

REQUEST FOR TECHNICAL CHANGE

AGENCY: NC Board of Agriculture

RULE CITATION: 02 NCAC 09B .0116

DEADLINE FOR RECEIPT: Monday, April 9, 2018

PLEASE NOTE: This request may extend to several pages. Please be sure you have reached the end of the document.

The Rules Review Commission staff has completed its review of this Rule prior to the Commission's next meeting. The Commission has not yet reviewed this Rule and therefore there has not been a determination as to whether the Rule will be approved. You may call our office to inquire concerning the staff recommendation.

In reviewing this Rule, the staff recommends that the following technical changes be made:

Page 7, lines 33-35 – delete these lines

Page 7, lines 36-37 – move this to the appropriate subparagraph in Paragraph (o) and renumber the lines that follow.

Page 8, lines 1-2 – move this to the appropriate subparagraph in Paragraph (o) and renumber the lines that follow.

Page 8, lines 3-4 – delete these lines

Page 8, lines 5-7 – renumber these paragraphs

Please retype the rule accordingly and resubmit it to our office at 1711 New Hope Church Road, Raleigh, North Carolina 27609.

Jason Thomas
Commission Counsel
Date submitted to agency: Friday, March 23, 2018

1 02 NCAC 09B .0116 is amended as published in 32:08 NCR 722-726 as follows:

2
3 **02 NCAC 09B .0116 ADOPTIONS BY REFERENCE**

4 (a) The Board incorporates by reference, including subsequent amendments and editions, "Official Methods of
5 Analysis of AOAC," published by the Association of Official Analytical Chemists. Copies of this document may be
6 obtained from the Association of Official Analytical Chemists International, Department 0742, 1970 Chain Bridge
7 Road, McLean, VA 22109-0742, at a cost of six hundred thirty dollars (\$630.00).

8 (b) The Board incorporates by reference, including subsequent amendments and editions, "U.S. Pharmacopeia
9 National Formulary USP XXXIII-NFXXVIII" and supplements, published by the U.S. Pharmacopeial Convention,
10 Inc. Copies of this document may be obtained from The United States Pharmacopeial Convention, Inc., Attention:
11 Customer Service, 12601 Twinbrook Parkway, Rockville, MD 20852, at a cost of eight-hundred fifty dollars
12 (\$850.00).

13 (c) The Board incorporates by reference, including subsequent amendments and editions, "ASTM Standards on
14 Engine Coolants," published by ASTM International. Copies of this document may be obtained from ASTM
15 International, 100 Bar Harbor Drive, West Conshohocken, PA 19428-2959, at a cost of two hundred eleven dollars
16 (\$211.00).

17 (d) The Board incorporates by reference, including subsequent amendments and editions, "EPA Manual of
18 Chemical Methods for Pesticides and Devices" and supplements, published by AOAC. Copies of this document
19 may be obtained online from the Environmental Protection Agency National Service Center for Environmental
20 Publications at <http://nepis.epa.gov/EXE/ZyPURL.cgi?Dockey=2000YS3Y.txt>.

21 (e) The Board incorporates by reference, including subsequent amendments and editions, "Pesticide Analytical
22 Manual," Volumes I and II, published by the United States Department of Health and Human Services, Food and
23 Drug Administration. Copies of this document may be obtained online at <http://www.fda.gov/Food/Science>
24 [Research/LaboratoryMethods/PesticideAnalysisManualPAM/default.htm](http://www.fda.gov/Food/Science).

25 (f) The Board incorporates by reference, including subsequent amendments and editions, "FDA Compliance Policy
26 Guides," published by the United States Department of Health and Human Services, Food and Drug Administration.
27 Copies of this document may be obtained online at <http://www.fda.gov/iceci/compliancemanuals/compliancepolicy>
28 [guidancemanual/default.htm](http://www.fda.gov/iceci/compliancemanuals/compliancepolicy) or from the State Information Branch (HFC-151), Division of Federal-State Relations,
29 US Food and Drug Administration, 5600 Fishers Lane, Room 12-07, Rockville, MD 20857.

