REFERENCE TITLE: hemp-derived manufactured impairing cannabinoids

State of Arizona Senate Fifty-sixth Legislature Second Regular Session 2024

SB 1401

Introduced by Senators Shamp: Carroll, Gowan, Kerr

AN ACT

AMENDING SECTIONS 13-3401, 15-712 AND 32-1901, ARIZONA REVISED STATUTES; RELATING TO DRUG OFFENSES.

(TEXT OF BILL BEGINS ON NEXT PAGE)

1 Be it enacted by the Legislature of the State of Arizona: 2 Section 1. Section 13-3401, Arizona Revised Statutes, is amended to 3 read: 4 13-3401. Definitions 5 In this chapter, unless the context otherwise requires: 6 1. "Administer" means to apply, inject or facilitate the inhalation 7 or ingestion of a substance to the body of a person. 8 2. "Amidone" means substance identified chemically any as 9 (4-4-diphenyl-6-dimethylamine-heptanone-3), or any salt of such substance, 10 by whatever trade name designated. 11 3. "Board" means the Arizona state board of pharmacy. 12 4. "Cannabis" means the following substances under whatever names 13 they may be designated: (a) The resin extracted from any part of a plant of the genus 14 cannabis, and every compound, manufacture, salt, derivative, mixture or 15 16 preparation of such plant, its seeds or its resin. Cannabis does not 17 include oil or cake made from the seeds of such plant, any fiber, 18 compound, manufacture, salt, derivative, mixture or preparation of the 19 mature stalks of such plant except the resin extracted from the stalks or 20 any fiber, oil or cake or the sterilized seed of such plant which is 21 incapable of germination. 22 (b) Every compound, manufacture, salt, derivative, mixture or 23 preparation of such resin or tetrahydrocannabinol. 24 5. "Coca leaves" means cocaine, its optical isomers and any compound, manufacture, salt, derivative, mixture or preparation of coca 25 26 leaves, except derivatives of coca leaves which do not contain cocaine, 27 ecgonine or substances from which cocaine or ecgonine may be synthesized 28 or made. 29 "Dangerous drug" means the following by whatever official, 6. 30 common, usual, chemical or trade name designated: 31 (a) Any material, compound, mixture or preparation that contains 32 any quantity of the following hallucinogenic substances and their salts, 33 isomers, whether optical, positional or geometric, and salts of isomers, 34 unless specifically excepted, whenever the existence of such salts, 35 isomers and salts of isomers is possible within the specific chemical 36 designation: 37 (i) Alpha-ethyltryptamine. 38 (ii) Alpha-methyltryptamine. 39 (iii) (2-aminopropyl) benzofuran (APB). (iv) (2-aminopropyl)-2, 3-dihydrobenzofuran (APDB). 40 41 (v) Aminorex. 42 (vi) 4-bromo-2, 5-dimethoxyphenethylamine. 43 (vii) 4-bromo-2, 5-dimethoxyamphetamine. 44 (viii) Bufotenine.

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1
           (ix) [3-(3-carbamoylphenyl)phenyl]N-cyclohexyl carbamate (URB-597).
 2
           (x) Diethyltryptamine.
 3
           (xi) 2, 5-dimethoxyamphetamine.
 4
           (xii) Dimethyltryptamine.
 5
           (xiii) (2-ethylaminopropyl)-benzofuran (EAPB).
 6
           (xiv) 5-methoxy-alpha-methyltryptamine.
 7
           (xv) 5-methoxy-3, 4-methylenedioxyamphetamine.
 8
           (xvi) 4-methyl-2, 5-dimethoxyamphetamine.
 9
           (xvii) (2-methylaminopropyl)-benzofuran (MAPB).
10
           (xviii) Ibogaine.
11
           (xix) Lysergic acid amide.
           (xx) Lysergic acid diethylamide.
12
13
           (xxi) Mescaline.
14
           (xxii) 4-methoxyamphetamine.
15
           (xxiii) Methoxymethylenedioxyamphetamine (MMDA).
16
           (xxiv) Methylenedioxyamphetamine (MDA).
17
           (xxv) 3, 4-methylenedioxymethamphetamine.
18
           (xxvi) 3, 4-methylenedioxy-N-ethylamphetamine.
19
           (xxvii) N-ethyl-3-piperidyl benzilate (JB-318).
20
           (xxviii) N-hydroxy-3, 4-methylenedioxyamphetamine.
21
           (xxix) N-methyl-3-piperidyl benzilate (JB-336).
22
           (xxx) N-methyltryptamine mimetic substances that are any substances
23
     derived from N-methyltryptamine by any substitution at the nitrogen, any
24
     substitution at the indole ring, any substitution at the alpha carbon, any
25
     substitution at the beta carbon or any combination of the above.
26
     N-methyltryptamine
                          mimetic
                                    substances
                                                 do
                                                      not
                                                            include
                                                                      melatonin
27
     (5-methoxy n-acetyltryptamine). Substances
                                                  in
                                                       the
                                                             N-methyltryptamine
     generic definition include AcO-DMT, Baeocystine, Bromo-DALT, DiPT, DMT,
28
29
     DPT, HO-DET, HO-DIPT, HO-DMT, HO-DPT, HO-MET, MeO-DALT, MeO-DET, MeO-DiPT,
30
     MeO-DMT, MeO-DPT, MeO-NMT, MET, NMT and Norbufotenin.
31
           (xxxi) N-(1-phenylcyclohexyl) ethylamine (PCE).
32
           (xxxii) Nabilone.
33
           (xxxiii) 1-(1-phenylcyclohexyl) pyrrolidine (PHP).
34
           (xxxiv) 1-(1-(2-thienyl)-cyclohexyl) piperidine (TCP).
35
           (xxxv) 1-(1-(2-thienyl)-cyclohexyl) pyrrolidine.
36
           (xxxvi) Para-methoxyamphetamine (PMA).
37
           (xxxvii) Psilacetin.
38
           (xxxviii) Psilocybin.
39
           (xxxix) Psilocyn.
40
           (x1) Synhexyl.
41
           (xli) Trifluoromethylphenylpiperazine (TFMPP).
42
           (xlii) Trimethoxyamphetamine (TMA).
43
           (xliii) 1-pentyl-3-(naphthoyl)indole (JWH-018 and isomers).
44
           (xliv) 1-butyl-3-(naphthoyl)indole (JWH-073 and isomers).
45
           (xlv) 1-hexyl-3-(naphthoyl)indole (JWH-019 and isomers).
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1 (xlvi) 1-pentyl-3-(4-chloro naphthoyl)indole (JWH-398 and isomers). 2 (xlvii) 1-(2-(4-(morpholinyl)ethyl))-3-(naphthoyl)indole (JWH-200 3 and isomers). 4 (xlviii) 1-pentyl-3-(methoxyphenylacetyl)indole (JWH-250 and 5 isomers). 6 (xlix) (2-methyl-1-propyl-1H-indol-3-YL)-1-naphthalenyl-methanone 7 (JWH-015 and isomers). 8 (1) (6AR, 10AR)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan2-9 YL)-6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol) (HU-210). 10 (li) 5-(1,1-dimethylheptyl)-2-(3-hydroxycyclohexyl)-phenol 11 (CP 47,497 and isomers). 12 (lii) 5-(1,1-dimethyloctyl)-2-(3-hydroxycyclohexyl)-phenol (cannabicyclohexanol, CP-47,497 C8 homologue and isomers). 13 (b) Any material, compound, mixture or preparation that contains 14 15 any quantity of cannabimimetic substances and their salts, isomers, 16 whether optical, positional or geometric, and salts of isomers, unless specifically excepted, whenever the existence of such salts, isomers and 17 18 salts of isomers is possible within the specific chemical designation. For the purposes of this subdivision, "cannabimimetic substances" means 19 20 any substances within the following structural classes: 21 (i) 2-(3-hydroxycyclohexyl)phenol with substitution the at 22 5-position of the phenolic ring by alkyl or alkenyl, whether or not 23 substituted on the cyclohexyl ring to any extent. Substances in the 24 2-(3-hydroxycyclohexyl)phenol generic definition include CP-47,497, CP-47,497 C8-Homolog, CP-55,940 and CP-56,667. 25 26 (ii) 3-(naphthoyl)indole or 3-(naphthylmethane)indole by substitution at the nitrogen atom of the indole ring, whether or not 27 28 further substituted on the indole ring to any extent, whether or not substituted on the naphthoyl or naphthyl ring to any extent. 29 Substances 30 in the 3-(naphthoyl)indole generic definition include AM-678, AM-2201, 31 JWH-004, JWH-007, JWH-009, JWH-015, JWH-016, JWH-018, JWH-019, JWH-020, 32 JWH-046, JWH-047, JWH-048, JWH-049, JWH-050, JWH-070, JWH-071, JWH-072, 33 JWH-073, JWH-076, JWH-079, JWH-080, JWH-081, JWH-082, JWH-094, JWH-096, 34 JWH-098, JWH-116, JWH-120, JWH-122, JWH-148, JWH-149, JWH-175, JWH-180, 35 JWH-181, JWH-182, JWH-184, JWH-185, JWH-189, JWH-192, JWH-193, JWH-194, 36 JWH-195, JWH-196, JWH-197, JWH-199, JWH-200, JWH-210, JWH-211, JWH-212, 37 JWH-213, JWH-234, JWH-235, JWH-236, JWH-239, JWH-240, JWH-241, JWH-242, 38 JWH-262, JWH-386, JWH-387, JWH-394, JWH-395, JWH-397, JWH-398, JWH-399, 39 JWH-400, JWH-412, JWH-413, JWH-414 and JWH-415. 40 (iii) 3-naphthoyl-indazole or 3-(naphthylmethane)-indazole by 41 substitution at one or both of the nitrogen atoms of the indazole ring, 42 whether or not further substituted on the indazole ring to any extent, 43 whether or not substituted on the naphthoyl ring to any extent. 44 Substances in the 3-naphthoyl-indazole or 3-(naphthylmethane)-indazole 45 generic definition include THJ2201 and THJ-018.

