

Journal of W. H. S.

XII A.

12

1925

20-
Amanik
N27.12A.12
196151

305

Price, As. 6.]

[Vol. 12A, Part XII, pp. 179 to 184.]

JOURNAL

OF THE

Indian Institute of Science.

CONTENTS.



OIL FROM THE SEEDS OF *SAPINDUS TRIFOLIATUS* (LINN.).

BY

D. R. Paranjpe and P. Ramaswami Ayyar.

DR. M. O. FORSTER, F.R.S., CHAIRMAN OF EDITORIAL BOARD.

JOURNAL
OF THE
INDIAN INSTITUTE OF SCIENCE

Parts I—XXI.	VOLUME I.—(1914-18).
Parts I—XV.	VOLUME II.—(1918-19).
Parts I—IX.	VOLUME III.—(1920).
Parts I—XVI.	VOLUME IV.—(1921).
Parts I—XIV.	VOLUME V.—(1922).
Parts I—XIII.	VOLUME VI.—(1923).
Parts I—XV.	VOLUME VII.—(1924).
Parts I—XVI.	VOLUME VIII (A)—(1925).
Parts I—II.	VOLUME VIII (B)—1925.
Parts I—X.	VOLUME IX (A)—1926.
Parts I—V.	VOLUME IX (B)—1926.
Parts I—X.	VOLUME X (A)—1927.
	VOLUME X (B)—1927.

- Part I.—Tests on Suspension Insulators after Ten Years' Service. By A. S. Venkateswaran and U. Ganguly. Price, Re. 1.
- Part II.—The Measurement of Voltage Gradient on a String of Suspension Insulators. By G. Yoganandam and R. K. Sen. Price, Re. 1.
- Part III.—Intensity Variations of Madras (Fort) Radio Station. By K. Sreenivasan. Price, As. 8.
- Part IV.—Suspension Insulator Testing. By G. Yoganandam. Price, As. 8.

VOLUME XI (A)—1928.

- Part I.—I. Studies on Invertase, Part I. Preparation and Purification of the Enzyme. By B. N. Sastri and Roland V. Norris. II. Note on a Simple Method for concentrating Enzyme Solutions. By B. N. Sastri. Price, Re. 1.
- Part II.—The Bleaching of Lac. By M. Venugopalan. Price, As. 8.
- Part III.—Contributions to the Study of Spike-Disease of Sandal (*Santalum Album*, Linn.). Part I. Diastatic Activity of the Leaves. By M. Sreenivasaya and B. N. Sastri. Price, As. 12.
- Part IV.—A Micro-Method for the Determination of Enzyme Activity. By B. N. Sastri and M. Sreenivasaya. Price, As. 12.
- Part V.—The Dielectric Constants of Ammonia, Phosphine and Arsine. By H. E. Watson. Price, Re. 1.

Price, As. 6.]

[Vol. 12A, Part XII, pp. 179 to 184.

JOURNAL

OF THE

Indian Institute of Science.

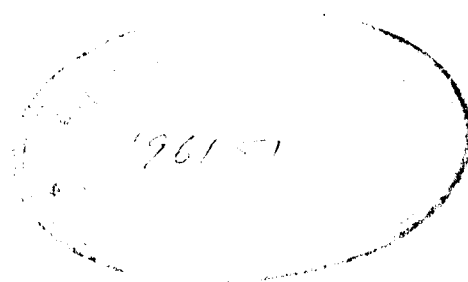
CONTENTS

OIL FROM SEEDS OF *SAPINDUS TRIFOLIATUS* (LINN.).

BY

D. R. Paranjpe and P. Ramaswami Ayyar.

DR. M. O. FORSTER, F.R.S., CHAIRMAN OF EDITORIAL BOARD.



OIL FROM THE SEEDS OF *SAPINDUS TRIFOLIATUS* (LINN.).

By D. R. Paranjpe and P. Ramaswami Ayyar.

Sapindus trifolius (N. O. *Sapindaceae*), the soapnut tree, is found in the villages of Southern India and Ceylon, and cultivated in Bengal for the fruit. It has long been used by the Hindus as a detergent, and is preferred to soap for certain purposes (*Pharm. Indica*, vol. 1, 368). The vernacular names of the tree are: Hindi and Mahratti, *Ritha*; Gujrathi, *Aritha*; Telugu, *Kunkudukayalu*; Kanarese, *Artela*; Tamil, *Poovandi*. While the fruit is valued principally for the high saponine content of its pericarp, it appeared of interest to examine also the oil contained in the seeds.

