

LEGISLATIVE ACTION

Senate Comm: RCS 04/19/2023

House

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The Appropriations Committee on Agriculture, Environment, and General Government (Gruters) recommended the following:

Senate Amendment (with title amendment)

Delete everything after the enacting clause and insert:

Section 1. Present subsections (9) through (24) of section 580.031, Florida Statutes, are redesignated as subsections (10) through (25), respectively, and a new subsection (9) is added to that section, to read:

9 580.031 Definitions of words and terms.—As used in this 10 chapter, the term:

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11	(9) "Dosage form animal product" means a feedstuff that
12	includes any product intended to affect the structure or
13	function of the animal's body other than by providing nutrition
14	to the animal.
15	(a) The term includes oils, tinctures, capsules, tablets,
16	liquids, and chewables.
17	(b) The term does not include:
18	1. Minerals or vitamins;
19	2. Products represented as a primary meal for the intended
20	animal species;
21	3. Products intended as a treat;
22	4. Dental products providing mechanical or abrasive action
23	or both; or
24	5. Drugs, biologics, parasiticides, medical devices, or
25	diagnostics used to treat, or administered to, animals pursuant
26	to:
27	a. The United States Food and Drug Administration Federal
28	Food, Drug, and Cosmetic Act, 21 U.S.C. ss. 301 et seq., as
29	amended;
30	b. The United States Department of Agriculture Federal
31	Virus-Serum-Toxin Act, 21 U.S.C. ss. 151 et seq., as amended; or
32	c. The United States Environmental Protection Agency
33	Federal Insecticide, Fungicide, and Rodenticide Act, 7 U.S.C.
34	ss. 136 et seq., as amended.
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36	Except as provided by law or rule, all terms used in connection
37	with commercial feed or feedstuff have the meanings ascribed to
38	them by the Association of American Feed Control Officials.
39	Section 2. Subsection (1) of section 580.051, Florida

601-03860-23

Statutes, is amended to read:



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580.051 Labels; requirements; penalty.-

42 (1) Any commercial feed or feedstuff distributed in this 43 state, except a customer-formula feed and feed distributed through an integrated poultry operation or by a cooperative to 44 45 its members, shall be accompanied by a legible label bearing all information required by the federal Food and Drug Administration 46 47 and the following information:

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(a) An accurate statement of the net weight.

(b) The name and principal address of the registrant.

(c) The brand name and product name, if any, under which the commercial feed is distributed. The word "medicated" shall be incorporated as part of the brand or product name if the commercial feed contains a drug.

54 1. The department may require feeding directions and 55 precautionary statements to be placed on the label for the safe 56 and effective use of medicated and other feed as deemed 57 necessary.

58 2. Labels on medicated feed shall include all of the 59 following:

60 a. Any feeding directions prescribed by the department to 61 ensure safe usage.

62 b. The stated purpose of the medication contained in the feed as stated in the claim statement. 63

c. The established name of each active drug ingredient. d. The level of each drug used in the final mixture 66 expressed in metric units as well as the required avoirdupois.

(d) The date of manufacture or expiration date of commercial feed sold at retail as the department may by rule

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(e) The guaranteed analysis stated in terms that advise the consumer of the composition of the feed or feedstuff or support claims made in the labeling. In all cases, the elements or compounds listed in the analysis must be determinable by laboratory methods approved by the department. <u>However, products</u> <u>sold solely as dosage form animal products and guaranteed as</u> <u>specified in this section need not show a guaranteed analysis.</u>

1. The guaranteed analysis, listing the minimum percentage of crude protein, minimum percentage of crude fat, and maximum percentage of crude fiber and, when more than 10 percent mineral ingredients are present, the minimum or maximum percentages of mineral elements or compounds as provided by rule.

82 2. Vitamin ingredients, when guaranteed, shall be shown in amounts and terms provided by rule. For mineral feed, the list 83 84 shall include the following: maximum or minimum percentages of calcium (Ca), phosphorus (P), salt (NaCl), iron (Fe), copper 85 86 (Cu), cobalt (Co), magnesium (Mg), manganese (Mn), potassium 87 (K), selenium (Se), zinc (Zn), and fluorine (F) if ingredients used as sources of any of these constituents are declared. All 88 89 mixtures that contain mineral or vitamin ingredients generally 90 regarded as dietary factors essential for the normal nutrition 91 of animals and that are sold or represented for the primary purpose of supplying these minerals or vitamins as additions to 92 93 rations in which these same mineral or vitamin factors may be 94 deficient shall be classified as mineral or vitamin supplements. 95 Products sold solely as mineral or vitamin supplements and 96 quaranteed as specified in this section need not show quarantees for protein, fat, and fiber. 97

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98	3. Other nutritional substances or elements determinable by
99	laboratory methods may be guaranteed by permission of, or shall
100	be guaranteed at the request of, the department as may be
101	provided by rule.
102	(f) The common or usual name of each ingredient used in the
103	manufacture of the commercial feed; however, for all commercial
104	feed except horse feed, the department by rule may permit the
105	use of collective terms for a group of ingredients which perform
106	a similar nutritional function.
107	(g) A label on a dosage form animal product must contain
108	all of the following:
109	1. An accurate statement of the net weight.
110	2. The name and principal address of the registrant.
111	3. The brand name and product name, if any, under which the
112	dosage form animal product is distributed.
113	4. The date of manufacture or expiration date of the dosage
114	form animal product sold at retail as the department may by rule
115	require.
116	5. The amount of each active ingredient per serving.
117	6. The common or usual name of each inactive ingredient
118	contained in the dosage form animal product.
119	7. A statement that identifies how the dosage form animal
120	product supports the structure or function of the animal.
121	8. Precautionary statements and warnings required to ensure
122	the safe and effective use of the dosage form animal product.
123	9. Recommended dosage by animal weight.
124	10. The statement "Not for human consumption."
125	Section 3. This act shall take effect October 1, 2023.
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128	And the title is amended as follows:
129	Delete everything before the enacting clause
130	and insert:
131	A bill to be entitled
132	An act relating to dosage form animal health products;
133	amending s. 580.031, F.S.; defining the term "dosage
134	formula animal product"; providing a definition;
135	amending s. 580.051, F.S.; providing an exception from
136	guaranteed analysis requirements for products sold
137	solely as dosage form animal products; providing
138	labeling requirements for dosage form animal products;
139	providing an effective date.

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