### As Passed by the House

## **136th General Assembly**

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Am. S. B. No. 152

# Senator Brenner

Cosponsors: Senators Lang, Cirino, DeMora, Hicks-Hudson, Liston, Patton, Reineke, Roegner

Representatives Mathews, T., Brennan, Brownlee, Deeter, Glassburn, Grim, Hall, D., Hiner, Hoops, Kishman, Miller, J., Mohamed, Piccolantonio, Rader, Richardson, Ritter, Robb Blasdel, Salvo, Schmidt, Sigrist, Tims, Williams, Willis

To	amend sections 4729.01, 4729.36, 4729.531,	1
	4729.532, 4729.54, and 4729.55 of the Revised	2
	Code to allow wild animal rehabilitation	3
	facilities to receive a limited license to	4
	administer euthanasia drugs and to modify the	5
	law regarding the use of terms that are limited	6
	to pharmacies and pharmacists.	7

#### BE IT ENACTED BY THE GENERAL ASSEMBLY OF THE STATE OF OHIO:

<b>Section 1</b> . That sections 4729.01, 4729.36, 4729.531,	8
4729.532, 4729.54, and 4729.55 of the Revised Code be amended to	9
read as follows:	10
Sec. 4729.01. As used in this chapter:	11
(A) "Pharmacy," except when used in a context that refers	12
to the practice of pharmacy, means any area, room, rooms, place	13
of business, department, or portion of any of the foregoing	14
where the practice of pharmacy is conducted.	15
(B) "Practice of pharmacy" means providing pharmacist care	16

requiring specialized knowledge, judgment, and skill derived	17
from the principles of biological, chemical, behavioral, social,	18
pharmaceutical, and clinical sciences. As used in this division,	19
"pharmacist care" includes the following:	20
(1) Interpreting prescriptions;	21
(2) Dispensing drugs and drug therapy related devices;	22
(3) Compounding drugs;	23
(4) Counseling individuals with regard to their drug	24
therapy, recommending drug therapy related devices, and	25
assisting in the selection of drugs and appliances for treatment	26
of common diseases and injuries and providing instruction in the	27
proper use of the drugs and appliances;	28
(5) Performing drug regimen reviews with individuals by	29
discussing all of the drugs that the individual is taking and	30
explaining the interactions of the drugs;	31
(6) Performing drug utilization reviews with licensed	32
health professionals authorized to prescribe drugs when the	33
pharmacist determines that an individual with a prescription has	34
a drug regimen that warrants additional discussion with the	35
prescriber;	36
(7) Advising an individual and the health care	37
professionals treating an individual with regard to the	38
<pre>individual's drug therapy;</pre>	39
(8) Acting pursuant to a consult agreement, if an	40
agreement has been established;	41
(9) Engaging in the administration of immunizations to the	42
extent authorized by section 4729.41 of the Revised Code;	43

(10) Engaging in the administration of drugs to the extent	44
authorized by section 4729.45 of the Revised Code.	45
(C) "Compounding" means the preparation, mixing,	46
assembling, packaging, and labeling of one or more drugs in any	47
of the following circumstances:	48
(1) Pursuant to a prescription issued by a licensed health	49
professional authorized to prescribe drugs;	50
(2) Pursuant to the modification of a prescription made in	51
accordance with a consult agreement;	52
(3) As an incident to research, teaching activities, or	53
chemical analysis;	54
(4) In anticipation of orders for drugs pursuant to	55
prescriptions, based on routine, regularly observed dispensing	56
patterns;	57
(5) Pursuant to a request made by a licensed health	58
professional authorized to prescribe drugs for a drug that is to	59
be used by the professional for the purpose of direct	60
administration to patients in the course of the professional's	61
practice, if all of the following apply:	62
(a) At the time the request is made, the drug is not	63
commercially available regardless of the reason that the drug is	64
not available, including the absence of a manufacturer for the	65
drug or the lack of a readily available supply of the drug from	66
a manufacturer.	67
(b) A limited quantity of the drug is compounded and	68
provided to the professional.	69
(c) The drug is compounded and provided to the	70
nrofessional as an occasional exception to the normal practice	71

