1		AN.	ACT relating to the Board of Pharmacy.
2	Be it	t enac	ted by the General Assembly of the Commonwealth of Kentucky:
3		→ S	ection 1. KRS 315.010 is amended to read as follows:
4	As u	ised in	this chapter, unless the context requires otherwise:
5	(1)	"Adı	minister" means the direct application of a drug to a patient or research subject
6		by in	njection, inhalation, or ingestion, whether topically or by any other means;
7	(2)	"Adı	ministrative activities of a pharmacy" means the following functions performed
8		by a	pharmacy adhering to all local, state, and federal patient privacy laws:
9		(a)	Investigating and researching a patient's insurance benefits and updating the
10			patient profile regarding insurance coverage;
11		(b)	Billing and collections activities, including:
12			1. Contacting patients for copayments and coinsurance payments; and
13			2. Communicating with insurance companies;
14		(c)	Performing patient financial assistance activities and updating patient records
15			accordingly;
16		(d)	Opening faxes and accessing electronic prescriptions for the purposes of
17			setting up patient demographic and insurance profiles, excluding height,
18			weight, and allergy information, so long as the activity does not involve the
19			entering of a prescription order into the dispensing or medication management
20			system;
21		(e)	Initiating insurance prior authorizations for submission to the licensed
22			pharmacy, including communications with the prescribing physician to
23			collect, record, and transmit information to insurance companies, so long as
24			the activity does not include the authorization or receipt of new or refill
25			prescription orders;
26		(f)	Answering and transferring telephone calls, whether or not such calls require

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accessing a patient record, so long as the call does not involve the

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1			interpretation, evaluation, or implementation of a drug order; and
2		(g)	Communicating with patients via telephone or electronically regarding refill
3			reminders, so long as the communication does not involve the interpretation,
4			evaluation, or implementation of a drug order and a pharmacist is readily
5			available for patient consultation;
6	(3)	"Ass	sociation" means the Kentucky Pharmacists Association;
7	(4)	"Bo	ard" means the Kentucky Board of Pharmacy;
8	(5)	"Co	llaborative care agreement" means a written agreement between a pharmacist or
9		phar	rmacists and a practitioner or practitioners that outlines a plan of cooperative
10		man	agement of patients' drug-related health care needs where:
11		(a)	Patients' drug-related health care needs fall within the practitioner's or
12			practitioners' statutory scope of practice;
13		(b)	Patients are referred by the practitioner or practitioners to the pharmacist or
14			pharmacists; and
15		(c)	The agreement:
16			1. Identifies the practitioner or practitioners and the pharmacist or
17			pharmacists who are parties to the agreement;
18			2. Specifies the drug-related regimen to be provided, and how drug therapy
19			is to be monitored; and
20			3. Stipulates the conditions for initiating, continuing, or discontinuing drug
21			therapy and conditions which warrant modifications to dose, dosage
22			regimen, dosage form, or route of administration;
23	(6)	"Co	mpound" or "compounding" means the preparation or labeling of a drug
24		purs	tuant to or in anticipation of a valid prescription drug order, including but not
25		limi	ted to packaging, intravenous admixture or manual combination of drug
26		ingr	edients. "Compounding," as used in this chapter, shall not preclude simple

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reconstitution, mixing, or modification of drug products prior to administration by

1		nong	pharmacists;
2	(7)	"Coı	nfidential information" means information which is accessed or maintained by a
3		phar	macist in a patient's record, or communicated to a patient as part of patient
4		cour	seling, whether it is preserved on paper, microfilm, magnetic media, electronic
5		med	ia, or any other form;
6	(8)	"Coı	ntinuing education unit" means ten (10) contact hours of board approved
7		cont	inuing pharmacy education. A "contact hour" means fifty (50) continuous
8		minı	ntes without a break period;
9	(9)	"Dis	pense" or "dispensing" means to deliver one (1) or more doses of a prescription
10		drug	in a suitable container, appropriately labeled for subsequent administration to
11		or us	se by a patient or other individual entitled to receive the prescription drug;
12	(10)	"Dru	g" means any of the following:
13		(a)	Articles recognized as drugs or drug products in any official compendium or
14			supplement thereto;
15		(b)	Articles, other than food, intended to affect the structure or function of the
16			body of man or other animals;
17		(c)	Articles, including radioactive substances, intended for use in the diagnosis,
18			cure, mitigation, treatment or prevention of disease in man or other animals;
19			or
20		(d)	Articles intended for use as a component of any articles specified in
21			paragraphs (a) to (c) of this subsection;
22	(11)	"Dru	ig regimen review" means retrospective, concurrent, and prospective review by
23		a ph	armacist of a patient's drug-related history, including but not limited to the
24		follo	owing areas:
25		(a)	Evaluation of prescription drug orders and patient records for:
26			1. Known allergies;

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Rational therapy contraindications;

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2.

