternative nicotine 648 product, or hemp product intended for smoking shall post in a conspicuous manner and place a sign or 649 signs indicating that the sale of tobacco products, nicotine vapor products, alternative nicotine products, or hemp products intended for smoking to any person under 21 years of age is prohibited by law. Any 650 attorney for the county, city, or town in which an alleged violation of this subsection occurred may 651 652 enforce this subsection by civil action to recover a civil penalty not to exceed \$50 \$500. The civil 653 penalty shall be paid into the local treasury. No filing fee or other fee or cost shall be charged to the 654 county, city, or town which instituted the action.

655 2. For the purpose of compliance with regulations of the Substance Abuse and Mental Health
656 Services Administration published at 61 Federal Register 1492, the Department of Agriculture and
657 Consumer Services may promulgate regulations which allow the Department to undertake the activities
658 necessary to comply with such regulations.

659 3. Any attorney for the county, city, or town in which an alleged violation of this subsection
occurred may enforce this subsection by civil action to recover a civil penalty not to exceed \$100 \$500.
661 The civil penalty shall be paid into the local treasury. No filing fee or other fee or cost shall be charged
to the county, city, or town which instituted the action.

663 G. Nothing in this section shall be construed to create a private cause of action.

H. Agents of the Virginia Alcoholic Beverage Control Authority designated pursuant to § 4.1-105may issue a summons for any violation of this section.

666 I. As used in this section:

"Alternative nicotine product" means any noncombustible product containing nicotine that is intended
for human consumption, whether chewed, absorbed, dissolved, or ingested by any other means.
"Alternative nicotine product" does not include any nicotine vapor product, tobacco product, or product
regulated as a drug or device by the U.S. Food and Drug Administration (FDA) under Chapter V (21
U.S.C. § 351 et seq.) of the Federal Food, Drug, and Cosmetic Act.

672 "Bidi" means a product containing tobacco that is wrapped in temburni leaf (diospyros melanoxylon)
673 or tendu leaf (diospyros exculpra), or any other product that is offered to, or purchased by, consumers as
674 a bidi or beedie.

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675 "Hemp product" means the same as that term is defined in § 3.2-4112.

676 "Nicotine vapor product" means any noncombustible product containing nicotine that employs a heating element, power source, electronic circuit, or other electronic, chemical, or mechanical means, 677 678 regardless of shape or size, that can be used to produce vapor from nicotine in a solution or other form. 679 "Nicotine vapor product" includes any electronic cigarette, electronic cigar, electronic cigarillo, electronic 680 pipe, or similar product or device and any cartridge or other container of nicotine in a solution or other 681 form that is intended to be used with or in an electronic cigarette, electronic cigar, electronic cigarillo, electronic pipe, or similar product or device. "Nicotine vapor product" does not include any product **682** regulated by the FDA under Chapter V (21 U.S.C. § 351 et seq.) of the Federal Food, Drug, and 683 **684** Cosmetic Act.

"Tobacco product" means any product made of tobacco and includes cigarettes, cigars, smokeless tobacco, pipe tobacco, bidis, and wrappings. "Tobacco product" does not include any nicotine vapor 685 686 **687** product, alternative nicotine product, or product that is regulated by the FDA under Chapter V (21 U.S.C. § 351 et seq.) of the Federal Food, Drug, and Cosmetic Act. 688

"Wrappings" includes papers made or sold for covering or rolling tobacco or other materials for 689 690 smoking in a manner similar to a cigarette or cigar.

691 § 54.1-3401. Definitions.

As used in this chapter, unless the context requires a different meaning:

693 "Administer" means the direct application of a controlled substance, whether by injection, inhalation, 694 ingestion, or any other means, to the body of a patient or research subject by (i) a practitioner or by his 695 authorized agent and under his direction or (ii) the patient or research subject at the direction and in the 696 presence of the practitioner.

697 "Advertisement" means all representations disseminated in any manner or by any means, other than 698 by labeling, for the purpose of inducing, or which are likely to induce, directly or indirectly, the 699 purchase of drugs or devices.

700 "Agent" means an authorized person who acts on behalf of or at the direction of a manufacturer, 701 distributor, or dispenser. It does not include a common or contract carrier, public warehouseman, or 702 employee of the carrier or warehouseman.

703 "Anabolic steroid" means any drug or hormonal substance, chemically and pharmacologically related 704 to testosterone, other than estrogens, progestins, corticosteroids, and dehydroepiandrosterone. 705

"Animal" means any nonhuman animate being endowed with the power of voluntary action.

706 "Automated drug dispensing system" means a mechanical or electronic system that performs operations or activities, other than compounding or administration, relating to pharmacy services, 707 708 including the storage, dispensing, or distribution of drugs and the collection, control, and maintenance of 709 all transaction information, to provide security and accountability for such drugs.

"Biological product" means a virus, therapeutic serum, toxin, antitoxin, vaccine, blood, blood 710 711 component or derivative, allergenic product, protein other than a chemically synthesized polypeptide, or analogous product, or arsphenamine or any derivative of arsphenamine or any other trivalent organic 712 713 arsenic compound, applicable to the prevention, treatment, or cure of a disease or condition of human 714 beings.

715 "Biosimilar" means a biological product that is highly similar to a specific reference biological 716 product, notwithstanding minor differences in clinically inactive compounds, such that there are no 717 clinically meaningful differences between the reference biological product and the biological product that has been licensed as a biosimilar pursuant to 42 U.S.C. § 262(k) in terms of safety, purity, and potency 718 719 of the product. 720

"Board" means the Board of Pharmacy.

721 "Bulk drug substance" means any substance that is represented for use, and that, when used in the compounding, manufacturing, processing, or packaging of a drug, becomes an active ingredient or a finished dosage form of the drug; however, "bulk drug substance" shall not include intermediates that 722 723 724 are used in the synthesis of such substances.

"Change of ownership" of an existing entity permitted, registered, or licensed by the Board means (i) 725 the sale or transfer of all or substantially all of the assets of the entity or of any corporation that owns 726 727 or controls the entity; (ii) the creation of a partnership by a sole proprietor, the dissolution of a partnership, or change in partnership composition; (iii) the acquisition or disposal of 50 percent or more 728 729 of the outstanding shares of voting stock of a corporation owning the entity or of the parent corporation 730 of a wholly owned subsidiary owning the entity, except that this shall not apply to any corporation the 731 voting stock of which is actively traded on any securities exchange or in any over-the-counter market; 732 (iv) the merger of a corporation owning the entity or of the parent corporation of a wholly-owned 733 subsidiary owning the entity with another business or corporation; or (v) the expiration or forfeiture of a 734 corporation's charter.

735 'Co-licensed partner" means a person who, with at least one other person, has the right to engage in 736 the manufacturing or marketing of a prescription drug, consistent with state and federal law.

737 "Compounding" means the combining of two or more ingredients to fabricate such ingredients into a 738 single preparation and includes the mixing, assembling, packaging, or labeling of a drug or device (i) by 739 a pharmacist, or within a permitted pharmacy, pursuant to a valid prescription issued for a medicinal or 740 therapeutic purpose in the context of a bona fide practitioner-patient-pharmacist relationship, or in 741 expectation of receiving a valid prescription based on observed historical patterns of prescribing and 742 dispensing; (ii) by a practitioner of medicine, osteopathy, podiatry, dentistry, or veterinary medicine as 743 an incident to his administering or dispensing, if authorized to dispense, a controlled substance in the 744 course of his professional practice; or (iii) for the purpose of, or as incident to, research, teaching, or 745 chemical analysis and not for sale or for dispensing. The mixing, diluting, or reconstituting of a 746 manufacturer's product drugs for the purpose of administration to a patient, when performed by a 747 practitioner of medicine or osteopathy licensed under Chapter 29 (§ 54.1-2900 et seq.), a person 748 supervised by such practitioner pursuant to subdivision A 6 or 19 of § 54.1-2901, or a person supervised 749 by such practitioner or a licensed nurse practitioner or physician assistant pursuant to subdivision A 4 of 750 § 54.1-2901 shall not be considered compounding.

751 "Controlled substance" means a drug, substance, or immediate precursor in Schedules I through VI of 752 this chapter. The term shall not include distilled spirits, wine, malt beverages, or tobacco as those terms 753 are defined or used in Title 3.2 or Title 4.1. The term "controlled substance" includes a controlled 754 substance analog that has been placed into Schedule I or II by the Board pursuant to the regulatory 755 authority in subsection D of § 54.1-3443.

756 "Controlled substance analog" means a substance the chemical structure of which is substantially 757 similar to the chemical structure of a controlled substance in Schedule I or II and either (i) which has a 758 stimulant, depressant, or hallucinogenic effect on the central nervous system that is substantially similar 759 to or greater than the stimulant, depressant, or hallucinogenic effect on the central nervous system of a 760 controlled substance in Schedule I or II or (ii) with respect to a particular person, which such person represents or intends to have a stimulant, depressant, or hallucinogenic effect on the central nervous 761 762 system that is substantially similar to or greater than the stimulant, depressant, or hallucinogenic effect 763 on the central nervous system of a controlled substance in Schedule I or II. "Controlled substance analog" does not include (a) any substance for which there is an approved new drug application as 764 defined under § 505 of the federal Food, Drug, and Cosmetic Act (21 U.S.C. § 355) or that is generally 765 recognized as safe and effective pursuant to §§ 501, 502, and 503 of the federal Food, Drug, and 766 Cosmetic Act (21 U.S.C. §§ 351, 352, and 353) and 21 C.F.R. Part 330; (b) with respect to a particular 767 768 person, any substance for which an exemption is in effect for investigational use for that person under 769 § 505 of the federal Food, Drug, and Cosmetic Act to the extent that the conduct with respect to that 770 substance is pursuant to such exemption; or (c) any substance to the extent not intended for human 771 consumption before such an exemption takes effect with respect to that substance.

"DEA" means the Drug Enforcement Administration, U.S. Department of Justice, or its successoragency.

"Deliver" or "delivery" means the actual, constructive, or attempted transfer of any item regulated by
this chapter, whether or not there exists an agency relationship, including delivery of a Schedule VI
prescription device to an ultimate user or consumer on behalf of a medical equipment supplier by a
manufacturer, nonresident manufacturer, wholesale distributor, nonresident wholesale distributor,
warehouser, nonresident warehouser, third-party logistics provider, or nonresident third-party logistics
provider at the direction of a medical equipment supplier in accordance with § 54.1-3415.1.

780 "Device" means instruments, apparatus, and contrivances, including their components, parts, and
781 accessories, intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in
782 man or animals or to affect the structure or any function of the body of man or animals.

783 "Dialysis care technician" or "dialysis patient care technician" means an individual who is certified
784 by an organization approved by the Board of Health Professions pursuant to Chapter 27.01
785 (§ 54.1-2729.1 et seq.) and who, under the supervision of a licensed physician, nurse practitioner,
786 physician assistant, or a registered nurse, assists in the care of patients undergoing renal dialysis
787 treatments in a Medicare-certified renal dialysis facility.

788 "Dialysis solution" means either the commercially available, unopened, sterile solutions whose
789 purpose is to be instilled into the peritoneal cavity during the medical procedure known as peritoneal
790 dialysis, or commercially available solutions whose purpose is to be used in the performance of
791 hemodialysis not to include any solutions administered to the patient intravenously.

792 "Dispense" means to deliver a drug to an ultimate user or research subject by or pursuant to the 793 lawful order of a practitioner, including the prescribing and administering, packaging, labeling, or 794 compounding necessary to prepare the substance for that delivery. However, dispensing shall not include 795 the transportation of drugs mixed, diluted, or reconstituted in accordance with this chapter to other sites 796 operated by such practitioner or that practitioner's medical practice for the purpose of administration of 797 such drugs to patients of the practitioner or that practitioner's medical practice at such other sites. For

798 practitioners of medicine or osteopathy, "dispense" shall only include the provision of drugs by a 799 practitioner to patients to take with them away from the practitioner's place of practice.

800 "Dispenser" means a practitioner who dispenses.

801 "Distribute" means to deliver other than by administering or dispensing a controlled substance.

802 "Distributor" means a person who distributes.

803 "Drug" means (i) articles or substances recognized in the official United States Pharmacopoeia 804 National Formulary or official Homeopathic Pharmacopoeia of the United States, or any supplement to 805 any of them; (ii) articles or substances intended for use in the diagnosis, cure, mitigation, treatment, or 806 prevention of disease in man or animals; (iii) articles or substances, other than food, intended to affect the structure or any function of the body of man or animals; (iv) articles or substances intended for use 807 as a component of any article specified in clause (i), (ii), or (iii); or (v) a biological product. "Drug" 808 809 does not include devices or their components, parts, or accessories.

810 "Drug product" means a specific drug in dosage form from a known source of manufacture, whether by brand or therapeutically equivalent drug product name. 811

812 "Electronic prescription" means a written prescription that is generated on an electronic application 813 and is transmitted to a pharmacy as an electronic data file; Schedule II through V prescriptions shall be 814 transmitted in accordance with 21 C.F.R. Part 1300.

"Facsimile (FAX) prescription" means a written prescription or order that is transmitted by an 815 816 electronic device over telephone lines that sends the exact image to the receiving pharmacy in hard copy 817 form. 818

"FDA" means the U.S. Food and Drug Administration.

819 "Immediate precursor" means a substance which the Board of Pharmacy has found to be and by 820 regulation designates as being the principal compound commonly used or produced primarily for use, 821 and which is an immediate chemical intermediary used or likely to be used in the manufacture of a 822 controlled substance, the control of which is necessary to prevent, curtail, or limit manufacture.

823 "Interchangeable" means a biosimilar that meets safety standards for determining interchangeability 824 pursuant to 42 U.S.C. § 262(k)(4).

825 "Label" means a display of written, printed, or graphic matter upon the immediate container of any 826 article. A requirement made by or under authority of this chapter that any word, statement, or other information appear on the label shall not be considered to be complied with unless such word, 827 828 statement, or other information also appears on the outside container or wrapper, if any, of the retail 829 package of such article or is easily legible through the outside container or wrapper.

830 "Labeling" means all labels and other written, printed, or graphic matter on an article or any of its 831 containers or wrappers, or accompanying such article.

832 "Manufacture" means the production, preparation, propagation, conversion, or processing of any item 833 regulated by this chapter, either directly or indirectly by extraction from substances of natural origin, or 834 independently by means of chemical synthesis, or by a combination of extraction and chemical synthesis, and includes any packaging or repackaging of the substance or labeling or relabeling of its 835 836 container. This term does not include compounding.

837 "Manufacturer" means every person who manufactures, a manufacturer's co-licensed partner, or a 838 repackager.