1 (iv) 3-(naphthoyl)pyrrole by substitution at the nitrogen atom of 2 the pyrrole ring, whether or not further substituted in the pyrrole ring 3 to any extent, whether or not substituted on the naphthoyl ring to any 4 extent. Substances in the 3-(naphthoyl)pyrrole generic definition include 5 JWH-030, JWH-145, JWH-146, JWH-147, JWH-150, JWH-156, JWH-243, JWH-244, 6 JWH-245, JWH-246, JWH-292, JWH-293, JWH-307, JWH-308, JWH-346, JWH-348, 7 JWH-363, JWH-364, JWH-365, JWH-367, JWH-368, JWH-369, JWH-370, JWH-371, 8 JWH-373 and JWH-392.

9 (v) 1-(naphthylmethylene)indene by substitution of the 3-position 10 of the indene ring, whether or not further substituted in the indene ring 11 to any extent, whether or not substituted on the naphthyl ring to any 12 extent. Substances in the 1-(naphthylmethylene)indene generic definition 13 include JWH-176.

(vi) 3-(phenylacetyl)indole or 3-(benzoyl)indole by substitution at 14 the nitrogen atom of the indole ring, whether or not further substituted 15 16 in the indole ring to any extent, whether or not substituted on the phenyl 17 ring to any extent. Substances in the 3-(phenylacetyl)indole generic definition include AM-694, AM-2233, JWH-167, JWH-201, JWH-202, JWH-203, 18 JWH-204, JWH-205, JWH-206, JWH-207, JWH-208, JWH-209, JWH--237, JWH-248, 19 20 JWH-250, JWH-251, JWH-253, JWH-302, JWH-303, JWH-304, JWH-305, JWH-306, 21 JWH-311, JWH-312, JWH-313, JWH-314, JWH-315, JWH-316, RCS-4, RCS-8, SR-18 22 and SR-19.

23 (vii) 3-(cyclopropylmethanone) indole or 3-(cyclobutylmethanone) 24 indole or 3-(cyclopentylmethanone) indole by substitution at the nitrogen 25 atom of the indole ring, whether or not further substituted in the indole 26 ring to any extent, whether or not substituted on the cyclopropyl, 27 cyclobutyl or cyclopentyl rings to any extent. Substances in the 28 3-(cyclopropylmethanone) indole generic definition include UR-144, 29 fluoro-UR-144 and XLR-11.

30 (viii) 3-adamantoylindole with substitution at the nitrogen atom of 31 the indole ring, whether or not further substituted on the indole ring to 32 any extent, whether or not substituted on the adamantyl ring to any 33 extent. Substances in the 3-adamantoylindole generic definition include 34 AB-001.

35 (ix) N-(adamantyl)-indole-3-carboxamide with substitution at the 36 nitrogen atom of the indole ring, whether or not further substituted on 37 the indole ring to any extent, whether or not substituted on the adamantyl 38 ring to any extent. Substances in the N-(adamantyl)-indole-3-carboxamide 39 generic definition include SDB-001.

40 with (x) Indole-3-carboxamide or indazole-3-carboxamide 41 substitution at the nitrogen atom of the indole ring or by substitution at 42 one or both of the nitrogen atoms of the indazole ring, whether or not 43 further substituted on the indole ring or the indazole ring to any extent, 44 whether or not substituted on the nitrogen of the carboxamide to any 45 Substances in the indole-3-carboxamide or indazole-3-carboxamide extent.

generic definition include AKB-48, fluoro-AKB-48, APINACA, AB-PINACA,
 AB-FUBINACA, ABICA and ADBICA.

3 (xi) 8-Quinolinyl-indole-3-carboxylate or 8-quinolinyl-indazole-3-4 carboxylate by substitution at the nitrogen atom of the indole ring or by 5 substitution at one or both of the nitrogen atoms of the indazole ring, 6 whether or not further substituted in the indole ring or indazole ring to 7 any extent, whether or not substituted on the quinoline ring to any 8 in the 8-quinolinyl-indole-3-carboxylate or extent. Substances the 9 8-quinolinyl-indazole-3-carboxylate generic definition include PB-22, fluoro-PB-22, NPB-22 and fluoro-NPB-22. 10

11 (xii) Naphthalenyl-indole-3-carboxylate or naphthalenyl-indazole-3-12 carboxylate by substitution at the nitrogen atom of the indole ring or by 13 substitution at one or both of the nitrogen atoms of the indazole ring, whether or not further substituted in the indole or indazole ring to any 14 15 extent, whether or not substituted on the naphthalenyl ring to any extent. 16 Substances in the naphthalenyl-indole-3-carboxylate or 17 naphthalenyl-indazole-3-carboxylate generic definition include NM2201, 18 FDU-PB-22, SDB-005 and fluoro SDB-005.

19 (c) Any material, compound, mixture or preparation that contains 20 any quantity of the following substances and their salts, isomers, whether 21 optical, positional or geometric, and salts of isomers having a potential 22 for abuse associated with a stimulant effect on the central nervous 23 system:

24

(i) Alpha-pyrrolidinobutiophenone (Alpha-PBP).

25

(ii) Alpha-pyrrolidinopropiophenone (Alpha-PPP).

26 27

(iii) Alpha-pyrrolidinovalerophenone (Alpha-PVP).(iv) Alpha-pyrrolidinovalerothiophenone (Alpha-PVT).

28 (v) Aminoindane mimetic substances that are derived from 29 aminoindane by any substitution at the indane ring, replacement of the 30 amino group with another N group or any combination of the above. 31 Substances in the aminoindane generic definition include MDAI, MMAI, IAI 32 and AMMI.

33 (vi) Amphetamine.

34 (vii) Benzphetamine.

35 36

37

(viii) Benzylpiperazine (BZP).

(ix) Beta-keto-n-methylbenzodioxolylbutanamine (Butylone).

(x) Beta-keto-n-methylbenzodioxolylpentanamine (Pentylone).

38 (xi) Butorphanol.

39 (xii) Cathine ((+)-norpseudoephedrine).

40 (xiii) Cathinomimetic substances that are any substances derived 41 from cathinone, (2-amino-1-phenyl-1-propanone) by any substitution at the 42 phenyl ring, any substitution at the 3 position, any substitution at the 43 nitrogen atom or any combination of the above substitutions.

44 (xiv) Cathinone.

45 (xv) 2-(4-Chloro-2,5-dimethoxyphenyl)ethanamine (2C-C).

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1
           (xvi) Chlorphentermine.
 2
           (xvii) Clortermine.
 3
           (xviii) Diethylpropion.
 4
           (xix) Dihydro-5H-indeno-(5,6-d)-1,3-dioxol-6-amine) (MDAI).
 5
           (xx) 2-(2,5-Dimethoxy-4-ethylphenyl)ethanamine (2C-E).
 6
           (xxi) 2-(2,5-Dimethoxy-4-methylphenyl)ethanamine (2C-D).
 7
           (xxii) 2-(2,5-Dimethoxy-4-nitro-phenyl)ethanamine (2C-N).
 8
           (xxiii) 2-(2,5-Dimethoxy-4-(n)-propylphenyl)ethanamine (2C-P).
 9
           (xxiv) 2-(2,5-Dimethoxyphenyl)ethanamine (2C-H).
10
           (xxv) Dimethylcathinone (Metamfepramone).
11
           (xxvi) Ethcathinone.
           (xxvii) 2-[4-(Ethylthio)-2,5-dimethoxyphenyl]ethanamine (2C-T-2).
12
13
           (xxviii) Fencamfamin.
           (xxix) Fenethylline.
14
15
           (xxx) Fenproporex.
16
           (xxxi) Fluoroamphetamine.
17
           (xxxii) Fluoromethamphetamine.
18
           (xxxiii) Fluoromethcathinone.
           (xxxiv) 2-(4-Iodo-2,5-dimethoxyphenyl)ethanamine (2C-I).
19
20
           (xxxv) 2-[4-(Isopropylthio)-2,5-dimethoxyphenyl]ethanamine(2C-T-4).
21
           (xxxvi) Mazindol.
22
           (xxxvii) Mefenorex.
23
           (xxxviii) Methamphetamine.
24
           (xxxix) Methcathinone.
25
           (x1) Methiopropamine.
26
           (xli) Methoxy-alpha-pyrrolidinopropiophenone (MOPPP).
27
           (xlii) Methoxymethcathinone (methedrone).
28
           (xliii) Methoxyphenethylamine mimetic substances that
                                                                       are
                                                                            any
29
     substances derived from 2, 5-dimethoxy-phenethylamine by any substitution
30
         the phenyl ring, any substitution at the nitrogen atom,
     at
                                                                            any
31
     substitutions at the carbon atoms of the ethylamine, or any combination of
32
     the above substitutions.
33
           (xliv) 4-methylaminorex.
34
           (xlv) Methyl-a-pyrrolidinobutiophenone (MPBP).
35
           (xlvi) Methylenedioxy-alphapyrrolidinopropiophenone (MDPPP).
36
           (xlvii) Methylenedioxyethcathinone (Ethylone).
37
           (xlviii) Methylenedioxymethcathinone (Methylone).
38
           (xlix) Methylenedioxypyrovalerone (MDPV).
39
           (1) Methylmethcathinone (Mephedrone).
40
           (li) Methylphenidate.
41
           (lii) Modafinil.
42
           (liii) Naphthylpyrovalerone (Naphyrone).
43
           (liv) N-ethylamphetamine.
44
           (lv) N, N-dimethylamphetamine.
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1 (lvi) Pemoline. 2 (lvii) Phendimetrazine. 3 (lviii) Phenmetrazine. 4 (lix) Phentermine. 5 (lx) Pipradol. 6 (lxi) Propylhexedrine. 7 (lxii) Pyrovalerone. 8 (lxiii) Sibutramine. 9 (lxiv) Spa ((-)-1-dimethylamino-1,2-diphenylethane). 10 (d) Any material, compound, mixture or preparation that contains 11 any quantity of the following substances having a potential for abuse 12 associated with a depressant effect on the central nervous system: 13 (i) Any substance which contains any quantity of a derivative of 14 barbituric acid, or any salt of a derivative of barbituric acid, unless 15 specifically excepted. 16 (ii) Alprazolam. 17 (iii) Bromazepam. 18 (iv) Camazepam. 19 (v) Carisoprodol. 20 (vi) Chloral betaine. 21 (vii) Chloral hydrate. 22 (viii) Chlordiazepoxide. 23 (ix) Chlorhexadol. 24 (x) Clobazam. 25 (xi) Clonazepam. 26 (xii) Clorazepate. 27 (xiii) Clotiazepam. (xiv) Cloxazolam. 28 29 (xv) Delorazepam. 30 (xvi) Diazepam. 31 (xvii) Dichloralphenazone. 32 (xviii) Estazolam. 33 (xix) Ethchlorvynol. 34 (xx) Ethinamate. 35 (xxi) Ethyl loflazepate. 36 (xxii) Etizolam. 37 (xxiii) Fenfluramine. 38 (xxiv) Fludiazepam. 39 (xxv) Flunitrazepam. 40 (xxvi) Flurazepam. 41 (xxvii) Gamma hydroxy butyrate. (xxviii) Glutethimide. 42 43 (xxix) Halazepam. (xxx) Haloxazolam. 44 45 (xxxi) Hydroxyphencyclidine (HO-PCP).

