1 AN ACT relating to controlled substances.

## 2 Be it enacted by the General Assembly of the Commonwealth of Kentucky:

- 3 → Section 1. KRS 218A.010 is amended to read as follows:
- 4 As used in this chapter:
- 5 (1) "Administer" means the direct application of a controlled substance, whether by
- 6 injection, inhalation, ingestion, or any other means, to the body of a patient or
- 7 research subject by:
- 8 (a) A practitioner or by his or her authorized agent under his or her immediate
- 9 supervision and pursuant to his or her order; or
- 10 (b) The patient or research subject at the direction and in the presence of the
- 11 practitioner;
- 12 (2) "Anabolic steroid" means any drug or hormonal substance chemically and
- pharmacologically related to testosterone that promotes muscle growth and includes
- those substances *classified as Schedule III controlled substances pursuant to*
- 15 Section 2 of this Act[listed in KRS 218A.090(5)] but does not include estrogens,
- progestins, and anticosteroids;
- 17 (3) "Cabinet" means the Cabinet for Health and Family Services;
- 18 (4) "Child" means any person under the age of majority as specified in KRS 2.015;
- 19 (5) "Cocaine" means a substance containing any quantity of cocaine, its salts, optical
- and geometric isomers, and salts of isomers;
- 21 (6) "Controlled substance" means methamphetamine, or a drug, substance, or
- 22 immediate precursor in Schedules I through V and includes a controlled substance
- analogue;
- 24 (7) (a) "Controlled substance analogue," except as provided in paragraph (b) of this
- subsection, means a substance:
- 26 1. The chemical structure of which is substantially similar to the structure
- of a controlled substance in Schedule I or II; and

1			2.	Which has a stimulant, depressant, or hallucinogenic effect on the
2				central nervous system that is substantially similar to or greater than the
3				stimulant, depressant, or hallucinogenic effect on the central nervous
4				system of a controlled substance in Schedule I or II; or
5			3.	With respect to a particular person, which such person represents or
6				intends to have a stimulant, depressant, or hallucinogenic effect on the
7				central nervous system that is substantially similar to or greater than the
8				stimulant, depressant, or hallucinogenic effect on the central nervous
9				system of a controlled substance in Schedule I or II.
10		(b)	Sucl	n term does not include:
11			1.	Any substance for which there is an approved new drug application;
12			2.	With respect to a particular person, any substance if an exemption is in
13				effect for investigational use for that person pursuant to federal law to
14				the extent conduct with respect to such substance is pursuant to such
15				exemption; or
16			3.	Any substance to the extent not intended for human consumption before
17				the exemption described in subparagraph 2. of this paragraph takes
18				effect with respect to that substance;
19	(8)	"Coı	unterf	eit substance" means a controlled substance which, or the container or
20		labe	ling o	of which, without authorization, bears the trademark, trade name, or other
21		iden	tifyin	g mark, imprint, number, or device, or any likeness thereof, of a
22		man	ufactı	arer, distributor, or dispenser other than the person who in fac-
23		man	ufactı	ared, distributed, or dispensed the substance;
24	(9)	"Dis	pense	" means to deliver a controlled substance to an ultimate user or research
25		subj	ect by	or pursuant to the lawful order of a practitioner, including the packaging

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labeling, or compounding necessary to prepare the substance for that delivery;

(10) "Dispenser" means a person who lawfully dispenses a Schedule II, III, IV, or V

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1	controlled	substance to	or for the	use of an	ultimate	iicer.
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- 2 (11) "Distribute" means to deliver other than by administering or dispensing a controlled
- 3 substance;
- (12) "Dosage unit" means a single pill, capsule, ampule, liquid, or other form of 4
- 5 administration available as a single unit;
- 6 (13) "Drug" means:
- 7 Substances recognized as drugs in the official United States Pharmacopoeia, (a)
- 8 official Homeopathic Pharmacopoeia of the United States, or official National
- 9 Formulary, or any supplement to any of them;
- 10 Substances intended for use in the diagnosis, care, mitigation, treatment, or (b)
- 11 prevention of disease in man or animals;
- 12 (c) Substances (other than food) intended to affect the structure or any function of
- 13 the body of man or animals; and
- 14 Substances intended for use as a component of any article specified in this
- 15 subsection.
- 16 It does not include devices or their components, parts, or accessories;
- 17 (14) "Good faith prior examination," as used in KRS Chapter 218A and for criminal
- 18 prosecution only, means an in-person medical examination of the patient conducted
- 19 by the prescribing practitioner or other health-care professional routinely relied
- 20 upon in the ordinary course of his or her practice, at which time the patient is
- 21 physically examined and a medical history of the patient is obtained. "In-person"
- 22 includes telehealth examinations. This subsection shall not be applicable to hospice
- 23 providers licensed pursuant to KRS Chapter 216B;
- 24 (15) "Hazardous chemical substance" includes any chemical substance used or intended
- 25 for use in the illegal manufacture of a controlled substance as defined in this section
- or the illegal manufacture of methamphetamine as defined in KRS 218A.1431, 26
- 27 which:

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1		(a)	Poses an explosion hazard;
2		(b)	Poses a fire hazard; or
3		(c)	Is poisonous or injurious if handled, swallowed, or inhaled;
4	(16)	"Hei	roin" means a substance containing any quantity of heroin, or any of its salts,
5		ison	ners, or salts of isomers;
6	(17)	"Hy	drocodone combination product" means a drug with:
7		(a)	Not more than three hundred (300) milligrams of dihydrocodeinone, or any of
8			its salts, per one hundred (100) milliliters or not more than fifteen (15)
9			milligrams per dosage unit, with a fourfold or greater quantity of an
10			isoquinoline alkaloid of opium; or
11		(b)	Not more than three hundred (300) milligrams of dihydrocodeinone, or any of
12			its salts, per one hundred (100) milliliters or not more than fifteen (15)
13			milligrams per dosage unit, with one (1) or more active, nonnarcotic
14			ingredients in recognized therapeutic amounts;
15	(18)	"Imr	mediate precursor" means a substance which is the principal compound
16		com	monly used or produced primarily for use, and which is an immediate chemical
17		inter	rmediary used or likely to be used in the manufacture of a controlled substance
18		or m	nethamphetamine, the control of which is necessary to prevent, curtail, or limit
19		man	ufacture;
20	(19)	"Inte	ent to manufacture" means any evidence which demonstrates a person's
21		cons	scious objective to manufacture a controlled substance or methamphetamine.
22		Such	n evidence includes but is not limited to statements and a chemical substance's
23		usag	ge, quantity, manner of storage, or proximity to other chemical substances or
24		equi	pment used to manufacture a controlled substance or methamphetamine;
25	(20)	"Iso	mer" means the optical isomer, except the Cabinet for Health and Family
26		Serv	rices may include the optical, positional, or geometric isomer to classify any

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substance pursuant to Section 2 of this Act[as used in KRS 218A.050(3) and

1		218.	A.070(1)(d). As used in KRS 218 $A.050(3)$ , the term "isomer" means the optical,
2		<del>posi</del>	tional, or geometric isomer. As used in KRS 218A.070(1)(d), the term "isomer"
3		mea	ns the optical or geometric isomer];
4	(21)	"Ma	nufacture," except as provided in KRS 218A.1431, means the production,
5		prep	paration, propagation, compounding, conversion, or processing of a controlled
6		subs	stance, either directly or indirectly by extraction from substances of natural
7		orig	in or independently by means of chemical synthesis, or by a combination of
8		extra	action and chemical synthesis, and includes any packaging or repackaging of the
9		subs	stance or labeling or relabeling of its container except that this term does not
10		inclu	ude activities:
11		(a)	By a practitioner as an incident to his or her administering or dispensing of a
12			controlled substance in the course of his or her professional practice;
13		(b)	By a practitioner, or by his or her authorized agent under his supervision, for
14			the purpose of, or as an incident to, research, teaching, or chemical analysis
15			and not for sale; or
16		(c)	By a pharmacist as an incident to his or her dispensing of a controlled
17			substance in the course of his or her professional practice;
18	(22)	"Ma	rijuana" means all parts of the plant Cannabis sp., whether growing or not; the
19		seed	Is thereof; the resin extracted from any part of the plant; and every compound,
20		man	ufacture, salt, derivative, mixture, or preparation of the plant, its seeds or resin
21		or a	my compound, mixture, or preparation which contains any quantity of these
22		subs	stances. The term "marijuana" does not include:
23		(a)	Industrial hemp as defined in KRS 260.850;
24		(b)	The substance cannabidiol, when transferred, dispensed, or administered
25			pursuant to the written order of a physician practicing at a hospital or
26			associated clinic affiliated with a Kentucky public university having a college
27			or school of medicine; or

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1		(c)	For persons participating in a clinical trial or in an expanded access program,
2			a drug or substance approved for the use of those participants by the United
3			States Food and Drug Administration;
4	(23)	"Me	dical history," as used in KRS Chapter 218A and for criminal prosecution only,
5		mea	ns an accounting of a patient's medical background, including but not limited to
6		prio	medical conditions, prescriptions, and family background;
7	(24)	"Me	dical order," as used in KRS Chapter 218A and for criminal prosecution only,
8		mea	ns a lawful order of a specifically identified practitioner for a specifically
9		iden	tified patient for the patient's health-care needs. "Medical order" may or may
10		not i	nclude a prescription drug order;
11	(25)	"Me	dical record," as used in KRS Chapter 218A and for criminal prosecution only,
12		mea	ns a record, other than for financial or billing purposes, relating to a patient,
13		kept	by a practitioner as a result of the practitioner-patient relationship;
14	(26)	"Me	thamphetamine" means any substance that contains any quantity of
15		metł	namphetamine, or any of its salts, isomers, or salts of isomers;
16	(27)	"Naı	cotic drug" means any of the following, whether produced directly or indirectly
17		by e	extraction from substances of vegetable origin, or independently by means of
18		chen	nical synthesis, or by a combination of extraction and chemical synthesis:
19		(a)	Opium and opiate, and any salt, compound, derivative, or preparation of
20			opium or opiate;
21		(b)	Any salt, compound, isomer, derivative, or preparation thereof which is
22			chemically equivalent or identical with any of the substances referred to in
23			paragraph (a) of this subsection, but not including the isoquinoline alkaloids
24			of opium;
25		(c)	Opium poppy and poppy straw;
26		(d)	Coca leaves, except coca leaves and extracts of coca leaves from which
27			cocaine, ecgonine, and derivatives of ecgonine or their salts have been