1		3. Appropriate dose and route of administration;
2		4. Appropriate directions for use; or
3		5. Duplicative therapies.
4		b) Evaluation of prescription drug orders and patient records for drug-drug, drug-
5		food, drug-disease, and drug-clinical laboratory interactions;
6		c) Evaluation of prescription drug orders and patient records for adverse drug
7		reactions; or
8		d) Evaluation of prescription drug orders and patient records for proper
9		utilization and optimal therapeutic outcomes;
10	(12)	Immediate supervision" means under the physical and visual supervision of a
11		pharmacist;
12	(13)	'Manufacturer" means any person, except a pharmacist compounding in the normal
13		course of professional practice,[within the Commonwealth] engaged in the
14		commercial production, preparation, propagation, compounding, conversion, or
15		processing of a drug, either directly or indirectly, by extraction from substances of
16		natural origin or independently by means of chemical synthesis, or both, and
17		ncludes any packaging or repackaging of a drug or the labeling or relabeling of its
18		container;
19	(14)	Medical order" means a lawful order of a specifically identified practitioner for a
20		pecifically identified patient for the patient's health care needs. "Medical order"
21		may or may not include a prescription drug order;
22	(15)	Nonprescription drugs" means nonnarcotic medicines or drugs which may be sold
23		without a prescription and are prepackaged and labeled for use by the consumer in
24		accordance with the requirements of the statutes and regulations of this state and the
25		ederal government;
26	(16)	'Outsourcing facility'' means a facility at one (1) geographic location or address
27		hat:

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1	<u>(a)</u>	Is engaged in the compounding of human sterile drugs without a patient-
2		specific prescription;
3	<u>(b)</u>	Has registered as an outsourcing facility with the secretary of the United
4		States Department of Health and Human Services, Food and Drug
5		Administration; and
6	<u>(c)</u>	Complies with all applicable state and federal requirements;
7	<u>(17)</u> "Pha	rmacist" means a natural person licensed by this state to engage in the practice
8	of the	e profession of pharmacy;
9	<u>(18)</u> [(17)]	"Pharmacist intern" means a natural person who is:
10	(a)	Currently certified by the board to engage in the practice of pharmacy under
11		the direction of a licensed pharmacist and who satisfactorily progresses
12		toward meeting the requirements for licensure as a pharmacist;
13	(b)	A graduate of an approved college or school of pharmacy or a graduate who
14		has established educational equivalency by obtaining a Foreign Pharmacy
15		Graduate Examination Committee (FPGEC) certificate, who is currently
16		licensed by the board for the purpose of obtaining practical experience as a
17		requirement for licensure as a pharmacist;
18	(c)	A qualified applicant awaiting examination for licensure as a pharmacist or
19		the results of an examination for licensure as a pharmacist; or
20	(d)	An individual participating in a residency or fellowship program approved by
21		the board for internship credit;
22	<u>(19)</u> [(18)]	"Pharmacy" means every place where:
23	(a)	Drugs are dispensed under the direction of a pharmacist;
24	(b)	Prescription drug orders are compounded under the direction of a pharmacist;
25		or
26	(c)	A registered pharmacist maintains patient records and other information for
27		the purpose of engaging in the practice of pharmacy, whether or not

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1	prescription drug orders are being dispensed;
2	(20)[(19)] "Pharmacy-related primary care" means the pharmacists' activities in patien
3	education, health promotion, and assistance in the selection and use of over-the
4	counter drugs and appliances for the treatment of common diseases and injuries, as
5	well as those other activities falling within their statutory scope of practice;
6	(21)[(20)] "Pharmacy technician" means a natural person who works under the
7	immediate supervision, or general supervision if otherwise provided for by statute
8	or administrative regulation, of a pharmacist for the purpose of assisting a
9	pharmacist with the practice of pharmacy;
10	(22)[(21)] "Practice of pharmacy" means interpretation, evaluation, and implementation
11	of medical orders and prescription drug orders; responsibility for dispensing
12	prescription drug orders, including radioactive substances; participation in drug and
13	drug-related device selection; administration of medications or biologics in the
14	course of dispensing or maintaining a prescription drug order; the administration or
15	adult immunizations pursuant to prescriber-approved protocols; the administration
16	of influenza vaccines to individuals nine (9) to thirteen (13) years of age pursuant to
17	prescriber-approved protocols with the consent of a parent or guardian; the
18	administration of immunizations to individuals fourteen (14) to seventeen (17) years
19	of age pursuant to prescriber-approved protocols with the consent of a parent of
20	guardian; the administration of immunizations to a child as defined in KRS
21	214.032, pursuant to protocols as authorized by KRS 315.500; drug evaluation
22	utilization, or regimen review; maintenance of patient pharmacy records; and
23	provision of patient counseling and those professional acts, professional decisions
24	or professional services necessary to maintain and manage all areas of a patient's
25	pharmacy-related care, including pharmacy-related primary care as defined in this
26	section;

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(23)[(22)] "Practitioner" has the same meaning given in KRS 217.015(35);