839 "Marijuana" means any part of a plant of the genus Cannabis whether growing or not, its seeds, or 840 its resin; and every compound, manufacture, salt, derivative, mixture, or preparation of such plant, its seeds, its resin, or any extract containing one or more cannabinoids. "Marijuana" does not include (i) the 841 842 mature stalks of such plant, fiber produced from such stalk, or oil or cake made from the seeds of such 843 plant, unless such stalks, fiber, oil, or cake is combined with other parts of plants of the genus Cannabis-Marijuana does not include (i); (ii) industrial hemp, as defined in § 3.2-4112, that is possessed by a person registered pursuant to subsection A of § 3.2-4115 or his agent, (iii) industrial hemp, as 844 845 846 defined in § 3.2-4112, that is possessed by a person who holds a hemp producer license issued by the 847 U.S. Department of Agriculture pursuant to 7 C.F.R. Part 990, or (iii); (iv) a hemp product, as defined 848 in § 3.2-4112, containing a tetrahydrocannabinol concentration of no greater than 0.3 percent that is 849 derived from industrial hemp, as defined in § 3.2-4112, that is grown, dealt, or processed in compliance 850 with state or federal law; (v) an industrial hemp extract, as defined in § 3.2-5145.1; or (vi) any 851 substance containing a tetrahydrocannabinol isomer, ester, ether, salt, or salts of such isomer, ester, or 852 ether that has been placed by the Board of Pharmacy into one of the schedules set forth in the Drug Control Act (§ 54.1-3400 et seq.) pursuant to § 54.1-3443. 853

"Medical equipment supplier" means any person, as defined in § 1-230, engaged in the delivery to 854 855 the ultimate consumer, pursuant to the lawful order of a practitioner, of hypodermic syringes and needles, medicinal oxygen, Schedule VI controlled devices, those Schedule VI controlled substances with 856 857 no medicinal properties that are used for the operation and cleaning of medical equipment, solutions for 858 peritoneal dialysis, and sterile water or saline for irrigation.

859 "Narcotic drug" means any of the following, whether produced directly or indirectly by extraction

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from substances of vegetable origin, or independently by means of chemical synthesis, or by a 860 combination of extraction and chemical synthesis: (i) opium, opiates, and any salt, compound, derivative, 861 862 or preparation of opium or opiates; (ii) any salt, compound, isomer, derivative, or preparation thereof which is chemically equivalent or identical with any of the substances referred to in clause (i), but not 863 864 including the isoquinoline alkaloids of opium; (iii) opium poppy and poppy straw; (iv) coca leaves and 865 any salt, compound, derivative, or preparation of coca leaves, and any salt, compound, isomer, 866 derivative, or preparation thereof which is chemically equivalent or identical with any of these 867 substances, but not including decocainized coca leaves or extraction of coca leaves which do not contain 868 cocaine or ecgonine.

869 "New drug" means (i) any drug, except a new animal drug or an animal feed bearing or containing a 870 new animal drug, the composition of which is such that such drug is not generally recognized, among 871 experts qualified by scientific training and experience to evaluate the safety and effectiveness of drugs, 872 as safe and effective for use under the conditions prescribed, recommended, or suggested in the labeling, 873 except that such a drug not so recognized shall not be deemed to be a "new drug" if at any time prior 874 to the enactment of this chapter it was subject to the Food and Drugs Act of June 30, 1906, as 875 amended, and if at such time its labeling contained the same representations concerning the conditions 876 of its use, or (ii) any drug, except a new animal drug or an animal feed bearing or containing a new 877 animal drug, the composition of which is such that such drug, as a result of investigations to determine 878 its safety and effectiveness for use under such conditions, has become so recognized, but which has not, 879 otherwise than in such investigations, been used to a material extent or for a material time under such 880 conditions.

881 "Nuclear medicine technologist" means an individual who holds a current certification with the 882 American Registry of Radiological Technologists or the Nuclear Medicine Technology Certification 883 Board.

884 "Official compendium" means the official United States Pharmacopoeia National Formulary, official 885 Homeopathic Pharmacopoeia of the United States, or any supplement to any of them.

886 "Official written order" means an order written on a form provided for that purpose by the U.S. Drug 887 Enforcement Administration, under any laws of the United States making provision therefor, if such 888 order forms are authorized and required by federal law, and if no such order form is provided then on 889 an official form provided for that purpose by the Board of Pharmacy.

890 "Opiate" means any substance having an addiction-forming or addiction-sustaining liability similar to 891 morphine or being capable of conversion into a drug having such addiction-forming or 892 addiction-sustaining liability. It does not include, unless specifically designated as controlled under 893 Article 4 (§ 54.1-3437 et seq.), the dextrorotatory isomer of 3-methoxy-n-methylmorphinan and its salts 894 (dextromethorphan). It does include its racemic and levorotatory forms.

895 "Opium poppy" means the plant of the species Papaver somniferum L., except the seeds thereof.

896 "Original package" means the unbroken container or wrapping in which any drug or medicine is 897 enclosed together with label and labeling, put up by or for the manufacturer, wholesaler, or distributor 898 for use in the delivery or display of such article.

899 "Outsourcing facility" means a facility that is engaged in the compounding of sterile drugs and is 900 currently registered as an outsourcing facility with the U.S. Secretary of Health and Human Services and 901 that complies with all applicable requirements of federal and state law, including the Federal Food, 902 Drug, and Cosmetic Act.

903 "Person" means both the plural and singular, as the case demands, and includes an individual, 904 partnership, corporation, association, governmental agency, trust, or other institution or entity.

905 "Pharmacist-in-charge" means the person who, being licensed as a pharmacist, signs the application 906 for a pharmacy permit and assumes full legal responsibility for the operation of the relevant pharmacy in a manner complying with the laws and regulations for the practice of pharmacy and the sale and dispensing of controlled substances; the "pharmacist-in-charge" shall personally supervise the pharmacy and the pharmacy's personnel as required by § 54.1-3432. 907 908 909 910

"Poppy straw" means all parts, except the seeds, of the opium poppy, after mowing.

911 "Practitioner" means a physician, dentist, licensed nurse practitioner pursuant to § 54.1-2957.01, 912 licensed physician assistant pursuant to § 54.1-2952.1, pharmacist pursuant to § 54.1-3300, TPA-certified 913 optometrist pursuant to Article 5 (§ 54.1-3222 et seq.) of Chapter 32, veterinarian, scientific investigator, 914 or other person licensed, registered, or otherwise permitted to distribute, dispense, prescribe and 915 administer, or conduct research with respect to a controlled substance in the course of professional 916 practice or research in the Commonwealth.

917 "Prescriber" means a practitioner who is authorized pursuant to §§ 54.1-3303 and 54.1-3408 to issue 918 a prescription.

919 "Prescription" means an order for drugs or medical supplies, written or signed or transmitted by word 920 of mouth, telephone, telegraph, or other means of communication to a pharmacist by a duly licensed 921 physician, dentist, veterinarian, or other practitioner authorized by law to prescribe and administer such 922 drugs or medical supplies.

923 "Prescription drug" means any drug required by federal law or regulation to be dispensed only 924 pursuant to a prescription, including finished dosage forms and active ingredients subject to § 503(b) of 925 the Federal Food, Drug, and Cosmetic Act (21 U.S.C. § 353(b)).

926 "Production" or "produce" includes the manufacture, planting, cultivation, growing, or harvesting of a 927 controlled substance or marijuana.

928 "Proprietary medicine" means a completely compounded nonprescription drug in its unbroken, 929 original package which does not contain any controlled substance or marijuana as defined in this chapter 930 and is not in itself poisonous, and which is sold, offered, promoted, or advertised directly to the general 931 public by or under the authority of the manufacturer or primary distributor, under a trademark, trade 932 name, or other trade symbol privately owned, and the labeling of which conforms to the requirements of 933 this chapter and applicable federal law. However, this definition shall not include a drug that is only 934 advertised or promoted professionally to licensed practitioners, a narcotic or drug containing a narcotic, 935 a drug that may be dispensed only upon prescription or the label of which bears substantially the 936 statement "Warning — may be habit-forming," or a drug intended for injection.

937 "Radiopharmaceutical" means any drug that exhibits spontaneous disintegration of unstable nuclei 938 with the emission of nuclear particles or photons and includes any non-radioactive reagent kit or 939 radionuclide generator that is intended to be used in the preparation of any such substance, but does not 940 include drugs such as carbon-containing compounds or potassium-containing salts that include trace quantities of naturally occurring radionuclides. The term also includes any biological product that is 941 942 labeled with a radionuclide or intended solely to be labeled with a radionuclide.

943 "Reference biological product" means the single biological product licensed pursuant to 42 U.S.C. 944 § 262(a) against which a biological product is evaluated in an application submitted to the U.S. Food 945 and Drug Administration for licensure of biological products as biosimilar or interchangeable pursuant to 946 42 U.S.C. § 262(k).

947 "Sale" includes barter, exchange, or gift, or offer therefor, and each such transaction made by any 948 person, whether as an individual, proprietor, agent, servant, or employee.

949 "Tetrahydrocannabinol" means any naturally occurring or synthetic tetrahydrocannabinol, including 950 its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of 951 isomers is possible within the specific chemical designation and any preparation, mixture, or substance 952 containing, or mixed or infused with, any detectable amount of tetrahydrocannabinol. 953 "Tetrahydrocannabinol" includes delta-6a(10a), delta-7, delta-8, delta-9, and delta-10 954 tetrahydrocannabinol. For the purposes of this definition, "isomer" means the optical, position, and 955 geometric isomers.

956 "Therapeutically equivalent drug products" means drug products that contain the same active 957 ingredients and are identical in strength or concentration, dosage form, and route of administration and that are classified as being therapeutically equivalent by the U.S. Food and Drug Administration 958 959 pursuant to the definition of "therapeutically equivalent drug products" set forth in the most recent 960 edition of the Approved Drug Products with Therapeutic Equivalence Evaluations, otherwise known as 961 the "Orange Book."

"Third-party logistics provider" means a person that provides or coordinates warehousing of or other 962 963 logistics services for a drug or device in interstate commerce on behalf of a manufacturer, wholesale 964 distributor, or dispenser of the drug or device but does not take ownership of the product or have 965 responsibility for directing the sale or disposition of the product.

966 "Total tetrahydrocannabinol" means the sum, after the application of any necessary conversion factor, of the percentage by weight of tetrahydrocannabinol and the percentage by weight of 967 968 tetrahvdrocannabinolic acid. 969

"USP-NF" means the current edition of the United States Pharmacopeia-National Formulary.

970 "Warehouser" means any person, other than a wholesale distributor, manufacturer, or third-party 971 logistics provider, engaged in the business of (i) selling or otherwise distributing prescription drugs or 972 devices to any person who is not the ultimate user or consumer and (ii) delivering Schedule VI 973 prescription devices to the ultimate user or consumer pursuant to § 54.1-3415.1. No person shall be 974 subject to any state or local tax by reason of this definition.

975 Wholesale distribution" means (i) distribution of prescription drugs to persons other than consumers 976 or patients and (ii) delivery of Schedule VI prescription devices to the ultimate user or consumer 977 pursuant to § 54.1-3415.1, subject to the exemptions set forth in the federal Drug Supply Chain Security 978 Act.

979 "Wholesale distributor" means any person other than a manufacturer, a manufacturer's co-licensed 980 partner, a third-party logistics provider, or a repackager that engages in wholesale distribution.

The words "drugs" and "devices" as used in Chapter 33 (§ 54.1-3300 et seq.) and in this chapter 981 982 shall not include surgical or dental instruments, physical therapy equipment, X-ray apparatus, or glasses **983** or lenses for the eyes.

984 The terms "pharmacist," "pharmacy," and "practice of pharmacy" as used in this chapter shall be defined as provided in Chapter 33 (§ 54.1-3300 et seq.) unless the context requires a different meaning.

986 § 54.1-3408.3. Certification for use of cannabis oil for treatment.

987 A. As used in this section:

988 "Botanical cannabis" means cannabis that is composed wholly of usable cannabis from the same parts989 of the same chemovar of cannabis plant.

990 "Cannabis oil" means any formulation of processed Cannabis plant extract, which may include 991 industrial hemp extracts, including isolates and distillates, acquired by a pharmaceutical processor 992 pursuant to § 54.1-3442.6, or a dilution of the resin of the Cannabis plant that contains no more than 10 993 milligrams of delta-9-tetrahydrocannabinol tetrahydrocannabinol per dose. "Cannabis oil" does not 994 include industrial hemp, as defined in § 3.2-4112, that is grown, dealt handled, or processed in 995 compliance with state or federal law, unless it has been grown and processed in the Commonwealth by 996 a registered industrial hemp processor and acquired and formulated by a pharmaceutical processor.

997 "Cannabis product" means a product that is (i) produced by a pharmaceutical processor, registered
998 with the Board, and compliant with testing requirements and (ii) composed of cannabis oil or botanical
999 cannabis.

1000 "Designated caregiver facility" means any hospice or hospice facility licensed pursuant to
1001 § 32.1-162.3, or home care organization as defined in § 32.1-162.7 that provides pharmaceutical services
1002 or home health services, private provider licensed by the Department of Behavioral Health and
1003 Developmental Services pursuant to Article 2 (§ 37.2-403 et seq.) of Chapter 4 of Title 37.2, assisted
1004 living facility licensed pursuant to § 63.2-1701, or adult day care center licensed pursuant to
1005 § 63.2-1701.

1006 "Practitioner" means a practitioner of medicine or osteopathy licensed by the Board of Medicine, a
1007 physician assistant licensed by the Board of Medicine, or a nurse practitioner jointly licensed by the
1008 Board of Medicine and the Board of Nursing.

1009 "Registered agent" means an individual designated by a patient who has been issued a written
1010 certification, or, if such patient is a minor or a vulnerable adult as defined in § 18.2-369, designated by
1011 such patient's parent or legal guardian, and registered with the Board pursuant to subsection G.

1012 "Usable cannabis" means any cannabis plant material, including seeds, but not (i) resin that has been 1013 extracted from any part of the cannabis plant, its seeds, or its resin; (ii) the mature stalks, fiber produced 1014 from the stalks, or any other compound, manufacture, salt, or derivative, mixture, or preparation of the 1015 mature stalks; or (iii) oil or cake made from the seeds of the plant.

1016 B. A practitioner in the course of his professional practice may issue a written certification for the 1017 use of cannabis products for treatment or to alleviate the symptoms of any diagnosed condition or 1018 disease determined by the practitioner to benefit from such use. The practitioner shall use his 1019 professional judgment to determine the manner and frequency of patient care and evaluation and may 1020 employ the use of telemedicine, provided that the use of telemedicine includes the delivery of patient 1021 care through real-time interactive audio-visual technology. If a practitioner determines it is consistent 1022 with the standard of care to dispense botanical cannabis to a minor, the written certification shall 1023 specifically authorize such dispensing. If not specifically included on the initial written certification, 1024 authorization for botanical cannabis may be communicated verbally or in writing to the pharmacist at 1025 the time of dispensing.

1026 C. The written certification shall be on a form provided by the Board of Pharmacy. Such written 1027 certification shall contain the name, address, and telephone number of the practitioner; the name and 1028 address of the patient issued the written certification; the date on which the written certification was 1029 made; and the signature or authentic electronic signature of the practitioner. Such written certification 1030 issued pursuant to subsection B shall expire no later than one year after its issuance unless the 1031 practitioner provides in such written certification an earlier expiration. A written certification shall not be 1032 issued to a patient by more than one practitioner during any given time period.

