SENATE BILL NO. 249

BY SENATOR CATHEY AND REPRESENTATIVE THOMPSON

1	AN ACT
2	To amend and reenact R.S. 3:1382(E), 1391(7) through (26), 1392(B)(2), 1393(A),
3	1396(A)(1) and (6) through (9), and 1398(A) and to enact R.S. 3:1391(27),
4	1396(A)(10) through (13), and 1400(A)(5), relative to the Louisiana Agricultural
5	Chemistry and Seed Commission; to provide relative to the state chemist's
6	responsibilities; to provide for definitions; to provide relative to the commission's
7	powers and authority; to provide relative to registration and labeling; to provide for
8	commercial feed adulteration; to provide relative to inspection, sampling, and
9	analysis regulations; to provide relative to deficiency assessments; and to provide for
10	related matters.
11	Be it enacted by the Legislature of Louisiana:
12	Section 1. R.S. 3:1382(E), 1391(7) through (26), 1392(B)(2), 1393(A), 1396(A)(1)
13	and (6) through (9), and 1398(A) are hereby amended and reenacted and R.S. 3:1391(27),
14	1396(A)(10) through (13), and 1400(A)(5) are hereby enacted to read as follows:
15	§1382. Commission; creation
16	* * *
17	E. The state chemist shall be responsible for making any chemical analysis
18	or other tests necessary for carrying out the provisions of this Chapter. He shall
19	determine annually the values per pound of nitrogen, available phosphoric acid,
20	potash, and any other substance claimed to have value as a fertilizer. The values so
21	determined shall be used in determining and assessing penalties. In addition to his
22	responsibilities, the following apply:
23	(1) The state chemist shall determine annually the values per pound of
24	nitrogen, available phosphoric acid, potash, and any other substance claimed

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1	to have value as a fertilizer.
2	(2) The state chemist may determine the value of protein and any other
3	substance guaranteed as a commercial feed.
4	(3) The values so determined shall be used in determining and assessing
5	penalties.
6	§1391. Definitions
7	For the purposes of this Part the following definitions shall apply:
8	* * *
9	(7) "Guaranteed feeding units" means the minimum crude protein, minimum
10	crude fat, maximum crude fiber, and minimum or maximum minerals expressed as
11	percentages or other required official units of measure, based on weight and
12	indicated on the label as being contained in the commercial feed.
13	(8) "Guarantor" means the entity listed on a commercial feed label or
14	package that guarantees quality, quantity, and safety of the product.
15	(8)(9) "Ingredient" or "ingredients" means any of the constituent materials
16	making up a commercial feed.
17	(9)(10) "Label" means a display of written, printed, or graphic matter upon
18	or affixed to the container in which a commercial feed is distributed or on the invoice
19	or delivery slip with which a commercial feed is distributed.
20	(10)(11) "Labeling" means all labels and other written, printed, or graphic
21	matter which is located upon a commercial feed or any of its containers or wrapper
22	or accompanying such commercial feed.
23	(11)(12) "Livestock" means cattle, buffalo, bison, oxen, and other bovine;
24	horses, mules, donkeys, and other equine; sheep; goats; swine; domestic rabbits; fish,
25	turtles, and other animals identified with aquaculture that are located in artificial
26	reservoirs or enclosures that are both on privately owned property and constructed
27	so as to prevent, at all times, the ingress and egress of fish life from public waters;
28	imported exotic deer and antelope, elk, farm-raised white-tailed deer, farm-raised
29	ratites, and other farm-raised exotic animals; chickens, turkeys, and other poultry;
30	and animals placed under the jurisdiction of the commissioner of agriculture and