of dispensing drugs pursuant to patient-specific prescriptions.	72
(D) "Consult agreement" means an agreement that has been	73
entered into under section 4729.39 of the Revised Code.	74
(E) "Drug" means:	75
(1) Any article recognized in the United States	76
pharmacopoeia and national formulary, or any supplement to them,	77
intended for use in the diagnosis, cure, mitigation, treatment,	78
or prevention of disease in humans or animals;	79
(2) Any other article intended for use in the diagnosis,	80
cure, mitigation, treatment, or prevention of disease in humans	81
or animals;	82
(3) Any article, other than food, intended to affect the	83
structure or any function of the body of humans or animals;	84
(4) Any article intended for use as a component of any	85
article specified in division (E)(1), (2), or (3) of this	86
section; but does not include devices or their components,	87
parts, or accessories.	88
"Drug" does not include "hemp" or a "hemp product" as	89
those terms are defined in section 928.01 of the Revised Code.	90
(F) "Dangerous drug" means any of the following:	91
(1) Any drug to which either of the following applies:	92
(a) Under the "Federal Food, Drug, and Cosmetic Act," 52	93
Stat. 1040 (1938), 21 U.S.C.A. 301, as amended, the drug is	94
required to bear a label containing the legend "Caution: Federal	95
law prohibits dispensing without prescription" or "Caution:	96
Federal law restricts this drug to use by or on the order of a	97
licensed veterinarian" or any similar restrictive statement, or	98

the drug may be dispensed only upon a prescription;	99
(b) Under Chapter 3715. or 3719. of the Revised Code, the	100
drug may be dispensed only upon a prescription.	101
(2) Any drug that contains a schedule V controlled	102
substance and that is exempt from Chapter 3719. of the Revised	103
Code or to which that chapter does not apply;	104
(3) Any drug intended for administration by injection into	105
the human body other than through a natural orifice of the human	106
body;	107
(4) Any drug that is a biological product, as defined in	108
section 3715.01 of the Revised Code.	109
(G) "Federal drug abuse control laws" has the same meaning	110
as in section 3719.01 of the Revised Code.	111
(H) "Prescription" means all of the following:	112
(1) A written, electronic, or oral order for drugs or	113
combinations or mixtures of drugs to be used by a particular	114
individual or for treating a particular animal, issued by a	115
licensed health professional authorized to prescribe drugs;	116
(2) For purposes of sections 4723.4810, 4729.282,	117
4730.432, and 4731.93 of the Revised Code, a written,	118
electronic, or oral order for a drug to treat chlamydia,	119
gonorrhea, or trichomoniasis issued to and in the name of a	120
patient who is not the intended user of the drug but is the	121
sexual partner of the intended user;	122
(3) For purposes of sections 3313.7110, 3313.7111,	123
3314.143, 3326.28, 3328.29, 4723.483, 4729.88, 4730.433,	124
4731.96, and 5101.76 of the Revised Code, a written, electronic,	125
or oral order for an epinephrine autoinjector issued to and in	126

the name of a school, school district, or camp;	127
(4) For purposes of Chapter 3728. and sections 4723.483,	128
4729.88, 4730.433, and 4731.96 of the Revised Code, a written,	129
electronic, or oral order for an epinephrine autoinjector issued	130
to and in the name of a qualified entity, as defined in section	131
3728.01 of the Revised Code;	132
(5) For purposes of sections 3313.7115, 3313.7116,	133
3314.147, 3326.60, 3328.38, 4723.4811, 4730.437, 4731.92, and	134
5101.78 of the Revised Code, a written, electronic, or oral	135
order for injectable or nasally administered glucagon in the	136
name of a school, school district, or camp.	137
(I) "Licensed health professional authorized to prescribe	138
drugs" or "prescriber" means an individual who is authorized by	139
law to prescribe drugs or dangerous drugs or drug therapy	140
related devices in the course of the individual's professional	141
practice, including only the following:	142
(1) A dentist licensed under Chapter 4715. of the Revised	143
Code;	144
(2) A clinical nurse specialist, certified nurse-midwife,	145
or certified nurse practitioner who holds a current, valid	146
license issued under Chapter 4723. of the Revised Code to	147
practice nursing as an advanced practice registered nurse;	148
(3) A certified registered nurse anesthetist who holds a	149
current, valid license issued under Chapter 4723. of the Revised	150
Code to practice nursing as an advanced practice registered	151
nurse, but only to the extent of the nurse's authority under	152
sections 4723.43 and 4723.434 of the Revised Code;	153
(4) An optometrist licensed under Chapter 4725. of the	154
Revised Code to practice optometry;	155