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1
           (xxxii) Ketamine.
 2
           (xxxiii) Ketazolam.
 3
           (xxxiv) Loprazolam.
 4
           (xxxv) Lorazepam.
 5
           (xxxvi) Lormetazepam.
 6
           (xxxvii) Lysergic acid.
 7
           (xxxviii) Mebutamate.
 8
           (xxxix) Meclogualone.
 9
           (x1) Medazepam.
10
           (xli) Meprobamate.
11
           (xlii) Methagualone.
12
           (xliii) Methohexital.
13
           (xliv) 2-(methoxyphenyl)-2-(ethylamino)cyclohexanone
14
     (Methoxetamine).
           (xlv) 2-(methoxyphenyl)-2-(methylamino)cyclohexanone
15
16
           (Methoxyketamine).
17
           (xlvi) Methoxyphencyclidine(MeO-PCP).
18
           (xlvii) Methyprylon.
19
           (xlviii) Midazolam.
20
           (xlix) Nimetazepam.
21
           (1) Nitrazepam.
22
           (li) Nordiazepam.
23
           (lii) Oxazepam.
24
           (liii) Oxazolam.
25
           (liv) Paraldehyde.
26
           (lv) Petrichloral.
27
           (lvi) Phencyclidine (PCP).
           (lvii) Phencyclidine mimetic substances that are any substances
28
29
     derived from phenylcyclohexylpiperidine by any substitution at the phenyl
30
     ring, any substitution at the piperidine ring, any substitution at the
31
     cyclohexyl ring, any replacement of the phenyl ring or any combination of
                                           phenylcyclohexylpiperidine
32
     the
           above.
                   Substances
                                in
                                     the
                                                                         generic
                            Amino-PCP,
33
     definition
                  include
                                         BCP,
                                               Bromo-PCP,
                                                             BTCP.
                                                                     Chloro-PCP.
34
     Fluoro-PCP, HO-PCP, MeO-PCP, Methyl-PCP, Nitro-PCP, Oxo-PCP, PCE, PCM,
35
     PCPY, TCP and TCPY.
36
           (lviii) Pinazepam.
37
           (lix) Prazepam.
38
           (lx) Scopolamine.
39
           (lxi) Sulfondiethylmethane.
40
           (lxii) Sulfonethylmethane.
41
           (lxiii) Sulfonmethane.
42
           (lxiv) Quazepam.
43
           (lxv) Temazepam.
44
           (lxvi) Tetrazepam.
45
           (lxvii) Tiletamine.
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1 (lxviii) Triazolam. 2 (lxix) Zaleplon. 3 (lxx) Zolazepam. 4 (lxxi) Zolpidem. 5 (lxxii) Zopiclone. 6 (e) Any material, compound, mixture or preparation that contains 7 any quantity of the following anabolic steroids and their salts, isomers 8 or esters: 9 (i) Boldenone. (ii) Clostebol (4-chlorotestosterone). 10 11 (iii) Dehydrochloromethyltestosterone. 12 (iv) Drostanolone. 13 (v) Ethylestrenol. (vi) Fluoxymesterone. 14 (vii) Formebulone (formebolone). 15 16 (viii) Mesterolone. 17 (ix) Methandriol. 18 (x) Methandrostenolone (methandienone). 19 (xi) Methenolone. 20 (xii) Methyltestosterone. 21 (xiii) Mibolerone. (xiv) Nandrolone. 22 23 (xv) Norethandrolon. 24 (xvi) Oxandrolone. 25 (xvii) Oxymesterone. 26 (xviii) Oxymetholone. 27 (xix) Stanolone (4-dihydrotestosterone). 28 (xx) Stanozolol. 29 (xxi) Testolactone. 30 (xxii) Testosterone. 31 (xxiii) Trenbolone. (f) ANY HEMP-DERIVED MANUFACTURED IMPAIRING CANNABINOIDS. 32 33 7. "Deliver" means the actual, constructive or attempted exchange 34 one person to another, whether or not there is an from agency 35 relationship. 36 8. "Director" means the director of the department of health 37 services. "Dispense" means distribute, leave with, give away, dispose of 38 9. or deliver. 39 40 10. "Drug court program" means a program that is established 41 pursuant to section 13-3422 by the presiding judge of the superior court 42 in cooperation with the county attorney in a county for the purpose of 43 prosecuting, adjudicating and treating drug dependent persons who meet the criteria and guidelines for entry into the program that are developed and 44 45 agreed on by the presiding judge and the prosecutor.

1 "Drug dependent person" means a person who is using a substance 11. 2 that is listed in paragraph 6, 19, 20, 21, 22 or 28 29 of this section and 3 who is in a state of psychological or physical dependence, or both, 4 arising from the use of that substance. 5 "Federal act" has the same meaning prescribed in section 12. 6 32-1901. 7 "HEMP-DERIVED MANUFACTURED IMPAIRING CANNABINOIDS": 13. 8 (a) MEANS ANY CANNABINOID THAT IS DERIVED FROM THE PLANT OF THE 9 GENUS CANNABIS OR ANY PART OF THE PLANT AND THAT IS ALTERED BY A CHEMICAL REACTION THAT CHANGES THE MOLECULAR STRUCTURE OF ANY NATURAL CANNABINOID 10 11 DERIVED FROM HEMP TO ANOTHER CANNABINOID WITH IMPAIRING PROPERTIES THAT IS 12 FOUND NATURALLY IN HEMP, INCLUDING ALL OF THE FOLLOWING: 13 (i) DELTA-8 TETRAHYDROCANNABINOL. 14 (ii) DELTA-9 TETRAHYDROCANNABINOL OVER 0.3 PERCENT ON A DRY-WEIGHT 15 BASIS, INCLUDING ANY PRODUCT MADE TO BE INGESTED. 16 (iii) DELTA-10 TETRAHYDROCANNABINOL. 17 (iv) HEXAHYDROCANNABINOL. 18 (v) TETRAHYDROCANNABIPHOROL. 19 (vi) TETRAHYDROCANNABINOL ACETATE ESTER. 20 (vii) TETRAHYDROCANNABIOCTYL. 21 (viii) TETRAHYDROCANNABIHEXOL. 22 (ix) TETRAHYDROCANNABINOLIC ACID. 23 (x) TETRAHYDROCANNABUTOL. 24 (xi) CANNABINOID THAT IS DERIVED FROM HEMP AND ALTERED BY A CHEMICAL REACTION THAT CHANGES THE MOLECULAR STRUCTURE OF ANY NATURAL 25 26 CANNABINOID DERIVED FROM HEMP TO ANOTHER CANNABINOID WITH IMPAIRING PROPERTIES WITH A SUBSTANTIALLY SIMILAR MOLECULAR STRUCTURE TO ANY 27 28 CANNABINOID THAT IS FOUND NATURALLY IN HEMP. 29 (b) DOES NOT INCLUDE ANY CANNABINOID DERIVED FROM HEMP THAT IS PRODUCED BY DECARBOXYLATION FROM NATURALLY OCCURRING CANNABINOID WITHOUT 30 31 THE USE OF A CHEMICAL CATALYST OR NONINTOXICATING CANNABINOIDS DERIVED 32 FROM HEMP, INCLUDING CANNABIDIOL, CANNABINOL, CANNABIGEROL. 33 CANNABICHROMENE, CANNABICYCLOL, CANNABIDIVARIN, CANNABIVARIN AND 34 TETRAHYDROCANNABIVARIN. 35 13. 14. "Isoamidone" means any substance identified chemically as 36 (4-4-diphenyl-5-methyl-6-dimethylaminohexanone-3), or any salt of such 37 substance, by whatever trade name designated. 14. 15. "Isonipecaine" means any substance identified chemically 38 39 as (1-methyl-4-phenyl-piperidine-4-carboxylic acid ethyl ester), or any 40 salt of such substance, by whatever trade name designated. 41 15. 16. "Ketobemidone" means any substance identified chemically 42 as (4-(3-hydroxyphenyl)-1-methyl-4-piperidylethyl ketone hydrochloride), 43 or any salt of such substance, by whatever trade name designated.