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1		removed;
2		(e) Cocaine, its salts, optical and geometric isomers, and salts of isomers;
3		(f) Ecgonine, its derivatives, their salts, isomers, and salts of isomers; and
4		(g) Any compound, mixture, or preparation which contains any quantity of any of
5		the substances referred to in paragraphs (a) to (f) of this subsection;
6	(28)	"Opiate" means any substance having an addiction-forming or addiction-sustaining
7		liability similar to morphine or being capable of conversion into a drug having
8		addiction-forming or addiction-sustaining liability. It does not include, unless
9		specifically designated as controlled under <b>Section 2 of this Act</b> [KRS 218A.030],
10		the dextrorotatory isomer of 3-methoxy-n-methylmorphinan and its salts
11		(dextromethorphan). It does include its racemic and levorotatory forms;
12	(29)	"Opium poppy" means the plant of the species papaver somniferum L., except its
13		seeds;
14	(30)	"Person" means individual, corporation, government or governmental subdivision
15		or agency, business trust, estate, trust, partnership or association, or any other legal
16		entity;
17	(31)	"Physical injury" has the same meaning it has in KRS 500.080;
18	(32)	"Poppy straw" means all parts, except the seeds, of the opium poppy, after mowing;
19	(33)	"Pharmacist" means a natural person licensed by this state to engage in the practice
20		of the profession of pharmacy;
21	(34)	"Practitioner" means a physician, dentist, podiatrist, veterinarian, scientific
22		investigator, optometrist as authorized in KRS 320.240, advanced practice
23		registered nurse as authorized under KRS 314.011, or other person licensed,
24		registered, or otherwise permitted by state or federal law to acquire, distribute,
25		dispense, conduct research with respect to, or to administer a controlled substance
26		in the course of professional practice or research in this state. "Practitioner" also

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includes a physician, dentist, podiatrist, veterinarian, or advanced practice registered

1		nurse authorized under KRS 314.011 who is a resident of and actively practicing in
2		a state other than Kentucky and who is licensed and has prescriptive authority for
3		controlled substances under the professional licensing laws of another state, unless
4		the person's Kentucky license has been revoked, suspended, restricted, or probated,
5		in which case the terms of the Kentucky license shall prevail;
6	(35)	"Practitioner-patient relationship," as used in KRS Chapter 218A and for criminal
7		prosecution only, means a medical relationship that exists between a patient and a
8		practitioner or the practitioner's designee, after the practitioner or his or her
9		designee has conducted at least one (1) good faith prior examination;
10	(36)	"Prescription" means a written, electronic, or oral order for a drug or medicine, or
11		combination or mixture of drugs or medicines, or proprietary preparation, signed or
12		given or authorized by a medical, dental, chiropody, veterinarian, optometric
13		practitioner, or advanced practice registered nurse, and intended for use in the
14		diagnosis, cure, mitigation, treatment, or prevention of disease in man or other
15		animals;
16	(37)	"Prescription blank," with reference to a controlled substance, means a document
17		that meets the requirements of KRS 218A.204 and 217.216;
18	(38)	"Presumptive probation" means a sentence of probation not to exceed the maximum
19		term specified for the offense, subject to conditions otherwise authorized by law,
20		that is presumed to be the appropriate sentence for certain offenses designated in
21		this chapter, notwithstanding contrary provisions of KRS Chapter 533. That
22		presumption shall only be overcome by a finding on the record by the sentencing
23		court of substantial and compelling reasons why the defendant cannot be safely and
24		effectively supervised in the community, is not amenable to community-based
25		treatment, or poses a significant risk to public safety;
26	(39)	"Production" includes the manufacture, planting, cultivation, growing, or harvesting

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of a controlled substance;

(40) "Recovery program" means an evidence-based, nonclinical service that assists individuals and families working toward sustained recovery from substance use and other criminal risk factors. This can be done through an array of support programs and services that are delivered through residential and nonresidential means;

- (41) "Salvia" means Salvia divinorum or Salvinorin A and includes all parts of the plant presently classified botanically as Salvia divinorum, whether growing or not, the seeds thereof, any extract from any part of that plant, and every compound, manufacture, derivative, mixture, or preparation of that plant, its seeds, or its extracts, including salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation of that plant, its seeds, or extracts. The term shall not include any other species in the genus salvia;
- (42) "Second or subsequent offense" means that for the purposes of this chapter an offense is considered as a second or subsequent offense, if, prior to his or her conviction of the offense, the offender has at any time been convicted under this chapter, or under any statute of the United States, or of any state relating to substances classified as controlled substances or counterfeit substances, except that a prior conviction for a nontrafficking offense shall be treated as a prior offense only when the subsequent offense is a nontrafficking offense. For the purposes of this section, a conviction voided under KRS 218A.275 or 218A.276 shall not constitute a conviction under this chapter;
- 22 (43) "Sell" means to dispose of a controlled substance to another person for consideration or in furtherance of commercial distribution;
- 24 (44) "Serious physical injury" has the same meaning it has in KRS 500.080;
- 25 (45) "Synthetic cannabinoids or piperazines" means any chemical compound which is 26 not approved by the United States Food and Drug Administration or, if approved, 27 which is not dispensed or possessed in accordance with state and federal law, that

1 contains Benzylpiperazine (BZP); Trifluoromethylphenylpiperazine (TFMPP); 1,12 Dimethylheptyl-11-hydroxytetrahydrocannabinol (HU-210); 1-Butyl-3-(13 naphthoyl)indole; 1-Pentyl-3-(1-naphthoyl)indole; dexanabinol (HU-211); or any
4 compound in the following structural classes:

- (a) Naphthoylindoles: Any compound containing a 3-(1-naphthoyl)indole structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group, whether or not further substituted in the indole ring to any extent and whether or not substituted in the naphthyl ring to any extent. Examples of this structural class include but are not limited to JWH-015, JWH-018, JWH-019, JWH-073, JWH-081, JWH-122, JWH-200, and AM-2201;
- (b) Phenylacetylindoles: Any compound containing a 3-phenylacetylindole structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group whether or not further substituted in the indole ring to any extent and whether or not substituted in the phenyl ring to any extent. Examples of this structural class include but are not limited to JWH-167, JWH-250, JWH-251, and RCS-8;
- (c) Benzoylindoles: Any compound containing a 3-(benzoyl)indole structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group whether or not further substituted in the indole ring to any extent and whether or not substituted in the phenyl ring to any extent. Examples of this structural class include but are not limited to AM-630, AM-2233, AM-694, Pravadoline (WIN 48,098), and RCS-4;
- (d) Cyclohexylphenols: Any compound containing a 2-(3-

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hydroxycyclohexyl)phenol structure with substitution at the 5-position of the phenolic ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group whether or not substituted in the cyclohexyl ring to any extent. Examples of this structural class include but are not limited to CP 47,497 and its C8 homologue (cannabicyclohexanol);

- (e) Naphthylmethylindoles: Any compound containing a 1H-indol-3-yl-(1-naphthyl)methane structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group whether or not further substituted in the indole ring to any extent and whether or not substituted in the naphthyl ring to any extent. Examples of this structural class include but are not limited to JWH-175, JWH-184, and JWH-185;
- (f) Naphthoylpyrroles: Any compound containing a 3-(1-naphthoyl)pyrrole structure with substitution at the nitrogen atom of the pyrrole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group whether or not further substituted in the pyrrole ring to any extent and whether or not substituted in the naphthyl ring to any extent. Examples of this structural class include but are not limited to JWH-030, JWH-145, JWH-146, JWH-307, and JWH-368;
- (g) Naphthylmethylindenes: Any compound containing a 1-(1-naphthylmethyl)indene structure with substitution at the 3-position of the indene ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group whether or not further substituted in the indene ring to any extent and whether or not substituted in the naphthyl ring to any extent. Examples of this structural class include but are not limited to JWH-176;

(h	) Tetramethylcyclopropanoylindoles: Any compound containing a 3-(1
	tetramethylcyclopropoyl)indole structure with substitution at the nitroge
	atom of the indole ring by an alkyl, haloalkyl, cycloalkylmethy
	cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethy
	group, whether or not further substituted in the indole ring to any extent an
	whether or not further substituted in the tetramethylcyclopropyl ring to an
	extent. Examples of this structural class include but are not limited to UR-14
	and XLR-11;

- (i) Adamantoylindoles: Any compound containing a 3-(1-adamantoyl)indole structure with substitution at the nitrogen atom of the indole ring by an alkyl, haloalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, or 2-(4-morpholinyl)ethyl group, whether or not further substituted in the indole ring to any extent and whether or not substituted in the adamantyl ring system to any extent. Examples of this structural class include but are not limited to AB-001 and AM-1248; or
- (j) Any other synthetic cannabinoid or piperazine which is not approved by the United States Food and Drug Administration or, if approved, which is not dispensed or possessed in accordance with state and federal law;
- (46) "Synthetic cathinones" means any chemical compound which is not approved by the United States Food and Drug Administration or, if approved, which is not dispensed or possessed in accordance with state and federal law (not including bupropion or compounds listed under a different schedule) structurally derived from 2-aminopropan-1-one by substitution at the 1-position with either phenyl, naphthyl, or thiophene ring systems, whether or not the compound is further modified in one (1) or more of the following ways:
  - (a) By substitution in the ring system to any extent with alkyl, alkylenedioxy, alkoxy, haloalkyl, hydroxyl, or halide substituents, whether or not further

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I			substituted in the ring system by one (1) or more other univalent substituents.
2			Examples of this class include but are not limited to 3,4-
3			Methylenedioxycathinone (bk-MDA);
4		(b)	By substitution at the 3-position with an acyclic alkyl substituent. Examples of
5			this class include but are not limited to 2-methylamino-1-phenylbutan-1-one
6			(buphedrone);
7		(c)	By substitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl, or
8			methoxybenzyl groups, or by inclusion of the 2-amino nitrogen atom in a
9			cyclic structure. Examples of this class include but are not limited to
10			Dimethylcathinone, Ethcathinone, and $\alpha$ -Pyrrolidinopropiophenone ( $\alpha$ -PPP);
11			or
12		(d)	Any other synthetic cathinone which is not approved by the United States
13			Food and Drug Administration or, if approved, is not dispensed or possessed
14			in accordance with state or federal law;
15	(47)	"Syn	thetic drugs" means any synthetic cannabinoids or piperazines or any synthetic
16		cath	inones;
17	(48)	"Tel	ehealth" has the same meaning it has in KRS 311.550;
18	(49)	"Tet	rahydrocannabinols" means synthetic equivalents of the substances contained in
19		the	plant, or in the resinous extractives of the plant Cannabis, sp. or synthetic
20		subs	tances, derivatives, and their isomers with similar chemical structure and
21		phar	macological activity such as the following:
22		(a)	Delta 1 cis or trans tetrahydrocannabinol, and their optical isomers;
23		(b)	Delta 6 cis or trans tetrahydrocannabinol, and their optical isomers; and
24		(c)	Delta 3, 4 cis or trans tetrahydrocannabinol, and its optical isomers;
25	(50)	"Tra	ffic," except as provided in KRS 218A.1431, means to manufacture, distribute,
26		disp	ense, sell, transfer, or possess with intent to manufacture, distribute, dispense,
27		or se	ell a controlled substance;