(5) A physician authorized under Chapter 4731. of the	156
Revised Code to practice medicine and surgery, osteopathic	157
medicine and surgery, or podiatric medicine and surgery;	158
(6) A physician assistant who holds a license to practice	159
as a physician assistant issued under Chapter 4730. of the	160
Revised Code, holds a valid prescriber number issued by the	161
state medical board, and has been granted physician-delegated	162
prescriptive authority;	163
(7) A veterinarian licensed under Chapter 4741. of the	164
Revised Code;	165
(8) A certified mental health assistant licensed under	166
Chapter 4772. of the Revised Code who has been granted	167
physician-delegated prescriptive authority by the physician	168
supervising the certified mental health assistant.	169
(J) "Sale" or "sell" includes any transaction made by any	170
person, whether as principal proprietor, agent, or employee, to	171
do or offer to do any of the following: deliver, distribute,	172
broker, exchange, gift or otherwise give away, or transfer,	173
whether the transfer is by passage of title, physical movement,	174
or both.	175
(K) "Wholesale sale" and "sale at wholesale" mean any sale	176
in which the purpose of the purchaser is to resell the article	177
purchased or received by the purchaser.	178
(L) "Retail sale" and "sale at retail" mean any sale other	179
than a wholesale sale or sale at wholesale.	180
(M) "Retail seller" means any person that sells any	181
dangerous drug to consumers without assuming control over and	182
responsibility for its administration. Mere advice or	183
instructions regarding administration do not constitute control	184

or establish responsibility.	185
(N) "Price information" means the price charged for a	186
prescription for a particular drug product and, in an easily	187
understandable manner, all of the following:	188
(1) The proprietary name of the drug product;	189
(2) The established (generic) name of the drug product;	190
(3) The strength of the drug product if the product	191
contains a single active ingredient or if the drug product	192
contains more than one active ingredient and a relevant strength	193
can be associated with the product without indicating each	194
active ingredient. The established name and quantity of each	195
active ingredient are required if such a relevant strength	196
cannot be so associated with a drug product containing more than	197
one ingredient.	198
(4) The dosage form;	199
(5) The price charged for a specific quantity of the drug	200
product. The stated price shall include all charges to the	201
consumer, including, but not limited to, the cost of the drug	202
product, professional fees, handling fees, if any, and a	203
statement identifying professional services routinely furnished	204
by the pharmacy. Any mailing fees and delivery fees may be	205
stated separately without repetition. The information shall not	206
be false or misleading.	207
(O) "Wholesale distributor of dangerous drugs" or	208
"wholesale distributor" means a person engaged in the sale of	209
dangerous drugs at wholesale and includes any agent or employee	210
of such a person authorized by the person to engage in the sale	211
of dangerous drugs at wholesale.	212

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(P) "Manufacturer of dangerous drugs" or "manufacturer" 213 means a person, other than a pharmacist or prescriber, who 214 manufactures dangerous drugs and who is engaged in the sale of 215 216 those dangerous drugs. (Q) "Terminal distributor of dangerous drugs" or "terminal 217 distributor" means a person who is engaged in the sale of 218 dangerous drugs at retail, or any person, other than a 219 manufacturer, repackager, outsourcing facility, third-party 220 logistics provider, wholesale distributor, or pharmacist, who 221 222 has possession, custody, or control of dangerous drugs for any 223 purpose other than for that person's own use and consumption. "Terminal distributor" includes pharmacies, hospitals, nursing 224 homes, and laboratories and all other persons who procure 225 dangerous drugs for sale or other distribution by or under the 226 supervision of a pharmacist, licensed health professional 227 authorized to prescribe drugs, or other person authorized by the 228 state board of pharmacy. 229 (R) "Promote to the public" means disseminating a 230 representation to the public in any manner or by any means, 231 other than by labeling, for the purpose of inducing, or that is 232 likely to induce, directly or indirectly, the purchase of a 233 234 dangerous drug at retail. (S) "Person" includes any individual, partnership, 235 association, limited liability company, or corporation, the 236 state, any political subdivision of the state, and any district, 237 department, or agency of the state or its political 238 subdivisions. 239 (T)(1) "Animal shelter" means a facility operated by a 240

