

1960. No. 187

[C]

FOOD AND DRUGS**Butter and Margarine**

REGULATIONS, DATED 23RD NOVEMBER, 1960, MADE BY THE MINISTRY OF HEALTH AND LOCAL GOVERNMENT UNDER SECTIONS 4, 48 AND 68 OF THE FOOD AND DRUGS ACT (NORTHERN IRELAND), 1958.

The Ministry of Health and Local Government, in exercise of the powers conferred upon it by Sections 4, 48 and 68 of the Food and Drugs Act (Northern Ireland), 1958(a), hereby makes the following regulations:—

Citation

1. These regulations may be cited as the Food Standards (Butter and Margarine) Regulations (Northern Ireland), 1960.

Interpretation

2. In these Regulations—

“the Act” means the Food and Drugs Act (Northern Ireland), 1958;

“butter” means the substance usually known as butter made exclusively from milk with or without salt and with or without the addition of colouring matter;

“catering business” includes the business or undertaking of an inn, public-house, hotel, restaurant, café, tea-shop, buffet, coffee-stall or any place of refreshment open to the public, or of a club, boarding-house, apartment-house, refreshment contractor, school feeding-centre, staff dining-room or canteen; and the word “caterer” shall be construed accordingly;

“margarine” means the food usually known as margarine being an emulsion of edible oils and fats with water or skimmed milk or other substances, with or without the addition of colouring matter, which is capable of being used for the same purposes as butter; and for the purpose of regulations 4(2), 5 and 6 any reference to “margarine” shall be construed as including a reference to any food, whether mixed with butter or not, which resembles butter;

“sale” includes an offer for sale and exposure for sale;

“sale by retail” means a sale to a person otherwise than for the purpose of re-sale and includes a sale of margarine as such by a caterer in the course of his catering business but does not include a sale to a manufacturer for the purposes of his manufacturing business or a sale to a caterer for the purposes of his catering business;

“vitamin D” means the anti-rachitic vitamins.

Standard for butter

3. The standard for butter as respects any water contained therein shall be as follows, that is to say; butter shall not contain more than 16 per cent. of water.

Standard for margarine

4.—(1) The standard for margarine as respects any water contained therein and as respects the fat content thereof shall be as follows, that is to say:—

(a) 1958. c. 27.

- (a) margarine shall not contain more than 16 per cent. of water;
- (b) the fat content of margarine shall not contain more than 10 per cent. of fat derived from milk.

(2) The standard for margarine, as respects vitamin A and vitamin D contained therein, shall be as specified in the First Schedule.

5. The standard for margarine prescribed in regulation 4(2) shall apply only as respects sales by retail.

6. No person shall sell or offer or expose for sale any food intended for human consumption under such a description as to lead an intending purchaser to believe he is purchasing butter or margarine unless the food complies with the said standards.

7. Where a person sells any food to a purchaser in response to a request for butter or margarine he shall be deemed to sell butter or margarine respectively unless he clearly notifies the purchaser at the time of sale that the food is not butter or margarine.

Penalties

8. If any person contravenes or fails to comply with any of the provisions of these regulations he shall be guilty of an offence and shall be liable on summary conviction—

- (a) to a fine not exceeding one hundred pounds or to imprisonment for a term not exceeding three months, or to both such fine and such imprisonment; and
- (b) in the case of a continuing offence, to a further fine not exceeding five pounds for each day during which the offence continues after conviction.

Legal Proceedings—Application of Act

9. In any prosecution for an offence under these regulations the relevant provisions of Sections 46, 48, 50 and 53 of the Act shall apply.

Revocation

10. The Sale of Butter (Ireland) Regulations, 1902, are hereby revoked.

Sealed with the Official Seal of the Ministry of Health and Local Government for Northern Ireland this 23rd day of November, nineteen hundred and sixty, in the presence of

(L.S.)

J. L. O. Andrews,
Minister of Health and Local Government.

FIRST SCHEDULE

The standard for margarine shall be as follows:—

Each ounce of margarine shall contain—

- (a) not less than 760 international units and not more than 940 international units of vitamin A determined in accordance with the method set forth in the Second Schedule. The vitamin A content shall be calculated as the sum of the vitamin A present as such or as its esters plus 0.8 times the beta-carotene equivalent of any carotenes present; any alpha-carotene being considered as equivalent in potency to half its weight of beta-carotene; and when red palm oil is used as a source of carotenes, the beta-carotene equivalent shall be taken as 53.5 per cent. of the total carotenes;
- (b) not less than 80 international units and not more than 100 international units of vitamin D.