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1	<u>(24)[(23)]</u>	"Prescription drug" means a drug which:
2	(a)	Under federal law is required to be labeled with either of the following
3		statements:
4		1. "Caution: Federal law prohibits dispensing without prescription";
5		2. "Caution: Federal law restricts this drug to use by, or on the order of, a
6		licensed veterinarian";
7		3. "Rx Only"; or
8		4. "Rx"; or
9	(b)	Is required by any applicable federal or state law or administrative regulation
10		to be dispensed only pursuant to a prescription drug order or is restricted to
11		use by practitioners;
12	<u>(25)</u> [(24)]	"Prescription drug order" means an original or new order from a practitioner
13	for d	rugs, drug-related devices or treatment for a human or animal, including orders
14	issue	d through collaborative care agreements or protocols authorized by the board.
15	Law	ful prescriptions result from a valid practitioner-patient relationship, are
16	inten	ded to address a legitimate medical need, and fall within the prescribing
17	pract	itioner's scope of professional practice;
18	<u>(26)[(25)]</u>	"Society" means the Kentucky Society of Health-Systems Pharmacists;
19	<u>(27)[(26)]</u>	"Supervision" means the presence of a pharmacist on the premises to which a
20	phar	macy permit is issued, who is responsible, in whole or in part, for the
21	profe	essional activities occurring in the pharmacy; and
22	<u>(28)</u> [(27)]	"Wholesaler" means any person who legally buys drugs for resale or
23	distri	bution to persons other than patients or consumers.
24	→ SI	ECTION 2. A NEW SECTION OF KRS CHAPTER 315 IS CREATED TO
25	READ AS	FOLLOWS:
26	(1) (a)	A person shall not operate an outsourcing facility within this

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Commonwealth, physically or by means of the Internet, facsimile, phone,

1		mail, or any other means, without first obtaining a permit from the board.
2	<u>(b)</u>	An application for a permit to operate an outsourcing facility shall be made
3		to the board upon forms provided by the board and shall contain any
4		information the board requires, which may include affirmative evidence of
5		the applicant's ability to comply with the reasonable standards and
6		regulations as may be prescribed by the board.
7	<u>(c)</u>	Each application shall be accompanied by a nonrefundable permit fee not
8		to exceed five hundred dollars (\$500) to be set by administrative regulation
9		promulgated by the board.
10	(2) As a	prerequisite to obtaining or renewing a permit from the board, the
11	<u>outso</u>	purcing facility shall:
12	<u>(a)</u>	Register as an outsourcing facility with the United States Secretary of
13		Health and Human Services in accordance with 21 U.S.C. sec. 353b; and
14	<u>(b)</u>	Submit a copy of a current inspection report resulting from an inspection
15		conducted by the United States Food and Drug Administration that
16		indicates compliance with the requirements of state and federal law and
17		regulations, including all applicable guidance documents and Current
18		Good Manufacturing Practices published by the United States Food and
19		Drug Administration.
20	(3) (a)	The inspection report required pursuant to subsection (2) of this section
21		shall be deemed current for the purposes of this section if the inspection
22		was conducted:
23		1. No more than one (1) year prior to the date of submission of an
24		application for a permit to the board; or
25		2. No more than two (2) years prior to the date of submission of an
26		application for renewal of a permit to the board.
27	(b)	If the outsourcing facility has not been inspected by the United States Food

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1		and Drug Administration within the period required under subparagraph 1.
2		of paragraph (a) of this subsection, the board may:
3		1. Accept an inspection report or other documentation from another
4		entity that is satisfactory to the board; or
5		2. Cause an inspection to be conducted by its duly authorized agent and
6		charge an inspection fee in an amount sufficient to cover the costs of
7		the inspection.
8	(4) (a)	Upon receipt of an application for a permit to operate an outsourcing
9		facility accompanied by the permit fee prescribed by administrative
10		regulation, the board shall:
11		1. Issue a permit if the outsourcing facility meets the standards and
12		requirements of this chapter and administrative regulations
13		promulgated by the board; or
14		2. Refuse to issue or renew any permit to operate if the outsourcing
15		facility fails to meet the standards and requirements of this chapter
16		and administrative regulations promulgated by the board.
17	<u>(b)</u>	The board shall act upon an application for a permit to operate within thirty
18		(30) days of the receipt of the application. The board may issue a temporary
19		permit to operate in any instance where it considers additional time
20		necessary for investigation and consideration before taking final action on
21		the application. The temporary permit shall be valid for a period of thirty
22		(30) days, unless extended.
23	(5) A se	eparate permit to operate shall be required for each outsourcing facility.
24	(6) (a)	Each permit to operate an outsourcing facility, unless suspended or
25		revoked, shall expire on June 30 following its date of issuance and be
26		renewable annually thereafter upon proper application accompanied by the
27		renewal fee as established by an administrative regulation of the board. The