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1	forestry and any hybrid, mixture, or mutation of any such animal.
2	(12)(13) "Manufacture" means to grind, mix, blend, or further process a
3	commercial feed for distribution.
4	(13)(14) "Manufacturer" means a person who manufactures a commercial
5	feed or a customer-formula feed.
6	(14)(15) "Medication" means any drug, antibiotic, or other substance
7	intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease
8	in animals other than man and any substance other than feed ingredients intended to
9	affect the structure or any function of the animal body.
10	(15)(16) "Official sample" means a sample of feed taken by the
11	commissioner or his agent in accordance with provisions of R.S. 3:1398.
12	(16)(17) "Package" means a parcel, bag, or other container.
13	(17)(18) "Percent" or "percentages" mean percentages by weights.
14	(18)(19) "Person" means any individual, partnership, corporation, and
15	association, or other legal entity.
16	(19)(20) "Pet" means any domesticated animal normally maintained in or
17	near the household of the owner thereof.
18	(20)(21) "Pet food" means any commercial feed prepared and distributed for
19	consumption by pets.
20	(21)(22) "Premises" means any place such as, but not exclusively,
21	warehouses, factories, stores, trucks, railroad cars, boats, etc.
22	(22)(23) "Protein derived from mammalian tissues" means any protein
23	containing a portion of mammalian animals, excluding: blood and blood products,
24	gelatin, inspected meat products which have been cooked and offered for human
25	food and further heat-processed for feed such as plate waste and used cellulosic food
26	casings; milk products including milk and milk proteins; and any product in which
27	the only mammalian protein consists entirely of porcine or equine protein.
28	(23)(24) "Registrant" means the person registering a feed with the
29	commission.
30	(24)(25) "Ruminant" includes any mammal of the suborder Ruminantia,

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1	which includes but is not limited to cattle, buffalo, sheep, goats, deer, elk, and
2	antelopes.
3	(25)(26) "Ton" means a net weight of two thousand pounds avoirdupois.
4	(26)(27) "Value of the protein deficiency" means the value of the crude
5	protein as set by the state chemist times the difference between the guaranteed
6	protein analysis and the actual protein analysis of the feed sample.
7	§1392. Commission; powers and authority
8	* * *
9	B. In the interest of uniformity, the commission by regulation may adopt,
10	unless it determines that they are inconsistent with the provisions of this Part or are
11	not appropriate to conditions which exist in this state, the following:
12	* * *
13	(2) Any federal regulation promulgated pursuant to the authority of the
14	Federal Food, Drug, and Cosmetic Act, the Food and Drug Administration, or the
15	Food Safety Modernization Act.
16	* * *
17	§1393. Registration and labeling
18	A. No person shall manufacture a commercial or customer-formula feed for
19	distribution in this state unless he has registered with the commission by filing on
20	forms provided by the commissioner his name, state of incorporation if incorporated,
21	the location of his principal place of business, and the location of each manufacturing
22	facility in this state when such facilities are so located. Registration shall be renewed
23	annually on July first. Renewal of registration may be denied by the commissioner
24	for cause. A distributor or guarantor may apply to the commission for registration
25	as a manufacturer and for authority to label feeds for sale in this state. All provisions
26	applicable to a manufacturer shall then apply to the distributor or guarantor.
27	* * *
28	§1396. Adulteration
29	A commercial feed shall be deemed to be adulterated:
30	(1) If it bears or contains any poisonous or deleterious substance which may