humane society or any society organized under Chapter 1717. of

the Revised Code or a dog pound operated pursuant to Chapter

955. of the Revised Code.	243
(2) "County dog warden" means a dog warden or deputy dog	244
warden appointed or employed under section 955.12 of the Revised	245
Code.	246
(3) "Wild animal rehabilitation facility" means a facility	247
that holds a permit issued by the chief of the division of	248
wildlife for rehabilitation purposes in accordance with section	249
1533.08 of the Revised Code or rules adopted by the chief.	250
(U) "Food" has the same meaning as in section 3715.01 of	251
the Revised Code.	252
(V) "Pain management clinic" has the same meaning as in	253
section 4731.054 of the Revised Code.	254
(W) "Investigational drug or product" means a drug or	255
product that has successfully completed phase one of the United	256
States food and drug administration clinical trials and remains	257
under clinical trial, but has not been approved for general use	258
by the United States food and drug administration.	259
"Investigational drug or product" does not include controlled	260
substances in schedule I, as defined in section 3719.01 of the	261
Revised Code.	262
(X) "Product," when used in reference to an	263
investigational drug or product, means a biological product,	264
other than a drug, that is made from a natural human, animal, or	265
microorganism source and is intended to treat a disease or	266
medical condition.	267
(Y) "Third-party logistics provider" means a person that	268
provides or coordinates warehousing or other logistics services	269
pertaining to dangerous drugs including distribution, on behalf	270
of a manufacturer, wholesale distributor, or terminal	271

Sec. 4729.36. (A) No place except a pharmacy licensed as a

terminal distributor of dangerous drugs and no person except a

licensed pharmacist shall knowingly display any sign or

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advertise in any fashion, using the words "pharmacy," "drugs,"	301
"drug store," "drug store supplies," "pharmacist," "druggist,"	302
"pharmaceutical chemist," "apothecary," "drug sundries," or	303
"medicine," or knowingly use any of these words or their	304
equivalent, of those words in any manner that would lead or tend	305
to lead the public to believe that the place is a pharmacy or	306
the person is a pharmacist.	307
(B) A pharmacy making retail sales may advertise by name	308
or therapeutic class the availability for sale or dispensing of	309
any dangerous drug provided that the advertising includes the	310
price information specified in the definition of that term in	311
section 4729.01 of the Revised Code.	312
Sec. 4729.531. (A) The state board of pharmacy may issue a	313
limited license to an animal shelter, a wild animal	314
rehabilitation facility, or county dog warden solely for the	315
purpose of purchasing, possessing, and administering drugs that	316
are distributed in a manufactured dosage form as described in	317
section 4729.532 of the Revised Code. Unless otherwise approved	318
by the board, no such license shall authorize or permit the	319
distribution of these drugs to any person other than the	320
originating wholesale distributor of the drugs. An application	321
for licensure shall include the information the board requires	322
by rule under this section. If the application meets the	323
requirements of the rules adopted under this section, the board	324
shall issue the license.	325
(B) The board, in accordance with Chapter 119. of the	326
Revised Code, shall adopt any rules necessary to administer and	327
enforce this section. The rules shall do all of the following:	328

(1) Require as a condition of licensure that an agent or

employee of an animal shelter or wild animal rehabilitation

<u>facility</u> or an agent or employee of a county dog warden, other	331
than a registered veterinary technician as defined in section	332
4741.01 of the Revised Code, has successfully completed a	333
euthanasia technician certification course described in section	334
4729.532 of the Revised Code;	335
(2) Specify the information the animal shelter, wild	336
animal rehabilitation facility, or county dog warden must	337
provide the board for issuance or renewal of a license;	338
(3) Address any other matters the board considers	339
necessary or appropriate for the administration and enforcement	340
of this section.	341
Sec. 4729.532. (A) No agent or employee of an animal	342
shelter, no agent or employee of a wild animal rehabilitation	343
<pre>facility, and no county dog warden or agent or employee of a</pre>	344
county dog warden shall perform euthanasia by means of lethal	345
injection on an animal by use of any substance other than a	346
substance in a manufactured dosage form that the state	347
veterinary medical licensing board, in consultation with the	348
state board of pharmacy, approves by rule adopted in accordance	349
with Chapter 119. of the Revised Code.	350
The agent or employee of an animal shelter or wild animal	351
rehabilitation facility, county dog warden, or agent or employee	352
of a county dog warden when using a lethal solution to perform	353
euthanasia on an animal shall use the solution in accordance	354
with the following methods:	355
(1) Intravenous injection by hypodermic needle;	356
(2) Intraperitoneal injection by hypodermic needle;	357
(3) Intracardial injection by hypodermic needle, but only	358

on an animal verified to be unconscious;

