

DECLARATION OF EMERGENCY

Department of Health Office of Public Health

Registration of Foods, Drugs, Cosmetics and
Prophylactic Devices
(LAC 49:Chapter 5 and LAC 51.VI.301)

The Louisiana Department of Health, Office of Public Health (LDH/OPH), pursuant to the emergency rulemaking authority granted by R.S. 40:4(A)(13), hereby adopts the following Emergency Rule for the protection of public health. This Emergency Rule is promulgated specifically in accordance with R.S. 49:953(B) of the Administrative Procedure Act (R.S. 49:950 et seq.).

The LDH/OPH finds it necessary to make changes to the Louisiana Administrative Code given the need for regulation of the cannabidiol-containing products made legal for sale to consumers under the provisions of Act No. 164 of the 2019 Louisiana Legislature. The following changes will authorize the LDH/OPH the ability to properly register these items, inspect firms that manufacture such items for human consumption, and conduct oversight of labelling, which could affect the health of Louisiana's citizens and visitors. Further, this Emergency Rule will provide the state health officer the ability to make critical decisions that protect human health. Accordingly, the following Emergency Rule, effective upon signature, shall remain in effect for a maximum of 120 days, or until the final Rule is promulgated, whichever occurs first.

This rule amends §501, §503, §509, and §515, repeals §511, and adds new §§517-529 of Chapter 5 of Title 49—Public Health—Food, Drugs, and Cosmetics. Changes to §501 amend typographical errors in the original language and add new definitions. Changes to §503 reflect changes to the name of the unit and the agency since the promulgation of the original language. Changes to §509 reflect the schedule actually being followed for registrations, which matches with the state's fiscal year (July 1 – June 30). Changes to §515 address the deletion of date language for February 1, 1986. §511 referenced a delinquent penalty schedule no longer in use or authorized by state law. §§517-529 are the new industrial-hemp-derived cannabidiol product registration rules.

Additionally, this rule amends §301 of Chapter 3 of Part VI of Title 51—Public Health—Sanitary Code. Changes to §301 update an adoption-by-reference of federal regulations and add a new rule regarding the inspection of manufacturers of cannabidiol-containing products for human consumption

Title 49

PUBLIC HEALTH—FOOD, DRUGS, AND COSMETICS

Chapter 5. Registration of Foods, Drugs, Cosmetics and Prophylactic Devices

§501. Definitions

[Formerly 49:2.2100]

A. Unless otherwise specifically provided herein, the following words and terms used in this Chapter of Title 49, and all other Chapters of Title 49 which are adopted or may be adopted, are defined for the purposes thereof as follows.

Accrediting Body—for the purposes of this Chapter, the International Organization for Standardization (ISO).

* * *

Cannabidiol—a nonpsychotropic cannabinoid found in *Cannabis sativa* L. and other conspecifics that can have a variety of physiological effects on the human body.

CBD—cannabidiol.

Certificate of Analysis—a document produced by an approved laboratory attesting to the composition of a product.

Certificate of Registration (FD-8)—certificate issued by the Food and Drug/Milk and Dairy Unit attesting that products produced or distributed by the holder's company have been registered with that entity.

* * *

Dietary Supplement—means a product other than tobacco intended to supplement the diet that is not represented for use as a conventional food, that is not a drug, and that is labeled as a dietary supplement and bears or contains one or more of the following dietary ingredients or a concentrate, metabolite, constituent, extract, or combination thereof: a vitamin, a mineral, a botanical, an amino acid, or a dietary substance for use by man to supplement the diet by increasing the total dietary intake.

* * *

Food—includes all substances and preparations used for or entering into the composition of food, drink, confectionery, chewing gum or condiment for man.

Industrial Hemp—the plant *Cannabis sativa* L. and any part of that plant, including the seeds thereof and all derivatives, extracts, cannabinoids, isomers, acids, salts, and salts of isomers, whether growing or not, with a delta-9 tetrahydrocannabinol (THC) concentration of not more than 0.3 percent on a dry weight basis.

Industrial-Hemp-Derived Cannabidiol Products (IHDCP)—any product intended for consumption and containing cannabidiol that was made from industrial hemp.

Industrial Hemp-Derived Cannabidiol Products Database—repository of information on products and firms that are registered with the Food and Drug/Milk and Dairy Unit of LDH/OPH that fall into the category of industrial-hemp-derived cannabidiol products.

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QR Code—Quick Response Code, a type of machine-readable, two-dimensional barcode that stores information about a product.

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THC—delta-9 tetrahydrocannabinol.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), R.S. 40:5(A)(8)(17) and R.S. 40:604.