1033 D. No practitioner shall be prosecuted under § 18.2-248 or 18.2-248.1 for the issuance of a 1034 certification for the use of cannabis products for the treatment or to alleviate the symptoms of a patient's 1035 diagnosed condition or disease pursuant to a written certification issued pursuant to subsection B. 1036 Nothing in this section shall preclude the Board of Medicine from sanctioning a practitioner for failing 1037 to properly evaluate or treat a patient's medical condition or otherwise violating the applicable standard 1038 of care for evaluating or treating medical conditions.

E. A practitioner who issues a written certification to a patient pursuant to this section shall register with the Board and shall hold sufficient education and training to exercise appropriate professional judgment in the certification of patients. The Board shall not limit the number of patients to whom a practitioner may issue a written certification. The Board may report information to the applicable licensing board on unusual patterns of certifications issued by a practitioner.

1044 F. No patient shall be required to physically present the written certification after the initial 1045 dispensing by any pharmaceutical processor or cannabis dispensing facility under each written 1046 certification, provided that the pharmaceutical processor or cannabis dispensing facility maintains an 1047 electronic copy of the written certification. Pharmaceutical processors and cannabis dispensing facilities 1048 shall electronically transmit, on a monthly basis, all new written certifications received by the 1049 pharmaceutical processor or cannabis dispensing facility to the Board.

1050 G. A patient, or, if such patient is a minor or a vulnerable adult as defined in § 18.2-369, such 1051 patient's parent or legal guardian, may designate an individual to act as his registered agent for the 1052 purposes of receiving cannabis products pursuant to a valid written certification. Such designated individual shall register with the Board. The Board may set a limit on the number of patients for whom 1053 1054 any individual is authorized to act as a registered agent.

H. Upon delivery of a cannabis product by a pharmaceutical processor or cannabis dispensing facility 1055 1056 to a designated caregiver facility, any employee or contractor of a designated caregiver facility, who is licensed or registered by a health regulatory board and who is authorized to possess, distribute, or 1057 1058 administer medications, may accept delivery of the cannabis product on behalf of a patient or resident 1059 for subsequent delivery to the patient or resident and may assist in the administration of the cannabis 1060 product to the patient or resident as necessary.

I. Information obtained under the registration process shall be confidential and shall not be subject to 1061 1062 the disclosure provisions of the Virginia Freedom of Information Act (§ 2.2-3700 et seq.). However, 1063 reasonable access to registry information shall be provided to (i) the Chairmen of the House Committee 1064 for Courts of Justice and the Senate Committee on the Judiciary, (ii) state and federal agencies or local law enforcement for the purpose of investigating or prosecuting a specific individual for a specific 1065 violation of law, (iii) licensed practitioners or pharmacists, or their agents, for the purpose of providing 1066 patient care and drug therapy management and monitoring of drugs obtained by a patient, (iv) a 1067 1068 pharmaceutical processor or cannabis dispensing facility involved in the treatment of a patient, or (v) a 1069 registered agent, but only with respect to information related to such patient.

1070 § 54.1-3423. Board to issue registration unless inconsistent with public interest; authorization to 1071 conduct research; application and fees.

1072 A. The Board shall register an applicant to manufacture or distribute controlled substances included 1073 in Schedules I through V unless it determines that the issuance of that registration would be inconsistent 1074 with the public interest. In determining the public interest, the Board shall consider the following 1075 factors:

1076 1. Maintenance of effective controls against diversion of controlled substances into other than 1077 legitimate medical, scientific, or industrial channels; 1078

2. Compliance with applicable state and local law;

3. Any convictions of the applicant under any federal and state laws relating to any controlled 1079 1080 substance;

1081 4. Past experience in the manufacture or distribution of controlled substances, and the existence in 1082 the applicant's establishment of effective controls against diversion;

1083 5. Furnishing by the applicant of false or fraudulent material in any application filed under this 1084 chapter:

6. Suspension or revocation of the applicant's federal registration to manufacture, distribute, or 1085 1086 dispense controlled substances as authorized by federal law; and 1087

7. Any other factors relevant to and consistent with the public health and safety.

1088 B. Registration under subsection A does not entitle a registrant to manufacture and distribute 1089 controlled substances in Schedule I or II other than those specified in the registration.

1090 C. Practitioners must be registered to conduct research or laboratory analysis with controlled substances in Schedules II through VI- tetrahydrocannabinol, or marijuana. Practitioners registered under 1091 1092 federal law to conduct research with Schedule I substances, other than tetrahydrocannabinol marijuana, 1093 may conduct research with Schedule I substances within this the Commonwealth upon furnishing the 1094 evidence of that federal registration.

1095 D. The Board may register other persons or entities to possess controlled substances listed on 1096 Schedules II through VI upon a determination that (i) there is a documented need, (ii) the issuance of 1097 the registration is consistent with the public interest, (iii) the possession and subsequent use of the 1098 controlled substances complies with applicable state and federal laws and regulations, and (iv) the 1099 subsequent storage, use, and recordkeeping of the controlled substances will be under the general 1100 supervision of a licensed pharmacist, practitioner of medicine, osteopathy, podiatry, dentistry, or veterinary medicine as specified in the Board's regulations. The Board shall consider, at a minimum, the 1101 factors listed in subsection A of this section in determining whether the registration shall be issued. 1102 Notwithstanding the exceptions listed in § 54.1-3422 A, the Board may mandate a controlled substances 1103 1104 registration for sites maintaining certain types and quantities of Schedules II through VI controlled substances as it may specify in its regulations. The Board shall promulgate regulations related to 1105

1106 requirements or criteria for the issuance of such controlled substances registration, storage, security, 1107 supervision, and recordkeeping.

1108 E. The Board may register a public or private animal shelter as defined in § 3.2-6500 to purchase, 1109 possess, and administer certain Schedule II through VI controlled substances approved by the State 1110 Veterinarian for the purpose of euthanizing injured, sick, homeless, and unwanted domestic pets and 1111 animals and to purchase, possess, and administer certain Schedule VI drugs and biological products for 1112 the purpose of preventing, controlling, and treating certain communicable diseases that failure to control 1113 would result in transmission to the animal population in the shelter. Controlled substances used for 1114 euthanasia shall be administered only in accordance with protocols established by the State Veterinarian 1115 and only by persons trained in accordance with instructions by the State Veterinarian. The list of 1116 Schedule VI drugs and biological products used for treatment and prevention of communicable diseases 1117 within the shelter shall be determined by the supervising veterinarian of the shelter and the drugs and 1118 biological products shall be administered only pursuant to written protocols established or approved by the supervising veterinarian of the shelter and only by persons who have been trained in accordance 1119 1120 with instructions established or approved by the supervising veterinarian. The shelter shall maintain a 1121 copy of the approved list of drugs and biological products, written protocols for administering, and 1122 training records of those persons administering drugs and biological products on the premises of the 1123 shelter.

1124 F. The Board may register a crisis stabilization unit established pursuant to § 37.2-500 or 37.2-601 1125 and licensed by the Department of Behavioral Health and Developmental Services to maintain a stock of 1126 Schedule VI controlled substances necessary for immediate treatment of patients admitted to the crisis 1127 stabilization unit, which may be accessed and administered by a nurse pursuant to a written or oral order 1128 of a prescriber in the absence of a prescriber. Schedule II through Schedule V controlled substances 1129 shall only be maintained if so authorized by federal law and Board regulations.

1130 G. The Board may register an entity at which a patient is treated by the use of instrumentation and 1131 diagnostic equipment through which images and medical records may be transmitted electronically for 1132 the purpose of establishing a bona fide practitioner-patient relationship and is prescribed Schedule II 1133 through VI controlled substances when such prescribing is in compliance with federal requirements for 1134 the practice of telemedicine and the patient is not in the physical presence of a practitioner registered 1135 with the U.S. Drug Enforcement Administration. In determining whether the registration shall be issued, 1136 the Board shall consider (i) the factors listed in subsection A, (ii) whether there is a documented need 1137 for such registration, and (iii) whether the issuance of the registration is consistent with the public 1138 interest.

1139 H. Applications for controlled substances registration certificates and renewals thereof shall be made 1140 on a form prescribed by the Board and such applications shall be accompanied by a fee in an amount to 1141 be determined by the Board.

1142 I. Upon (i) any change in ownership or control of a business, (ii) any change of location of the 1143 controlled substances stock, (iii) the termination of authority by or of the person named as the 1144 responsible party on a controlled substances registration, or (iv) a change in the supervising practitioner, 1145 if applicable, the registrant or responsible party shall immediately surrender the registration. The 1146 registrant shall, within 14 days following surrender of a registration, file a new application and, if 1147 applicable, name the new responsible party or supervising practitioner. 1148

§ 54.1-3442.6. Permit to operate pharmaceutical processor or cannabis dispensing facility.

1149 A. No person shall operate a pharmaceutical processor or a cannabis dispensing facility without first 1150 obtaining a permit from the Board. The application for such permit shall be made on a form provided 1151 by the Board and signed by a pharmacist who will be in full and actual charge of the pharmaceutical 1152 processor's dispensing area or cannabis dispensing facility. The Board shall establish an application fee 1153 and other general requirements for such application.

1154 B. Each permit shall expire annually on a date determined by the Board in regulation. The number of 1155 permits that the Board may issue or renew in any year is limited to one pharmaceutical processor and 1156 up to five cannabis dispensing facilities for each health service area established by the Board of Health. 1157 Permits shall be displayed in a conspicuous place on the premises of the pharmaceutical processor and 1158 cannabis dispensing facility.

1159 C. The Board shall adopt regulations establishing health, safety, and security requirements for 1160 pharmaceutical processors and cannabis dispensing facilities. Such regulations shall include requirements 1161 for (i) physical standards; (ii) location restrictions; (iii) security systems and controls; (iv) minimum 1162 equipment and resources; (v) recordkeeping; (vi) labeling, including the potency of each botanical 1163 cannabis product and the amounts recommended by the practitioner or dispensing pharmacist, and 1164 packaging; (vii) routine inspections no more frequently than once annually; (viii) processes for safely 1165 and securely dispensing and delivering in person cannabis products to a patient, his registered agent, or, if such patient is a minor or a vulnerable adult as defined in § 18.2-369, such patient's parent or legal 1166

1228

1167 guardian; (ix) dosage limitations for cannabis oil that provide that each dispensed dose of cannabis oil 1168 not exceed 10 milligrams of delta-9-tetrahydrocannabinol tetrahydrocannabinol; (x) a process for the 1169 wholesale distribution of and the transfer of usable cannabis, botanical cannabis, cannabis oil, and 1170 cannabis products between pharmaceutical processors, between a pharmaceutical processors and a cannabis dispensing facility, and between cannabis dispensing facilities; (xi) an allowance for the sale of 1171 1172 devices for administration of dispensed cannabis products and hemp-based CBD products that meet the 1173 applicable standards set forth in state and federal law, including the laboratory testing standards set forth 1174 in subsection M; (xii) an allowance for the use and distribution of inert product samples containing no cannabinoids for patient demonstration exclusively at the pharmaceutical processor or cannabis 1175 1176 dispensing facility, and not for further distribution or sale, without the need for a written certification; 1177 (xiii) a process for acquiring industrial hemp extracts and formulating such extracts into cannabis 1178 products; and (xiv) an allowance for the advertising and promotion of the pharmaceutical processor's 1179 products and operations, which shall not limit the pharmaceutical processor from the provision of 1180 educational material to practitioners who issue written certifications and patients. The Board shall also 1181 adopt regulations for pharmaceutical processors that include requirements for (a) processes for safely and 1182 securely cultivating Cannabis plants intended for producing cannabis products, (b) the secure disposal of 1183 agricultural waste, and (c) a process for registering cannabis oil products.

1184 D. The Board shall require that, after processing and before dispensing any cannabis products, a 1185 pharmaceutical processor shall make a sample available from each batch of cannabis product for testing 1186 by an independent laboratory located in Virginia meeting Board requirements. A valid sample size for 1187 testing shall be determined by each laboratory and may vary due to sample matrix, analytical method, and laboratory-specific procedures. A minimum sample size of 0.5 percent of individual units for 1188 dispensing or distribution from each homogenized batch of cannabis oil is required to achieve a 1189 representative cannabis oil sample for analysis. A minimum sample size, to be determined by the 1190 1191 certified testing laboratory, from each batch of botanical cannabis is required to achieve a representative 1192 botanical cannabis sample for analysis. Botanical cannabis products shall only be tested for the 1193 following: total cannabidiol (CBD); total tetrahydrocannabinol (THC); terpenes; pesticide chemical 1194 residue; heavy metals; mycotoxins; moisture; and microbiological contaminants. Testing thresholds shall 1195 be consistent with generally accepted cannabis industry thresholds. The pharmaceutical processor may 1196 remediate botanical cannabis or cannabis oil that fails any quality testing standard except pesticides. 1197 Following remediation, all remediated botanical cannabis or cannabis oil shall be subject to laboratory 1198 testing and approved upon satisfaction of applicable testing standards, which shall not be more stringent 1199 than initial testing prior to remediation. If a batch of botanical cannabis fails retesting after remediation, 1200 it shall be considered usable cannabis and may be processed into cannabis oil. Stability testing shall not 1201 be required for any cannabis product with an expiration date assigned by the pharmaceutical processor 1202 of six months or less from the date of the cannabis product registration approval. Stability testing 1203 required for assignment of an expiration date longer than six months shall be limited to microbial 1204 testing, on a pass/fail basis, and potency testing, on a 10 percent deviation basis, of active ingredients.

E. A laboratory testing samples for a pharmaceutical processor shall obtain a controlled substances
 registration certificate pursuant to § 54.1-3423 and shall comply with quality standards established by the
 Board in regulation.

F. Every pharmaceutical processor's dispensing area or cannabis dispensing facility shall be under the personal supervision of a licensed pharmacist on the premises of the pharmaceutical processor or cannabis dispensing facility. The pharmaceutical processor shall ensure that security measures are adequate to protect the cannabis from diversion at all times, and the pharmacist-in-charge shall have concurrent responsibility for preventing diversion from the dispensing area.

Every pharmaceutical processor shall designate a person who shall have oversight of the cultivation
and production areas of the pharmaceutical processor and shall provide such information to the Board.
The Board shall direct all communications related to enforcement of requirements related to cultivation
and production of cannabis oil products by the pharmaceutical processor to such designated person.

1217 G. The Board shall require the material owners of an applicant for a pharmaceutical processor or 1218 cannabis dispensing facility permit to submit to fingerprinting and provide personal descriptive information to be forwarded along with his fingerprints through the Central Criminal Records Exchange 1219 1220 to the Federal Bureau of Investigation for the purpose of obtaining criminal history record information 1221 regarding the applicant's material owners. The cost of fingerprinting and the criminal history record 1222 search shall be paid by the applicant. The Central Criminal Records Exchange shall forward the results 1223 of the criminal history background check to the Board or its designee, which shall be a governmental 1224 entity. A pharmaceutical processor shall maintain evidence of criminal background checks for all 1225 employees and delivery agents of the pharmaceutical processor. Criminal background checks of 1226 employees and delivery agents may be conducted by any service sufficient to disclose any federal and 1227 state criminal convictions.