(4) Oral administration of solution or powder.	360
(B) Before euthanasia, a euthanasia technician may	361
administer a solution of one or more drugs exclusively for the	362
purpose of inducing anesthesia, sedation, or unconsciousness	363
prior to euthanasia. Only those drugs that have been approved by	364
rule adopted in accordance with Chapter 119. of the Revised Code	365
by the state board of pharmacy, in consultation with the state	366
veterinary medical licensing board, may be used.	367
(C) No agent or employee of an animal shelter or wild	368
animal rehabilitation facility and no county dog warden or agent	369
or employee of a county dog warden, other than a registered	370
veterinary technician as defined in section 4741.01 of the	371
Revised Code, shall perform euthanasia by means of lethal	372
injection on an animal or administer pre-euthanasia drugs that	373
induce anesthesia, sedation, or unconsciousness unless the agent	374
or employee or county dog warden has received certification	375
after successfully completing a euthanasia technician	376
certification course as described in this division.	377
The curriculum for a euthanasia technician certification	378
course shall be one that has been approved by the state	379
veterinary medical licensing board, shall be at least sixteen	380
hours in length, and shall include information in at least all	381
of the following areas:	382
(1) The pharmacology, proper administration, and storage	383
of euthanasia, sedation, and anesthesia solutions;	384
(2) Federal and state laws regulating the storage and	385
accountability of euthanasia, sedation, and anesthesia	386
solutions;	387

(3) Euthanasia technician stress management;

(4) Proper disposal of euthanized animals.	389
(D)(1) No agent or employee of an animal shelter or wild	390
animal rehabilitation facility shall perform euthanasia by means	391
of lethal injection on animals or administer pre-euthanasia	392
drugs that induce anesthesia, sedation, or unconsciousness under	393
this section unless the facility in which the agent or employee	394
works or is employed is licensed with the state board of	395
pharmacy under section 4729.531 of the Revised Code. No agent or	396
employee of a county dog warden shall perform euthanasia by	397
means of lethal injection on animals or administer pre-	398
euthanasia drugs that induce anesthesia, sedation, or	399
unconsciousness under this section unless the county dog warden	400
is licensed under section 4729.531 of the Revised Code.	401
(2) Any agent or employee of an animal shelter or wild	402
animal rehabilitation -facility or county dog warden performing	403
euthanasia by means of lethal injection or administering pre-	404
euthanasia drugs that induce anesthesia, sedation, or	405
unconsciousness shall do so only in a humane and proficient	406
manner that is in conformity with the methods described in	407
divisions (A) and (B) of this section and not in violation of	408
Chapter 959. of the Revised Code.	409
(E) Nothing in this section precludes a licensed	410
veterinarian or registered veterinary technician as defined in	411
section 4741.01 of the Revised Code from engaging in the	412
practice of veterinary medicine as authorized in Chapter 4741.	413
of the Revised Code.	414
Sec. 4729.54. (A) As used in this section:	415
(1) "Category II" means any dangerous drug that is not	416
included in category III.	417

(2) "Category III" means any controlled substance that is	418
contained in schedule I, II, III, IV, or V.	419
(3) "Emergency medical service organization" has the same	420
meaning as in section 4765.01 of the Revised Code.	421
(4) "Emergency medical service organization satellite"	422
means a location where dangerous drugs are stored that is	423
separate from, but associated with, the headquarters of an	424
emergency medical service organization. "Emergency medical	425
service organization satellite" does not include the units under	426
the control of the emergency medical service organization.	427
(5) "Person" includes an emergency medical service	428
organization or an emergency medical service organization	429
satellite.	430
(6) "Schedule I," "schedule II," "schedule III," "schedule	431
IV," and "schedule V" have the same meanings as in section	432
3719.01 of the Revised Code.	433
(B)(1) A person seeking to be licensed as a terminal	434
distributor of dangerous drugs shall file with the executive	435
director of the state board of pharmacy a verified application.	436
After it is filed, the application may not be withdrawn without	437
approval of the board.	438
(2) An application shall contain all the following that	439
apply in the applicant's case:	440
(a) Information that the board requires relative to the	441
qualifications of a terminal distributor of dangerous drugs set	442
forth in section 4729.55 of the Revised Code;	443
(b) A statement as to whether the person is seeking to be	444
licensed as a category II. category III. limited category II. or	445