44 **16.** 17. "Licensed" or "permitted" means authorized by the laws of 45 this state to do certain things. 1 17. 18. "Manufacture" means produce, prepare, propagate, compound, 2 mix or process, directly or indirectly, by extraction from substances of 3 natural origin or independently by means of chemical synthesis, or by a 4 combination of extraction and chemical synthesis. Manufacture includes 5 any packaging or repackaging or labeling or relabeling of containers. 6 Manufacture does not include any producing, preparing, propagating, 7 compounding, mixing, processing, packaging or labeling done in conformity 8 with applicable state and local laws and rules by a licensed practitioner 9 incident to and in the course of his licensed practice.

10 18. 19. "Manufacturer" means a person who manufactures a narcotic 11 or dangerous drug or other substance controlled by this chapter.

12 19. 20. "Marijuana" means all parts of any plant of the genus 13 cannabis, from which the resin has not been extracted, whether growing or 14 not, and the seeds of such plant. Marijuana does not include the mature 15 stalks of such plant or the sterilized seed of such plant which is 16 incapable of germination.

17 20. 21. "Narcotic drugs" means the following, whether of natural 18 or synthetic origin and any substance neither chemically nor physically 19 distinguishable from them:

- 20 (a) Acetyl-alpha-methylfentanyl.
- 21 (b) Acetylmethadol.
- 22 (c) Alfentanil.
- 23 (d) Allylprodine.
- 24 (e) Alphacetylmethadol.
- 25 (f) Alphameprodine.
- 26 (g) Alphamethadol.
- 27 (h) Alpha-methylfentanyl.
- 28 (i) Alpha-methylthiofentanyl.
- 29 (j) Alphaprodine.
- 30 (k) Amidone (methadone).
- 31 (l) Anileridine.
- 32 (m) Benzethidine.
- 33 (n) Benzylfentanyl.
- 34 (o) Betacetylmethadol.
- 35 (p) Beta-hydroxyfentanyl.
- 36 (q) Beta-hydroxy-3-methylfentanyl.
- 37 (r) Betameprodine.
- 38 (s) Betamethadol.
- 39 (t) Betaprodine.
- 40 (u) Bezitramide.
- 41 (v) Buprenorphine and its salts.
- 42 (w) Cannabis.
- 43 (x) Carfentanil.
- 44 (y) 4-chloro-n-[-1-[2-(4-nitrophenyl)ethyl]-2-piperidinylidene]benz
- 45 enesulfonamide (W-18).