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1	render it injurious to human or animal health. If the substance is not an added
2	substance, the commercial feed shall not be considered adulterated under this
3	Paragraph if the quantity of the substance in the commercial feed does not ordinarily
4	render it injurious to health.
5	* * *
6	(6) If it is, or it bears or contains any new animal drug which is unsafe
7	within the meaning of Section 512 of the Federal Food, Drug, and Cosmetic Act.
8	(7) If it consists in whole or in part of any filthy, putrid, or decomposed
9	substance, or if it is otherwise unfit for feed.
10	(8) If it is, in whole or in part, the product of a diseased animal or of an
11	animal which has died otherwise than by slaughter which is unsafe within the
12	meaning of Section 402(a)(1) or (2) of the Federal Food, Drug, and Cosmetic
13	<u>Act.</u>
14	(9) If any valuable constituent has been in whole or part omitted or abstracted
15	therefrom or any less valuable substance substituted therefor.
16	(7)(10) If its composition or quality falls below or differs from that which it
17	is purported or is represented to possess by its labeling.
18	(11) If the manufacture, processing, packaging, distribution and use do
19	not comply with the requirements of Title 21, Code of Federal Regulations, Part
20	507, Subparts A, B, C, E, and F, except when the commission determines these
21	federal regulations are not appropriate to the conditions which exist in this
22	state.
23	(8)(12) If it contains a drug, as defined by the Act, or antibiotic and the
24	methods used in or the facilities or controls used for its manufacture, processing, or
25	packaging, or distribution and use do not conform to good manufacturing practice
26	regulations promulgated by the commission to assure that the drug meets the
27	requirement of this Part as to safety and has the identity and strength and meets the
28	quality and purity characteristics which it purports or is represented to possess. In
29	promulgating such regulations, the commission shall adopt the good manufacturing
30	practice regulations for Type A medicated articles , medicated feed premixes and

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1	for medicated feeds in accordance with the Federal Food, Drug, and Cosmetic Act
2	and 21 CFR Parts 225 and 507 226, except when the commission determines these
3	federal regulations are not appropriate to the conditions which exist in this state.
4	(9)(13) If it contains viable or poisonous weed seeds in amounts exceeding
5	the limits which the commission shall establish by rule or regulation.
6	* * *
7	§1398. Inspection, sampling, and analysis
8	A. For the purpose of enforcement of this Part and in order to determine
9	whether its provisions have been complied with including whether or not an
10	operation may be subject to such provisions, officers or employees duly designated
11	by the commissioner upon presenting appropriate credentials to the owner, operator,
12	employee in charge, are authorized to enter, during normal business hours, any
13	premises within the state in which commercial feeds are manufactured, processed,
14	packed, held for distribution, or sold or to enter any vehicle being used to
15	commercially transport or hold such feeds; and to obtain official samples and to
16	inspect at reasonable times and within reasonable limits and in a reasonable manner
17	such premises or vehicle and all pertinent equipment, finished and unfinished
18	materials, containers, and labeling thereof. The inspection may include the
19	verification of such records and production and control procedures as may be
20	necessary to determine compliance with the good manufacturing practice regulations
21	for medicated feeds by regulation of the commission. In promulgating such
22	regulations, the commission may adopt the good manufacturing practice
23	regulations in accordance with Title 21, Code of Federal Regulations Part 225,
24	Part 226, and Part 507 Subparts A, B, C, E, and F, except when the commission
25	determines these federal regulations are not appropriate to the conditions which
26	exist in this state. Each such inspection shall be commenced and completed with
27	reasonable promptness. Upon completion of the inspection, the person in charge of
28	the facility or vehicle shall be notified.
29	* * *

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§1400. Deficiency assessments; enforcement

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1	A. If a given lot or shipment of feed is found by official sample and analysis
2	to be deficient in one or more of the guaranteed feeding units, a deficiency
3	assessment of no less than ten dollars shall be assessed against the registrant with
4	respect to the lot or shipment of feed in question in accordance with the following
5	provisions:
6	* * *
7	(5) All other guarantees: A deficiency assessment, not to exceed ten
8	percent of the purchase price of the feed, if the deficiency or excess, where
9	applicable, is greater than the tolerances established by the commission by rule.
10	* * *
11	Section 2. This Act shall become effective upon signature by the governor or, if not
12	signed by the governor, upon expiration of the time for bills to become law without signature
13	by the governor, as provided by Article III, Section 18 of the Constitution of Louisiana. If
14	vetoed by the governor and subsequently approved by the legislature, this Act shall become
15	effective on the day following such approval.

PRESIDENT OF THE SENATE

SPEAKER OF THE HOUSE OF REPRESENTATIVES

GOVERNOR OF THE STATE OF LOUISIANA

APPROVED: _____