limited category III terminal distributor of dangerous drugs;	446
(c) If the person is seeking to be licensed as a limited	447
category II or limited category III terminal distributor of	448
dangerous drugs, a list of the dangerous drugs that the person	449
is seeking to possess, have custody or control of, and	450
distribute, which list shall also specify the purpose for which	451
those drugs will be used and their source;	452
(d) If the person is an emergency medical service	453
organization, the information that is specified in divisions (C)	454
(1) and (2) of this section, and if the person is an emergency	455
medical service organization satellite, the information required	456
under division (D) of this section;	457
(e) Except with respect to the units under the control of	458
an emergency medical service organization, the identity of the	459
one establishment or place at which the person intends to engage	460
in the sale or other distribution of dangerous drugs at retail,	461
and maintain possession, custody, or control of dangerous drugs	462
for purposes other than the person's own use or consumption;	463
(f) If the application pertains to a pain management	464
clinic, information that demonstrates, to the satisfaction of	465
the board, compliance with division (A) of section 4729.552 of	466
the Revised Code.	467
(C)(1) Each emergency medical service organization that	468
applies for a terminal distributor of dangerous drugs license	469
shall submit with its application all of the following:	470
(a) A copy of its standing orders or protocol, which	471
orders or protocol shall be signed by a physician;	472
(b) A list of the dangerous drugs that the units under its	473
control may carry, expressed in standard dose units, which shall	474

be signed by a physician;	475
(c) A list of the personnel employed or used by the	476
organization to provide emergency medical services in accordance	477
with Chapter 4765. of the Revised Code.	478
In accordance with Chapter 119. of the Revised Code, the	479
board shall adopt rules specifying when an emergency medical	480
service organization that is licensed as a terminal distributor	481
must notify the board of any changes in its documentation	482
submitted pursuant to division (C)(1) of this section.	483
(2) An emergency medical service organization seeking to	484
be licensed as a terminal distributor of dangerous drugs shall	485
list in its application for licensure the following additional	486
information:	487
(a) The units under its control that the organization	488
determines will possess dangerous drugs for the purpose of	489
administering emergency medical services in accordance with	490
Chapter 4765. of the Revised Code;	491
(b) With respect to each such unit, whether the dangerous	492
drugs that the organization determines the unit will possess are	493
in category II or III.	494
(3) An emergency medical service organization that is	495
licensed as a terminal distributor of dangerous drugs shall file	496
a new application for such licensure if there is any change in	497
the number or location of any of its units or if there is any	498
change in the category of the dangerous drugs that any unit will	499
possess.	500
(4) A unit listed in an application for licensure pursuant	501
to division (C)(2) of this section may obtain the dangerous	502
drugs it is authorized to possess from its emergency medical	503

service organization or, on a replacement basis, from a hospital	504
pharmacy. If units will obtain dangerous drugs from a hospital	505
pharmacy, the organization shall file, and maintain in current	506
form, the following items with the pharmacist who is responsible	507
for the hospital's terminal distributor of dangerous drugs	508
license:	509
(a) A copy of its standing orders or protocol;	510
(b) A list of the personnel employed or used by the	511
organization to provide emergency medical services in accordance	512
with Chapter 4765. of the Revised Code, who are authorized to	513
possess the drugs, which list also shall indicate the personnel	514
who are authorized to administer the drugs.	515
(D) Each emergency medical service organization satellite	516
that applies for a terminal distributor of dangerous drugs	517
license shall submit with its application all of the information	518
that the board requires to be submitted with the application, as	519
specified in rules the board shall adopt in accordance with	520
Chapter 119. of the Revised Code.	521
(E) There shall be four categories of terminal distributor	522
of dangerous drugs licenses. The categories are as follows:	523
(1) Category II license. A person who obtains this license	524
may possess, have custody or control of, and distribute only the	525
dangerous drugs described in category II.	526
(2) Limited category II license. A person who obtains this	527
license may possess, have custody or control of, and distribute	528
only the dangerous drugs described in category II that were	529
listed in the application for licensure.	530
(3) Category III license, which may include a pain	531

management clinic classification issued under section 4729.552

f the Revised Code. A person who obtains this license may	533
possess, have custody or control of, and distribute the	534
dangerous drugs described in category II and category III. If the license includes a pain management clinic classification,	535
	536
the person may operate a pain management clinic.	537