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1		<u>nonrefundable renewal fee shall not exceed five hundred dollars (\$500).</u>
2		(b) An additional nonrefundable fee not to exceed the annual renewal fee may
3		be assessed and set by administrative regulation as a delinquent renewal
4		penalty for failure to renew by June 30 of each year.
5	<u>(7)</u>	Permits to operate shall be issued only for the premises and persons named in the
6		application and shall not be transferable, except that a buyer may operate the
7		outsourcing facility under the permit of the seller pending a decision by the board
8		on an application which shall be filed by the buyer with the board at least five (5)
9		days prior to the date of sale.
10	<u>(8)</u>	The board may promulgate administrative regulations to ensure:
11		(a) That proper equipment and reference material is on hand considering the
12		nature of the pharmaceutical practice conducted at the particular
13		outsourcing facility: and
14		(b) Health and sanitation standards for areas within outsourcing facilities that
15		adhere to Current Good Manufacturing Practices published by the United
16		States Food and Drug Administration.
17	<u>(9)</u>	Each outsourcing facility shall comply with KRS 218A.202.
18	<u>(10)</u>	Each outsourcing facility shall compound in compliance with the requirements
19		of state and federal law and regulations, including all applicable guidance
20		documents and Current Good Manufacturing Practices published by the United
21		States Food and Drug Administration.
22	<u>(11)</u>	A pharmacist may temporarily operate an outsourcing facility in an area not
23		designated on the permit as authorized in KRS 315.500.
24		→SECTION 3. A NEW SECTION OF KRS CHAPTER 315 IS CREATED TO
25	REA	AD AS FOLLOWS:
26	<u>(1)</u>	(a) Each out-of-state outsourcing facility that does business, physically or by
27		means of the Internet, facsimile, phone, mail, or any other means, inside

1		this Commonwealth shall hold a current outsourcing facility permit issued
2		by the board.
3	<u>(b)</u>	An application for a permit to operate an out-of-state outsourcing facility
4		shall be made to the board upon forms provided by it and shall contain
5		information the board requires, which may include affirmative evidence of
6		the applicant's ability to comply with reasonable standards and regulations
7		as may be prescribed by the board.
8	<u>(c)</u>	Each application shall be accompanied by a nonrefundable permit fee to be
9		set by administrative regulation promulgated by the board. The fee shall
10		<u>not:</u>
11		1. Exceed five hundred dollars (\$500); or
12		2. Exceed the current fee for an in-state outsourcing facility permit.
13	(2) As a	prerequisite to obtaining or renewing a permit from the board, the out-of-
14	<u>state</u>	outsourcing facility shall:
15	<u>(a)</u>	Register as an outsourcing facility with the United States Secretary of
16		Health and Human Services in accordance with 21 U.S.C. sec. 353b; and
17	<u>(b)</u>	Submit a copy of a current inspection report resulting from an inspection
18		conducted by the United States Food and Drug Administration that
19		indicates compliance with the requirements of state and federal law and
20		regulations, including all applicable guidance documents and Current
21		Good Manufacturing Practices published by the United States Food and
22		Drug Administration.
23	(3) (a)	The inspection report required pursuant to paragraph (b) of subsection (2)
24		of this section shall be deemed current for the purposes of this section if the
25		inspection was conducted:
26		1. No more than one (1) year prior to the date of submission of an
27		application for a permit to the board; or

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1			2. No more than two (2) years prior to the date of submission of an
2			application for renewal of a permit to the board.
3		<u>(b)</u>	If the out-of-state outsourcing facility has not been inspected by the United
4			States Food and Drug Administration within the period required under
5			subparagraph 1. of paragraph (a) of this subsection, the board may:
6			1. Accept an inspection report or other documentation from another
7			entity that is satisfactory to the board; or
8			2. Cause an inspection to be conducted by its duly authorized agent and
9			charge a nonrefundable inspection fee in an amount sufficient to
10			cover the costs of the inspection.
11	<u>(4)</u>	(a)	Upon receipt of an application to operate an out-of-state outsourcing
12			facility, accompanied by the permit fee required by subsection (1) of this
13			section, the board shall:
14			1. Issue a permit if the out-of-state outsourcing facility meets the
15			standards and requirements of this chapter and administrative
16			regulations promulgated by the board; or
17			2. Refuse to renew any permit to operate unless the out-of-state
18			outsourcing facility meets the standards and requirements of this
19			chapter and administrative regulations promulgated by the board.
20		<u>(b)</u>	The board shall act upon an application for a permit to operate within thirty
21			(30) days after the receipt thereof. The board may issue a temporary permit
22			to operate in any instance where it considers additional time necessary for
23			investigation and consideration before taking final action upon the
24			application, and the temporary permit shall be valid for a period of thirty
25			(30) days, unless extended.
26	<u>(5)</u>	A seg	parate permit to operate shall be required for each out-of-state outsourcing
27		<u>facili</u>	ity.