SECOND SCHEDULE

Method for Determination of Vitamin A Content

Principle

The margarine is saponified, the unsaponifiable matter is extracted with diethyl ether and passed through a process of double column chromatography, which separates carotene from vitamin A. The optical density of the carotene solution is measured over the spectral range 440-450 $m\mu$. and the vitamin A fractions, after being identified by the Carr-Price reaction, are combined in light petroleum. The optical density of the solution is measured at the wavelength of maximum absorption, normally at 324 $m\mu$. All operations should be carried out under subdued lighting conditions.

Apparatus.

1. Chromatographic apparatus, (see sketch of assembly and of components). It consists of an upper and lower column. The lower column is fitted with a side arm which can be adjusted so that the eluate from the upper column can either by-pass, or pass through, the lower column. The assembly is fitted with three-way taps to enable air pressure to be applied either to the top column alone or to both columns.

2. Graduated tubes.—1 ml. (see sketch).

3. Spectrophotometer.—For measuring ultra-violet absorption of vitamin A solution.

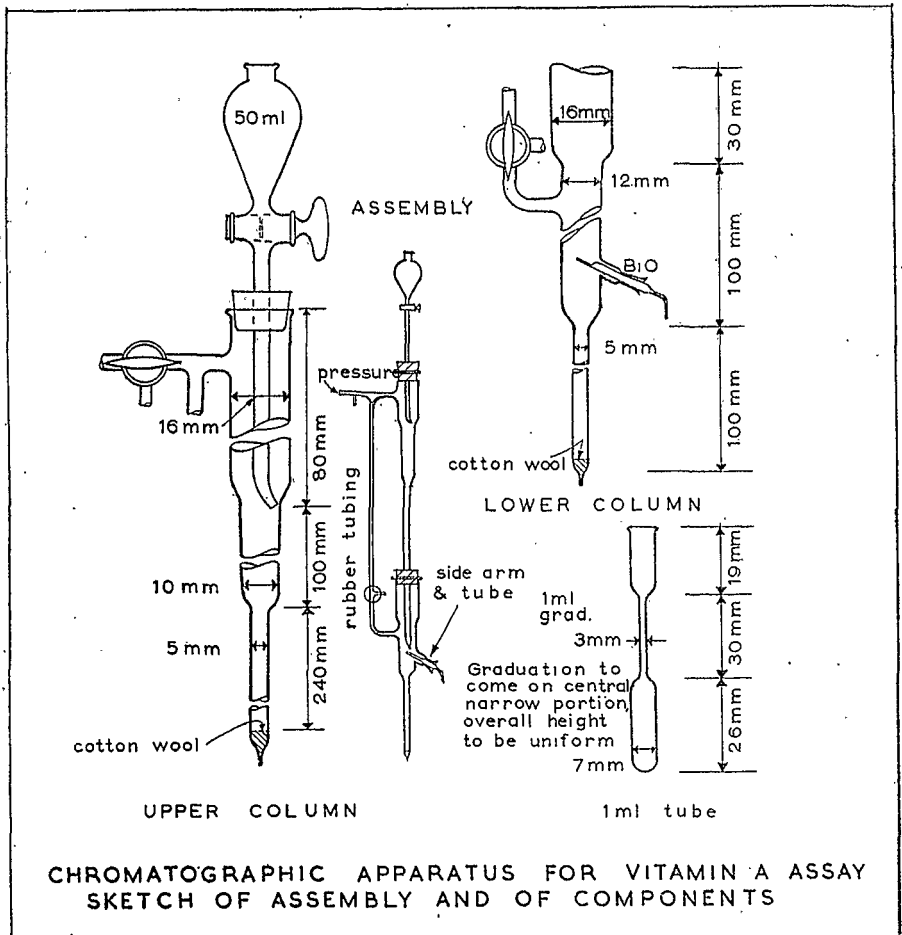
4. Quartz cuvettes.—Two; each of 1 cm.

5. Saponification flasks.—300 ml. resistance glass.

6. Separators.—500 ml. capacity.

7. Graduated flasks ("A" grade).—10 ml., 50 ml.

8. Pipette ("A" grade).—1 ml., the tip drawn out to a capillary to fit graduated tubes.



Reagents

All reagents must be of the quality required for quantitative chemical analysis.