- (4) Limited category III license. A person who obtains this license may possess, have custody or control of, and distribute only the dangerous drugs described in category II or category III that were listed in the application for licensure.
- (F) Except for an application made by a county dog warden or on behalf of an animal shelter or wild animal rehabilitation facility, if an applicant for a limited category II license or limited category III license intends to administer dangerous drugs to a person or animal, the applicant shall submit, with the application, a copy of its protocol or standing orders. The protocol or orders shall be signed by a licensed health professional authorized to prescribe drugs, specify the dangerous drugs to be administered, and list personnel who are authorized to administer the dangerous drugs in accordance with federal law or the law of this state.

An application made by a county dog warden or on behalf of an animal shelter or wild animal rehabilitation facility shall include a list of the dangerous drugs to be administered to animals and the personnel who are authorized to administer the drugs to animals in accordance with section 4729.532 of the Revised Code.

In accordance with Chapter 119. of the Revised Code, the board shall adopt rules specifying when a licensee must notify the board of any changes in its documentation submitted pursuant to this division.

(G)(1) Except as provided in division (G)(3) of this	563
section, each applicant for licensure as a terminal distributor	564
of dangerous drugs shall submit, with the application, a license	565
fee. The amount assessed shall not be returned to the applicant	566
if the applicant fails to qualify for the license.	567
(2) The following fees apply under division (G)(1) of this	568
section:	569
(a) Except as provided in division (G)(2)(b) of this	570
section:	571
(i) Three hundred twenty dollars for a category II or	572
limited category II license;	573
(ii) Four hundred forty dollars for a category III	574
license, including a license with a pain management clinic	575
classification issued under section 4729.552 of the Revised	576
Code, or a limited category III license.	577
(b) One hundred twenty dollars for all of the following:	578
(i) A person who is required to hold a license as a	579
terminal distributor of dangerous drugs pursuant to division (C)	580
of section 4729.541 of the Revised Code;	581
(ii) A professional association, corporation, partnership,	582
or limited liability company organized for the purpose of	583
practicing veterinary medicine that is not included in division	584
(G)(2)(b)(i) of this section;	585
(iii) An emergency medical service organization satellite.	586
(3) No fee applies for a license issued to a charitable	587
pharmacy, as defined in section 3719.811 of the Revised Code, if	588
the charitable pharmacy is participating in the drug repository	589
program established under section 3715.87 of the Revised Code.	590

(H)(1) The board shall issue a terminal distributor of	591
dangerous drugs license to each person who submits an	592
application for such licensure in accordance with this section,	593
pays the required license fee, is determined by the board to	594
meet the requirements set forth in section 4729.55 of the	595
Revised Code, and satisfies any other applicable requirements of	596
this section.	597

(2) Except for the license of a county dog warden, the 598 license shall describe the one establishment or place at which 599 the licensee may engage in the sale or other distribution of 600 dangerous drugs at retail and maintain possession, custody, or 601 control of dangerous drugs for purposes other than the 602 licensee's own use or consumption. The one establishment or 603 place shall be that which is identified in the application for 604 licensure. 605

No such license shall authorize or permit the terminal 606 distributor of dangerous drugs named in it to engage in the sale 607 or other distribution of dangerous drugs at retail or to 608 maintain possession, custody, or control of dangerous drugs for 609 any purpose other than the distributor's own use or consumption, 610 at any establishment or place other than that described in the 611 license, except that an agent or employee of an animal shelter 612 or wild animal rehabilitation facility or county dog warden may 613 possess and use dangerous drugs in the course of business as 614 provided in section 4729.532 of the Revised Code. 615

- (3) The license of an emergency medical service 616 organization shall cover the organization's headquarters and, in 617 addition, shall cover and describe all the units of the 618 organization listed in its application for licensure. 619
  - (I) (1) All licenses issued or renewed pursuant to this

section shall be effective for a period specified by the board	621
in rules adopted under section 4729.26 of the Revised Code. The	622
effective period for an initial or renewed license shall not	623
exceed twenty-four months unless the board extends the period in	624
rules to adjust license renewal schedules. A license shall be	625
renewed by the board according to the provisions of this	626
section, the standard renewal procedure of Chapter 4745. of the	627
Revised Code, and rules adopted by the board under section	628
4729.26 of the Revised Code. A person seeking to renew a license	629
shall submit an application for renewal and pay the required fee	630
on or before the date specified in the rules adopted by the	631
board. The fee required for the renewal of a license shall be	632
the same as the license fee that applies under division (G)(2)	633
of this section.	634