H. In addition to other employees authorized by the Board, a pharmaceutical processor may employ

individuals who may have less than two years of experience (i) to perform cultivation-related duties
under the supervision of an individual who has received a degree in a field related to the cultivation of
plants or a certification recognized by the Board or who has at least two years of experience cultivating
plants, (ii) to perform extraction-related duties under the supervision of an individual who has a degree
in chemistry or pharmacology or at least two years of experience extracting chemicals from plants, and
(iii) to perform duties at the pharmaceutical processor and cannabis dispensing facility upon certification
as a pharmacy technician.

1236 I. A pharmaceutical processor to whom a permit has been issued by the Board may establish up to
1237 five cannabis dispensing facilities for the dispensing of cannabis products that have been cultivated and
1238 produced on the premises of a pharmaceutical processor permitted by the Board. Each cannabis
1239 dispensing facility shall be located within the same health service area as the pharmaceutical processor.

1240 J. No person who has been convicted of a felony under the laws of the Commonwealth or another 1241 jurisdiction within the last five years shall be employed by or act as an agent of a pharmaceutical 1242 processor or cannabis dispensing facility.

1243 K. Every pharmaceutical processor or cannabis dispensing facility shall adopt policies for 1244 pre-employment drug screening and regular, ongoing, random drug screening of employees.

L. A pharmacist at the pharmaceutical processor's dispensing area and the cannabis dispensing facility shall determine the number of pharmacy interns, pharmacy technicians, and pharmacy technician trainees who can be safely and competently supervised at one time; however, no pharmacist shall supervise more than six persons performing the duties of a pharmacy technician at one time in the pharmaceutical processor's dispensing area or cannabis dispensing facility.

M. A pharmaceutical processor may acquire industrial hemp extracts grown and processed in 1250 1251 Virginia, and in compliance with state or federal law, from a registered industrial hemp dealer or 1252 processor. A pharmaceutical processor may process and formulate such extracts into an allowable dosage 1253 of cannabis product. Industrial hemp extracts acquired and formulated by a pharmaceutical processor are 1254 subject to the same third-party testing requirements that may apply to cannabis plant extract. Testing 1255 shall be performed by a laboratory located in Virginia and in compliance with state law governing the 1256 testing of cannabis products. The industrial hemp dealer or processor shall provide such third-party 1257 testing results to the pharmaceutical processor before industrial hemp extracts may be acquired.

1258 N. With the exception of § 2.2-4031, neither the provisions of the Administrative Process Act 1259 (§ 2.2-4000 et seq.) nor public participation guidelines adopted pursuant thereto shall apply to the 1260 adoption of any regulation pursuant to this section. Prior to adopting any regulation pursuant to this 1261 section, the Board of Pharmacy shall publish a notice of opportunity to comment in the Virginia 1262 Register of Regulations and post the action on the Virginia Regulatory Town Hall. Such notice of 1263 opportunity to comment shall contain (i) a summary of the proposed regulation; (ii) the text of the 1264 proposed regulation; and (iii) the name, address, and telephone number of the agency contact person 1265 responsible for receiving public comments. Such notice shall be made at least 60 days in advance of the last date prescribed in such notice for submittals of public comment. The legislative review provisions of 1266 subsections A and B of § 2.2-4014 shall apply to the promulgation or final adoption process for regulations pursuant to this section. The Board of Pharmacy shall consider and keep on file all public 1267 1268 1269 comments received for any regulation adopted pursuant to this section. 1270

O. The Board shall register all cannabis products that meet testing, labeling, and packaging standards. § 54.1-3442.7. Dispensing cannabis products; report.

1271 A. A pharmaceutical processor or cannabis dispensing facility shall dispense or deliver cannabis 1272 1273 products only in person to (i) a patient who is a Virginia resident or temporarily resides in Virginia and 1274 has been issued a valid written certification; (ii) such patient's registered agent; or (iii) if such patient is 1275 a minor or a vulnerable adult as defined in § 18.2-369, such patient's parent or legal guardian who is a 1276 Virginia resident or temporarily resides in Virginia. A companion may accompany a patient into a 1277 pharmaceutical processor's dispensing area or cannabis dispensing facility. Prior to the initial dispensing 1278 of cannabis products pursuant to each written certification, a pharmacist or pharmacy technician 1279 employed by the pharmaceutical processor or cannabis dispensing facility shall make and maintain, on 1280 site or remotely by electronic means, for two years a paper or electronic copy of the written certification 1281 that provides an exact image of the document that is clearly legible; shall view, in person or by 1282 audiovisual means, a current photo identification of the patient, registered agent, parent, or legal 1283 guardian; and shall verify current board registration of the practitioner and the corresponding registered 1284 agent if applicable. Thereafter, an initial dispensing may be delivered to the patient, registered agent, 1285 parent, legal guardian, or designated caregiver facility. Prior to any subsequent dispensing of cannabis 1286 products pursuant to each written certification, an employee or delivery agent shall view a current photo 1287 identification of the patient, registered agent, parent, or legal guardian and the current board registration 1288 issued to the registered agent if applicable. No pharmaceutical processor or cannabis dispensing facility shall dispense more than a 90-day supply, as determined by the dispensing pharmacist or certifying 1289

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1290 practitioner, for any patient during any 90-day period. A pharmaceutical processor or cannabis 1291 dispensing facility may dispense less than a 90-day supply of a cannabis product for any patient during 1292 any 90-day period; however, a pharmaceutical processor or cannabis dispensing facility may dispense 1293 more than one cannabis product to a patient at one time. No more than four ounces of botanical 1294 cannabis shall be dispensed for each 30-day period for which botanical cannabis is dispensed. The Board 1295 shall establish in regulation an amount of cannabis oil that constitutes a 90-day supply to treat or 1296 alleviate the symptoms of a patient's diagnosed condition or disease. In determining the appropriate 1297 amount of a cannabis product to be dispensed to a patient, a pharmaceutical processor or cannabis 1298 dispensing facility shall consider all cannabis products dispensed to the patient and adjust the amount 1299 dispensed accordingly.

1300 B. A pharmaceutical processor or cannabis dispensing facility shall dispense only cannabis products produced on the premises of a pharmaceutical processor permitted by the Board or cannabis products 1301 1302 that have been formulated with extracts from industrial hemp acquired by a pharmaceutical processor 1303 from a registered industrial hemp dealer or processor pursuant to § 54.1-3442.6. A pharmaceutical 1304 processor may begin cultivation upon being issued a permit by the Board.

1305 C. The Board shall report annually by December 1 to the Chairmen of the House Committee for 1306 Health, Welfare and Institutions and the Senate Committee on Education and Health on the operation of 1307 pharmaceutical processors and cannabis dispensing facilities issued a permit by the Board.

1308 D. The concentration of delta-9-tetrahydrocannabinol tetrahydrocannabinol in any cannabis product 1309 on site may be up to 10 percent greater than or less than the level of delta 9-tetrahydrocannabinol 1310 tetrahydrocannabinol measured for labeling. A pharmaceutical processor and cannabis dispensing facility shall ensure that such concentration in any cannabis product on site is within such range. A 1311 1312 pharmaceutical processor producing cannabis products shall establish a stability testing schedule of 1313 cannabis products. 1314

§ 54.1-3443. Board to administer article.

1315 A. The Board shall administer this article and may add substances to or deschedule or reschedule all substances enumerated in the schedules in this article pursuant to the procedures of the Administrative 1316 1317 Process Act (§ 2.2-4000 et seq.). In making a determination regarding a substance, the Board shall 1318 consider the following: 1319

- 1. The actual or relative potential for abuse;
- 1320 2. The scientific evidence of its pharmacological effect, if known;
- 1321 3. The state of current scientific knowledge regarding the substance;
- 1322 4. The history and current pattern of abuse;
- 1323 5. The scope, duration, and significance of abuse;
- 1324 6. The risk to the public health; 1325
 - 7. The potential of the substance to produce psychic or physical dependence; and

1326 8. Whether the substance is an immediate precursor of a substance already controlled under this 1327 article.

1328 B. After considering the factors enumerated in subsection A, the Board shall make findings and issue 1329 a regulation controlling the substance if it finds the substance has a potential for abuse.

1330 C. If the Board designates a substance as an immediate precursor, substances which are precursors of 1331 the controlled precursor shall not be subject to control solely because they are precursors of the 1332 controlled precursor.

1333 D. If the Board, in consultation with the Department of Forensic Science, determines the substance 1334 shall be placed into Schedule I or II pursuant to § 54.1-3445 or 54.1-3447, the Board may amend its regulations pursuant to Article 2 (§ 2.2-4006 et seq.) of the Administrative Process Act. Prior to making 1335 1336 such amendments, the Board shall conduct a public hearing. At least 30 days prior to conducting such hearing, it shall post notice of the hearing on the Virginia Regulatory Town Hall and shall send notice 1337 1338 of the hearing to any persons requesting to be notified of a regulatory action. In the notice, the Board 1339 shall include a list of all substances it intends to schedule by regulation. The Board shall notify the 1340 House Committee for Courts of Justice and the Senate Committee on the Judiciary of any new substance 1341 added to Schedule I or II pursuant to this subsection. Any substance added to Schedule I or II pursuant 1342 to this subsection shall remain on Schedule I or II for a period of 18 months. Upon expiration of such 1343 18-month period, such substance shall be descheduled unless a general law is enacted adding such 1344 substance to Schedule I or II. Nothing in this subsection shall preclude the Board from adding 1345 substances to or descheduling or rescheduling all substances enumerated in the schedules pursuant to the 1346 provisions of subsections A, B, and E.

1347 E. If any substance is designated, rescheduled, or descheduled as a controlled substance under federal 1348 law and notice of such action is given to the Board, the Board may similarly control the substance under 1349 this chapter after the expiration of 30 days from publication in the Federal Register of a final or interim 1350 final order or rule designating a substance as a controlled substance or rescheduling or descheduling a 1351 substance by amending its regulations in accordance with the requirements of Article 2 (§ 2.2-4006 et 1352 seq.) of the Administrative Process Act. Prior to making such amendments, the Board shall post notice 1353 of the hearing on the Virginia Regulatory Town Hall and shall send notice of the hearing to any persons 1354 requesting to be notified of a regulatory action. The Board shall include a list of all substances it intends 1355 to schedule by regulation in such notice.

1356 F. Authority to control under this section does not extend to distilled spirits, wine, malt beverages, or 1357 tobacco as those terms are defined or used in Title 4.1.

1358 G. The Board shall exempt any nonnarcotic substance from a schedule if such substance may, under 1359 the provisions of the federal Food, Drug, and Cosmetic Act (21 U.S.C. § 301 et seq.) or state law, be 1360 lawfully sold over the counter without a prescription.

1361 H. Any tetrahydrocannabinol isomer, ester, ether, salt, or salts of such isomer, ester, or ether 1362 scheduled pursuant to this section shall not be included in the definition of marijuana set forth in 1363 § 4.1-600, 18.2-247, or 54.1-3401. 1364

§ 54.1-3446. Schedule I.

The controlled substances listed in this section are included in Schedule I:

1366 1. Any of the following opiates, including their isomers, esters, ethers, salts, and salts of isomers, 1367 esters, and ethers, unless specifically excepted, whenever the existence of these isomers, esters, ethers 1368 and salts is possible within the specific chemical designation:

1369 1-{1-[1-[4-bromophenyl]ethyl]-4-piperidinyl}-1,3-dihydro-2H-benzimidazol-2-one (other name: 1370 Brorphine);

- 1371 1-[2-methyl-4-(3-phenyl-2-propen-1-yl)-1-piperazinyl]-1-butanone (other name: 2-methyl AP-237);
- 1372 1-(2-phenylethyl)-4-phenyl-4-acetyloxypiperidine (other name: PEPAP);
- 1373 1-methyl-4-phenyl-4-propionoxypiperidine (other name: MPPP);
- 1374 2-[(4-methoxyphenyl)methyl]-N,N-diethyl-5-nitro-1H-benzimidazole-1-ethanamine (other name: 1375 Metonitazene);
- 1376 2-methoxy-N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-acetamide (other name: Methoxyacetyl 1377 fentanyl);
- 1378 3,4-dichloro-N-[2-(dimethylamino)cyclohexyl]-N-methyl-benzamide (other name: U-47700);
- 1379 3,4-dichloro-N-{[1-(dimethylamino)cyclohexyl]methyl}benzamide (other name: AH-7921);
- 1380 Acetyl fentanyl (other name: desmethyl fentanyl);
- 1381 Acetylmethadol;
- 1382 Allylprodine;

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1383 Alphacetylmethadol (except levo-alphacetylmethadol, also known as levo-alpha-acetylmethadol, 1384 levomethadyl acetate, or LAAM);

- 1385 Alphameprodine;
- 1386 Alphamethadol;
- 1387 Benzethidine;
- 1388 Betacetylmethadol;
- 1389 Betameprodine;
- 1390 Betamethadol;
- 1391 Betaprodine;
- 1392 Clonitazene;
- 1393 Dextromoramide:
- 1394 Diampromide;
- 1395 Diethylthiambutene;
- 1396 Difenoxin;
- 1397 Dimenoxadol:
- 1398 Dimepheptanol;
- 1399 Dimethylthiambutene;
- 1400 Dioxaphetylbutyrate;
- 1401 Dipipanone;
- 1402 Ethylmethylthiambutene:
- 1403 Etonitazene;
- 1404 Etoxeridine;
- 1405 Furethidine;
- 1406 Hydroxypethidine;
- 1407 Ketobemidone;
- 1408 Levomoramide: 1409
- Levophenacylmorphan;
- 1410 Morpheridine;
- 1411 MT-45 (1-cvclohexyl-4-(1,2-diphenylethyl)piperazine);
- 1412 N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopropanecarboxamide (other name: Cyclopropyl fentanyl);