30 (g) The Board incorporates by reference, including subsequent amendments and editions, "Bergey's Manual of
31 Determinative Bacteriology," Lippincott, Williams & Wilkins Company, Baltimore. Copies of this document may
32 be obtained from the Lippincott, Williams & Wilkins Company, P.O. Box 1620, Hagerstown, MD 21741 at a cost of
33 one hundred thirty-seven dollars and ninety-nine cents (\$137.99).

34 (h) The Board incorporates by reference, including subsequent amendments and editions, "Microbiology
35 Laboratory Guidebook," published by the United States Department of Agriculture, Food Safety and Inspection
36 Service, Washington, DC. Copies of this document may be obtained online from [http://www.fsis.usda.gov/science/](http://www.fsis.usda.gov/science/microbiological_Lab_Guidebook/)
37 [microbiological_Lab_Guidebook/](http://www.fsis.usda.gov/science/microbiological_Lab_Guidebook/) at no charge.

1 (i) The Board incorporates by reference, including subsequent amendments and editions, "FDA Bacteriological
2 Analytical Manual," published by the United States Department of Health and Human Services, Food and Drug
3 Administration. Copies of this document may be obtained online at <http://www.fda.gov/Food/FoodScience>
4 [Research/LaboratoryMethods/ucm114664.htm](http://www.fda.gov/Food/FoodScience) at no charge.

5 (j) The Board incorporates by reference, including subsequent amendments and editions, "Standard Methods for the
6 Examination of Dairy Products," published by the American Public Health Association. Copies of this document
7 may be obtained from the American Public Health Association Publication Sales, P.O. Box 933019, Atlanta, GA at
8 a cost of eighty-five dollars (\$85.00).

9 (k) The Board incorporates by reference, including subsequent amendments and editions, "Compendium of
10 Methods for the Microbiological Examination of Foods," published by the American Public Health Association.
11 Copies of this document may be obtained from the American Public Health Association Publication Sales, P.O. Box
12 933019, Atlanta, GA at a cost of one hundred fifty dollars (\$150.00).

13 (l) The Board incorporates by reference, including subsequent amendments and editions, "Bergey's Manual of
14 Systematic Bacteriology," Springer Publishing, New York, NY. Copies of this document may be obtained from
15 Springer Publishing, 233 Spring Street, New York, NY, 10013 at a cost of one hundred fifty-nine dollars (\$159.00).

16 (m) The Board incorporates by reference, including subsequent amendments and editions, "Manual of Clinical
17 Microbiology," published by the American Society for Microbiology. Copies of this document may be obtained
18 from the American Society for Microbiology Press, PO Box 605, Herndon, VA 22070, at a cost of two hundred
19 sixty-nine dollars and ninety-five cents (\$269.95).

20 (n) The Board incorporates by reference, including subsequent amendments and editions, "Standard Methods for
21 the Examination of Water and Waste Water," published by American Public Health Association, American Water
22 Works Association, and Water Pollution Control Federation. Copies of this document may be obtained from the
23 American Public Health Association Publication Sales, P.O. Box 933019, Atlanta, GA at a cost of two hundred
24 ninety-five dollars (\$295.00).

25 (o) The Board incorporates by reference, including subsequent amendments and editions, the following parts or
26 sections of the Code of Federal Regulations, Title 21, Chapter I, as promulgated by the Commissioner of the Food
27 and Drug Administration under the authority of the Federal Food, Drug, and Cosmetic Act:

28	Part or	
29	Section	Description of Part or Section
30	(1)	1.1 General
31	(2)	1.3 Labeling - Definitions
32	(3)	1.20 Presence of Mandatory Label Information
33	(4)	1.21 Failure to Reveal Material Facts
34	(5)	1.24 Exemptions from Required Label Statements
35	(6)	1.326 Who is Subject to this Subpart?
36	(7)	1.327 Who is Excluded from All or Part of the Regulations in this Subpart?
37	(8)	1.328 What Definitions Apply to this Subpart?