1	<u>(0)</u>	Each out-of-state outsourcing facility grantea an out-of-state outsourcing facility
2		permit by the board shall disclose to the board the location, names, and titles of
3		all principal corporate officers and all pharmacists who are dispensing
4		prescription drugs to entities within the Commonwealth. A report containing this
5		information shall be made to the board on an annual basis and within thirty (30)
6		days after any change of office, corporate officer, or pharmacist.
7	<u>(7)</u>	(a) An out-of-state outsourcing facility granted an out-of-state outsourcing
8		facility permit shall comply with all requests for information within three
9		(3) business days of a written request by the board or its agents.
10		(b) An out-of-state outsourcing facility shall maintain at all times a valid
11		unexpired permit, license, or registration to conduct the outsourcing facility
12		in compliance with the laws of the jurisdiction in which it is a resident.
13		(c) As a prerequisite to seeking a permit from the board, the out-of-state
14		outsourcing facility shall submit a copy of the most recent inspection report
15		resulting from an inspection conducted by the regulatory or licensing
16		agency of the jurisdiction in which it is located. Thereafter, the out-of-state
17		outsourcing facility granted a permit shall submit to the board a copy of any
18		subsequent inspection report of the outsourcing facility conducted by the
19		regulatory or licensing body of the jurisdiction in which it is located.
20	<u>(8)</u>	Each out-of-state outsourcing facility granted an out-of-state outsourcing facility
21		permit by the board shall maintain records of any controlled substances or
22		dangerous drugs.
23	<u>(9)</u>	Each out-of-state outsourcing facility shall, during its regular hours of operation,
24		but not less than five (5) days per week and for a minimum of forty (40) hours per
25		week, provide a toll-free telephone service directly to the pharmacist in charge of
26		the out-of-state outsourcing facility for the purpose of facilitating
27		communication. A toll-free number shall be placed on a label affixed to each

1	container of drugs dispensed to an entity within the Commonwealth.
2	(10) An out-of-state outsourcing facility shall comply with KRS 218A.202.
3	(11) An out-of-state outsourcing facility doing business within the Commonwealth of
4	Kentucky shall use the address on file with the board as the return address on the
5	labels of any package shipped into or within the Commonwealth. The return
6	address shall be placed on the package in a clear and prominent manner.
7	(12) (a) A permit to operate an out-of-state outsourcing facility, unless suspended or
8	revoked, shall expire on June 30 following its date of issuance and be
9	renewable annually thereafter upon proper application accompanied by the
10	nonrefundable renewal fee established by subsection (1) of this section.
11	(b) An additional nonrefundable fee not to exceed the annual renewal fee may
12	be assessed and set by administrative regulation as a delinquent renewal
13	penalty for failure to renew by June 30 of each year.
14	(13) Permits to operate shall be issued only for the premises and persons named in the
15	application and shall not be transferable, except that a buyer may operate the
16	out-of-state outsourcing facility under the permit of the seller pending a decision
17	by the board on an application which shall be filed by the buyer with the board at
18	least five (5) days prior to the date of sale.
19	(14) The board may promulgate administrative regulations to ensure that proper
20	equipment and reference material is on hand considering the nature of the
21	pharmaceutical practice conducted at the particular out-of-state outsourcing
22	facility.
23	(15) Each out-of-state outsourcing facility shall compound in compliance with the
24	requirements of state and federal law and regulations, to include all applicable
25	guidance documents and Current Good Manufacturing Practices published by
26	the United States Food and Drug Administration.
27	→ Section 4. KRS 315.400 is amended to read as follows:

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1 As used in KRS 315,400 to 315,412:

distribution in effect; and

- 2 (1) "Authorized distributor of record" means a wholesale distributor that:
- Has established an ongoing relationship with a manufacturer to distribute the manufacturer's prescription drug. An ongoing relationship exists between a wholesale distributor and a manufacturer if the wholesale distributor, including any affiliated group of the wholesale distributor as defined in Section 1504 of the Internal Revenue Code, has a written agreement for
- 9 (b) Is listed on the manufacturer's current list of authorized distributors of record;
- 10 (2) "Co-licensed partner" means two (2) or more entities that have the right to engage in 11 the manufacturing or marketing or both of a prescription drug consistent with the
- Federal Drug Administration's implementation of the federal Prescription Drug
- 13 Marketing Act;

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- 14 (3) "Co-licensed product" means a prescription drug manufactured by two (2) or more co-licensed partners;
 - (4) "Counterfeit prescription drug" means a drug which, or the container or labeling of which, without authorization, bears the trademark, trade name, or other identifying mark, imprint, or device, or any likeness thereof, of a drug manufacturer, processor, packer, or distributor other than the person or persons who in fact manufactured, processed, packed, or distributed the drug and which thereby falsely purports or is represented to be the product of, or to have been packed or distributed by, the other
- drug manufacturer, processor, packer, or distributor;

 23 (5) "Dispenser" means a retail pharmacy, hospital
 - (5) "Dispenser" means a retail pharmacy, hospital pharmacy, a group of chain pharmacies under common ownership and control that do not act as a wholesale distributor, or any other person authorized by law to dispense or administer prescription drugs, and the affiliated warehouses or distribution centers of such entities under common ownership and control that do not act as a wholesale