The fruits are fleshy, two or three united, and each is the size of a cherry. The seeds resemble the fruit in shape, are black, subspheroid, smooth, and have the size of a large pea. The fruits used in the present investigation were obtained locally and were separated into pericarp (65.7 per cent.) and seeds (34.3 per cent.). The latter were further divided into one part of yellowish green kernels and two parts of outer shell. The finely powdered seed-kernels on extraction with petroleum ether gave 44.7 per cent. of a dark yellow oil which deposits a small quantity of stearin on standing.

An examination of the oil in the usual manner has revealed the remarkable fact that in addition to the glycerides of palmitic, stearic, oleic and lignoceric acids, the oil contains a high proportion (22 per cent.) of *n*-eicosanic acid. The oil may be classed as a non-drying oil from the low iodine value (58.5).

EXPERIMENTAL.

The analytical constants of the oil given in Table I were determined in the usual way and compare well with those of A. K. Menon (*J. Soc. Chem. Ind.*, 1910, 1431), whose results appear in the second column.

TABLE I.

Sp. gr. $\frac{100^{\circ}}{100^{\circ}}$	...	...	0.8540	0.8542
$n_D^{25^{\circ}}$	...	...	1.4764	1.4748
Saponification value	...	...	194.1	191.8
Iodine value	...	...	58.5	58.6
Reichert-Meisel value	...	...	1.5	1.6
Acetyl value	...	...	0.0	0.0
Unsaponifiable matter	...	...	1.2 per cent.	1.1 per cent.
Hehner value	...	...	93.5	93.9

THE MIXED FATTY ACIDS.

For the preparation of the fatty acids, the oil was saponified with an alcoholic solution of sodium hydroxide, the dried soap extracted with ether to remove unsaponifiable matter, and the fatty acids regenerated from the soap by boiling with 1 : 1 diluted hydrochloric acid. The acids obtained after taking up in ether and removing the solvent had the constants given in Table II, the second column showing those of A. K. Menon.

TABLE II.

Saponification value	...	...	195.4	188.6
Iodine value	...	...	61.3	57.0
Equivalent weight	...	...	287.0	297.3
Titre	...	...	47°	54.4° (m. p.)

The mixed acids were separated into the saturated and unsaturated acids by the method of Twitchell (*J. Ind. Eng. Chem.*, 1921, 13, 806), the process being repeated twice. The separated acids were analysed with results summarised in Table III.

TABLE III.

	Saturated acids (38.5 per cent.)	Unsaturated acids (61.5 per cent.)
Iodine value	...	...
Equivalent weight	...	...
n_D^{60}	...	...
	1.1	86.2
	292.5	283.0
	1.4400	1.4450

IDENTIFICATION OF THE UNSATURATED ACIDS.

The additive bromo-compounds were prepared by the method of Jamieson and Baughman (*J. Amer. Chem. Soc.*, 1920, 42, 2398). No hexabromides separated at zero nor was any crystalline tetrabromide formed from petroleum solution. The solvent was therefore distilled and bromine estimated in the residual oily bromide (Br, 36.2. Calc. for dibromide of oleic acid, Br, 36.2 per cent.). The unsaturated acid was therefore assumed to be oleic acid.

IDENTIFICATION OF THE SATURATED ACIDS.

The saturated acids recovered from the insoluble lead salts were converted into the methyl esters by boiling with ten times their volume of anhydrous methyl alcohol containing 4 per cent. of hydrogen chloride. The purified esters were twice fractionated under a pressure of 6 mm. and twelve fractions were obtained. The main results of the fractionation are summarised in Table IV, and the detailed examination of each fraction is given below.

TABLE IV.

Fraction No.	B. p. 6 mm.	Yield in grams	Titre of ester	M. W. of ester	M. p. of acid
I	below 190°	9.7	25.4°	285.5	54.5°
II	191-196	11.7	28.4	289.1	53-54
III	197-198	12.0	31.2	297.0	55-56
IV	199-206	14.5	33.6	305.5	57-58
V	206-207	20.1	36.8	315.2	67
VI	207-208	17.1	38.5	320.2	70
VII	208-209	10.4	39.8	322.3	...
VIII	209-210	10.0	41.2	325.3	73
IX	211-217	11.8	42.0	329.0	74
X	218-225	13.4	43.0	336.7	71
XI	226-235	5.2	44.6	335.2	69-70
XII	...	1.1	...	...	80.5

Fraction I was a liquid. The acids obtained from it were crystallised from 70 per cent. alcohol, when an acid melting at 73° separated; and from the mother liquor, a solid acid melting at 53-54° was obtained having an equivalent weight of 268. Assuming that the acid melting at 73° was *n*-eicosanic acid which was identified in the later fractions and that the yield was approximately quantitative, it was possible to calculate from the molecular weight that the fraction was a ternary mixture of palmitic ester (5.1 g.), stearic ester (3.8 g.) and *n*-eicosanic ester (0.8 g.).