1	(9)	1.329	Do Other Statutory Provisions and Regulations Apply?
2	(10)	1.330	Can Existing Records Satisfy the Requirements of this Subpart?
3	(11)	1.337	What Information Must Non-transporters Establish and Maintain to Identify the
4			Nontransporter and Transporter Immediate Previous Sources of Food?
5	(12)	1.345	What Information Must Non-transporters Establish and Maintain to Identify the
6			Nontransporter and Transporter Immediate Subsequent Recipients of Food?
7	(13)	1.352	What Information Must Transporters Establish and Maintain?
8	(14)	1.360	What are the Record Retention Requirements?
9	(15)	1.361	What are the Record Availability Requirements?
10	(16)	1.362	What Records are Excluded from this Subpart?
11	(17)	1.363	What are the Consequences of Failing to Establish, or Maintain Records or Make Them
12			Available to FDA as Required by this Subpart?
13	(18)	1.368	What are the Compliance Dates for this Subpart?
14	(19)	2.25	Grain Seed Treated with Poisonous Substances; Color Identification to Prevent
15			Adulteration of Human and Animal Food
16	(20)	2.35	Use of Secondhand Containers for the Shipment or Storage of Food and Animal Feed
17	(21)	7.1	Scope
18	(22)	7.3	Definitions
19	(23)	7.12	Guaranty
20	(24)	7.13	Suggested Forms of Guaranty
21	(25)	7.40	Recall Policy
22	(26)	7.41	Health Hazard Evaluation and Recall Classification
23	(27)	7.42	Recall Strategy
24	(28)	7.45	Food and Drug Administration - Requested Recall
25	(29)	7.46	Firm-initiated Recall
26	(30)	7.49	Recall Communications
27	(31)	7.50	Public Notification of Recall
28	(32)	7.53	Recall Status Reports
29	(33)	7.55	Termination of a Recall
30	(34)	7.59	General Industry Guidance
31	(35)	70	Color Additives
32	(36)	73	Listing of Color Additives Exempt from Certification
33	(37)	74	Listing of Color Additives Subject to Certification
34	(38)	81	General Specifications and General Restrictions for Provisioned Color Additives for Use
35			in Foods, Drugs and Cosmetics
36	(39)	82	Listing of Certified Provisionally Listed Colors and Specifications
37	(40)	100	General

1	(41)	101	Food Labeling
2	(42)	102	Common or Usual Name for Nonstandardized Foods
3	(43)	104	Nutritional Quality Guidelines for Foods
4	(44)	105	Foods for Special Dietary Use
5	(45)	106	Infant Formula Quality Control Procedures
6	(46)	107	Infant Formula
7	(47)	108	Emergency Permit Control
8	(48)	109	Unavoidable Contaminants in Food for Human Consumption and Food-Packaging
9			Material
10	(49)	110	Current Good Manufacturing Practice in Manufacturing, Packing, or Holding Human
11			Food
12	(50)	111	Current Good Manufacturing Practice in Manufacturing, Packaging, Labeling, or Holding
13			Operations for Dietary Supplements
14	(51)	113	Thermally Processed Low-Acid Foods Packaged in Hermetically Sealed Containers
15	(52)	114	Acidified Foods
16	(53)	115	Shell Eggs
17	(54)	118	Production, Storage, and Transportation of Shell Eggs
18	(55)	120	Hazard Analysis and Critical Control Point (HACCP) Systems
19	(56)	123	Fish and Fishery Products
20	(57)	129	Processing and Bottling of Bottled Drinking Water (Except as amended by 02 NCAC 09C
21			.0700 - Bottled Water)
22	(58)	130	Food Standards: General
23	(59)	131	Milk and Cream
24	(60)	133	Cheeses and Related Cheese Products
25	(61)	135	Frozen Desserts
26	(62)	136	Bakery Products
27	(63)	137	Cereal Flours and Related Products
28	(64)	139	Macaroni and Noodle Products
29	(65)	145	Canned Fruits
30	(66)	146	Canned Fruit Juices
31	(67)	150	Fruit Butters, Jellies, Preserves, and Related Products
32	(68)	152	Fruit Pies
33	(69)	155	Canned Vegetables
34	(70)	156	Vegetable Juices
35	(71)	158	Frozen Vegetables
36	(72)	160	Eggs and Egg Products
37	(73)	161	Fish and Shellfish (Except Section 161.30 and 161.130 through 161.145)