1413	N-(1-phenethylpiperidin-4-yl)-N-phenyltetrahydrofuran-2-carboxamide (other name: Tetrahydrofuranyl
1414	fentanyl);
1415 1416	N-[1-[1-methyl-2-(2-thienyl)ethyl]-4-piperidyl]-N-phenylpropanamide (other name: alpha-methylthiofentanyl);
1417	N-[1-(1-methyl-2-phenylethyl)-4-piperidyl]-N-phenylacetamide (other name:
1418	acetyl-alpha-methylfentanyl);
1419	N-{1-[2-hydroxy-2-(2-thienyl)ethyl]-4-piperidinyl}-N-phenylpropanamide (other name:
1420	beta-hydroxythiofentanyl);
1421	N-[1-(2-hydroxy-2-phenyl)ethyl-4-piperidyl]-N-phenylpropanamide (other name:
1422	beta-hydroxyfentanyl);
1423	N-[1-(alpha-methyl-beta-phenyl)ethyl-4-piperidyl]propionanilide (other names:
1424	1-(1-methyl-2-phenylethyl)-4-(N-propanilido) piperidine, alpha-methylfentanyl);
1425	N-(2-fluorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-propanamide (other names: 2-fluorofentanyl, ortho-fluorofentanyl);
1426 1427	N-(3-fluorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-propanamide (other name: 3-fluorofentanyl);
1428	N-[3-methyl-1-(2-hydroxy-2-phenylethyl)4-piperidyl]-N-phenylpropanamide (other name:
1429	beta-hydroxy-3-methylfentanyl);
1430	N-[3-methyl-1-(2-phenylethyl)-4-piperidyl]-N-phenylpropanamide (other name: 3-methylfentanyl);
1431	N-[3-methyl-1-(2-thienyl)ethyl-4-piperidinyl]-N-phenylpropanamide (other name:
1432	3-methylthiofentanyl);
1433	N-(4-chlorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-propanamide (other names:
1434	para-chlorofentanyl, 4-chlorofentanyl);
1435 1436	N-(4-fluorophenyl)-2-methyl-N-[1-(2-phenylethyl)-4-piperidinyl]-propanamide (other name:
1430 1437	para-fluoroisobutyryl fentanyl); N-(4-fluorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-butanamide (other name:
1438	para-fluorobutyrylfentanyl);
1439	N-(4-fluorophenyl)-N-1-(2-phenylethyl)-4-piperidinyl]-propanamide (other name: para-fluorofentanyl);
1440	N,N-diethyl-2-(2-(4-isopropoxybenzyl)-5-nitro-1H-benzimidazol-1-yl)ethan-1-amine (other name:
1441	Isotonitazene);
1442	N,N-diethyl-2-{[(4-ethoxyphenyl) methyl]-1H-benzimidazol-1-yl}-ethan-1-amine (other names:
1443	Etazene, Desnitroetonitazene);
1444 1445	N,N-diethyl-2-[(4-methoxyphenyl)methyl]-1H-benzimidazole-1-ethanamine (other name: Metodesnitazene);
1446	N-phenyl-N-[1-(2-phenylmethyl)-4-piperidinyl]-2-furancarboxamide (other name: N-benzyl Furanyl
1447	norfentanyl);
1448	N-phenyl-N-(4-piperidinyl)-propanamide (other name: Norfentanyl);
1449	Noracymethadol;
1450	Norlevorphanol;
1451 1452	Normethadone;
1452 1453	Norpipanone; N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-2-furancarboxamide (other name: Furanyl fentanyl);
1454	N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-2-propenamide (other name: Acryl fentanyl);
1455	N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-butanamide (other name: butyryl fentanyl);
1456	N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-pentanamide (other name: Pentanoyl fentanyl);
1457	N-phenyl-N-[1-(2-thienyl)ethyl-4-piperidinyl]-propanamide (other name: thiofentanyl);
1458	Phenadoxone;
1459	Phenampromide;
1460 1461	Phenomorphan; Phenomorphical
1401	Phenoperidine; Piritramide;
1463	Proheptazine;
1464	Properidine;
1465	Propiram;
1466	Racemoramide;
1467	Tilidine;
1468	Trimeperidine;
1469 1470	N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-1,3-benzodioxole-5-carboxamide (other name:
1470 1471	Benzodioxole fentanyl); 3,4-dichloro-N-[2-(diethylamino)cyclohexyl]-N-methylbenzamide (other name: U-49900);
14/1	2-(2,4-dichlorophenyl)-N-[2-(dimethylamino)cyclohexyl]-N-methyl acetamide (other name: U-48800);
14/2	2-(2,4-dichorophenyl)-N-[2-(dimethylamino)cyclohesyl] N-methyl acatamide (other name, U-51754).

1473 2-(3,4-dichlorophenyl)-N-[2-(dimethylamino)cyclohexyl]-N-methyl acetamide (other name: U-51754);
1474 N-(2-fluorophenyl)-2-methoxy-N-[1-(2-phenylethyl)-4-piperidinyl]-acetamide (other name: Ocfentanil);

names:

- 1475 N-(4-methoxyphenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-butanamide (other name:
 1476 4-methoxybutyrylfentanyl);
- 1477 N-phenyl-2-methyl-N-[1-(2-phenylethyl)-4-piperidinyl]-propanamide (other name: Isobutyryl fentanyl);
- 1478 N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-cyclopentanecarboxamide (other name: Cyclopentyl 1479 fentanyl);
- **1480** N-phenyl-N-(1-methyl-4-piperidinyl)-propanamide (other name: N-methyl norfentanyl);
- 1481 N-[2-(dimethylamino)cyclohexyl]-N-methyl-1,3-benzodioxole-5-carboxamide (other names:
 1482 3,4-methylenedioxy U-47700 or 3,4-MDO-U-47700);
- **1483** N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-2-butenamide (other name: Crotonyl fentanyl);
- 1484 N-phenyl-N-[4-phenyl-1-(2-phenylethyl)-4-piperidinyl]-propanamide (other name: 4-phenylfentanyl);
- **1485** N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-benzamide (other names: Phenyl fentanyl, Benzoyl **1486** fentanyl);
- **1487** N-[2-(dimethylamino)cyclohexyl]-N-phenylfuran-2-carboxamide (other name: Furanyl UF-17);
- **1488** N-[2-(dimethylamino)cyclohexyl]-N-phenylpropionamide (other name: UF-17);
- **1489** 3,4-dichloro-N-[2-(dimethylamino)cyclohexyl]-N-isopropyl-benzamide (other name: Isopropyl **1490** U-47700).
- 1491 2. Any of the following opium derivatives, their salts, isomers and salts of isomers, unless
 1492 specifically excepted, whenever the existence of these salts, isomers and salts of isomers is possible
 1493 within the specific chemical designation:
- 1494 Acetorphine;
- 1495 Acetyldihydrocodeine;
- **1496** Benzylmorphine;
- 1497 Codeine methylbromide;
- 1498 Codeine-N-Oxide;
- 1499 Cyprenorphine;
- **1500** Desomorphine;
- 1501 Dihydromorphine;
- 1502 Drotebanol;
- 1503 Etorphine;
- 1504 Heroin;
- **1505** Hydromorphinol;
- 1506 Methyldesorphine;
- **1507** Methyldihydromorphine;
- **1508** Morphine methylbromide;
- **1509** Morphine methylsulfonate;
- 1510 Morphine-N-Oxide;
- 1511 Myrophine;
- 1512 Nicocodeine;
- 1513 Nicomorphine;
- 1514 Normorphine;
- 1515 Pholcodine;
- **1516** Thebacon.
- 1517 3. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation, which contains any quantity of the following hallucinogenic substances, or which contains any of its salts, isomers, and salts of isomers, whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation (for purposes of this subdivision only, the term "isomer" includes the optical, position, and geometric isomers):
- **1522** Alpha-ethyltryptamine (some trade or other names: Monase; a-ethyl-1H-indole-3-ethanamine; **1523** 3-2-aminobutyl] indole; a-ET; AET);
- 1524 4-Bromo-2,5-dimethoxyphenethylamine (some trade or other
 - 1525 2-4-bromo-2,5-dimethoxyphenyl]-1-aminoethane;alpha-desmethyl DOB; 2C-B; Nexus);
 - **1526** 3,4-methylenedioxy amphetamine;
 - **1527** 5-methoxy-3,4-methylenedioxy amphetamine;
 - **1528** 3,4,5-trimethoxy amphetamine;
 - 1529 Alpha-methyltryptamine (other name: AMT);
 - 1530 Bufotenine;
 - **1531** Diethyltryptamine;
 - **1532** Dimethyltryptamine;
 - **1533** 4-methyl-2,5-dimethoxyamphetamine;
 - **1534** 2,5-dimethoxy-4-ethylamphetamine (DOET);
 - **1535** 4-fluoro-N-ethylamphetamine;

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names:

- 1536 2.5-dimethoxy-4-(n)-propylthiophenethylamine (other name: 2C-T-7);
- 1537 Ibogaine:
- 1538 5-methoxy-N,N-diisopropyltryptamine (other name: 5-MeO-DIPT);
- 1539 Lysergic acid diethylamide;
- 1540 Mescaline;
- 1541 Parahexvl (some trade other or 1542

3-Hexyl-1-hydroxy-7.8,9,10-tetrahydro-6,6,9-trimethyl-6H-dibenzo [b,d] pyran; Synhexyl);

- 1543 Pevote:
- 1544 N-ethyl-3-piperidyl benzilate;
- 1545 N-methyl-3-piperidyl benzilate;
- 1546 Psilocybin;
- 1547 Psilocvn:
- 1548 Salvinorin A;

1549 Tetrahydrocannabinols, except as present in (i) industrial hemp, as defined in § 3.2-4112, that is possessed by a person registered pursuant to subsection A of § 3.2-4115 or his agent; (ii) a hemp 1550 1551 product, as defined in § 3.2-4112, containing a tetrahydrocannabinol concentration of no greater than 0.3 percent that is derived from industrial hemp, as defined in § 3.2-4112, that is grown, dealt, or processed 1552 in compliance with state or federal law; (iii) marijuana; (iv) dronabinol in sesame oil and encapsulated 1553 1554 in a soft gelatin capsule in a drug product approved by the U.S. Food and Drug Administration; or (v) 1555 industrial hemp, as defined in § 3.2-4112, that is possessed by a person who holds a hemp producer license issued by the U.S. Department of Agriculture pursuant to 7 C.F.R. Part 990; 1556

- 1557 2,5-dimethoxyamphetamine (some trade or other names: 2,5-dimethoxy-a-methylphenethylamine; 1558 2,5-DMA);
- 1559 3,4-methylenedioxymethamphetamine (MDMA), its optical, positional and geometric isomers, salts 1560 and salts of isomers;
- 1561 3,4-methylenedioxy-N-ethylamphetamine (also known as N-ethyl-alpha-methyl-3,4 1562 (methylenedioxy)phenethylamine, N-ethyl MDA, MDE, MDEA);
- 1563 N-hydroxy-3, 4-methylenedioxyamphetamine(some other names: N-hydroxy-alpha-methyl-3,4(methylenedioxy)phenethylamine, and N-hydroxy MDA); 1564
- 4-bromo-2,5-dimethoxyamphetamine 1565 (some trade other or names: 1566 4-bromo-2,5-dimethoxy-a-methylphenethylamine; 4-bromo-2,5-DMA);
- 4-methoxyamphetamine (some trade or other names: 4-methoxy-a-methylphenethylamine; 1567 1568 paramethoxyamphetamine; PMA);
- 1569 Ethylamine analog of phencyclidine (some other names: N-ethyl-1-phenylcyclohexylamine, 1570 (1-phenylcyclohexyl) ethylamine, N-(1-phenylcyclohexyl) ethylamine, cyclohexamine, PCE);
- 1571 Pyrrolidine analog of phencyclidine (some other names: 1-(1-phenylcyclohexyl)-pyrrolidine, PCPy, 1572 PHP);
- Thiophene analog of phencyclidine (some other names: 1-[1-(2-thienyl)-cyclohexyl]-piperidine, 1573 2-thienyl analog of phencyclidine, TPCP, TCP); 1574
- 1575 1-1-(2-thienyl)cyclohexyl]pyrrolidine (other name: TCPy);
- 1576 3,4-methylenedioxypyrovalerone (other name: MDPV);
- 4-methylmethcathinone (other names: mephedrone, 4-MMC); 1577
- 1578 3,4-methylenedioxymethcathinone (other name: methylone);
- Naphthylpyrovalerone (other name: naphyrone); 1579
- 1580 4-fluoromethcathinone (other names: flephedrone, 4-FMC);
- 1581 4-methoxymethcathinone (other names: methedrone; bk-PMMA);
- Ethcathinone (other name: N-ethylcathinone); 1582
- 1583 3,4-methylenedioxyethcathinone (other name: ethylone);
- 1584 Beta-keto-N-methyl-3,4-benzodioxolylbutanamine (other name: butylone);
- 1585 N,N-dimethylcathinone (other name: metamfepramone);
- 1586 Alpha-pyrrolidinopropiophenone (other name: alpha-PPP);
- 1587 4-methoxy-alpha-pyrrolidinopropiophenone (other name: MOPPP);
- 1588 3.4-methylenedioxy-alpha-pyrrolidinopropiophenone (other name: MDPPP);
- Alpha-pyrrolidinovalerophenone (other name: alpha-PVP); 1589
- 1590 6,7-dihydro-5H-indeno-(5,6-d)-1,3-dioxol-6-amine (other name: MDAI);
- 1591 3-fluoromethcathinone (other name: 3-FMC);
- 4-Ethyl-2,5-dimethoxyphenethylamine (other name: 2C-E); 1592
- 1593 4-Iodo-2,5-dimethoxyphenethylamine (other name: 2C-I);
- 1594 4-Methylethcathinone (other name: 4-MEC);
- 1595 4-Ethylmethcathinone (other name: 4-EMC);
- 1596 N,N-diallyl-5-methoxytryptamine (other name: 5-MeO-DALT);
- 1597 Beta-keto-methylbenzodioxolylpentanamine (other names: Pentylone, bk-MBDP);