HISTORICAL NOTE: Adopted by the Louisiana State Board of Health, September 1968, amended by the Louisiana Department of Health, Office of Public Health, LR 45:

§503. Registration Provisions [Formerly 49:2.2110]

A. In accordance with the provisions of R.S. 40:627, each manufacturer, packer or proprietor of processed foods, proprietary or patent medicines, prophylactic devices and cosmetics in packaged form shall register each separate and distinct product annually with the Louisiana Food and Drug/Milk and Dairy Unit/ LDH/OPH.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), R.S. 40:5(A)(8)(17) and R.S. 40:604.

HISTORICAL NOTE: Adopted by the Louisiana State Board of Health, September 1968, amended by the Louisiana Department of Health, Office of Public Health, LR 45:

§509. Product Registration Procedure
[Formerly 49:2.2140]

A. In accordance with the provisions of R.S. 40:627 and 628 and in order to establish revised procedures for the annual registration of products, manufacturers, packers, processors and distributors of all processed foods, proprietary or patent medicines, prophylactic devices and cosmetics in packaged form, whose names appear on the labels, must submit an application for registration of such products on or before July 1 of each year. Certificates of registration will be issued to each firm for a period of one year expiring on June 30 of each year.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), R.S. 40:5(A)(8)(17) and R.S. 40:604.

HISTORICAL NOTE: Adopted by the Louisiana State Board of Health, September 1968, amended by the Department of Health and Human Resources, Office of Health Services and Environmental Quality, LR 10:9 (January 1984), LR 9:562 (August 1983), amended by the Department of Health and Human Resources, Office of Preventive and Public Health Services LR 11:1161 (December 1985), amended by the Louisiana Department of Health, Office of Public Health 45:

§511. Late Registration Penalty Fees
[Formerly 49:2.2150]

Repealed.

AUTHORITY NOTE: Promulgated in accordance with R.S. 40:4(A)(13), R.S. 40:604, and R.S. 40:627(D).

HISTORICAL NOTE: Adopted by the Department of Health and Human Resources, Office of Preventive and Public Health Services, September 1968, amended by the Department of Health and Human Resources, Office of Health Services and Environmental Quality, LR 10:9 (January 1984), LR 9:562 (August 1983), amended by the Department of Health and Human Resources, Office of Preventive and Public Health Services LR 11:1161 (December 1985), repealed by the Louisiana Department of Health, Office of Public Health, LR 45:

§515. Late Registration Penalty Fee Assessment
[Formerly 49:2.2170]

A. The late registration penalty fees as established by Act 344 of the 1985 Louisiana Legislature will assess each manufacturer, packer, or proprietor a penalty of \$10 for failure to register each separate and distinct product annually. The penalty assessed shall be in addition to the examination and investigation charge (registration fee). No manufacturer, packer, or proprietor shall be assessed a late registration penalty fee of more than \$100 in any calendar year.

B. ...

C. Late registration penalty fees will be imposed on those firms which fail to submit an application for registration and registration fees on or before July 1 of each year.

AUTHORITY NOTE: Promulgated in accordance with R.S. 40:4(A)(13), R.S. 40:604, and R.S. 40:627(D).

HISTORICAL NOTE: Adopted by the Louisiana State Board of Health, September 1968, amended by the Department of Health and Human Resources, Office of Health Services and Environmental Quality, LR 10:9 (January 1984), LR 9:562 (August 1983), amended by the Department of Health and Human Resources, Office of Preventive and Public Health Services LR 11:1161 (December 1985), amended by the Louisiana Department of Health, Office of Public Health, LR 45:

§517. Registration of Industrial-Hemp-Derived Cannabidiol Products

A. In accordance with the provisions of R.S. 3:1470 as promulgated by the 2019 Legislature, manufacturers or distributors of industrial-hemp-derived cannabidiol products

must register each separate and distinct product with the Food and Drug/Milk and Dairy Unit of LDH/OPH annually and initially within 90 days of the effective date of these regulations or prior to marketing the products in the state of Louisiana, whichever comes first.

B. The manufacturer of any product that is not registered within the specified timeframe will be deemed to be in violation of these rules with respect to such product(s).

C. In lieu of the annual examination and administration charge normally collected under R.S. 40:628(B), the applicant for an industrial-hemp-derived cannabidiol product registration must provide (both initially and on or before July 1 of each year) the Food and Drug/Milk and Dairy Unit of LDH/OPH with an application form, a cashier's check or money order made payable to the department in the amount of \$50 per each separate and distinct CBD product, specimen copies of labeling in paper or electronic format, and a list of all products the applicant wishes to register with the department. If the packet meets these regulatory requirements, the department will issue to the applicant an FD-8a Certificate of IHDCP (Industrial Hemp-Derived Cannabidiol Products Database) Registration and the application information will be entered into the LDH/OPH Industrial-Hemp-Derived Cannabidiol Products Database.