1	(74)	163	Cacao Products
2	(75)	164	Tree Nut and Peanut Products
3	(76)	165	Beverages
4	(77)	166	Margarine
5	(78)	168	Sweeteners and Table Syrups
6	(79)	169	Food Dressings and Flavorings
7	(80)	170	Food Additives
8	(81)	172	Food Additives Permitted for Direct Addition to Food for Human Consumption
9	(82)	173	Secondary Direct Food Additives Permitted in Food for Human Consumption
10	(83)	174	Indirect Food Additives: General
11	(84)	175	Indirect Food Additives: Adhesives and Components of Coatings
12	(85)	176	Indirect Food Additives: Paper and Paperboard Components
13	(86)	177	Indirect Food Additives: Indirect Food Additives: Polymers
14	(87)	178	Indirect Food Additives: Adjuvants, Production Aids, and Sanitizers
15	(88)	179	Irradiation in the Production, Processing and Handling of Food
16	(89)	180	Food Additives Permitted in Food or in Contact with Food on an Interim Basis Pending
17			Additional Study
18	(90)	181	Prior-Sanctioned Food Ingredients
19	(91)	182	Substances Generally Recognized as Safe
20	(92)	184	Direct Food Substances Affirmed as Generally Recognized as Safe
21	(93)	186	Indirect Food Substances Affirmed as Generally Recognized as Safe
22	(94)	189	Substances Prohibited from Use in Human Food
23	(95)	190	Dietary Supplements
24	(96)	200	General
25	(97)	201	Labeling
26	(98)	202	Prescription Drug Advertising
27	(99)	210	Current Good Manufacturing Practice in Manufacturing, Processing, Packing or Holding
28			of Drugs; General
29	(100)	211	Current Good Manufacturing Practice for Finished Pharmaceuticals
30	(101)	225	Current Good Manufacturing Practice for Medicated Feeds
31	(102)	226	Current Good Manufacturing Practice for Type A Medicated Articles
32	(103)	250	Special Requirements for Specific Human Drugs
33	(104)	290	Controlled Drugs
34	(105)	299	Drugs; Official Names and Established Names
35	(106)	300	General
36	(107)	310	New Drugs
37	(108)	312	Investigational New Drug Application

1	(109)	314	Applications for FDA Approval to Market New Drug
2	(110)	320	Bioavailability and Bioequivalence Requirements
3	(111)	330	Over-the-Counter (OTC) Human Drugs Which Are Generally Recognized as Safe and
4			Effective and Not Misbranded
5	(112)	331	Antacid Products for Over-the-Counter (OTC) Human Use
6	(113)	332	Antiflatulent Products for Over-the-Counter Human Use
7	(114)	361	Prescription Drugs for Human Use Generally Recognized as Safe and Effective and Not
8			Misbranded: Drugs Used in Research
9	(115)	369	Interpretive Statements Re: Warnings on Drugs and Devices for Over-the-Counter Sale
10	(116)	809	In Vitro Diagnostic Products for Human Use
11	(117)	812	Investigational Device Exemptions
12	(118)	820	Quality System Regulation
13	(119)	860	Medical Device Classification Procedures
14	(120)	861	Procedures for Performance Standards Development
15	(121)	870	Cardiovascular Devices
16	(122)	882	Neurological Devices
17	(123)	884	Obstetrical and Gynecological Devices
18	(124)	895	Banned Devices
19	(125)	500	General
20	(126)	501	Animal Food Labeling
21	(127)	502	Common or Usual Names for Nonstandardized Animal Foods
22	(128)	509	Unavoidable Contaminants in Animal Food and Food-Packaging Material
23	(129)	510	New Animal Drugs
24	(130)	511	New Animal Drugs for Investigational Use
25	(131)	514	New Animal Drug Applications
26	(132)	520	Oral Dosage Form New Animal Drugs
27	(133)	522	Implantation or Injectable Dosage Form New Animal Drugs
28	(134)	524	Ophthalmic and Topical Dosage Form New Animal Drugs
29	(135)	526	Intramammary Dosage Form New Animal Drugs
30	(136)	529	Certain Other Dosage Form New Animal Drugs
31	(137)	556	Tolerances for Residues of New Animal Drugs in Food
32	(138)	558	New Animal Drugs for Use in Animal Feeds
33	(139)	570	Food Additives
34	(140)	573	Food Additives Permitted in Feed and Drinking Water of Animals
35	(141)	582	Substances Generally Recognized as Safe
36	(142)	584	Food Substances Affirmed as Generally Recognized as Safe in Feed and Drinking Water
37			of Animals