1	<u> </u>	distr	ibutor. "Dispenser" shall not include a person who dispenses only products
2	<u>t</u>	to be	used in animals in accordance with 21 U.S.C. sec. 360b(a)(4) and (5);
3	<u>(6)</u> '	'Dro	p shipment" means the sale of a prescription drug to a wholesale distributor by
4	t	he c	drug's manufacturer, the manufacturer's co-licensed partner, the manufacturer's
5	t	hird	-party logistics provider, the manufacturer's exclusive distributor, or by an
6	8	autho	orized distributor of record that purchased the product directly from the
7	1	manı	ufacturer, the manufacturer's co-licensed partner, the manufacturer's third-party
8	1	logis	tics provider, or the manufacturer's exclusive distributor, and:
9	((a)	The wholesale distributor takes title to but not physical possession of the drug;
10	((b)	The wholesale distributor invoices the pharmacy, pharmacy warehouse, or
11			other person authorized by law to dispense or administer a prescription drug;
12			and
13	((c)	The pharmacy, pharmacy warehouse, or other person authorized by law to
14			dispense or administer a prescription drug receives delivery directly from the
15			manufacturer, the manufacturer's co-licensed partner, the manufacturer's third-
16			party logistics provider, the manufacturer's exclusive distributor, or an
17			authorized distributor of record;
18	<u>(7)</u> [(6))]	"Emergency medical reasons" includes but is not limited to:
19	((a)	Transfers of a prescription drug between health-care entities or between a
20			health-care entity and a retail pharmacy to alleviate a temporary shortage of a
21			prescription drug arising from delays in or interruptions of the regular
22			distribution schedules;
23	((b)	Sales of drugs for use in the treatment of acutely ill or injured persons to
24			nearby emergency medical services providers, firefighting organizations, or
25			licensed health-care practitioners in the same marketing or service area;
26	((c)	The provision of emergency supplies of drugs to nearby nursing homes, home
27			health agencies, or hospice organizations for emergency use when necessary

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1		drugs cannot be obtained; or
2	(d)	Transfers of prescription drugs by a retail pharmacy to another retail pharmacy
3		to alleviate a temporary shortage;
4	<u>(8)</u> [(7)]	"End user" means a patient or consumer that uses a prescription drug as
5	presc	cribed by an authorized health-care professional;
6	<u>(9)[(8)]</u>	"FDA" means the United States Food and Drug Administration and any
7	succe	essor agency;
8	<u>(10)</u> [(9)]	"Manufacturer" means the same as defined in KRS 315.010;
9	<u>(11)</u> [(10)]	"Manufacturer's exclusive distributor" means a distributor who:
10	(a)	Contracts with a manufacturer to provide or coordinate the warehousing,
11		distributing, or other similar services on behalf of a manufacturer;
12	(b)	Takes title of the prescription drug but does not have responsibility to direct
13		the sale of the manufacturer's prescription drug;
14	(c)	Is licensed under KRS 315.402; and
15	(d)	Is an authorized distributor of record;
16	<u>(12)</u> [(11)]	"Normal distribution channel" means a chain of custody for a prescription
17	drug	from a manufacturer, a manufacturer's co-licensed partner, a manufacturer's
18	third	-party logistics provider, or a manufacturer's exclusive distributor that goes
19	direc	tly, by drop shipment or by intracompany transfer, to:
20	(a)	A pharmacy or other designated person authorized by law to distribute a
21		prescription drug to an end user;
22	(b)	A pharmacy warehouse that performs intracompany sales or transfers of
23		prescription drugs to a group of pharmacies under common ownership and
24		control to a patient, pursuant to a prescription for a patient, or to a person
25		authorized by law to administer a prescription drug for use by a patient;
26	(c)	An authorized distributor of record:

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Then to a pharmacy or other designated person authorized by law to

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1	distribute a prescription drug to an end user;
2	2. Then to a pharmacy warehouse as specified in paragraph (b) of this
3	subsection; or
4	3. Then to another authorized distributor of record to a licensed health-care
5	facility or pharmacy, or a practitioner authorized by law to distribute a
6	prescription drug to an end user; or
7	(d) A nonprofit organization under state contract to distribute prescription drugs
8	to pharmacies pursuant to the state's emergency response plan and the
9	subsequent distribution of those prescription drugs to pharmacies;
10	(13)[(12)] "Pedigree" means a document or electronic file containing information that
11	records each distribution of a prescription drug;
12	(14)[(13)] "Pharmacy warehouse" means a physical location for prescription drugs that
13	acts as a central warehouse and performs intracompany sales or transfers of
14	prescription drugs to a group of pharmacies under common ownership and control;
15	(15)[(14)] "Prescription drug" means the same as defined in KRS 315.010;
16	(16) "Repackager" means a person who owns or operates an establishment that
17	repacks and relabels a product or package for further sale or distribution without
18	a further transaction;
19	(17)[(15)] "Reverse distributor" means every person who acts as an agent for
20	pharmacies, drug wholesalers, manufacturers, or other entities by receiving, taking
21	inventory, and managing the disposition of outdated or nonsalable drugs;
22	(18)[(16)] "Third-party logistics provider" means an entity that contracts with a
23	manufacturer, wholesale distributor, repackager, or dispenser to provide and [or]
24	coordinate[the] warehousing[, distribution,] or other <u>logistics</u> [similar] services on
25	behalf of a manufacturer, wholesale distributor, repackager, or dispenser but does
26	not take title to the drug or have responsibility to direct the sale of the
27	manufacturer's] drug. A third-party logistics provider[who is a licensed wholesale