1	(z) A oblama n $\begin{bmatrix} 1 & (2 & nbay a b + b + 1) & 2 & ninanidiny a bidana \end{bmatrix}$
1 2	<pre>(z) 4-chloro-n-[1-(2-pheylethyl)-2-piperidinylidene] benzenesulfonamide (W-15).</pre>
2	(aa) Clonitazene.
4	(bb) Coca leaves.
5	(cc) 1-cyclohexyl-4-(1,2-diphenylethyl)piperazine (MT-45).
6	(dd) Dextromoramide.
7	(ee) Dextropropoxyphene.
8	(ff) Diampromide.
9	(gg) 3,4-dichloro-n-(-[1-(dimethylamino)cyclohexyl]methyl)-benzamid
10	e (AH-7921).
11	(hh) 3,4-dichloro-n-[2-(dimethylamino)cyclohexyl]-N-methylbenzamide
12	(U-47700).
13	(ii) Diethylthiambutene.
14	(jj) Difenoxin.
15	(kk) Dihydrocodeine.
16	(11) Dimenoxadol.
17	(mm) Dimepheptanol.
18	(nn) Dimethylthiambutene.
19	(oo) Dioxaphetyl butyrate.
20	(pp) Diphenidine (DEP).
21	(qq) Diphenoxylate.
22	(rr) Dipipanone.
23	(ss) Ephenidine.
24	(tt) Ethylmethylthiambutene.
25	(uu) Etonitazene.
26	(vv) Etoxeridine.
27	(ww) Fentanyl.
28	(xx) Fentanyl mimetic substances that are any substances derived
29	from fentanyl by any substitution in the phenethyl group, any substitution
30	in the piperidine ring, any substitution in the aniline ring, any
31	replacement of the phenyl portion of the phenethyl group, any replacement
32	of the N-propionyl group or any combination of the above.
33	(yy) Furethidine.
34	(zz) Hydroxypethidine.
35	(aaa) Isoamidone (isomethadone).
36	(bbb) Isophenidine.
37	(ccc) Pethidine (meperidine).
38	(ddd) Ketobemidone.
39	(eee) Lefetamine.
40	(fff) Levomethorphan.
41	(ggg) Levomoramide.
42	(hhh) Levophenacylmorphan.
43	(iii) Levorphanol.
44 4 F	(jjj) Metazocine.
45	(kkk) Methoxphenidine (MXP).

```
1
           (111) 3-methylfentanyl.
 2
                  1-methyl-4-phenyl-4-propionoxypiperidine (MPPP).
           (mmm)
 3
                 3-methylthiofentanyl.
           (nnn)
 4
           (000)
                  Morpheridine.
 5
                  Noracymethadol.
           (ppp)
 6
           (ppp)
                  Norlevorphanol.
 7
           (rrr) Normethadone.
 8
           (sss)
                  Norpipanone.
 9
           (ttt) Opium.
10
           (uuu) Para-fluorofentanyl.
11
           (vvv) Pentazocine.
12
           (www) Phenadoxone.
13
           (xxx) Phenampromide.
14
           (yyy) Phenazocine.
           (zzz) 1-(2-phenethyl)-4-phenyl-4-acetoxypiperidine (PEPAP).
15
16
           (aaaa) Phenomorphan.
17
           (bbbb) Phenoperidine.
18
           (cccc)
                   Piminodine.
19
           (dddd) Piritramide.
20
           (eeee) Proheptazine.
21
           (ffff)
                   Properidine.
22
           (gggg)
                   Propiram.
23
           (hhhh) Racemethorphan.
24
           (iiii)
                   Racemoramide.
25
           (jjjj)
                   Racemorphan.
26
           (kkkk) Remifentanil.
27
           (1111) Sufentanil.
28
           (mmmm)
                  Thenylfentanyl.
29
           (nnnn)
                  Thiofentanyl.
30
           (0000)
                  Tilidine.
31
                  Tramadol.
                                  2-[(dimethylamino)methyl]-1-(3-methoxyphenyl)
           (pppp)
     cyclohexanol, and its salts, optical and geometric isomers, and its salts
32
33
     of isomers.
34
           (qqqq) Trimeperidine.
35
                     "Opium" means any compound, manufacture, salt, isomer,
           <del>21.</del> 22.
36
     salt of isomer, derivative, mixture or preparation of the following, but
37
     does not include apomorphine or any of its salts:
38
           (a) Acetorphine.
39
           (b) Acetyldihydrocodeine.
40
           (c) Benzylmorphine.
41
           (d) Codeine.
42
           (e) Codeine methylbromide.
43
           (f) Codeine-N-oxide.
44
           (g) Cyprenorphine.
45
           (h)
                Desomorphine.
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1	(i) Dihydromorphine.
2	(j) Drotebanol.
3	(k) Ethylmorphine.
4	(1) Etorphine.
5	(m) Heroin.
6	(n) Hydrocodone.
7	(o) Hydromorphinol.
8	(p) Hydromorphone.
9	(q) Levo-alphacetylmethadol.
10	(r) Methyldesorphine.
11	(s) Methyldihydromorphine.
12	(t) Metopon.
13	(u) Morphine.
14	(v) Morphine methylbromide.
15	(w) Morphine methylsulfonate.
16	(x) Morphine-N-oxide.
17	(y) Myrophine.
18	(z) Nalorphine.
19	(aa) Nicocodeine.
20	(bb) Nicomorphine.
21	(cc) Normorphine.
22	(dd) Oxycodone.
23	(ee) Oxymorphone.
24	(ff) Pholcodine.
25	(gg) Thebacon.
26	(hh) Thebaine.
27	22. 23. "Ordinary ephedrine, pseudoephedrine,
28	(-)-norpseudoephedrine or phenylpropanolamine product" means a product
29	that contains ephedrine, pseudoephedrine, (-)-norpseudoephedrine or
30	phenylpropanolamine and that is all of the following:
31	(a) Approved for sale under the federal act.
32	(b) Labeled, advertised and marketed only for an indication that is
33	approved by the federal food and drug administration.
34	(c) Either:
35	(i) A nonliquid that is sold in package sizes of not more than
36	three grams of ephedrine, pseudoephedrine, (-)-norpseudoephedrine or
37	phenlypropanolamine and that is packaged in blister packs containing not
38	more than two dosage units or, if the use of blister packs is technically
39	infeasible, that is packaged in unit dose packets or pouches.
40	(ii) A liquid that is sold in package sizes of not more than three
41	grams of ephedrine, pseudoephedrine, (-)-norpseudoephedrine or
42	phenylpropanolamine.
43	23. 24. "Peyote" means any part of a plant of the genus
44	lophophora, known as the mescal button.

1 24. 25. "Pharmacy" means a licensed business where drugs are 2 compounded or dispensed by a licensed pharmacist. 3 25. 26. "Practitioner" means a person licensed to prescribe and 4 administer drugs. 5 26. 27. "Precursor chemical I" means any material, compound. 6 mixture or preparation which contains any quantity of the following 7 substances and their salts, optical isomers or salts of optical isomers: 8 (a) N-acetylanthranilic acid. 9 (b) Anthranilic acid. 10 (c) Ephedrine. 11 (d) Ergotamine. 12 (e) Isosafrole. 13 (f) Lysergic acid. 14 (g) Methylamine. 15 (h) N-ethylephedrine. 16 (i) N-ethylpseudoephedrine. 17 (j) N-methylephedrine. 18 (k) N-methylpseudoephedrine. 19 (1) Norephedrine. 20 (m) (-)-Norpseudoephedrine. 21 (n) Phenylacetic acid. 22 (o) Phenylpropanolamine. 23 (p) Piperidine. 24 (q) Pseudoephedrine. 27. 28. "Precursor chemical II" means any material, compound, 25 26 mixture or preparation which contains any quantity of the following 27 substances and their salts, optical isomers or salts of optical isomers: 28 (a) 4-cyano-2-dimethylamino-4, 4-diphenyl butane. 29 (b) 4-cyano-1-methyl-4-phenylpiperidine. (c) Chlorephedrine. 30 31 (d) Chlorpseudoephedrine. 32 (e) Ethyl-4-phenylpiperidine-4-carboxylate. 2-methyl-3-morpholino-1, 1-diphenylpropane-carboxylic acid. 33 (f) 34 (g) 1-methyl-4-phenylpiperidine-4-carboxylic acid. 35 (h) N-formyl amphetamine. 36 (i) N-formyl methamphetamine. 37 (j) Phenyl-2-propanone. 38 (k) 1-piperidinocyclohexane carbonitrile. 39 (1) 1-pyrrolidinocyclohexane carbonitrile. 40 28. 29. "Prescription-only drug" does not include a dangerous drug 41 or narcotic drug but means: (a) Any drug which because of its toxicity or other potentiality 42 43 for harmful effect, or the method of its use, or the collateral measures 44 necessary to its use, is not generally recognized among experts, qualified 45 by scientific training and experience to evaluate its safety and efficacy,

1 as safe for use except by or under the supervision of a medical 2 practitioner. (b) Any drug that is limited by an approved new drug application 3 4 under the federal act or section 32-1962 to use under the supervision of a 5 medical practitioner. 6 (c) Every potentially harmful drug, the labeling of which does not 7 bear or contain full and adequate directions for use by the consumer. 8 (d) Any drug required by the federal act to bear on its label the 9 legend "Caution: Federal law prohibits dispensing without prescription" 10 or "Rx only". 11 29. 30. "Produce" means grow, plant, cultivate, harvest, dry, 12 process or prepare for sale. 13 30. 31. "Regulated chemical" means the following substances in 14 bulk form that are not a useful part of an otherwise lawful product: 15 (a) Acetic anhydride. 16 (b) Hypophosphorous acid. 17 (c) Iodine. 18 (d) Sodium acetate. 19 (e) Red phosphorus. 20 (f) Gamma butyrolactone (GBL). 21 (g) 1, 4-butanediol. 22 (h) Butyrolactone. 23 (i) 1, 2 butanolide. 24 (j) 2-oxanalone. 25 (k) Tetrahydro-2-furanone. 26 Dihydro-2(3H)-furanone. 27 (m) Tetramethylene glycol. 31. 32. "Retailer" means either: 28 29 (a) A person other than a practitioner who sells any precursor chemical or regulated chemical to another person for purposes of 30 31 consumption and not resale, whether or not the person possesses a permit 32 issued pursuant to title 32, chapter 18. 33 (b) A person other than a manufacturer or wholesaler who purchases, receives or acquires more than twenty-four grams of a precursor chemical. 34 32. 33. "Sale" or "sell" means an exchange for anything of value 35 36 or advantage, present or prospective. 33. 34. "Sale for personal use" means the retail sale for a 37 legitimate medical use in a single transaction to an individual customer, 38 39 to an employer for dispensing to employees from first aid kits or medicine 40 chests or to a school for administration pursuant to section 15-344. 41 34. 35. "Scientific purpose" means research, teaching or chemical 42 analysis. 43 35. 36. "Suspicious transaction" means a transaction to which any 44 of the following applies: 45 (a) A report is required under the federal act.

1 (b) The circumstances would lead a reasonable person to believe 2 that any person is attempting to possess a precursor chemical or regulated 3 chemical for the purpose of unlawful manufacture of a dangerous drug or 4 narcotic drug, based on such factors as the amount involved, the method of 5 payment, the method of delivery and any past dealings with any 6 participant.

7 (c) The transaction involves payment for precursor or regulated 8 chemicals in cash or money orders in a total amount of more than \$200.

9 (d) The transaction involves a sale, a transfer or furnishing to a 10 retailer for resale without a prescription of ephedrine, pseudoephedrine, 11 (-)-norpseudoephedrine or phenylpropanolamine that is not an ordinary 12 ephedrine, pseudoephedrine, (-)-norpseudoephedrine or phenylpropanolamine 13 product.

14 36. 37. "Threshold amount" means a weight, market value or other 15 form of measurement of an unlawful substance as follows:

16

(a) One gram of heroin.(b) Nine grams of cospins

17 18 (b) Nine grams of cocaine.

18 (c) Seven hundred fifty milligrams of cocaine base or hydrolyzed 19 cocaine.

20

(d) Four grams or 50 milliliters of PCP.

(e) Nine grams of methamphetamine, including methamphetamine inliquid suspension.

23 (f) Nine grams of amphetamine, including amphetamine in liquid 24 suspension.

25 (g) One-half milliliter of lysergic acid diethylamide, or in the 26 case of blotter dosage units fifty dosage units.

(h) Two pounds of marijuana.

27 28

(i) Nine grams of fentanyl or fentanyl mimetic substances.

(j) For any combination consisting solely of those unlawful substances listed in subdivisions (a) through (i) of this paragraph, an amount equal to or in excess of the threshold amount, as determined by the application of section 13-3420.

33 (k) For any unlawful substance not listed in subdivisions (a) 34 through (i) of this paragraph or any combination involving any unlawful 35 substance not listed in subdivisions (a) through (i) of this paragraph, a 36 value of at least \$1,000.

37

37. 38. "Transfer" means furnish, deliver or give away.

38 38. "Vapor-releasing substance containing a toxic substance" 39 means a material which releases vapors or fumes containing any of the 40 following:

41 (a) Ketones, including acetone, methyl ethyl ketone, mibk, miak, 42 isophorone and mesityl oxide.

43 (b) Hydrocarbons, including propane, butane, pentane, hexane,44 heptane and halogenated hydrocarbons.

45 (c) Ethylene dichloride.

- 1 (d) Pentachlorophenol.
- 2 (e) Chloroform.
- 3 (f) Methylene chloride.
- 4 (g) Trichloroethylene.
- 5 (h) Difluoroethane.
- 6 (i) Tetrafluoroethane.
- 7 (j) Aldehydes, including formaldehyde.
 - (k) Acetates, including ethyl acetate and butyl acetate.
- 9 (1) Aromatics, including benzene, toluene, xylene, ethylbenzene and 10 cumene.
- 11 (m) Alcohols, including methyl alcohol, ethyl alcohol, isopropyl 12 alcohol, butyl alcohol and diacetone alcohol.