1	(143)	589	Substances Prohibited from Use in Animal Food or Feed
2	(144)	700	General
3	(145)	701	Cosmetic Labeling
4	(146)	720	Voluntary Filing of Cosmetic Product Ingredient Composition Statements
5	(147)	740	Cosmetic Product Warning Statements

6 Copies of the Code of Federal Regulations may be obtained at no cost by accessing the website of the U.S.
7 Government Printing Office at <http://www.gpoaccess.gov/cfr/index.html>.

8 (p) The Board incorporates by reference, including subsequent amendments and editions, "Tolerances and
9 Exemptions from Tolerances for Pesticide Chemicals in or on Raw Agricultural Commodities," 40 C.F.R. Part 180.
10 Copies of the Code of Federal Regulations may be obtained at no cost by accessing the website of the U.S.
11 Government Printing Office at <http://www.gpoaccess.gov/cfr/index.html>.

12 (q) The Board incorporates by reference, including subsequent amendments and editions, "Definitions and
13 Standards of Identity or Composition for Meats, Meat By-products, and Meat Food Products," 9 C.F.R. Part 319.
14 Copies of the Code of Federal Regulations may be obtained at no cost by accessing the website of the U.S.
15 Government Printing Office at <http://www.gpoaccess.gov/cfr/index.html>.

16 (r) The Board incorporates by reference, including subsequent amendments and editions, "Definitions and
17 Standards of Identity or Composition for Poultry and Poultry Products," 9 C.F.R. Sections 381.155 through 381.170.
18 Copies of the Code of Federal Regulations may be obtained at no cost by accessing the website of the U.S.
19 Government Printing Office at <http://www.gpoaccess.gov/cfr/index.html>.

20 (s) The Board incorporates by reference, including subsequent amendments and editions, Title 9, Part 317.2(1) of
21 the Code of Federal Regulations. Copies of Title 9 of the Code of Federal Regulations may be obtained from the
22 Superintendent of Documents, Government Printing Office, Washington, DC 20402, at a cost of sixty-four dollars
23 (\$64.00).

24 (t) The Board incorporates by reference, including subsequent amendments and editions, Title 9, Part 381.125(b) of
25 the Code of Federal Regulations. Copies of the Code of Federal Regulations may be obtained at no cost by
26 accessing the website of the U.S. Government Printing Office at <http://www.gpoaccess.gov/cfr/index.html>.

27 (u) The Board incorporates by reference, including subsequent amendments and editions, a document entitled,
28 "Fresh Air '2000' - A Look At FDA's Medical Gas Requirements," published by the United States Department of
29 Health and Human Services, Food and Drug Administration. A copy of this material may be obtained at no cost
30 from the Food and Drug Protection Division of the North Carolina Department of Agriculture and Consumer
31 Services.

32 (v) The Board incorporates by reference the definition of "dietary supplement" found at 21 USC 321(ff).