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1	distr	ibutor under KRS 315.402 and is a manufacturer's authorized distributor of
2	reco	rd] shall be considered as part of the normal distribution channel;
3	<u>(19)</u> [(17)]	"Wholesale distribution" means the distribution of a prescription drug to
4	perso	ons other than an end user, but does not include:
5	(a)	Intracompany sales or transfers;
6	(b)	The sale, purchase, distribution, trade, or transfer of a prescription drug for
7		emergency medical reasons;
8	(c)	The distribution of prescription drug samples by a manufacturer or authorized
9		distributor;
10	(d)	Drug returns or transfers to the original manufacturer, original wholesale
11		distributor, or transfers to a reverse distributor or third-party returns processor;
12	(e)	The sale, purchase, or trade of a drug pursuant to a prescription;
13	(f)	The delivery of a prescription drug by a common carrier;
14	(g)	The purchase or acquisition by a health-care entity or pharmacy that is a
15		member of a group purchasing organization of a drug for its own use from the
16		group purchasing organization, or health-care entities or pharmacies that are
17		members of the group organization;
18	(h)	The sale, purchase, distribution, trade, or transfer of a drug by a charitable
19		health-care entity to a nonprofit affiliate of the organization as otherwise
20		permitted by law;
21	(i)	The sale, transfer, merger, or consolidation of all or part of the business of a
22		pharmacy with another pharmacy or pharmacies; or
23	(j)	The distribution of a prescription drug to a health-care practitioner or to
24		another pharmacy if the total number of units transferred during a twelve (12)
25		month period does not exceed five percent (5%) of the total number of all
26		units dispensed by the pharmacy during the immediate twelve (12) month
27		period; and

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1	(20)[(18)] "Wholesale distributor" means an entity engaged in the wholesale distribution
2	of prescription drugs, including but not limited to manufacturers, manufacturers
3	exclusive distributors, authorized distributors of record, drug wholesalers or
4	distributors, [third-party logistics providers,] third-party returns processors, reverse
5	distributors, and pharmacy warehouses and retail pharmacies that engage in the
6	wholesale distribution of a prescription drug.
7	→SECTION 5. A NEW SECTION OF KRS 315.400 TO 315.412 IS CREATED
8	TO READ AS FOLLOWS:
9	(1) Each facility of a third-party logistics provider located within Kentucky shall be
10	licensed by the board prior to shipping a prescription drug:
11	(a) Within the borders of Kentucky; or
12	(b) To a location outside the borders of Kentucky.
13	(2) A license issued under subsection (1) of this section shall be renewed annually
14	<u>upon:</u>
15	(a) Completion of an application; and
16	(b) Payment of a nonrefundable renewal fee established by the board through
17	the promulgation of an administrative regulation.
18	(3) A third-party logistics provider located in another state seeking to ship a
19	prescription drug into Kentucky shall provide documentation upon request by the
20	board or its staff that the third-party logistics provider is licensed as a third-party
21	logistics provider by:
22	(a) The state from which the third-party logistics provider ships, if that state
23	licenses third-party logistics providers; or
24	(b) The United States Food and Drug Administration.
25	(4) A third-party logistics provider license shall be valid only for the name,
26	ownership, and location listed on the license. Changes of name, ownership, or
27	location shall require a new third-party logistics provider license.

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1	(5) Changes in information required for licensure shall be reported to the board, in
2	writing, within ten (10) days after the change.
3	(6) A third-party logistics provider shall not operate from a place of residence.
4	(7) A third-party logistics provider facility shall be located apart and separate from
5	any retail pharmacy licensed by the board.
6	(8) A third-party logistics provider shall publicly display all licenses and have the
7	most recent state and federal inspection reports readily available.
8	→SECTION 6. A NEW SECTION OF KRS 315.400 TO 315.412 IS CREATED
9	TO READ AS FOLLOWS:
10	(1) An applicant for licensure as a third-party logistics provider shall submit a
11	satisfactorily completed board-approved application together with the required
12	fee. New applicants shall provide, at minimum, the following:
13	(a) The applicant's full name, all trade or business names used, full business
14	address, and telephone number;
15	(b) Type of ownership, whether individual, partnership, limited liability
16	company, or corporation;
17	(c) Name of the owner or owners, including:
18	1. If a person, the name, address, Social Security number, and date of
19	<u>birth;</u>
20	2. If other than a person, the name, address, and Social Security number
21	and date of birth of each partner, limited liability company member, or
22	corporate officer and corporate director and the federal employer
23	identification number;
24	3. If a corporation, the state of incorporation;
25	4. If a publicly traded corporation, the information described in
26	subparagraph 2. of this paragraph is not required for corporate
27	officers and corporate directors; and