- (2) (a) Subject to division (I) (2) (b) of this section, a license that has not been renewed by the date specified in rules adopted by the board may be reinstated only upon payment of the required renewal fee and a penalty fee of one hundred ten dollars.
- (b) If an application for renewal has not been submitted by the sixty-first day after the renewal date specified in rules adopted by the board, the license is considered void and cannot be renewed, but the license holder may reapply for licensure.
- (3) A terminal distributor of dangerous drugs that fails to renew licensure in accordance with this section and rules adopted by the board is prohibited from engaging in the retail sale, possession, or distribution of dangerous drugs until a valid license is issued by the board.
- (J) (1) No emergency medical service organization that is 649 licensed as a terminal distributor of dangerous drugs shall fail 650

to comply with division (C)(1), (3), or (4) of this section.	651
(2) No licensed terminal distributor of dangerous drugs	652
shall possess, have custody or control of, or distribute	653
dangerous drugs that the terminal distributor is not entitled to	654
possess, have custody or control of, or distribute by virtue of	655
its category of licensure.	656
(3) No licensee that is required by division (F) of this	657
section to notify the board of changes in its protocol or	658
standing orders, or in personnel, shall fail to comply with that	659
division.	660
(K) The board may enter into agreements with other states,	661
federal agencies, and other entities to exchange information	662
concerning licensing and inspection of terminal distributors of	663
dangerous drugs located within or outside this state and to	664
investigate alleged violations of the laws and rules governing	665
distribution of drugs by terminal distributors. Any information	666
received pursuant to such an agreement is subject to the same	667
confidentiality requirements applicable to the agency or entity	668
from which it was received and shall not be released without	669
prior authorization from that agency or entity.	670
Sec. 4729.55. No license shall be issued to an applicant	671
for licensure as a terminal distributor of dangerous drugs	672
unless the applicant has furnished satisfactory proof to the	673
state board of pharmacy that:	674
(A) The applicant is equipped as to land, buildings, and	675
equipment to properly carry on the business of a terminal	676
distributor of dangerous drugs within the category of licensure	677
approved by the board.	678
(B) One of the following will maintain supervision and	679

control over the possession and custody of dangerous drugs and	680
controlled substances that may be acquired by or on behalf of	681
<pre>the applicant:</pre>	682
(1) A pharmacist, licensed health professional authorized	683
to prescribe drugs, or other person authorized by the board $\tau$ :	684
(2) An animal shelter, wild animal rehabilitation	685
<pre>facility, or county dog warden licensed under section 4729.531</pre>	686
of the Revised Code, or :	687
(3) A laboratory will maintain supervision and control	688
over the possession and custody of dangerous drugs and	689
controlled substances that may be acquired by or on behalf of	690
the applicant.	691
(C) Adequate safeguards are assured to prevent the sale or	692
other distribution of dangerous drugs by any person other than a	693
pharmacist or licensed health professional authorized to	694
prescribe drugs.	695
(D) Adequate safeguards are assured that the applicant	696
will carry on the business of a terminal distributor of	697
dangerous drugs in a manner that allows pharmacists and pharmacy	698
interns employed by the terminal distributor to practice	699
pharmacy in a safe and effective manner.	700
(E) If the applicant, or any agent or employee of the	701
applicant, has been found guilty of violating section 4729.51 of	702
the Revised Code, the "Federal Food, Drug, and Cosmetic Act," 52	703
Stat. 1040 (1938), 21 U.S.C.A. 301, the federal drug abuse	704
control laws, Chapter 2925., 3715., 3719., or 4729. of the	705
Revised Code, or any rule of the board, adequate safeguards are	706
assured to prevent the recurrence of the violation.	707
(F) If the application is made on behalf of an animal	708

Section 2. That existing sections 4729.01, 4729.36,

4729.531, 4729.532, 4729.54, and 4729.55 of the Revised Code are

Am. S. B. No. 152

hereby repealed.

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