- 1598 Alpha-methylamino-butyrophenone (other name: Buphedrone);
- 1599 Alpha-methylamino-valerophenone (other name: Pentedrone);
- 1600 3.4-Dimethylmethcathinone (other name: 3.4-dmmc);
- 1601 4-methyl-alpha-pyrrolidinopropiophenone (other name: MPPP);
- 1602 4-Iodo-2,5-dimethoxy-N-[(2-methoxyphenyl)methyl]-benzeneethanamine (other names: 25-I, 1603 25I-NBOMe, 2C-I-NBOMe);
- 1604 Methoxetamine (other names: MXE, 3-MeO-2-Oxo-PCE);
- 1605 4-Fluoromethamphetamine (other name: 4-FMA);
- 1606 4-Fluoroamphetamine (other name: 4-FA);
- 1607 2-(2,5-Dimethoxy-4-methylphenyl)ethanamine (other name: 2C-D);
- 1608 2-(4-Chloro-2,5-dimethoxyphenyl)ethanamine (other name: 2C-C);
- 1609 2-[4-(Ethylthio)-2,5-dimethoxyphenyl]ethanamine (other name: 2C-T-2);
- 2-[4-(Isopropylthio)-2,5-dimethoxyphenyl]ethanamine (other name: 2C-T-4); 1610
- 2-(2,5-Dimethoxyphenyl)ethanamine (other name: 2C-H); 1611
- 1612 2-(2,5-Dimethoxy-4-nitro-phenyl)ethanamine (other name: 2C-N);
- 1613 2-(2,5-Dimethoxy-4-(n)-propylphenyl)ethanamine (other name: 2C-P);
- (2-aminopropyl)benzofuran (other name: APB); 1614
- 1615 (2-aminopropyl)-2,3-dihydrobenzofuran (other name: APDB);
- 1616 4-chloro-2,5-dimethoxy-N-[(2-methoxyphenyl)methyl]-benzeneethanamine (other names:
- 1617 2C-C-NBOMe, 25C-NBOMe, 25C);
- 1618 4-bromo-2,5-dimethoxy-N-[(2-methoxyphenyl)methyl]-benzeneethanamine (other names: 1619 2C-B-NBOMe, 25B-NBOMe, 25B);
- 1620 Acetoxydimethyltryptamine (other names: AcO-Psilocin, AcO-DMT, Psilacetin);
- 1621 Benocyclidine (other names: BCP, BTCP);
- 1622 Alpha-pyrrolidinobutiophenone (other name: alpha-PBP);
- 1623 3.4-methylenedioxy-N,N-dimethylcathinone (other names: Dimethylone, bk-MDDMA);
- 1624 4-bromomethcathinone (other name: 4-BMC);
- 1625 4-chloromethcathinone (other name: 4-CMC);
- 1626 4-Iodo-2,5-dimethoxy-N-[(2-hydroxyphenyl)methyl]-benzeneethanamine (other name: 25I-NBOH);
- 1627 Alpha-Pyrrolidinohexiophenone (other name: alpha-PHP);
- 1628 Alpha-Pyrrolidinoheptiophenone (other name: PV8);
- 1629 5-methoxy-N.N-methylisopropyltryptamine (other name: 5-MeO-MIPT);
- 1630 Beta-keto-N,N-dimethylbenzodioxolylbutanamine (other names: Dibutylone, bk-DMBDB);
- 1631 Beta-keto-4-bromo-2,5-dimethoxyphenethylamine (other name: bk-2C-B);
- 1632 1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-1-pentanone (other name: N-ethylpentylone);
- 1633 1-[1-(3-methoxyphenyl)cyclohexyl]piperidine (other name: 3-methoxy PCP);
- 1634 1-[1-(4-methoxyphenyl)cyclohexyl]piperidine (other name: 4-methoxy PCP);
- 1635 4-Chloroethcathinone (other name: 4-CEC);
- 1636 3-Methoxy-2-(methylamino)-1-(4-methylphenyl)-1-propanone (other name: Mexedrone);
- 1637 1-propionyl lysergic acid diethylamide (other name: 1P-LSD);
- 1638 (2-Methylaminopropyl)benzofuran (other name: MAPB);
- 1639 1-(1,3-benzodioxol-5-yl)-2-(dimethylamino)-1-pentanone (other N,N-Dimethylpentylone, names: 1640 Dipentylone);
- 1641 1-(4-methoxyphenyl)-2-(pyrrolidin-1-yl)octan-1-one (other name: 4-methoxy-PV9);
- 1642 3.4-tetramethylene-alpha-pyrrolidinovalerophenone (other name: TH-PVP);
- 1643 4-allyloxy-3,5-dimethoxyphenethylamine (other name: Allylescaline);
- 1644 4-Bromo-2,5-dimethoxy-N-[(2-hydroxyphenyl)methyl]-benzeneethanamine (other name: 25B-NBOH);
- 1645 4-chloro-alpha-methylamino-valerophenone (other name: 4-chloropentedrone);
- 1646 4-chloro-alpha-Pyrrolidinovalerophenone (other name: 4-chloro-alpha-PVP);
- 1647 4-fluoro-alpha-Pyrrolidinoheptiophenone (other name: 4-fluoro-PV8);
- 1648 4-hydroxy-N.N-diisopropyltryptamine (other name: 4-OH-DIPT);
- 1649 4-methyl-alpha-ethylaminopentiophenone;
- 1650 4-methyl-alpha-Pyrrolidinohexiophenone (other name: MPHP);
- 1651 5-methoxy-N,N-dimethyltryptamine (other name: 5-MeO-DMT);
- 1652 5-methoxy-N-ethyl-N-isopropyltryptamine (other name: 5-MeO-EIPT);
- 6-ethyl-6-nor-lysergic acid diethylamide (other name: ETH-LAD); 1653
- 1654 6-allyl-6-nor-lysergic acid diethylamide (other name: AL-LAD);
- 1655 (N-methyl aminopropyl)-2,3-dihydrobenzofuran (other name: MAPDB);
- 1656 2-(methylamino)-2-phenyl-cyclohexanone (other name: Deschloroketamine);
- 1657 2-(ethylamino)-2-phenyl-cyclohexanone (other name: deschloro-N-ethyl-ketamine);
- 2-methyl-1-(4-(methylthio)phenyl)-2-morpholinopropiophenone (other name: MMMP); 1658

- 1659 Alpha-ethylaminohexanophenone (other name: N-ethylhexedrone);
- **1660** N-ethyl-1-(3-methoxyphenyl)cyclohexylamine (other name: 3-methoxy-PCE);
- **1661** 4-fluoro-alpha-pyrrolidinohexiophenone (other name: 4-fluoro-alpha-PHP);
- **1662** N-ethyl-1,2-diphenylethylamine (other name: Ephenidine);
- **1663** 2,5-dimethoxy-4-chloroamphetamine (other name: DOC);
- **1664** 3,4-methylenedioxy-N-tert-butylcathinone;
- 1665 Alpha-pyrrolidinoisohexiophenone (other name: alpha-PiHP);
- 1666 1-[1-(3-hydroxyphenyl)cyclohexyl]piperidine (other name: 3-hydroxy PCP);
- **1667** 4-acetyloxy-N,N-diallyltryptamine (other name: 4-AcO-DALT);
- **1668** 4-hydroxy-N,N-methylisopropyltryptamine (other name: 4-hydroxy-MiPT);
- **1669** 3,4-Methylenedioxy-alpha-pyrrolidinohexanophenone (other name: MDPHP);
- **1670** 5-methoxy-N,N-dibutyltryptamine (other name: 5-methoxy-DBT);
- **1671** 1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-1-butanone (other names: Eutylone, bk-EBDB);
- **1672** 1-(1,3-benzodioxol-5-yl)-2-(butylamino)-1-pentanone (other name: N-butylpentylone);
- **1673** N-benzyl-3,4-dimethoxyamphetamine (other name: N-benzyl-3,4-DMA);
- 1674 1-(benzo[d][1,3]dioxol-5-yl)-2-(sec-butylamino)pentan-1-one (other name: N-sec-butyl Pentylone);
- 1675 1-cyclopropionyl lysergic acid diethylamide (other name: 1cP-LSD);
- **1676** 2-(ethylamino)-1-phenylheptan-1-one (other name: N-ethylheptedrone);
- 1677 (2-ethylaminopropyl)benzofuran (other name: EAPB);
- **1678** 4-ethyl-2,5-dimethoxy-N-[(2-hydroxyphenyl)methyl]-benzeneethanamine (other name: 25E-NBOH);
- **1679** 2-fluoro-Deschloroketamine (other name: 2-(2-fluorophenyl)-2-(methylamino)-cyclohexanone);
- **1680** 4-hydroxy-N-ethyl-N-propyltryptamine (other name: 4-hydroxy-EPT);
- 1681 2-(isobutylamino)-1-phenylhexan-1-one (other names: N-Isobutyl Hexedrone, alpha-isobutylaminohexanphenone);
- 1683 1-(4-methoxyphenyl)-N-methylpropan-2-amine (other names: para-Methoxymethamphetamine, 1684 PMMA);
- 1685 N-ethyl-1-(3-hydroxyphenyl)cyclohexylamine (other name: 3-hydroxy-PCE);
- 1686 N-heptyl-3,4-dimethoxyamphetamine (other name: N-heptyl-3,4-DMA);
- 1687 N-hexyl-3,4-dimethoxyamphetamine (other name: N-hexyl-3,4-DMA);
- **1688** 4-fluoro-3-methyl-alpha-pyrrolidinovalerophenone (other name: 4-fluoro-3-methyl-alpha-PVP);
- **1689** 4-fluoro-alpha-methylamino-valerophenone (other name: 4-fluoropentedrone);
- **1690** N-(1,4-dimethylpentyl)-3,4-dimethoxyamphetamine (other name: N-(1,4-dimethylpentyl)-3,4-DMA);
- **1691** 4,5-methylenedioxy-N,N-diisopropyltryptamine (other name: 4,5-MDO-DiPT);
- 1692 Alpha-pyrrolidinocyclohexanophenone (other name: alpha-PCYP);
- **1693** 3,4-methylenedioxy-alpha-pyrrolidinoheptiophenone (other name: MDPV8);
- **1694** 4-chloro-alpha-methylaminobutiophenone (other name: 4-chloro Buphedrone).
- 4. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture
 or preparation which contains any quantity of the following substances having a depressant effect on the
 central nervous system, including its salts, isomers and salts of isomers whenever the existence of such
 salts, isomers and salts of isomers is possible within the specific chemical designation:
- 1699 5-(2-chlorophenyl)-1,3-dihydro-3-methyl-7-nitro-2H-1,4-benzodiazepin-2-one (other name:
 1700 Meclonazepam);
- **1701** 7-chloro-5-(2-fluorophenyl)-1,3-dihydro-1,4-benzodiazepin-2-one (other name: Norfludiazepam);
- **1702** Bromazolam;
- 1703 Clonazolam;
- 1704 Deschloroetizolam;
- 1705 Etizolam;
- 1706 Flualprazolam;
- 1707 Flubromazepam;
- 1708 Flubromazolam;
- 1709 Gamma hydroxybutyric acid (some other names include GHB; gamma hydroxybutyrate;
 1710 4-hydroxybutyrate; 4-hydroxybutanoic acid; sodium oxybate; sodium oxybutyrate);
- 1711 Mecloqualone;
- 1712 Methaqualone.
- 5. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture
 or preparation which contains any quantity of the following substances having a stimulant effect on the
 central nervous system, including its salts, isomers and salts of isomers:
- **1716** 2-(3-fluorophenyl)-3-methylmorpholine (other name: 3-fluorophenmetrazine);
- 1717 Aminorex (some trade or other names; aminoxaphen; 2-amino-5-phenyl-2-oxazoline;
 1718 4,5-dihydro-5-phenyl-2-oxazolamine);
- 1719 Cathinone (some trade or other names: 2-amino-1-phenyl-1-propanone, alpha-aminopropiophenone, 1720 2-aminopropiophenone, normalistic state state state of the state of t
- 1720 2-aminopropiophenone, norephedrone), and any plant material from which Cathinone may be derived;

- 1721 Cis-4-methylaminorex (other name: cis-4,5-dihydro-4-methyl-5-phenyl-2-oxazolamine);
- 1722 Ethylamphetamine;
- 1723 Ethyl phenyl(piperidin-2-yl)acetate (other name: Ethylphenidate);
- 1724 Fenethylline;

1725 Methcathinone (some other names: 2-(methylamino)-propiophenone;
1726 alpha-(methylamino)-propiophenone; 2-(methylamino)-1-phenylpropan-1-one;

alpha-N-methylaminopropiophenone; monomethylpropion; ephedrone; N-methylcathinone;
 methylcathinone; AL-464; AL-422; AL-463 and UR 1432);

- **1729** N-Benzylpiperazine (some other names: BZP, 1-benzylpiperazine);
- 1730 N,N-dimethylamphetamine (other names: N,N-alpha-trimethyl-benzeneethanamine,
 1731 N,N-alpha-trimethylphenethylamine);
- 1732 Methyl 2-(4-fluorophenyl)-2-(2-piperidinyl)acetate (other name: 4-fluoromethylphenidate);

1733 Isopropyl-2-phenyl-2-(2-piperidinyl)acetate (other name: Isopropylphenidate);

- 4-chloro-N,N-dimethylcathinone;
- 1735 3,4-methylenedioxy-N-benzylcathinone (other name: BMDP).

6. Any substance that contains one or more cannabimimetic agents or that contains their salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation, and any preparation, mixture, or substance containing, or mixed or infused with, any detectable amount of one or more cannabimimetic agents.

a. "Cannabimimetic agents" includes any substance that is within any of the following structuralclasses:

1742 2-(3-hydroxycyclohexyl)phenol with substitution at the 5-position of the phenolic ring by alkyl or 1743 alkenyl, whether or not substituted on the cyclohexyl ring to any extent;

3-(1-naphthoyl)indole or 1H-indol-3-yl-(1-naphthyl)methane with substitution at the nitrogen atom of
the indole ring, whether or not further substituted on the indole ring to any extent, whether or not
substituted on the naphthoyl or naphthyl ring to any extent;

- 1747 3-(1-naphthoyl)pyrrole with substitution at the nitrogen atom of the pyrrole ring, whether or not further substituted in the pyrrole ring to any extent, whether or not substituted on the naphthoyl ring to any extent;
- 1750 1-(1-naphthylmethyl)indene with substitution of the 3-position of the indene ring, whether or not further substituted in the indene ring to any extent, whether or not substituted on the naphthyl ring to any extent;
- 3-phenylacetylindole or 3-benzoylindole with substitution at the nitrogen atom of the indole ring,
 whether or not further substituted in the indole ring to any extent, whether or not substituted on the
 phenyl ring to any extent;
- 3-cyclopropoylindole with substitution at the nitrogen atom of the indole ring, whether or not further
 substituted on the indole ring to any extent, whether or not substituted on the cyclopropyl ring to any
 extent;
- 3-adamantoylindole with substitution at the nitrogen atom of the indole ring, whether or not further
 substituted on the indole ring to any extent, whether or not substituted on the adamantyl ring to any
 extent;

1762 N-(adamantyl)-indole-3-carboxamide with substitution at the nitrogen atom of the indole ring,
1763 whether or not further substituted on the indole ring to any extent, whether or not substituted on the
1764 adamantyl ring to any extent; and

1765 N-(adamantyl)-indazole-3-carboxamide with substitution at a nitrogen atom of the indazole ring,
1766 whether or not further substituted on the indazole ring to any extent, whether or not substituted on the
1767 adamantyl ring to any extent.