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1		(a) Upon the board's written request, a list of all manufacturers, wholesale
2		distributors, repackagers, and dispensers for whom the third-party logistics
3		provider provides services.
4	<u>(2)</u>	Renewal applicants shall provide the items listed in subsection (1) of this section
5		and any other information the board deems necessary to protect the public health
6		and safety as promulgated by administrative regulation.
7	<u>(3)</u>	The board may use a board-approved outside agency, if permitted by federal law,
8		to inspect third-party logistics providers.
9		→ SECTION 7. A NEW SECTION OF KRS 315.400 TO 315.412 IS CREATED
10	TO	READ AS FOLLOWS:
11	<u>(1)</u>	The board shall consider, at a minimum, the following factors in determining the
12		eligibility for initial licensure and renewal of third-party logistics providers:
13		(a) A finding by a law enforcement agency or regulatory agency that the
14		applicant or any of its owners has violated federal, state, or local laws;
15		(b) Suspension, revocation, or any other sanction against a license currently or
16		previously held by the applicant or any of its owners for a violation of
17		federal or state law;
18		(c) A finding that the applicant or any of its owners is guilty of or pleaded
19		guilty or nolo contendere to violating federal, state, or local laws;
20		(d) The furnishing by the applicant of false or fraudulent material in any
21		application;
22		(e) Failure to maintain or make available to the board or to federal, state, or
23		local law enforcement officials those records required to be maintained by
24		third-party logistics providers; and
25		(f) Any other factors or qualifications that the board considers relevant to and
26		consistent with public health and safety. Any factors inconsistent with
27		federal standards shall not be applied.

1	<u>(2)</u>	A licensee who has no record of providing third-party logistics services involving
2		prescription drugs during a routine inspection may have its subsequent renewal
3		application referred to the board for review and possible discipline, and the board
4		may require the licensee to appear before the board at the review.
5	<u>(3)</u>	A third-party logistics provider shall have and follow a diversion detection and
6		loss prevention plan that includes all prescription drugs, which shall be
7		immediately available to the board or its agents upon request.
8	<u>(4)</u>	The board shall have the right to deny licensure if it determines that granting the
9		license would not be consistent with public health and safety.
10		→SECTION 8. A NEW SECTION OF KRS 315.400 TO 315.412 IS CREATED
11	TO I	READ AS FOLLOWS:
12	<u>(1)</u>	Third-party logistics providers shall establish and maintain for board inspection
13		a list of each partner, limited liability company member, corporate officer, and
14		director, including a description of the duties and the qualifications of each.
15	<u>(2)</u>	A third-party logistics provider shall not have as an owner or designated
16		representative anyone convicted of a felony for conduct relating to:
17		(a) Providing third-party logistics services involving prescription drugs;
18		(b) A violation of 21 U.S.C. sec. 331(i) or (k); or
19		(c) A violation of 18 U.S.C. sec. 1365, relating to product tampering.
20	<u>(3)</u>	A third-party logistics provider shall not have as an owner or designated
21		representative anyone who has violated federal or state requirements for third-
22		party logistics provider licensure and presented a threat of serious adverse health
23		consequences or death to humans.
24		→SECTION 9. A NEW SECTION OF KRS 315.400 TO 315.412 IS CREATED
25	TO	READ AS FOLLOWS:
26	<u>(1)</u>	A third-party logistics provider shall operate in compliance with all applicable
27		federal state and local laws and regulations, including but not limited to:

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1	(a) The Drug Supply Chain Security Act of 2013 and rules promulgated
2	thereunder; and
3	(b) The storage practices set out in 21 U.S.C. sec. 360eee-3(d)(2)(C).
4	(2) A third-party logistics provider shall allow the board and authorized federal
5	state, and local law enforcement officials to enter and inspect its premises and
6	delivery vehicles, to audit its records and written operating procedures, and to
7	confiscate prescription drugs and records to the extent authorized by law, rule, or
8	regulation.
9	(3) Failure to operate in compliance with all applicable federal, state, and local law
10	and regulations shall constitute unprofessional conduct pursuant to KRS
11	315.121(1)(a).
12	→SECTION 10. A NEW SECTION OF KRS 315.400 TO 315.412 IS CREATED
13	TO READ AS FOLLOWS:
14	(1) A medical gas wholesaler, whether located within the Commonwealth of
15	operating within the Commonwealth from a location outside the Commonwealth
16	shall be licensed by the board. Each license application shall include a
17	nonrefundable fee which shall:
18	(a) Be prescribed by administrative regulation promulgated by the board in an
19	amount not to exceed two hundred fifty dollars (\$250); and
20	(b) Not be increased by more than twenty-five dollars (\$25) per year.
21	(2) A medical gas wholesaler shall be required to maintain accurate records of all
22	drugs handled. Records shall be made available to agents of the board for
23	inspection upon request.
24	(3) Failure to report to the board or willful submission of inaccurate information
25	shall be grounds for disciplinary action under KRS 315.131.
26	(4) The board shall promulgate administrative regulations to specify the criteria for
27	licensure and discipline of a medical gas wholesaler.

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- Section 11. KRS 315.205 is amended to read as follows:
- 2 Upon the request of an individual or his or her parent or guardian, a pharmacist who
- 3 administers an immunization to an individual who is fourteen (14) to seventeen (17) years
- 4 of age or an influenza vaccine to an individual who is nine (9) to thirteen (13) years of
- 5 age, as authorized in KRS 315.010(22)[(21)], shall provide notification of the
- 6 immunization to the individual's primary care provider.