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1	(51)	"Transfer" means to dispose of a controlled substance to another person without
2		consideration and not in furtherance of commercial distribution; and
3	(52)	"Ultimate user" means a person who lawfully possesses a controlled substance for
4		his or her own use or for the use of a member of his or her household or for
5		administering to an animal owned by him or her or by a member of his or her
6		household.
7		→ Section 2. KRS 218A.020 is amended to read as follows:
8	(1)	The Cabinet for Health and Family Services shall administer this chapter and may
9		by <u>administrative</u> regulation add substances to or delete or reschedule all
10		substances enumerated in the schedules <u>authorized under</u> [set forth in] this chapter.
11		In making a determination regarding a substance, the Cabinet for Health and Family
12		Services may consider the following:
13		(a) The actual or relative potential for abuse;
14		(b) The scientific evidence of its pharmacological effect, if known;
15		(c) The state of current scientific knowledge regarding the substance;
16		(d) The history and current pattern of abuse;
17		(e) The scope, duration, and significance of abuse;
18		(f) The risk to the public health;
19		(g) The potential of the substance to produce psychic or physiological dependence
20		liability; and
21		(h) Whether the substance is an immediate precursor of a substance already
22		controlled under this chapter.
23	(2)	After considering the factors enumerated in subsection (1) of this section, the
24		Cabinet for Health and Family Services may adopt a regulation controlling the
25		substance if it finds the substance has a potential for abuse.

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If any substance is designated  $\underline{or}_{\{,\}}$  rescheduled[, or deleted] as a controlled

substance under the federal Controlled Substances Act, the drug shall be

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(3)

1	considered to be controlled at the state level in the same numerical schedule
2	corresponding to the federal schedule.
3	(b) Notwithstanding paragraph (a) of this subsection, the Cabinet for Health
4	and Family Services may file an amendment to the administrative
5	regulations promulgated pursuant to this section to control the substance in
6	a more restrictive numerical schedule than the federal schedule as
7	permitted by subsection (1) of this section [law and notice thereof is given to
8	the Cabinet for Health and Family Services, the Cabinet for Health and Family
9	Services may similarly control the substance under this chapter by regulation].
10	(4) The Cabinet for Health and Family Services shall exclude any nonnarcotic
11	substance from a schedule if the substance may be lawfully sold over the counter
12	without prescription under the provisions of the Federal Food, Drug and Cosmetic
13	Act, or the Federal Comprehensive Drug Abuse Prevention and Control Act of
14	1970, or the Kentucky Revised Statutes (for the purposes of this section the
15	Kentucky Revised Statutes shall not include any regulations issued thereunder).
16	(5) The Office of Drug Control Policy may request that the Cabinet for Health and
17	Family Services schedule a substance substantially similar to a synthetic
18	cannabinoid or piperazine or a synthetic cathinone. The cabinet shall consider the
19	request utilizing the criteria established by this section and shall issue a writter
20	response within sixty (60) days of the scheduling request delineating the cabinet's
21	decision to schedule or not schedule the substance and the basis for the cabinet's
22	decision. The cabinet's response shall be provided to the Legislative Research
23	Commission and shall be a public record.
24	→ Section 3. KRS 243.100 is amended to read as follows:
25	A natural person shall not become a licensee under KRS 243.020 to 243.670 if he or she:
26	(1) (a) Has been convicted of any felony until five (5) years have passed from the
27	date of conviction, release from custody or incarceration, parole, or

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1			termination of probation, whichever is later;
2		(b)	Has been convicted of any misdemeanor involving a controlled substance
3			that is described in or classified pursuant to Section 2 of this Act or [ under]
4			KRS <del>[ 218A.050,]</del> 218A.040, 218A.060, <del>[ 218A.070,]</del> 218A.080, <del>[ 218A.090,]</del>
5			218A.100, $\underline{or}$ [218A.110,] 218A.120[, or 218A.130] in the two (2) years
6			immediately preceding the application;
7		(c)	Has been convicted of any misdemeanor directly or indirectly attributable to
8			the use of alcoholic beverages in the two (2) years immediately preceding the
9			application;
10		(d)	Is under the age of twenty-one (21) years;
11		(e)	Has had any license issued under this statute relating to the regulation of the
12			manufacture, sale, and transportation of alcoholic beverages revoked for cause
13			or has been convicted of a violation of any such statute, until the expiration of
14			two (2) years from the date of the revocation or conviction; or
15		(f)	Is not a citizen of the United States and has not had an actual, bona fide
16			residence in this state for at least one (1) year before the date on which his or
17			her application for a license is made. This subsection shall not apply to
18			applicants for manufacturers' licenses, to applicants that are corporations
19			authorized to do business in this state, or to persons licensed on March 7,
20			1938.
21	(2)	A p	partnership, limited partnership, limited liability company, corporation, or
22		gove	ernmental agency shall not be licensed if:
23		(a)	Each member of the partnership or each of the directors, principal officers, or
24			managers does not qualify under subsection (1)(a), (b), (c), and (d) of this
25			section;
26		(b)	It has had any license issued under this statute relating to the regulation of the

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manufacture, sale, and transportation of alcoholic beverages revoked for cause

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or has been convicted of a violation of any such statute, until the expiration of two (2) years from the date of the revocation or conviction; or