- 13

8

(n) Ether, including Diethyl ether and petroleum ether.(o) Nitrous oxide.

- 14 (o) Nitrous oxide15 (p) Amyl nitrite.
- 16 (q) Isobutyl nitrite.

17 39. 40. "Weight" unless otherwise specified includes the entire 18 weight of any mixture or substance that contains a detectable amount of an unlawful substance. If a mixture or substance contains more than one 19 20 unlawful substance, the weight of the entire mixture or substance is 21 assigned to the unlawful substance that results in the greater offense. 22 If a mixture or substance contains lysergic acid diethylamide, the offense that results from the unlawful substance shall be based on the greater 23 24 offense as determined by the entire weight of the mixture or substance or the number of blotter dosage units. For the purposes of this paragraph, 25 26 "mixture" means any combination of substances from which the unlawful 27 substance cannot be removed without a chemical process.

40. 41. "Wholesaler" means a person who in the usual course of 28 29 business lawfully supplies narcotic drugs, dangerous drugs, precursor 30 chemicals or regulated chemicals that he himself has not produced or 31 prepared, but not to a person for the purpose of consumption by the person, whether or not the wholesaler has a permit that is issued pursuant 32 33 to title 32, chapter 18. Wholesaler includes a person who sells, delivers or dispenses a precursor chemical in an amount or under circumstances that 34 35 would require registration as a distributor of precursor chemicals under 36 the federal act.

37 Sec. 2. Section 15-712, Arizona Revised Statutes, is amended to 38 read:

- 39
- 40

41 42 15-712. <u>Instruction on alcohol, tobacco, narcotic drugs,</u> <u>marijuana, date rape drugs and other dangerous</u> <u>drugs: chemical abuse prevention programs:</u> <u>definitions</u>

A. Instruction on the nature and harmful effects of alcohol,
tobacco, narcotic drugs, marijuana, date rape drugs and other dangerous
drugs on the human system and instruction on the laws related to the

1 control of these substances and the nonuse and prevention of use and abuse 2 of alcohol, tobacco, narcotic drugs, marijuana, date rape drugs and other 3 dangerous drugs may be included in the courses of study in common and high 4 schools, with emphasis on grades four through nine. Instruction on the 5 nature and harmful effects of alcohol, tobacco, narcotic drugs, marijuana, 6 date rape drugs and other dangerous drugs on a human fetus may be included 7 in the courses of study in grades six through twelve. The instruction may 8 be integrated into existing health, science, citizenship or similar 9 studies and shall meet the criteria for chemical abuse prevention education programs developed pursuant to subsection C of this section. 10

11 B. At the request of a school district, the department of education 12 shall provide technical assistance to school districts that choose to 13 implement programs to prevent chemical abuse.

14 C. The department of education and the department of health 15 services, in consultation with the committee established pursuant to 16 section 41-617, shall establish an interagency committee to coordinate 17 their assistance to school districts.

18 D. The state board of education may accept gifts and grants and 19 distribute them and monies appropriated for chemical shall abuse 20 prevention programs to school districts to assist with the costs of 21 programs designed to prevent chemical abuse by pupils in kindergarten 22 programs and grades one through twelve. School districts which THAT have 23 approved chemical abuse prevention policies and procedures as prescribed 24 in section 15–345 are eligible for a maximum of one dollar \$1 for each pupil or one thousand dollars \$1,000, whichever is more. If sufficient 25 26 monies are not available to meet all requests, the state board shall determine which school districts to fund based on need, availability of 27 other programs or sources of revenue and the likelihood of the school 28 29 district's proposed program successfully meeting needs identified by the school district. A school district shall include the monies it receives 30 31 for chemical abuse prevention programs under this section in the special 32 projects section of the budget as provided in section 15-903. 33 subsection F.

34

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E. For the purpose of this section:

35 1. "Date rape drug" means a drug prescribed LISTED in section 36 13-3401, paragraph 30 31, subdivisions (f) through (m).

37 2. "Narcotic drug", "marijuana" and "dangerous drug" have the same
 38 meaning MEANINGS prescribed in section 13-3401.

39 Sec. 3. Section 32–1901, Arizona Revised Statutes, is amended to 40 read:

32-1901. Definitions

In this chapter, unless the context otherwise requires:

1. "Administer" means directly applying a controlled substance,
 prescription-only drug, dangerous drug or narcotic drug, whether by
 injection, inhalation, ingestion or any other means, to the body of a

1 patient or research subject by a practitioner or by the practitioner's 2 authorized agent or the patient or research subject at the direction of 3 the practitioner.

2. "Advertisement" means all representations that are disseminated in any manner or by any means other than by labeling for the purpose of inducing, or that are likely to induce, directly or indirectly, the purchase of drugs, devices, poisons or hazardous substances.

8 3. "Advisory letter" means a nondisciplinary letter to notify a 9 licensee or permittee that either:

10 (a) While there is insufficient evidence to support disciplinary 11 action, the board believes that continuation of the activities that led to 12 the investigation may result in further board action against the licensee 13 or permittee.

14 (b) The violation is a minor or technical violation that is not of 15 sufficient merit to warrant disciplinary action.

16 (c) While the licensee or permittee has demonstrated substantial 17 compliance through rehabilitation, remediation or reeducation that has 18 mitigated the need for disciplinary action, the board believes that 19 repeating the activities that led to the investigation may result in 20 further board action against the licensee or permittee.

4. "Antiseptic", if a drug is represented as such on its label, means a representation that it is a germicide, except in the case of a drug purporting to be, or represented as, an antiseptic for inhibitory use as a wet dressing, ointment or dusting powder or other use that involves prolonged contact with the body.

5. "Authorized officers of the law" means legally empowered peace officers, compliance officers of the board of pharmacy and agents of the division of narcotics enforcement and criminal intelligence of the department of public safety.

6. "Automated prescription-dispensing kiosk" means a mechanical system that is operated as an extension of a pharmacy, that maintains all transaction information within the pharmacy operating system, that is separately permitted from the pharmacy and that performs operations that either:

(a) Accept a prescription or refill order, store prepackaged or
 repackaged medications, label and dispense patient-specific prescriptions
 and provide counseling on new or refilled prescriptions.

38 (b) Dispense or deliver a prescription or refill that has been 39 prepared by or on behalf of the pharmacy that oversees the automated 40 prescription-dispensing kiosk.

41 7. "Board" or "board of pharmacy" means the Arizona state board of 42 pharmacy.

43 8. "Certificate of composition" means a list of a product's44 ingredients.

1 9. "Certificate of free sale" means a document that authenticates a 2 product that is generally and freely sold in domestic or international 3 channels of trade.

4

10. "Color additive" means a material that either:

5 (a) Is any dye, pigment or other substance that is made by a 6 process of synthesis or similar artifice or that is extracted, isolated or 7 otherwise derived, with or without intermediate or final change of identity, from any vegetable, animal, mineral or other source. 8

9 (b) If added or applied to a drug, or to the human body or any part 10 of the human body, is capable of imparting color, except that color 11 additive does not include any material that has been or may be exempted 12 under the federal act. Color includes black, white and intermediate 13 grays.

14 "Compounding" means preparing, mixing, assembling, packaging or 11. labeling a drug by a pharmacist or an intern or pharmacy technician under 15 16 the pharmacist's supervision, for the purpose of dispensing to a patient 17 based on a valid prescription order. Compounding includes preparing drugs 18 in anticipation of prescription orders prepared on routine, regularly 19 observed prescribing patterns and preparing drugs as an incident to 20 research, teaching or chemical analysis or for administration by a medical 21 practitioner to the medical practitioner's patient and not for sale or 22 dispensing. Compounding does not include preparing commercially available products from bulk compounds or preparing drugs for sale to pharmacies, 23 24 practitioners or entities for the purpose of dispensing or distribution.

25 12. "Compressed medical gas distributor" means a person that holds 26 a current permit issued by the board to distribute compressed medical 27 gases to compressed medical gas suppliers and other entities that are 28 registered, licensed or permitted to use, administer or distribute 29 compressed medical gases.

30 "Compressed medical gases" means gases and liquid oxygen that a 13. 31 compressed medical gas distributor or manufacturer has labeled in 32 compliance with federal law.

14. "Compressed medical gas order" means an order for compressed 33 34 medical gases that is issued by a medical practitioner.

35 15. "Compressed medical gas supplier" means a person that holds a 36 current permit issued by the board to supply compressed medical gases 37 pursuant to a compressed medical gas order and only to the consumer or the 38 patient.

39 "Controlled substance" means a drug, substance or immediate 16. 40 precursor that is identified, defined or listed in title 36, chapter 27, 41 article 2 or the rules adopted pursuant to title 36, chapter 27, article 2. 42

1 17. "Corrosive" means any substance that when it comes in contact 2 with living tissue will cause destruction of the tissue by chemical 3 action.

4 18. "Counterfeit drug" means a drug that, or the container or 5 labeling of which, without authorization, bears the trademark, trade name 6 or other identifying mark, imprint, number or device, or any likeness of 7 these, of a manufacturer, distributor or dispenser other than the person 8 that in fact manufactured, distributed or dispensed that drug.

9 19. "Dangerous drug" has the same meaning prescribed in section 10 13-3401.

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20. "Day" means a business day.

12 21. "Decree of censure" means an official action that is taken by 13 the board and that may include a requirement for restitution of fees to a 14 patient or consumer.

15 22. "Deliver" or "delivery" means the actual, constructive or 16 attempted transfer from one person to another whether or not there is an 17 agency relationship.

18 23. "Deputy director" means a pharmacist who is employed by the 19 board and selected by the executive director to perform duties as 20 prescribed by the executive director.

24. "Device", except as used in paragraph 18 of this section, 22 section 32-1965, paragraph 4 and section 32-1967, subsection A, paragraph 23 15 and subsection C, means an instrument, apparatus or contrivance, 24 including its components, parts and accessories, including all such items 25 under the federal act, that is intended either:

26 (a) For use in diagnosing, curing, mitigating, treating or 27 preventing disease in the human body or other animals.

(b) To affect the structure or any function of the human body or other animals.

25. "Director" means the director of the division of narcotics
 enforcement and criminal investigation INTELLIGENCE of the department of
 public safety.

26. "Direct supervision of a pharmacist" means that the pharmacist is present. If relating to the sale of certain items, direct supervision of a pharmacist means that a pharmacist determines the legitimacy or advisability of a proposed purchase of those items.