- b. The term "cannabimimetic agents" includes:
- **1769** 5-(1,1-Dimethylheptyl)-2-[3-hydroxycyclohexyl]-phenol (other name: CP 47,497);
- 1770 5-(1,1-Dimethylhexyl)-2-[3-hydroxycyclohexyl]-phenol (other name: CP 47,497 C6 homolog);
- 1771 5-(1,1-Dimethyloctyl)-2-[3-hydroxycyclohexyl]-phenol (other name: CP 47,497 C8 homolog);
- 1772 5-(1,1-Dimethylnonyl)-2-[3-hydroxycyclohexyl]-phenol (other name: CP 47,497 C9 homolog);
- 1773 1-pentyl-3-(1-naphthoyl)indole (other names: JWH-018, AM-678);
- 1774 1-butyl-3-(1-naphthoyl)indole (other name: JWH-073);
- 1775 1-pentyl-3-(2-methoxyphenylacetyl)indole (other name: JWH-250);
- 1776 1-hexyl-3-(naphthalen-1-oyl)indole (other name: JWH-019);
- 1777 1-[2-(4-morpholinyl)ethyl]-3-(1-naphthoyl)indole (other name: JWH-200);
- **1778** (6aR, 10aR) 9 (hydroxymethyl) 6, 6 dimethyl 3 (2 methyloctan 2 yl) 6a, 7, 10, 10a tet
- 1779 rahydrobenzo[c]chromen-1-ol (other name: HU-210);
- 1780 1-pentyl-3-(4-methoxy-1-naphthoyl)indole (other name: JWH-081);
- 1781 1-pentyl-3-(4-methyl-1-naphthoyl)indole (other name: JWH-122);

- 1782 1-pentyl-3-(2-chlorophenylacetyl)indole (other name: JWH-203);
- 1783 1-pentyl-3-(4-ethyl-1-naphthoyl)indole (other name: JWH-210);
- 1784 1-pentyl-3-(4-chloro-1-naphthoyl)indole (other name: JWH-398);
- 1785 1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole (other name: AM-694);
- 1786 1-((N-methylpiperidin-2-yl)methyl)-3-(1-naphthoyl)indole (other name: AM-1220);
- 1787 1-(5-fluoropentyl)-3-(1-naphthoyl)indole (other name: AM-2201);
- 1788 1-[(N-methylpiperidin-2-yl)methyl]-3-(2-iodobenzoyl)indole (other name: AM-2233);
- **1789** Pravadoline (4-methoxyphenyl)-[2-methyl-1-(2-(4-morpholinyl)ethyl)indol-3-yl]methanone (other 1790 name: WIN 48,098);
- 1791 1-pentyl-3-(4-methoxybenzoyl)indole (other names: RCS-4, SR-19);
- 1792 1-(2-cyclohexylethyl)-3-(2-methoxyphenylacetyl)indole (other names: RCS-8, SR-18);
- 1793 1-pentyl-3-(2,2,3,3-tetramethylcyclopropylmethanone)indole (other name: UR-144);
- 1794 1-(5-fluoropentyl)-3-(2,2,3,3-tetramethylcyclopropylmethanone)indole (other names: XLR-11,
- **1795** 5-fluoro-UR-144);
- 1796 N-adamantyl-1-fluoropentylindole-3-carboxamide (other name: STS-135);
- 1797 N-adamantyl-1-pentylindazole-3-carboxamide (other names: AKB48, APINACA);
- 1798 1-pentyl-3-(1-adamantoyl)indole (other name: AB-001);
- (8-quinolinyl)(1-pentylindol-3-yl)carboxylate (other name: PB-22);
- **1800** (8-quinolinyl)(1-(5-fluoropentyl)indol-3-yl)carboxylate (other name: 5-fluoro-PB-22);
- **1801** (8-quinolinyl)(1-cyclohexylmethyl-indol-3-yl)carboxylate (other name: BB-22);
- 1802 N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-pentylindazole-3-carboxamide (other name: AB-PINACA);
- 1803 N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(4-fluorobenzyl)indazole-3-carboxamide (other name:
 1804 AB-FUBINACA);
- **1805** 1-(5-fluoropentyl)-3-(1-naphthoyl)indazole (other name: THJ-2201);
- 1806 N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentylindazole-3-carboxamide (other name:
 1807 ADB-PINACA);
- **1808** N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(cyclohexylmethyl)indazole-3-carboxamide (other name: **1809** AB-CHMINACA);
- 1810 N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(5-fluoropentyl)indazole-3-carboxamide (other name:
 1811 5-fluoro-AB-PINACA);
- 1812 N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(cyclohexylmethyl)indazole-3-carboxamide (other names:
 1813 ADB-CHMINACA, MAB-CHMINACA);
- 1814 Methyl-2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3-methylbutanoate (other name:
 1815 5-fluoro-AMB);
- **1816** 1-naphthalenyl 1-(5-fluoropentyl)-1H-indole-3-carboxylate (other name: NM-2201);
- **1817** 1-(4-fluorobenzyl)-3-(2,2,3,3-tetramethylcyclopropylmethanone)indole (other name: FUB-144);
- **1818** 1-(5-fluoropentyl)-3-(4-methyl-1-naphthoyl)indole (other name MAM-2201);
- 1819 N-(1-Amino-3,3-dimethyl-1-oxobutan-2-yl)-1-[(4-fluorophenyl)methyl]-1H-indazole-3-carboxamide
 1820 (other name: ADB-FUBINACA);
- Methyl 2-[1-[(4-fluorophenyl)methyl]-1H-indazole-3-carboxamido]-3,3-dimethylbutanoate (other name:
 MDMB-FUBINACA);
- 1823 Methyl 2-[1-(5-fluoropentyl)-1H-indazole-3-carboxamido]-3,3-dimethylbutanoate (other names:
 1824 5-fluoro-ADB, 5-Fluoro-MDMB-PINACA);
- **1825** Methyl 2-({1-[(4-fluorophenyl)methyl]-1H-indazole-3-carbonyl}amino)-3-methylbutanoate (other **1826** names: AMB-FUBINACA, FUB-AMB);
- **1827** N-(adamantan-1-yl)-1-(4-fluorobenzyl)-1H-indazole-3-carboxamide (other names: FUB-AKB48, **1828** 5F-APINACA);
- 1829 N-(adamantan-1-yl)-1-(5-fluoropentyl)-1H-indazole-3-carboxamide (other name: 5F-AKB48);
- 1830 N-(adamantanyl)-1-(5-chloropentyl) indazole-3-carboxamide (other name: 5-chloro-AKB48);
- 1831 Naphthalen-1-yl 1-pentyl-1H-indazole-3-carboxylate (other name: SDB-005);
- 1832 N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(cyclohexylmethyl)indole-3-carboxamide (other name:
 1833 AB-CHMICA);
- **1834** 1-pentyl-N-(phenylmethyl)-1H-indole-3-carboxamide (other name: SDB-006);
- 1835 Quinolin-8-yl 1-(4-fluorobenzyl)-1H-indole-3-carboxylate (other name: FUB-PB-22);
- 1836 Methyl N-[1-(cyclohexylmethyl)-1H-indole-3-carbonyl]valinate (other name: MMB-CHMICA);
- 1837 N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(5-fluoropentyl)indazole-3-carboxamide (other name: 1838 5-fluoro-ADB-PINACA);
- **1839** 1-(4-cyanobutyl)-N-(1-methyl-1-phenylethyl)-1H-indazole-3-carboxamide (other name: 4-cyano **1840** CUMYL-BUTINACA);
- 1841 Methyl 2-[1-(5-fluoropentyl)-1H-indole-3-carboxamido]-3,3-dimethylbutanoate (other names: 5-fluoro 1842 MDMB-PICA, 5F-MDMB-PICA);
- **1843** Ethyl 2-({1-[(4-fluorophenyl)methyl]-1H-indazole-3-carbonyl}amino)-3-methylbutanoate (other name:

1844 EMB-FUBINACA);

- 1845 Methyl 2-[1-4-fluorobutyl)-1H-indazole-3-carboxamido]-3,3-dimethylbutanoate (other name: 1846 4-fluoro-MDMB-BUTINACA);
- 1847 1-(5-fluoropentyl)-N-(1-methyl-1-phenylethyl)-1H-indole-3-carboxamide (other name: 5-fluoro 1848 CUMYL-PICA);
- 1849 Methyl 2-[1-(pent-4-enyl)-1H-indazole-3-carboxamindo]-3,3-dimethylbutanoate (other name: 1850 MDMB-4en-PINAČA);
- 1851 Methyl 2-({1-[(4-fluorophenyl)methyl]-1H-indole-3-carbonyl}amino)-3-methylbutanoate (other names: 1852 MMB-FUBICA, AMB-FUBICA);
- 1853 Methyl 2-[1-(4-penten-1-yl)-1H-indole-3-carboxamido]-3-methylbutanoate (other names: MMB022, 1854 MMB-4en-PICA);
- 1855 Methyl 2-[1-(5-fluoropentyl)-1H-indole-3-carboxamido]-3-methylbutanoate (other name: MMB 2201); Methyl 2-[1-(5-fluoropentyl)-1H-indole-3-carboxamido]-3-phenylpropanoate (other name: 1856
- 1857 5-fluoro-MPP-PICA);
- 1858 N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-butylindazole-3-carboxamide (other name: ADB-BUTINACA); 1859
- 1860 N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(5-chloropentyl)indazole-3-carboxamide (other name: 1861 5-chloro-AB-PINACA);
- 1862 1-(5-fluoropentyl)-N-(2-phenylpropan-2-yl)-1H-indazole-3-carboxamide (other names: 1863 5F-CUMYL-PINACA, 5-fluoro CUMYL-PINACA, CUMYL-5F-PINACA);
- 1864 Ethyl 2-[1-(5-fluoropentyl)-1H-indazole-3-carboxamido]-3,3-dimethylbutanoate (other names: 1865 5F-EDMB-PINACA, 5-fluoro EDMB-PINACA);
- 1866 Ethyl-2-[1-(5-fluoropentyl)-1H-indazole-3-carboxamido]-3-methylbutanoate (other names: 1867 5-fluoro-EMB-PINACA, 5F-AEB);
- 1868 Ethyl 2-[1-(5-fluoropentyl)-1H-indole-3-carboxamido]-3-methylbutanoate (other name: 1869 5-fluoro-EMB-PICA);
- 1870 Ethyl-2-[1-(5-fluoropentyl)-1H-indole-3-carboxamido]-3.3-dimethylbutanoate (other name: 5-fluoro 1871 EDMB-PICA);
- 1872 Methyl 2-[1-(4-fluorobutyl)-1H-indole-3-carboxamido]-3,3-dimethylbutanoate (other name: 1873 4-fluoro-MDMB-BUTICA);
- 1874 Methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-3,3-dimethylbutanoate (other names: 1875 MDMB-CHMICA, MMB-CHMINACA);
- 1876 N-(1-amino-3.3-dimethyl-1-oxobutan-2-yl)-1-(pent-4-enyl)indazole-3-carboxamide (other name: 1877 ADB-4en-PINACA).
- 1878 § 59.1-200. Prohibited practices.
- 1879 A. The following fraudulent acts or practices committed by a supplier in connection with a consumer 1880 transaction are hereby declared unlawful: 1881
 - 1. Misrepresenting goods or services as those of another;
- 1882 2. Misrepresenting the source, sponsorship, approval, or certification of goods or services;
- 1883 3. Misrepresenting the affiliation, connection, or association of the supplier, or of the goods or 1884 services, with another; 1885
 - 4. Misrepresenting geographic origin in connection with goods or services;
- 1886 5. Misrepresenting that goods or services have certain quantities, characteristics, ingredients, uses, or 1887 benefits; 1888
 - 6. Misrepresenting that goods or services are of a particular standard, quality, grade, style, or model;
- 1889 7. Advertising or offering for sale goods that are used, secondhand, repossessed, defective, 1890 blemished, deteriorated, or reconditioned, or that are "seconds," irregulars, imperfects, or "not first 1891 class," without clearly and unequivocally indicating in the advertisement or offer for sale that the goods 1892 are used, secondhand, repossessed, defective, blemished, deteriorated, reconditioned, or are "seconds," 1893 irregulars, imperfects or "not first class";
- 1894 8. Advertising goods or services with intent not to sell them as advertised, or with intent not to sell 1895 at the price or upon the terms advertised.
- 1896 In any action brought under this subdivision, the refusal by any person, or any employee, agent, or servant thereof, to sell any goods or services advertised or offered for sale at the price or upon the terms 1897 1898 advertised or offered, shall be prima facie evidence of a violation of this subdivision. This paragraph 1899 shall not apply when it is clearly and conspicuously stated in the advertisement or offer by which such 1900 goods or services are advertised or offered for sale, that the supplier or offeror has a limited quantity or 1901 amount of such goods or services for sale, and the supplier or offeror at the time of such advertisement 1902 or offer did in fact have or reasonably expected to have at least such quantity or amount for sale;
- 1903 9. Making false or misleading statements of fact concerning the reasons for, existence of, or amounts 1904 of price reductions;

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1905 10. Misrepresenting that repairs, alterations, modifications, or services have been performed or parts 1906 installed;

1907 11. Misrepresenting by the use of any written or documentary material that appears to be an invoice 1908 or bill for merchandise or services previously ordered;

1909 12. Notwithstanding any other provision of law, using in any manner the words "wholesale," 1910 "wholesaler," "factory," or "manufacturer" in the supplier's name, or to describe the nature of the 1911 supplier's business, unless the supplier is actually engaged primarily in selling at wholesale or in 1912 manufacturing the goods or services advertised or offered for sale;

1913 13. Using in any contract or lease any liquidated damage clause, penalty clause, or waiver of 1914 defense, or attempting to collect any liquidated damages or penalties under any clause, waiver, damages, 1915 or penalties that are void or unenforceable under any otherwise applicable laws of the Commonwealth, 1916 or under federal statutes or regulations;

1917 13a. Failing to provide to a consumer, or failing to use or include in any written document or 1918 material provided to or executed by a consumer, in connection with a consumer transaction any 1919 statement, disclosure, notice, or other information however characterized when the supplier is required 1920 by 16 C.F.R. Part 433 to so provide, use, or include the statement, disclosure, notice, or other 1921 information in connection with the consumer transaction;

14. Using any other deception, fraud, false pretense, false promise, or misrepresentation in connection 1922 1923 with a consumer transaction;

15. Violating any provision of § 3.2-6509, 3.2-6512, 3.2-6513, 3.2-6513.1, 3.2-6514, 3.2-6515, 3.2-6516, or 3.2-6519 is a violation of this chapter; 1924 1925 1926

16. Failing to disclose all conditions, charges, or fees relating to:

1927 a. The return of goods for refund, exchange, or credit. Such disclosure shall be by means of a sign 1928 attached to the goods, or placed in a conspicuous public area of the premises of the supplier, so as to be 1929 readily noticeable and readable by the person obtaining the goods from the supplier. If the supplier does 1930 not permit a refund, exchange, or credit for return, he shall so state on a similar sign. The provisions of 1931 this subdivision shall not apply to any retail merchant who has a policy of providing, for a period of not 1932 less than 20 days after date of purchase, a cash refund or credit to the purchaser's credit card account 1933 for the return of defective, unused, or undamaged merchandise upon presentation of proof of purchase. 1934 In the case of merchandise paid for by check, the purchase shall be treated as a cash purchase and any 1935 refund may be delayed for a period of 10 banking days to allow for the check to clear. This subdivision 1936 does not apply to sale merchandise that is obviously distressed, out of date, post season, or otherwise 1937 reduced for clearance; nor does this subdivision apply to special order purchases where the purchaser 1938 has requested the supplier to order merchandise of a specific or unusual size, color, or brand not 1939 ordinarily carried in the store or the store's catalog; nor shall this subdivision apply in connection with a 1940 transaction for the sale or lease of motor vehicles, farm tractors, or motorcycles as defined in 1941 § 46.2-100;

1942 b. A layaway agreement. Such disclosure shall be furnished to the consumer (i) in writing at the time 1943 of the layaway agreement, or (ii) by means of a sign placed in a conspicuous public area of the 1944 premises of the supplier, so as to be readily noticeable and readable by the consumer, or (iii) on the bill 1945 of sale. Disclosure shall include the conditions, charges, or fees in the event that a consumer breaches 1946 the agreement:

1947 16a. Failing to provide written notice to a consumer of an existing open-end credit balance in excess 1948 of \$5 (i) on an account maintained by the supplier and (ii) resulting from such consumer's overpayment 1949 on such account. Suppliers shall give consumers written notice of such credit balances within 60 days of 1950 receiving overpayments. If the credit balance information is incorporated into statements of account 1951 furnished consumers by suppliers within such 60-day period, no separate or additional notice is required;

1952 17. If a supplier enters into a written agreement with a consumer to resolve a dispute that arises in 1953 connection with a consumer transaction, failing to adhere to the terms and conditions of such an 1954 agreement; 1955

18. Violating any provision of the Virginia Health Club Act, Chapter 24 (§ 59.1-294 et seq.);

1956 19. Violating any provision of the Virginia Home Solicitation Sales Act, Chapter 2.1 (§ 59.1-21.1 et 1957 seq.);