- (c) It is a partnership or corporation, if any member of the partnership or any director, manager, or principal officer of the corporation has had any license issued under any statute relating to the regulation of the manufacture, sale, and transportation of alcoholic beverages, revoked for cause or has been convicted of a violation of any such statute, until the expiration of the later of two (2) years from the date of the revocation or two (2) years from the date of conviction.
- (3) The provisions of subsection (1)(a) and (b) shall apply to anyone applying for a new license under this chapter after July 15, 1998, but shall not apply to those who renew a license that was originally issued prior to July 15, 1998, or an application for a supplemental license where the original license was issued prior to July 15, 1998.
  - → Section 4. KRS 243.390 is amended to read as follows:
- 16 (1) In addition to other information as the board may by administrative regulation 17 require, every application for a license under KRS 243.020 to 243.670 shall contain 18 the following information, given under oath:
  - (a) The name, age, Social Security number, address, residence, and citizenship of each applicant;
    - (b) If the applicant is a partner, the name, age, Social Security number, address, residence, and citizenship of each partner and the name and address of the partnership;
    - (c) The name, age, Social Security number, address, residence, and citizenship of each person interested in the business for which the license is sought, together with the nature of that interest, and, if the applicant is a corporation, limited partnership company, or limited liability company, the name, age, Social

Security number, address, and residence of each officer, director, member, partner, and managerial employee and the citizenship of each, and the state under the laws of which the corporate applicant is incorporated or organized. The department may require the names of all the stockholders and the percentage of stock held by each;

- (d) The premises to be licensed, stating the street and number, if the premises has a street number, and otherwise such a description that will reasonably indicate the location of the premises;
- (e) A statement that neither the applicant nor any other person referred to in this section has been convicted of [;] any misdemeanor directly or indirectly attributable to alcoholic beverages; any violation involving a controlled substance that is described in or classified pursuant to Section 2 of this Act or [of] KRS[218A.050,] 218A.040, 218A.060,[218A.070,] 218A.080,[218A.090,] 218A.100, or [218A.110,] 218A.120[, or 218A.130] within the two (2) years immediately preceding the application; any felony, within five (5) years from the later of the date of parole or the date of conviction; or providing false information to the department preceding the application; and that the applicant or any other person referred to in this section has not had any license that has been issued to him under any alcoholic beverage statute revoked for cause within two (2) years prior to the date of the application; and
- (f) A statement that the applicant will in good faith abide by every state and local statute, regulation, and ordinance relating to the manufacture, sale, use of, and trafficking in alcoholic beverages.
- (2) If, after a license has been issued, there is a change in any of the facts required to be set forth in the application, a verified supplemental statement in writing giving notice of the change shall be filed with the board within ten (10) days after the change.

1	(3)	In giving any notice or taking any action in reference to a license, the board may
2		rely upon the information furnished in the application or in the supplemental
3		statement connected with the application. This information, as against the licensee
4		or applicant, shall be conclusively presumed to be correct. The information required
5		to be furnished in the application or supplemental statement shall be deemed
6		material in any prosecution for perjury.

- 7 → Section 5. KRS 243.500 is amended to read as follows:
- 8 Any license issued under KRS 243.020 to 243.670 may be revoked or suspended for the
- 9 following causes:
- 10 (1) Conviction of the licensee or his agent or employee for selling any illegal beverages 11 on the licensed premises.
- 12 (2) Making any false, material statements in an application for a license or supplemental license.
- 14 (3) Violation of the provisions of KRS 243.670.
- 15 (4) Conviction of the licensee or any of his clerks, servants, agents, or employees of:
- 16 (a) Two (2) violations of the terms and provisions of KRS Chapter 241, 243, or
  17 244 or any act regulating the manufacture, sale, and transportation of alcoholic
  18 beverages within two (2) consecutive years;
- 19 (b) Two (2) misdemeanors directly or indirectly attributable to the use of 20 intoxicating liquors within two (2) consecutive years; or
- (c) Any felony.
- 22 (5) Failure or default of a licensee to pay an excise tax or any part of the tax or any penalties imposed by or under the provisions of any statutes, ordinances, or Acts of Congress relative to taxation, or for a violation of any administrative regulations
- promulgated by the Department of Revenue made in pursuance thereof.
- 26 (6) Revocation of any license or permit provided in KRS 243.060, 243.070, 243.600, and 243.610, or granted under any Act of Congress relative to the regulation of the

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1		manufacture, sale, and transportation of alcoholic beverages. Any license issued
2		under KRS 243.020 to 243.670 shall be revoked or suspended if the licensee sells
3		the alcoholic beverages at a price in excess of the price set by federal or state
4		regulations.
5	(7)	Setting up, conducting, operating, or keeping, on the licensed premises, any
6		gambling game, device, machine, contrivance, lottery, gift enterprise, handbook, or

- gambling game, device, machine, contrivance, lottery, gift enterprise, handbook, or facility for betting or transmitting bets on horse races; or permitting to be set up, conducted, operated, kept, or engaged in, on the licensed premises, any such game, device, machine, contrivance, lottery, gift enterprise, handbook, or facility. This 10 section shall not apply to contests in which eligibility to participate is determined by chance and the ultimate winner is determined by skill and the licensee has no direct 12 interest, or to the sale of lottery tickets sold under the provisions of KRS Chapter 13 154A.
- 14 Conviction of the licensee, his agents, servants, or employees for:
- 15 (a) The sale or use upon the licensed premises of those items *classified pursuant* 16 to Section 2 of this Act described in KRS 218A.050 to 218A.130 as 17 controlled substances, including synthetic drugs;
  - (b) Knowingly permitting the sale or use by patrons upon the licensed premises of those items classified pursuant to Section 2 of this Act described in KRS 218A.050 to 218A.130] as controlled substances, including synthetic drugs; or
- 21 Knowingly receiving stolen property upon the licensed premises.
- 22 → Section 6. KRS 314.011 is amended to read as follows:
- 23 As used in this chapter, unless the context thereof requires otherwise:
- 24 (1) "Board" means Kentucky Board of Nursing;