1. Alumina (Type 1).—Prepare from alumina trihydrate.

Activate the fraction passing through a 150 mesh British Standard sieve by heating at 800°C. for 7 hours. After cooling, add distilled water, 2 g. per 98 g. activated alumina, and mix well. Keep in glass stoppered bottle.

2. Alumina (Type 2).—Place 10 g. of activated alumina (Type 1) in a 50 ml. round bottom flask, add 10 ml. of sodium hydroxide solution, and mix to a thin paste. Attach the flask to a vacuum pump and evacuate for 30 minutes at room temperature at 20 mm. pressure or lower. While still under this reduced pressure, place the flask in an oil bath at 135°C. for 30 minutes. Cool under reduced pressure and shake out the alumina-sodium hydroxide into a small mortar, ignoring any which adheres to flask. Add 1.2 ml. water slowly while stirring gently with a pestle. Immediately transfer the smooth powder to a small well stoppered bottle.

3. Antimony trichloride.—A saturated solution in chloroform (Carr-Price reagent).

4. Diethyl ether.—Freshly distilled over sodium hydroxide pellets.

5. Ethyl alcohol.—Absolute.
6. Light petroleum (40 to 60°C).—To obtain an optically pure fraction it may be necessary to distil and use only that portion boiling below 45°C.
7. Potassium hydroxide solution.—Dissolve 60 g. of potassium hydroxide pellets in distilled water and make up to 100 ml.
8. Sodium hydroxide solution.—Dissolve 10 g. of sodium hydroxide pellets in distilled water and make up to 100 ml.
9. Carbon dioxide or hydrogen supply.
10. Hydroquinone.

Saponification and Extraction of Unsaponifiable Matter

Weigh 10.0 g. margarine into a 300 ml. flat bottom flask, add 20 mg. hydroquinone, 40 ml. absolute alcohol, 10 ml. potassium hydroxide solution and boil under reflux for 20 minutes. Transfer quantitatively to a 500 ml. separator using two 40 ml. quantities of distilled water and extract once with 100 ml. and then three times with 50 ml. quantities of diethyl ether. Combine the ether extracts and wash with four successive portions, each of 50 ml. of distilled water. Transfer the ethereal solution in suitable aliquots quantitatively to a 300 ml. wide-mouth flat-bottom flask. All subsequent operations are conducted in an inert atmosphere e.g. carbon dioxide. Distil off the ether, add to the residue about 5 ml. absolute alcohol and reheat to remove the last traces of water. Make sure that no trace of alcohol remains, as otherwise the chromatogram will not develop properly. Dissolve the residue in 5 to 10 ml. of light petroleum.

Chromatography and Measurement of Optical Density

Place a small wad of cotton wool in the lower tip of each of the two columns of the chromatographic apparatus. Fill the upper column to the middle of the tube of 10 mm. diameter with light petroleum and pour in sufficient alumina (Type 1) to fill the tube of 5 mm. diameter. Prepare the lower column in a similar manner with alumina (Type 2), half filling the 5 mm. diameter section. Close the tip of the lower column with rubber tubing plugged with a short piece of glass rod. Connect the lower column to the upper column so that the outflow passes through the side arm.

Apply air pressure to the top column. When the excess of light petroleum has been forced through the column, release the pressure and transfer quantitatively the light petroleum solution of unsaponifiable matter to the column. Develop under pressure, first with 5 ml. of light petroleum, and then successively with 5 ml. quantities of light petroleum and diethyl ether mixtures containing respectively 4, 8, 12, 16, 20, 24 and 36 per cent. of diethyl ether. During these operations the level of liquid should not fall below the upper surface of the alumina.

Carotene

Carotene passes quickly down the column and is eluted before the 16 per cent. diethyl ether solvent mixture is introduced, the eluate being by-passed through the side tube into a receiver. Reduce the volume of the carotene solution, if necessary, by evaporation in an inert atmosphere, transfer to a 50 ml. graduated flask and make up to volume with light petroleum.

Measure the optical density of the solution in a 1 cm. quartz cuvette, against a light petroleum blank over the spectral range 440-450 m μ . at 2 m μ . intervals. From the optical density at λ max.* calculate the E (1 per cent., 1 cm.) value and multiply this by 358 to obtain the equivalent β -carotene potency in International Units per g. This value multiplied again by 0.8 will give the equivalent vitamin A potency of the carotenoids in International Units per g.

* λ max. means the wavelength of light at which the optical density reaches its maximum value.