37 27. "Dispense" means to deliver to an ultimate user or research 38 subject by or pursuant to the lawful order of a practitioner, including 39 prescribing, administering, packaging, labeling or compounding as 40 necessary to prepare for that delivery.

28. "Dispenser" means a practitioner who dispenses.

42 29. "Distribute" means to deliver, other than by administering or 43 dispensing.

30. "Distributor" means a person who distributes.

1	31. "Drug" means:
2	(a) Articles that are recognized, or for which standards or
3	specifications are prescribed, in the official compendium.
4	(b) Articles that are intended for use in the diagnosis, cure,
5	mitigation, treatment or prevention of disease in the human body or other
6	animals.
7	(c) Articles other than food that are intended to affect the
8	structure or any function of the human body or other animals.
9	(d) Articles that are intended for use as a component of any
10	articles specified in subdivision (a), (b) or (c) of this paragraph but
11	does not include devices or their components, parts or accessories.
12	32. "Drug enforcement administration" means the drug enforcement
13	administration of the United States department of justice or its successor
14	agency.
15	33. "Drug or device manufacturing" means producing, preparing,
16	propagating or processing a drug or device, either directly or indirectly,
17	by extraction from substances of natural origin or independently by means
18	of chemical synthesis and includes any packaging or repackaging of
19	substances or labeling or relabeling of its container and promoting and
20	marketing the same. Drug or device manufacturing does not include
21	compounding.
22	34. "Durable medical equipment" means technologically sophisticated
23	medical equipment as prescribed by the board in rule that a patient or
24	consumer may use in a home or residence and that may be a
25	prescription-only device.
26	35. "Durable medical equipment distributor":
27	(a) Means a person that stores or distributes durable medical
28	equipment other than to the patient or consumer.
29	(b) Includes a virtual durable medical equipment distributor as
30	prescribed in rule by the board.
31	36. "Durable medical equipment supplier":
32	(a) Means a person that sells, leases or supplies durable medical
33	equipment to the patient or consumer.
34	(b) Includes a virtual durable medical equipment supplier as
35	prescribed in rule by the board.
36	37. "Economic poison" means any substance that alone, in chemical
37	combination with or in formulation with one or more other substances is a
38	pesticide within the meaning of the laws of this state or the federal
39	insecticide, fungicide and rodenticide act and that is used in producing,
40	storing or transporting raw agricultural commodities.
41	38. "Enteral feeding" means nourishment that is provided by means
42	of a tube inserted into the stomach or intestine.

1 39. "Established name", with respect to a drug or ingredient of a 2 drug, means any of the following:

3

(a) The applicable official name.

4 (b) If there is no such name and the drug or ingredient is an 5 article recognized in an official compendium, the official title in an 6 official compendium.

7 8

(c) If neither subdivision (a) nor (b) of this paragraph applies, the common or usual name of the drug.

9 40. "Executive director" means the executive director of the board 10 of pharmacy.

11 41. "Federal act" means the federal laws and regulations that 12 pertain to drugs, devices, poisons and hazardous substances and that are 13 official at the time any drug, device, poison or hazardous substance is 14 affected by this chapter.

15

42. "Full-service wholesale permittee":

16 (a) Means a permittee who may distribute prescription-only drugs 17 and devices, controlled substances and over-the-counter drugs and devices 18 to pharmacies or other legal outlets from a place devoted in whole or in 19 part to wholesaling these items.

20

(b) Includes a virtual wholesaler as defined in rule by the board.

21 43. "Good manufacturing practice" means a system for ensuring that 22 products are consistently produced and controlled according to quality 23 standards and covering all aspects of design, monitoring and control of 24 manufacturing processes and facilities to ensure that products do not pose 25 any risk to the consumer or public.

26 44. "Highly toxic" means any substance that falls within any of the 27 following categories:

(a) Produces death within fourteen days in half or more than half 28 29 of a group of ten or more laboratory white rats each weighing between two 30 hundred and three hundred grams, at a single dose of fifty milligrams or 31 less per kilogram of body weight, when orally administered.

(b) Produces death within fourteen days in half or more than half 32 33 of a group of ten or more laboratory white rats each weighing between two hundred and three hundred grams, if inhaled continuously for a period of 34 35 one hour or less at an atmospheric concentration of two hundred parts per 36 million by volume or less of gas or vapor or two milligrams per liter by 37 volume or less of mist or dust, provided the concentration is likely to be 38 encountered by humans if the substance is used in any reasonably 39 foreseeable manner.

40 (c) Produces death within fourteen days in half or more than half 41 of a group of ten or more rabbits tested in a dosage of two hundred 42 milligrams or less per kilogram of body weight, if administered by 43 continuous contact with the bare skin for twenty-four hours or less. Ιf 44 the board finds that available data on human experience with any substance 45 indicate results different from those obtained on animals in the dosages

1 or concentrations prescribed in this paragraph, the human data shall take 2 precedence.

45. "Hospital" means any institution for the care and treatment of the sick and injured that is approved and licensed as a hospital by the department of health services.

6

46. "Intern" means a pharmacy intern.

47. "Internship" means the practical, experiential, hands-ontraining of a pharmacy intern under the supervision of a preceptor.

9 48. "Irritant" means any substance, other than a corrosive, that on 10 immediate, prolonged or repeated contact with normal living tissue will 11 induce a local inflammatory reaction.

12 49. "Jurisprudence examination" means a board-approved pharmacy law 13 examination that is written and administered in cooperation with the 14 national association of boards of pharmacy or another board-approved 15 pharmacy law examination.

16 50. "Label" means a display of written, printed or graphic matter 17 on the immediate container of any article that, unless easily legible 18 through the outside wrapper or container, also appears on the outside 19 wrapper or container of the article's retail package. For the purposes of 20 this paragraph, the immediate container does not include package liners.

21 51. "Labeling" means all labels and other written, printed or 22 graphic matter that either:

23

(a) Is on any article or any of its containers or wrappers.

24

(b) Accompanies that article.

52. "Letter of reprimand" means a disciplinary letter that is a public document issued by the board and that informs a licensee or permittee that the licensee's or permittee's conduct violates state or federal law and may require the board to monitor the licensee or permittee.

30 53. "Limited service pharmacy" means a pharmacy that is approved by 31 the board to practice a limited segment of pharmacy as indicated by the 32 permit issued by the board.

33

54. "Manufacture" or "manufacturer":

(a) Means every person who prepares, derives, produces, compounds,
 processes, packages or repackages or labels any drug in a place, other
 than a pharmacy, that is devoted to manufacturing the drug.

37 (b) Includes a virtual manufacturer as defined in rule by the 38 board.

39

55. "Marijuana" has the same meaning prescribed in section 13-3401.

56. "Medical practitioner" means any medical doctor, doctor of osteopathic medicine, dentist, podiatrist, veterinarian or other person who is licensed and authorized by law to use and prescribe drugs and devices to treat sick and injured human beings or animals or to diagnose or prevent sickness in human beings or animals in this state or any state, territory or district of the United States. 1 57. "Medication order" means a written or verbal order from a 2 medical practitioner or that person's authorized agent to administer a 3 drug or device.

4 58. "Narcotic drug" has the same meaning prescribed in section 5 13-3401.

6

59. "New drug" means either:

7 (a) Any drug of which the composition is such that the drug is not 8 generally recognized among experts qualified by scientific training and 9 experience to evaluate the safety and effectiveness of drugs as safe and 10 effective for use under the conditions prescribed, recommended or 11 suggested in the labeling.

(b) Any drug of which the composition is such that the drug, as a result of investigations to determine its safety and effectiveness for use under such conditions, has become so recognized, but that has not, other than in the investigations, been used to a material extent or for a material time under those conditions.

17 60. "Nonprescription drug" or "over-the-counter drug" means any 18 nonnarcotic medicine or drug that may be sold without a prescription and that is prepackaged and labeled for use by the consumer in accordance with 19 20 the requirements of the laws of this state and federal law. 21 Nonprescription drug does not include:

(a) A drug that is primarily advertised and promoted professionally
 to medical practitioners and pharmacists by manufacturers or primary
 distributors.

25 26 (b) A controlled substance.

(c) A drug that is required to bear a label that states "Rx only".

27

(d) A drug that is intended for human use by hypodermic injection.61. "Nonprescription drug wholesale permittee":

61. "Nonprescription drug wholesale permittee":
(a) Means a permittee who may distribute only over-the-counter
drugs and devices to pharmacies or other lawful outlets from a place
devoted in whole or in part to wholesaling these items.

32

(b) Includes a virtual wholesaler as defined in rule by the board.

62. "Notice" means personal service or the mailing of a copy of the notice by certified mail and email addressed either to the person at the person's latest address of record in the board office or to the person and the person's attorney using the most recent information provided to the board in the board's licensing database.

38 63. "Nutritional supplementation" means vitamins, minerals and
 39 caloric supplementation. Nutritional supplementation does not include
 40 medication or drugs.

41 64. "Official compendium" means the latest revision of the United 42 States pharmacopeia and the national formulary or any current supplement.

65. "Other jurisdiction" means one of the other forty-nine states,
the District of Columbia, the Commonwealth of Puerto Rico or a territory
of the United States of America.

1 66. "Package" means a receptacle that is defined or described in 2 the United States pharmacopeia and the national formulary as adopted by 3 the board.

67. "Packaging" means the act or process of placing a drug item or below device in a container for the purpose or intent of dispensing or below distributing the item or device to another.

68. "Parenteral nutrition" means intravenous feeding that provides
an individual with fluids and essential nutrients the individual needs
while the individual is unable to receive adequate fluids or feedings by
mouth or by enteral feeding.

11 69. "Person" means an individual, partnership, corporation and 12 association, and their duly authorized agents.

13 70. "Pharmaceutical care" means the provision of drug therapy and14 other pharmaceutical patient care services.

15 71. "Pharmacist" means an individual who is currently licensed by 16 the board to practice the profession of pharmacy in this state.

17 72. "Pharmacist in charge" means the pharmacist who is responsible 18 to the board for a licensed establishment's compliance with the laws and 19 administrative rules of this state and of the federal government 20 pertaining to the practice of pharmacy, the manufacturing of drugs and the 21 distribution of drugs and devices.