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"Delegation" means directing a competent person to perform a selected nursing 25 (2)26 activity or task in a selected situation under the nurse's supervision and pursuant to 27 administrative regulations promulgated by the board in accordance with the

1		provisions of KRS Chapter 13A;
2	(3)	"Nurse" means a person who is licensed or holds the privilege to practice under the
3		provisions of this chapter as a registered nurse or as a licensed practical nurse;
4	(4)	"Nursing process" means the investigative approach to nursing practice utilizing a
5		method of problem-solving by means of:
6		(a) Nursing diagnosis, a systematic investigation of a health concern, and an
7		analysis of the data collected in order to arrive at an identifiable problem; and
8		(b) Planning, implementation, and evaluation based on nationally accepted
9		standards of nursing practice;
10	(5)	"Registered nurse" means one who is licensed or holds the privilege under the
11		provisions of this chapter to engage in registered nursing practice;
12	(6)	"Registered nursing practice" means the performance of acts requiring substantial
13		specialized knowledge, judgment, and nursing skill based upon the principles of
14		psychological, biological, physical, and social sciences in the application of the
15		nursing process in:
16		(a) The care, counsel, and health teaching of the ill, injured, or infirm;
17		(b) The maintenance of health or prevention of illness of others;
18		(c) The administration of medication and treatment as prescribed by a physician,

- physician assistant, dentist, or advanced practice registered nurse and as further authorized or limited by the board, and which are consistent either with American Nurses' Association Scope and Standards of Practice or with standards of practice established by nationally accepted organizations of registered nurses. Components of medication administration include but are not limited to:
  - 1. Preparing and giving medications in the prescribed dosage, route, and frequency, including dispensing medications only as defined in subsection (17)(b) of this section;

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1		2	. Observing, recording, and reporting desired effects, untoward reactions,
2			and side effects of drug therapy;
3		3	. Intervening when emergency care is required as a result of drug therapy;
4		4	. Recognizing accepted prescribing limits and reporting deviations to the
5			prescribing individual;
6		5	. Recognizing drug incompatibilities and reporting interactions or
7			potential interactions to the prescribing individual; and
8		$\epsilon$	. Instructing an individual regarding medications;
9		(d) T	The supervision, teaching of, and delegation to other personnel in the
10		ŗ	erformance of activities relating to nursing care; and
11		(e) T	The performance of other nursing acts which are authorized or limited by the
12		t	oard, and which are consistent either with American Nurses' Association
13		S	tandards of Practice or with Standards of Practice established by nationally
14		а	ccepted organizations of registered nurses;
15	(7)	"Adva	nced practice registered nurse" or "APRN" means a certified nurse
16		practit	oner, certified registered nurse anesthetist, certified nurse midwife, or
17		clinica	nurse specialist, who is licensed to engage in advance practice registered
18		nursin	g pursuant to KRS 314.042 and certified in at least one (1) population focus;
19	(8)	"Adva	nced practice registered nursing" means the performance of additional acts by
20		registe	red nurses who have gained advanced clinical knowledge and skills through
21		an acc	redited education program that prepares the registered nurse for one (1) of the
22		four (4	APRN roles; who are certified by the American Nurses' Association or
23		other	nationally established organizations or agencies recognized by the board to
24		certify	registered nurses for advanced practice registered nursing as a certified nurse
25		practit	oner, certified registered nurse anesthetist, certified nurse midwife, or
26		clinica	I nurse specialist; and who certified in at least one (1) population focus. The

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additional acts shall, subject to approval of the board, include but not be limited to

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prescribing treatment, drugs, devices, and ordering diagnostic tests. Advanced practice registered nurses who engage in these additional acts shall be authorized to issue prescriptions for and dispense nonscheduled legend drugs as defined in KRS 217.905 and to issue prescriptions for but not to dispense Schedules II through V controlled substances <u>described in or</u> as classified <u>pursuant to Section 2 of this Act or [in]</u> KRS 218A.060, 218A.070, 218A.080, 218A.090, 218A.090, 218A.100, and 218A.110, 218A.120, and 218A.130, under the conditions set forth in KRS 314.042 and regulations promulgated by the Kentucky Board of Nursing on or before August 15, 2006.

- (a) 1. Prescriptions issued by advanced practice registered nurses for Schedule II controlled substances classified under KRS 218A.060, except hydrocodone combination products as defined in KRS 218A.010, shall be limited to a seventy-two (72) hour supply without any refill.
  - 2. Prescriptions issued by advanced practice registered nurses for hydrocodone combination products as defined in KRS 218A.010 shall be limited to a thirty (30) day supply without any refill.
  - 3. Prescriptions issued under this subsection for psychostimulants may be written for a thirty (30) day supply only by an advanced practice registered nurse certified in psychiatric-mental health nursing who is providing services in a health facility as defined in KRS Chapter 216B or in a regional services program for mental health or individuals with an intellectual disability as defined in KRS Chapter 210.
- (b) Prescriptions issued by advanced practice registered nurses for Schedule III controlled substances classified under KRS 218A.080 shall be limited to a thirty (30) day supply without any refill. Prescriptions issued by advanced practice registered nurses for Schedules IV and V controlled substances classified under KRS 218A.100 and 218A.120 shall be limited to the original

prescription and refills not to exceed a six (6) month supply.