33 (w) The Board incorporates by reference, including subsequent amendments and editions, the following parts or
34 sections of the Code of Federal Regulations, Title 21, Chapter I, as promulgated by the Commissioner of the Food
35 and Drug Administration under the authority of the Federal Food, Drug, and Cosmetic Act:

36	<u>(1)</u>	<u>112</u>	<u>Standards for the Growing, Harvesting, Packing, and Holding of Produce for Human</u>
37			<u>Consumption</u>

1 (2) 117 Current Good Manufacturing Practice, Hazard Analysis, and Risk-Based Preventive
2 Controls for Human Food

3 Copies of the Code of Federal Regulations may be obtained at no cost by accessing the website of the U.S.
4 Government Printing Office at <http://www.gpoaccess.gov/cfr/index.html>.

5 (x) The Board incorporates by reference the definition of “processed food” found at 21 USC 321(gg).

6 (y) The Board incorporates by reference the definition of “major food allergen” found at 21 USC 321(qq).

7 (z) The Board incorporates by reference the definition of “knowingly” or “knew” found at 21 USC 321(bb).

8
9 *History Note: Authority G.S. 106-139; 106-245.16; 106-245.22; 106-245.32; 106-267;*

10 *Eff. December 14, 1981;*

11 *Amended Eff. May 1, 2013; January 1, 2011; June 1, 2004; April 1, 2003; June 1, 1995; April 1,*
12 *1992; June 1, 1988; October 1, 1987; May 1, 2018;*

13 *Pursuant to G.S. 150B-21.3A, rule is necessary without substantive public interest Eff. March 22,*
14 *2015.*

15

1 02 NCAC 09B .0134 is adopted as published in 32:08 NCR 726 as follows:

2

3 **02 NCAC 09B .0134** **DEFINING ESTABLISHMENT**

4 The term “establishment” under the North Carolina Food, Drugs and Cosmetics Act, N.C. Gen. Stat. § 106-120 et seq.
5 shall include farms as defined under 21 CFR § 112.3, which is hereby incorporated by reference including later
6 amendments or editions and can be accessed free of cost at <http://www.gpoaccess.gov/cfr/index.html>.

7

8 *History Note:* Authority G.S. 106-139;

9 *Eff. May 1, 2018.*

REQUEST FOR TECHNICAL CHANGE

AGENCY: NC Board of Agriculture

RULE CITATION: 02 NCAC 09B .0135

DEADLINE FOR RECEIPT: Monday, April 9, 2018

PLEASE NOTE: This request may extend to several pages. Please be sure you have reached the end of the document.

The Rules Review Commission staff has completed its review of this Rule prior to the Commission's next meeting. The Commission has not yet reviewed this Rule and therefore there has not been a determination as to whether the Rule will be approved. You may call our office to inquire concerning the staff recommendation.

In reviewing this Rule, the staff recommends that the following technical changes be made:

This rule does not impose any requirement, establish any benefit, or otherwise affect the legal rights or obligations of any person. Instead, it appears to hypothesize about legal effect of non-compliance with various rules and statutes. Withdraw this rule.

Please retype the rule accordingly and resubmit it to our office at 1711 New Hope Church Road, Raleigh, North Carolina 27609.

Jason Thomas
Commission Counsel
Date submitted to agency: Friday, March 23, 2018

1 02 NCAC 09B .0135 is adopted as published in 32:08 NCR 726 as follows:

2

3 **02 NCAC 09B .0135 ADULTERATION AND MISBRANDING**

4 (a) Failure to comply with 21 C.F.R. Part 112 as adopted under 02 NCAC 09B .0116(w)(1) may render food
5 adulterated or misbranded, or both, under N.C. Gen. Stat. § 106-129 and N.C. Gen. Stat. § 106-130 for purposes of
6 N.C. Gen. Stat. § 106-122(1)-(3).

7 (b) Failure to comply with 21 C.F.R. Part 117 as adopted under 02 NCAC 09B .0116(w)(2) may render food
8 adulterated or misbranded, or both, under N.C. Gen. Stat. § 106-129 and N.C. Gen. Stat. § 106-130 for purposes of
9 N.C. Gen Stat. § 106-122(1)-(3).

10

11 *History Note: Authority G.S. 106-122; 106-129; 106-130; 106-139;*
12 *Eff. May 1, 2018.*

Withdrawn