20. Violating any provision of the Automobile Repair Facilities Act, Chapter 17.1 (§ 59.1-207.1 et 1958 1959 seq.);

1960 Violating any provision of the Virginia Lease-Purchase Agreement Act, Chapter 17.4 21. 1961 (§ 59.1-207.17 et seq.);

1962 22. Violating any provision of the Prizes and Gifts Act, Chapter 31 (§ 59.1-415 et seq.);

1963 23. Violating any provision of the Virginia Public Telephone Information Act, Chapter 32 1964 (§ 59.1-424 et seq.);

24. Violating any provision of § 54.1-1505; 1965

1966 25. Violating any provision of the Motor Vehicle Manufacturers' Warranty Adjustment Act, Chapter

- **1967** 17.6 (§ 59.1-207.34 et seq.);
- **1968** 26. Violating any provision of § 3.2-5627, relating to the pricing of merchandise;
- 1969 27. Violating any provision of the Pay-Per-Call Services Act, Chapter 33 (§ 59.1-429 et seq.);
- 1970 28. Violating any provision of the Extended Service Contract Act, Chapter 34 (§ 59.1-435 et seq.);
- 1971 29. Violating any provision of the Virginia Membership Camping Act, Chapter 25 (§ 59.1-311 et 1972 seq.);
- **1973** 30. Violating any provision of the Comparison Price Advertising Act, Chapter 17.7 (§ 59.1-207.40 et seq.);
- 1975 31. Violating any provision of the Virginia Travel Club Act, Chapter 36 (§ 59.1-445 et seq.);
- **1976** 32. Violating any provision of §§ 46.2-1231 and 46.2-1233.1;
- **1977** 33. Violating any provision of Chapter 40 (§ 54.1-4000 et seq.) of Title 54.1;
- **1978** 34. Violating any provision of Chapter 10.1 (§ 58.1-1031 et seq.) of Title 58.1;
- 1979 35. Using the consumer's social security number as the consumer's account number with the supplier,
- **1980** if the consumer has requested in writing that the supplier use an alternate number not associated with **1981** the consumer's social security number;
- **1982** 36. Violating any provision of Chapter 18 (§ 6.2-1800 et seq.) of Title 6.2;
- **1983** 37. Violating any provision of § 8.01-40.2;
- **1984** 38. Violating any provision of Article 7 (§ 32.1-212 et seq.) of Chapter 6 of Title 32.1;
- **1985** 39. Violating any provision of Chapter 34.1 (§ 59.1-441.1 et seq.);
- 1986 40. Violating any provision of Chapter 20 (§ 6.2-2000 et seq.) of Title 6.2;
- **1987** 41. Violating any provision of the Virginia Post-Disaster Anti-Price Gouging Act, Chapter 46 (§ 59.1-525 et seq.);
- **1989** 42. Violating any provision of Chapter 47 (§ 59.1-530 et seq.);
- **1990** 43. Violating any provision of § 59.1-443.2;
- **1991** 44. Violating any provision of Chapter 48 (§ 59.1-533 et seq.);
- 1992 45. Violating any provision of Chapter 25 (§ 6.2-2500 et seq.) of Title 6.2;
- **1993** 46. Violating the provisions of clause (i) of subsection B of § 54.1-1115;
- **1994** 47. Violating any provision of § 18.2-239;
- **1995** 48. Violating any provision of Chapter 26 (§ 59.1-336 et seq.);

49. Selling, offering for sale, or manufacturing for sale a children's product the supplier knows or has
reason to know was recalled by the U.S. Consumer Product Safety Commission. There is a rebuttable
presumption that a supplier has reason to know a children's product was recalled if notice of the recall
has been posted continuously at least 30 days before the sale, offer for sale, or manufacturing for sale
on the website of the U.S. Consumer Product Safety Commission. This prohibition does not apply to
children's products that are used, secondhand or "seconds";

- **2002** 50. Violating any provision of Chapter 44.1 (§ 59.1-518.1 et seq.);
- 2003 51. Violating any provision of Chapter 22 (§ 6.2-2200 et seq.) of Title 6.2;
- **2004** 52. Violating any provision of § 8.2-317.1;
- **2005** 53. Violating subsection A of § 9.1-149.1;

54. Selling, offering for sale, or using in the construction, remodeling, or repair of any residential dwelling in the Commonwealth, any drywall that the supplier knows or has reason to know is defective drywall. This subdivision shall not apply to the sale or offering for sale of any building or structure in which defective drywall has been permanently installed or affixed;

- 55. Engaging in fraudulent or improper or dishonest conduct as defined in § 54.1-1118 while engaged in a transaction that was initiated (i) during a declared state of emergency as defined in § 44-146.16 or (ii) to repair damage resulting from the event that prompted the declaration of a state of emergency, regardless of whether the supplier is licensed as a contractor in the Commonwealth pursuant
- **2014** to Chapter 11 (§ 54.1-1100 et seq.) of Title 54.1;
- **2015** 56. Violating any provision of Chapter 33.1 (§ 59.1-434.1 et seq.);
- **2016** 57. Violating any provision of § 18.2-178, 18.2-178.1, or 18.2-200.1;
- 2017 58. Violating any provision of Chapter 17.8 (§ 59.1-207.45 et seq.);
- **2018** 59. Violating any provision of subsection E of § 32.1-126;
- **2019** 60. Violating any provision of § 54.1-111 relating to the unlicensed practice of a profession licensed **2020** under Chapter 11 (§ 54.1-1100 et seq.) or Chapter 21 (§ 54.1-2100 et seq.) of Title 54.1;
- **2021** 61. Violating any provision of § 2.2-2001.5;
- 2022 62. Violating any provision of Chapter 5.2 (§ 54.1-526 et seq.) of Title 54.1;
- **2023** 63. Violating any provision of § 6.2-312;
- 2024 64. Violating any provision of Chapter 20.1 (§ 6.2-2026 et seq.) of Title 6.2;
- 2025 65. Violating any provision of Chapter 26 (§ 6.2-2600 et seq.) of Title 6.2;
- 2026 66. Violating any provision of Chapter 54 (§ 59.1-586 et seq.);
- **2027** 67. Knowingly violating any provision of § 8.01-27.5;

2028 68. Failing to make available a conspicuous online option to cancel a recurring purchase of a good 2029 or service as required by § 59.1-207.46;

2030 69. Selling or offering for sale any substance intended for human consumption, orally or by 2031 inhalation, that contains a synthetic derivative of tetrahydrocannabinol. As used in this subdivision, 2032 "synthetic derivative" means a chemical compound produced by man through a chemical transformation 2033 to turn a compound into a different compound by adding or subtracting molecules to or from the 2034 original compound. This subdivision shall not (i) apply to products that are approved for marked by the U.S. Food and Drug Administration and scheduled in the Drug Control Act (§ 54.1-3400 et seq.) or (ii) 2035 be construed to prohibit any conduct permitted under Article 4.2 (§ 54.1-3442.5 et seq.) of Chapter 34 2036 2037 of Title 54.1.

2038 70. Selling or offering for sale to a person younger than 21 years of age any substance intended for human consumption, orally or by inhalation, that contains tetrahydrocannabinol. This subdivision shall 2039 2040 not (i) apply to products that are approved for marketing by the U.S. Food and Drug Administration and scheduled in the Drug Control Act (§ 54.1-3400 et seq.) or (ii) be construed to prohibit any conduct 2041 2042 permitted under Article 4.2 of Chapter 34 of Title 54.1 of the Code of Virginia;

2043 70. 71. Selling or offering for sale any substance intended for human consumption, orally or by 2044 inhalation, that contains tetrahydrocannabinol, unless such substance is (i) contained in child-resistant 2045 packaging, as defined in § 4.1-600; (ii) equipped with a label that states, in English and in a font no less 2046 than 1/16 of an inch, (a) that the substance contains tetrahydrocannabinol and may not be sold to 2047 persons younger than 21 years of age, (b) all ingredients contained in the substance, (c) the amount of 2048 such substance that constitutes a single serving, and (d) the total percentage and milligrams of 2049 tetrahydrocannabinol included in the substance and the number of milligrams of tetrahydrocannabinol 2050 that are contained in each serving; and (iii) accompanied by a certificate of analysis, produced by an independent laboratory that is registered with the U.S. Drug Enforcement Administration and accredited 2051 2052 pursuant to standard ISO/IEC 17025 of the International Organization of Standardization by a third-party accrediting body, that states the tetrahydrocannabinol concentration of the substance or the 2053 2054 tetrahydrocannabinol concentration of the batch from which the substance originates. This subdivision 2055 shall not (i) apply to products that are approved for marketing by the U.S. Food and Drug 2056 Administration and scheduled in the Drug Control Act (§ 54.1-3400 et seq.) or (ii) be construed to prohibit any conduct permitted under Article 4.2 of Chapter 34 of Title 54.1 of the Code of Virginia; 2057

2058 71. 72. Manufacturing, offering for sale at retail, or selling at retail an industrial hemp extract, as 2059 defined in § 3.2-5145.1, a food containing an industrial hemp extract, or a substance containing 2060 tetrahydrocannabinol that depicts or is in the shape of a human, animal, vehicle, or fruit; and

2061 72. 73. Selling or offering for sale any substance intended for human consumption, orally or by 2062 inhalation, that contains tetrahydrocannabinol and, without authorization, bears, is packaged in a 2063 container or wrapper that bears, or is otherwise labeled to bear the trademark, trade name, famous mark 2064 as defined in 15 U.S.C. § 1125, or other identifying mark, imprint, or device, or any likeness thereof, of 2065 a manufacturer, processor, packer, or distributor of a product intended for human consumption other 2066 than the manufacturer, processor, packer, or distributor that did in fact so manufacture, process, pack, or 2067 distribute such substance; and

2068 74. Selling or offering for sale a topical hemp product that does not contain a bittering agent that 2069 renders the product unpalatable. As used in this subdivision, "topical hemp product" means a hemp 2070 product, as defined in § 3.2-4112, that (i) is intended to be rubbed, poured, sprinkled, or sprayed on, 2071 introduced into, or otherwise applied to the human body and (ii) is not intended to be consumed orally 2072 or by inhalation. This subdivision shall not (a) apply to products that are approved for marketing by the U.S. Food and Drug Administration and scheduled in the Drug Control Act (§ 54.1-3400 et seq.) or (b) 2073 be construed to prohibit any conduct permitted under Article 4.2 (§ 54.1-3442.5 et seq.) of Chapter 34 2074 2075 of Title 54.1.

2076 B. Nothing in this section shall be construed to invalidate or make unenforceable any contract or 2077 lease solely by reason of the failure of such contract or lease to comply with any other law of the 2078 Commonwealth or any federal statute or regulation, to the extent such other law, statute, or regulation 2079 provides that a violation of such law, statute, or regulation shall not invalidate or make unenforceable 2080 such contract or lease. 2081

§ 59.1-203. Restraining prohibited acts.

2082 A. Notwithstanding any other provisions of law to the contrary, the Attorney General, any attorney 2083 for the Commonwealth, or the attorney for any city, county, or town may cause an action to be brought 2084 in the appropriate circuit court in the name of the Commonwealth, or of the county, city, or town to enjoin any violation of § 59.1-200 or 59.1-200.1. The circuit court having jurisdiction may enjoin such 2085 2086 violations notwithstanding the existence of an adequate remedy at law. In any action under this section, 2087 it shall not be necessary that damages be proved.

2088 B. Unless the Attorney General, any attorney for the Commonwealth, or the attorney for any county, 2089 city, or town determines that a person subject to the provisions of this chapter intends to depart from

this Commonwealth or to remove his property herefrom, or to conceal himself or his property herein, or on a reasonable determination that irreparable harm may occur if immediate action is not taken, he shall, before initiating any legal proceedings as provided in this section, give notice in writing that such proceedings are contemplated, and allow such person a reasonable opportunity to appear before said attorney and show that a violation did not occur or execute an assurance of voluntary compliance, as provided in § 59.1-202.

2096 C. The circuit courts are authorized to issue temporary or permanent injunctions to restrain and prevent violations of § 59.1-200 or 59.1-200.1.

D. The Commissioner of the Department of Agriculture and Consumer Services, or his duly authorized representative, shall have the power to inquire into possible violations of subdivisions A 18, 28, 29, 31, 39, and 41, as it relates to motor fuels, 69, 70, 71, 72, 73, and 74 of § 59.1-200 and § 59.1-335.12, and, if necessary, to request, but not to require, an appropriate legal official to bring an action to enjoin such violation.

§ 59.1-206. Civil penalties; attorney's fees.

2104 A. In any action brought under this chapter, if the court finds that a person has willfully engaged in 2105 an act or practice in violation of § 59.1-200 or 59.1-200.1, the Attorney General, the attorney for the 2106 Commonwealth, or the attorney for the county, city, or town may recover for the Literary Fund, upon 2107 petition to the court, a civil penalty of not more than \$2,500 per violation. If the court finds that a 2108 person has willfully committed a second or subsequent violation of subdivision A 69, 70, 71, 72, 73, or 2109 74 of § 59.1-200, the Attorney General, the attorney for the Commonwealth, or the attorney for the 2110 county, city, or town may recover for the Literary Fund, upon petition to the court, a civil penalty of not more than \$5,000 per violation. 2111

B. For purposes of this section, prima facie evidence of a willful violation may be shown when the Attorney General, the attorney for the Commonwealth, or the attorney for the county, city, or town notifies the alleged violator by certified mail that an act or practice is a violation of § 59.1-200 or 59.1-200.1, and the alleged violator, after receipt of said notice, continues to engage in the act or practice.

B. C. Any person who willfully violates the terms of an assurance of voluntary compliance or an injunction issued under § 59.1-203 shall forfeit and pay to the Literary Fund a civil penalty of not more than \$5,000 per violation. For purposes of this section, the circuit court issuing an injunction shall retain jurisdiction, and the cause shall be continued, and in such cases the Attorney General, the attorney for the Commonwealth, or the attorney for the county, city, or town may petition for recovery of civil penalties.

 C. D. In any action pursuant to subsection A Θr , B, or C and in addition to any other amount awarded, the Attorney General, the attorney for the Commonwealth, or the attorney for the county, city, or town may recover any applicable civil penalty or penalties, costs, reasonable expenses incurred by the state or local agency in investigating and preparing the case not to exceed \$1,000 per violation, and attorney's fees. Such civil penalty or penalties, costs, reasonable expenses, and attorney's fees shall be paid into the general fund of the Commonwealth or of the county, city, or town which such attorney represented.

2130 D. E. Nothing in this section shall be construed as limiting the power of the court to punish as
2131 contempt the violation of any order issued by the court, or as limiting the power of the court to enter
2132 other orders under § 59.1-203 or 59.1-205.

2133 E. F. The right of trial by jury as provided by law shall be preserved in actions brought under this section.

2135 2. That the provisions of this act may result in a net increase in periods of imprisonment or

2136 commitment. Pursuant to § 30-19.1:4 of the Code of Virginia, the estimated amount of the 2137 necessary appropriation is \$0 for periods of imprisonment in state adult correctional facilities and

2137 necessary appropriation is so for periods of imprisonment in state adult correctional facilities and 2138 cannot be determined for periods of commitment to the custody of the Department of Juvenile

2139 Justice.

2103