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Limitations for specific controlled substances which are identified as having the greatest potential for abuse or diversion, based on the best available scientific and law enforcement evidence, shall be established in an administrative regulation promulgated by the Kentucky Board of Nursing. The regulation shall be based on recommendations from the Controlled Substances Formulary Development Committee, which is hereby created. The committee shall be composed of two (2) advanced practice registered nurses appointed by the Kentucky Board of Nursing, one (1) of whom shall be designated as a committee co-chair; two (2) physicians appointed by the Kentucky Board of Medical Licensure, one (1) of whom shall be designated as a committee cochair; and one (1) pharmacist appointed by the Kentucky Board of Pharmacy. The initial regulation shall be promulgated on or before August 15, 2006, and shall be reviewed at least annually thereafter by the committee.

Nothing in this chapter shall be construed as requiring an advanced practice registered nurse designated by the board as a certified registered nurse anesthetist to obtain prescriptive authority pursuant to this chapter or any other provision of law in order to deliver anesthesia care. The performance of these additional acts shall be consistent with the certifying organization or agencies' scopes and standards of practice recognized by the board by administrative regulation;

- "Licensed practical nurse" means one who is licensed or holds the privilege under (9) the provisions of this chapter to engage in licensed practical nursing practice;
- 23 (10) "Licensed practical nursing practice" means the performance of acts requiring 24 knowledge and skill such as are taught or acquired in approved schools for practical 25 nursing in:
  - (a) The observing and caring for the ill, injured, or infirm under the direction of a registered nurse, advanced practice registered nurse, physician assistant,

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1		licensed physician, or dentist;
2	(b)	The giving of counsel and applying procedures to safeguard life and health, as
3		defined and authorized by the board;

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- (c) The administration of medication or treatment as authorized by a physician, physician assistant, dentist, or advanced practice registered nurse and as further authorized or limited by the board which is consistent with the National Federation of Licensed Practical Nurses or with Standards of Practice established by nationally accepted organizations of licensed practical nurses;
- Teaching, supervising, and delegating except as limited by the board; and (d)
- The performance of other nursing acts which are authorized or limited by the (e) board and which are consistent with the National Federation of Practical Nurses' Standards of Practice or with Standards of Practice established by nationally accepted organizations of licensed practical nurses;
- 15 (11) "School of nursing" means a nursing education program preparing persons for 16 licensure as a registered nurse or a practical nurse;
- 17 (12) "Continuing education" means offerings beyond the basic nursing program that 18 present specific content planned and evaluated to meet competency based 19 behavioral objectives which develop new skills and upgrade knowledge;
- 20 (13) "Nursing assistance" means the performance of delegated nursing acts by unlicensed 21 nursing personnel for compensation under supervision of a nurse;
- 22 (14) "Sexual assault nurse examiner" means a registered nurse who has completed the 23 required education and clinical experience and maintains a current credential from 24 the board as provided under KRS 314.142 to conduct forensic examinations of 25 victims of sexual offenses under the medical protocol issued by the Justice and 26 Public Safety Cabinet in consultation with the Sexual Assault Response Team

27 Advisory Committee pursuant to KRS 216B.400(4);

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1	(15)	"Competency" means the application of knowledge and skills in the utilization of
2		critical thinking, effective communication, interventions, and caring behaviors
3		consistent with the nurse's practice role within the context of the public's health,
4		safety, and welfare;
5	(16)	"Credential" means a current license, registration, certificate, or other similar
6		authorization that is issued by the board;
7	(17)	"Dispense" means:
8		(a) To receive and distribute noncontrolled legend drug samples from
9		pharmaceutical manufacturers to patients at no charge to the patient or any
10		other party; or
11		(b) To distribute noncontrolled legend drugs from a local, district, and
12		independent health department, subject to the direction of the appropriate
13		governing board of the individual health department;
14	(18)	"Dialysis care" means a process by which dissolved substances are removed from a
15		patient's body by diffusion, osmosis, and convection from one (1) fluid
16		compartment to another across a semipermeable membrane;
17	(19)	"Dialysis technician" means a person who is not a nurse, a physician assistant, or a
18		physician and who provides dialysis care in a licensed renal dialysis facility under
19		the direct, on-site supervision of a registered nurse or a physician;
20	(20)	"Population focus" means the section of the population within which the advanced
21		practice registered nurse has targeted to practice. The categories of population foci

- 23 (a) Family and individual across the lifespan;
- 24 (b) Adult gerontology;
- (c) Neonatal;

are:

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- 26 (d) Pediatrics;
- 27 (e) Women's health and gender-related health; and

- 1 (f) Psychiatric mental health; and
- 2 (21) "Conviction" means but is not limited to:
- 3 (a) An unvacated adjudication of guilt;
- 4 (b) Pleading no contest or nolo contendere or entering an Alford plea; or
- 5 (c) Entering a guilty plea pursuant to a pretrial diversion order;
- 6 Regardless of whether the penalty is rebated, suspended, or probated.
- 7 → Section 7. The following KRS sections are repealed:
- 8 218A.030 Controlled substances -- How scheduled.
- 9 218A.050 Schedule I controlled substances.
- 10 218A.070 Schedule II controlled substances.
- 11 218A.090 Schedule III controlled substances.
- 12 218A.110 Schedule IV controlled substances.
- 13 218A.130 Schedule V controlled substances.