D. No person is authorized to distribute any industrial-hemp-derived cannabidiol products in the state of Louisiana unless that person has first obtained a Certificate of IHDCP Registration from the Food and Drug/Milk and Dairy Unit.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

§519. Industrial-Hemp-Derived Cannabidiol Products Labeling Requirements: Certificate of Analysis

A. In addition to the requirements enumerated in R.S. 40:608, industrial-hemp-derived cannabidiol products must bear labeling that includes a scannable bar code, QR code, or a web address linked to a document or website containing the certificate of analysis for that product.

B. The certificate of analysis must be from a laboratory that is accredited by LDH/OPH.

C. The certificate of analysis must include, at a minimum, the following information:

1. the batch number of the product;
2. the date the batch was received by the laboratory;
3. the date the testing was completed;
4. the laboratory methodology used for each analysis referenced in the report;
5. the amount of THC by dry weight in milligrams;
6. the amount of CBD by dry weight in milligrams;
7. the amount of any detected residual solvent in the product in parts per million;
8. the amount of any detected pesticide residues in the product in parts per million;
9. the amount of any microbiological contaminants in the product in colony-forming units (CFU) per gram; and
10. the amount of any detected heavy metal traces in the product in parts per million.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

**§521. Industrial-Hemp-Derived Cannabidiol Products
Labeling Requirements: Disclaimer**

A. Each primary container of industrial-hemp-derived cannabidiol product must bear the following statement: "This product has not been evaluated by the Food and Drug Administration and is not intended to diagnose, treat, cure, or prevent any disease."

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

**§523. Industrial-Hemp-Derived Cannabidiol Products
Labeling Requirements: Health Claims
Prohibited**

A. No product labeling or advertising material for any industrial-hemp-derived cannabidiol product sold or otherwise distributed in the state of Louisiana may bear any implicit or explicit health claims.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

**§525. Industrial-Hemp-Derived Cannabidiol Products
Labeling Requirements: Dietary Supplements
Prohibited**

A. No industrial-hemp-derived cannabidiol product may be marketed as a dietary supplement.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

**§527. Penalties for Violations of Requirements to
Register Industrial-Hemp-Derived Cannabidiol
Products**

A. Any person who violates the provisions requiring registration of industrial-hemp-derived cannabidiol products is subject to the penalties provided for by R.S. 3:1484 and other sanctions as provided for by the State Food, Drug, and Cosmetic Law.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

§529. Exemptions

A. Industrial-hemp-derived cannabidiol products that have been produced in accordance with R.S. 40: 1046 or that are FDA-approved pharmaceuticals are not subject to the requirements of this regulation.

AUTHORITY NOTE: Promulgated in accordance with R.S. 3:1482(J), R.S. 40:4(A)(13), and R.S. 40:604.

HISTORICAL NOTE: Promulgated by the Louisiana Department of Health, Office of Public Health, LR 45:

Title 51

PUBLIC HEALTH—SANITARY CODE

**Part VI. Manufacturing, Processing, Packing and
Holding of Food, Drugs and Cosmetics**

**Chapter 3. Current Good Manufacturing Practices
in Manufacturing, Processing, Packing or
Holding Human Food**

**§301. General Provisions; Code of Federal Regulations
[formerly paragraph 6:039]**

A. The criteria in 21 CFR 117 Subpart B and Subpart F (Code of Federal Regulations) shall apply in determining whether the facilities, methods, practices, and controls used in the manufacturing, processing, packing or holding of food

are in conformance with or are operated or administered in conformity with good manufacturing practices to assure that food for human consumption is safe and has been prepared, packed and held under sanitary conditions.

B. In accordance with R.S. 3:1468, facilities producing industrial-hemp-derived cannabidiol products intended for human consumption will be inspected under the provisions of this Chapter.

AUTHORITY NOTE: Promulgated in accordance with the provisions of R.S. 40:4(A)(1)(a). Also see R.S. 40:601 et seq., and R.S. 3:1468.

HISTORICAL NOTE: Promulgated by the Department of Health and Hospitals, Office of Public Health, LR 28:1234 (June 2002), amended by the Louisiana Department of Health, Office of Public Health, LR 45:

Interested persons may submit written comments to Michael Vidrine, Director, Sanitarian Services, Office of Public Health, Louisiana Department of Health, P.O. Box 4489, Baton Rouge, LA 70821-4489. He is responsible for responding to inquiries regarding this Emergency Rule.

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State Health Officer
and
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