73. "Pharmacist licensure examination" means a board-approved examination that is written and administered in cooperation with the national association of boards of pharmacy or any other board-approved pharmacist licensure examination.

26

74. "Pharmacy" means:

(a) Any place where drugs, devices, poisons or related hazardous
 substances are offered for sale at retail or where prescription orders are
 dispensed by a licensed pharmacist.

30 (b) Any place that displays on or in the place or that displays a 31 sign on the place the words "pharmaceutical chemist", "apothecary", 32 "druggist", "pharmacy", "drugstore", "drugs" or "drug sundries", any 33 combination of these words, or any words of similar meaning in any 34 language.

35 (c) Any place where the characteristic symbol of pharmacy or the 36 characteristic prescription sign "Rx" is exhibited.

37 (d) Any building or other structure or portion of a building or 38 other structure that is leased, used or controlled by a permittee to 39 conduct the business authorized by the board at the address specified on 40 the permit issued to the permittee.

41

(e) A remote dispensing site pharmacy.

42 43

- (f) A remote hospital-site pharmacy.
- (g) A satellite pharmacy.

44 75. "Pharmacy intern" means a person who has all of the 45 qualifications and experience prescribed in section 32-1923. 1 76. "Pharmacy technician" means a person who is licensed pursuant 2 to this chapter.

3 77. "Pharmacy technician trainee" means a person who is licensed 4 **REGISTERED** pursuant to this chapter.

5

"Poison" or "hazardous substance" includes any of the following 78. 6 if intended and suitable for household use or use by children:

7 (a) Any substance that, according to standard works on medicine, 8 pharmacology, pharmacognosy or toxicology, if applied to, introduced into 9 or developed within the body in relatively small quantities by its inherent action uniformly produces serious bodily injury, disease or 10 11 death.

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(b) A toxic substance.

- (c) A highly toxic substance.
- (d) A corrosive substance.
- 15 (e) An irritant.
 - (f) A strong sensitizer.

17 (g) A mixture of any of the substances described in this paragraph, 18 if the substance or mixture of substances may cause substantial personal injury or substantial illness during or as a proximate result of any 19 20 customary or reasonably foreseeable handling or use, including reasonably 21 foreseeable ingestion by children.

22 (h) A substance that is designated by the board to be a poison or hazardous substance. This subdivision does not apply to radioactive 23 24 substances, economic poisons subject to the federal insecticide, fungicide and rodenticide act or the state pesticide act, foods, drugs and cosmetics 25 26 subject to state laws or the federal act or substances intended for use as fuels when stored in containers and used in the heating, cooking or 27 refrigeration system of a house. This subdivision applies to any 28 29 substance or article that is not itself an economic poison within the meaning of the federal insecticide, fungicide and rodenticide act or the 30 31 state pesticide act, but that is a poison or hazardous substance within 32 the meaning of this paragraph by reason of bearing or containing an 33 economic poison or hazardous substance.

34

79. "Practice of pharmacy":

35 (a) Means furnishing the following health care services as a 36 medical professional:

37 (i) Interpreting, evaluating and dispensing prescription orders in 38 the patient's best interests.

39 (ii) Compounding drugs pursuant to or in anticipation of a 40 prescription order.

41 (iii) Labeling drugs and devices in compliance with state and 42 federal requirements.

43 (iv) Participating in drug selection and drug utilization reviews, 44 drug administration, drug or drug-related research and drug therapy 45 monitoring or management.

1 (v) Providing patient counseling necessary to provide 2 pharmaceutical care.

3 (vi) Properly and safely storing drugs and devices in anticipation 4 of dispensing.

5

(vii) Maintaining required records of drugs and devices.

6 (viii) Offering or performing acts, services, operations or 7 transactions that are necessary to conduct, operate, manage and control a 8 pharmacy.

9 (ix) Providing patient care services pursuant to a collaborative 10 practice agreement with a provider as outlined in section 32-1970.

11 (x) Initiating and administering immunizations or vaccines pursuant 12 to section 32-1974.

13 (b) Does not include initiating a prescription order for any 14 medication, drug or other substance used to induce or cause a medication 15 abortion as defined in section 36-2151.

16 80. "Practitioner" means any physician, dentist, veterinarian, 17 scientific investigator or other person who is licensed, registered or 18 otherwise permitted to distribute, dispense, conduct research with respect to or administer a controlled substance in the course of professional 19 20 practice or research in this state, or any pharmacy, hospital or other 21 institution that is licensed, registered or otherwise permitted to 22 distribute, dispense, conduct research with respect to or administer a controlled substance in the course of professional practice or research in 23 24 this state.

25 81. "Preceptor" means a pharmacist who is serving as the practical
 26 instructor of an intern and who complies with section 32-1923.

27

82. "Precursor chemical" means a substance that is:

(a) The principal compound that is commonly used or that is produced primarily for use and that is an immediate chemical intermediary used or likely to be used in the manufacture of a controlled substance, the control of which is necessary to prevent, curtail or limit manufacture.

33

(b) Listed in section 13–3401, paragraph 26 or 27 OR 28.

34 83. "Prescription" means either a prescription order or a 35 prescription medication.

36 84. "Prescription medication" means any drug, including label and 37 container according to context, that is dispensed pursuant to a 38 prescription order.

39

85. "Prescription-only device" includes:

40 (a) Any device that is limited by the federal act to use under the 41 supervision of a medical practitioner.

42 (b) Any device required by the federal act to bear on its label 43 essentially the legend "Rx only".

44 86. "Prescription-only drug" does not include a controlled 45 substance but does include: 1 (a) Any drug that because of its toxicity or other potentiality for 2 harmful effect, the method of its use, or the collateral measures 3 necessary to its use is not generally recognized among experts, qualified 4 by scientific training and experience to evaluate its safety and efficacy, 5 as safe for use except by or under the supervision of a medical 6 practitioner.

7 (b) Any drug that is limited by an approved new drug application 8 under the federal act or section 32-1962 to use under the supervision of a 9 medical practitioner.

10 (c) Every potentially harmful drug, the labeling of which does not 11 bear or contain full and adequate directions for use by the consumer.

12 (d) Any drug, other than a controlled substance, that is required 13 by the federal act to bear on its label the legend "Rx only".

14

87. "Prescription order" means any of the following:

15 (a) An order to a pharmacist for drugs or devices that is issued 16 and signed by a duly licensed medical practitioner in the authorized 17 course of the practitioner's professional practice.

18 (b) An order that is transmitted to a pharmacist through word of 19 mouth, telephone or other means of communication directed by that medical 20 practitioner. Prescription orders received by word of mouth, telephone or 21 other means of communication shall be maintained by the pharmacist 22 pursuant to section 32-1964, and the record so made by the pharmacist constitutes the original prescription order to be dispensed by the 23 24 pharmacist. This paragraph does not alter or affect laws of this state or 25 any federal act requiring a written prescription order.

26 (c) An order that is initiated by a pharmacist pursuant to a 27 collaborative practice agreement with a provider as outlined in section 28 32–1970, or immunizations or vaccines administered by a pharmacist 29 pursuant to section 32–1974.

30 (d) A diet order or an order for enteral feeding, nutritional 31 supplementation or parenteral nutrition that is initiated by a registered 32 dietitian or other qualified nutrition professional in a hospital pursuant 33 to section 36-416.

34

88. "Professionally incompetent" means:

35 (a) Incompetence based on a variety of factors, including a lack of 36 sufficient pharmaceutical knowledge or skills or experience to a degree 37 likely to endanger the health of patients.

38 (b) When considered with other indications of professional 39 incompetence, a pharmacist or pharmacy intern who fails to obtain a 40 passing score on a board-approved pharmacist licensure examination or a 41 pharmacy technician or pharmacy technician trainee who fails to obtain a 42 score on а board-approved pharmacy technician licensure passing 43 examination.

44 89. "Radioactive substance" means a substance that emits ionizing 45 radiation. 1 90. "Remote dispensing site pharmacy" means a pharmacy where a 2 pharmacy technician or pharmacy intern prepares, compounds or dispenses 3 prescription medications under remote supervision by a pharmacist.

91. "Remote hospital-site pharmacy" means a pharmacy located in a satellite facility that operates under the license issued by the department of health services to the hospital of which it is a satellite.

92. "Remote supervision by a pharmacist" means that a pharmacist
directs and controls the actions of pharmacy technicians and pharmacy
interns through the use of audio and visual technology.

93. "Revocation" or "revoke" means the official cancellation of a license, permit, registration or other approval authorized by the board for a period of two years unless otherwise specified by the board. A request or new application for reinstatement may be presented to the board for review before the conclusion of the specified revocation period upon review of the executive director.

94. "Safely engage in employment duties" means that a permittee or the permittee's employee is able to safely engage in employment duties related to the manufacture, sale, distribution or dispensing of drugs, devices, poisons, hazardous substances, controlled substances or precursor chemicals.

21 95. "Satellite facility" has the same meaning prescribed in section 22 36-422.

23 96. "Satellite pharmacy" means a work area located within a 24 hospital or on a hospital campus that is not separated by other commercial 25 property or residential property, that is under the direction of a 26 pharmacist, that is a remote extension of a centrally licensed hospital 27 pharmacy, that is owned by and dependent on the centrally licensed 28 hospital pharmacy for administrative control, staffing and drug 29 procurement and that is not required to be separately permitted.

30 97. "Symbol" means the characteristic symbols that have 31 historically identified pharmacy, including show globes and mortar and 32 pestle, and the sign "Rx".

98. "Third-party logistics provider" means an entity that provides or coordinates warehousing or other logistics services for the following items, but that does not take ownership of the items, and that distributes those items as directed by a manufacturer, wholesaler, dispenser or durable medical equipment supplier that is permitted by the board:

(a) Narcotic drugs or other controlled substances.(b) Dangerous drugs as defined in section 13-3401.

Prescription-only drugs and devices.

- 38 39
- 40 41
- (d) Nonprescription drugs and devices.
- 42 (e) Precursor chemicals.

(c)

43 (f) Regulated chemicals as defined in section 13-3401.

1 99. "Toxic substance" means a substance, other than a radioactive 2 substance, that has the capacity to produce injury or illness in humans 3 through ingestion, inhalation or absorption through any body surface.

4 100. "Ultimate user" means a person who lawfully possesses a drug 5 or controlled substance for that person's own use, for the use of a member 6 of that person's household or for administering to an animal owned by that 7 person or by a member of that person's household.