Fraction II.—The ester (6.2 g.) was saponified with alcoholic potash, neutralised with dilute hydrochloric acid so as to make about an 80 per cent. alcohol and allowed to stand overnight. This

precipitated the acid of high molecular weight as potassium soap, from which the acid was regenerated and crystallised from acetone; 0.76 g. was obtained, and having m.p. 73–74° with equivalent, 310 was proved to be different from Kahlbaum's arachidic acid (mixed melting point, 63–65°), but identical with a synthetic specimen of *n*-eicosanic acid (mixed m. p., 73–75°). Acid (m. p. 50° with equivalent, 270) from the soluble potassium soap was resolved by fractional precipitation with lead acetate in 90 per cent. alcoholic solution, into (a) an acid (equivalent, 282; m. p. 69–70°, unchanged on mixing with stearic acid), (b) an acid m. p. 53–54° and (c) an acid m. p. 59–60° (unchanged on mixing with palmitic acid) having an equivalent weight of 251 (palmitic acid, 256). Assuming that the acid melting at 73–74°, is a mixture of *n*-eicosanic and stearic acids, and that all the former acid was precipitated, the composition of this fraction is palmitic ester (5.1 g.), stearic ester (5.1 g.) and *n*-eicosanic ester (1.5 g.).

Fraction III.—The ester (7.1 g.) was saponified with potash, and resolved into five fractions by fractional precipitation with an alcoholic solution of magnesium acetate (10 grams in 100 c.c. of 95 per cent. alcohol). The following acids were thus obtained:—

(a) 2.4 g., m.p. 72–73° (equivalent, 308.5) after crystallisation from acetone melted at 73–75° (unchanged on mixing with *n*-eicosanic acid, but lowered by 6° on mixing with Kahlbaum's arachidic acid).

(b) 0.2 g., m.p. 66–69° (with stearic acid, at 65–66°).

(c) 0.5 g., m.p. 65–68° (with stearic acid at 65–66°).

(d) 0.7 g., m.p. 60–62° (with palmitic acid at 55–57° and with stearic acid at 60–67°).

(e) 2.5 g., m.p. 54–55° with an equivalent of 270; on crystallising this from alcohol, ether and acetone consecutively, two crops of crystals were obtained melting at 58–59° and 59–61°, respectively. By taking mixed melting points of these with specimens of pure palmitic, stearic and synthetic heptadecoic (margaric) acids they were respectively found to be a palmitic-stearic-mixture, and pure palmitic acid. Fraction III therefore consists of palmitic ester (3.3 g.), stearic ester (4.9 g.) and *n*-eicosanic ester (3.8 g.).

Fraction IV.—This was further separated into four portions by refractionation under reduced pressure as shown in Table V. From the molecular weights and melting points of the acids obtained from three of the refractionated portions the mixture should consist of palmitic ester (4.9 g.), stearic (8.4 g.) and *n*-eicosanic ester (1.2 g.).

TABLE V.

Fraction No.	Yield in grams	Titre of ester	M. W. of ester	M. p. of acid
1	2.9	28.2°	278	56°
2	4.0	31.5°	...	...
3	3.9	36.5°	298	67°
4	1.1	40.4°	...	73°

Fraction V.—By careful refractionation and hydrolysis this gave 1.5 g. of pure palmitic acid, nearly 1 g. of stearic acid, and a residual mixture of stearic and *n*-eicosanic acids from which the latter was obtained pure (m.p. 73–75°) by crystallisation from acetone. Its composition is therefore palmitic ester (1.6 g.), stearic ester (2.7 g.) and *n*-eicosanic ester (15.8 g.).

Fractions VI–VIII constitute a mixture of stearic ester (5.2 g.), and *n*-eicosanic ester (32.3 g.).

Fraction IX is principally *n*-eicosanic ester, contaminated with a small proportion of lignoceric ester.

Fractions X and XI are undoubtedly mixtures of *n*-eicosanic and lignoceric esters, because from the residual fraction XII, an acid was prepared which after crystallisation from 98 per cent. alcohol melted at 80–80.5° (unchanged on mixing with pure lignoceric acid, but lowered by 3° on mixing with behenic acid).

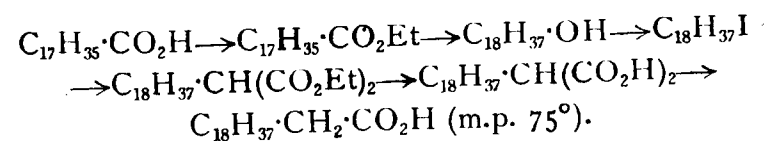
Detailed composition of the saturated esters (twelve fractions), summarised in Table VI (grams), shows totals of palmitic ester 14.6 per cent., stearic 21.9 per cent., *n*-eicosanic 56.8 per cent. and lignoceric ester 6.6 per cent.

TABLE VI.

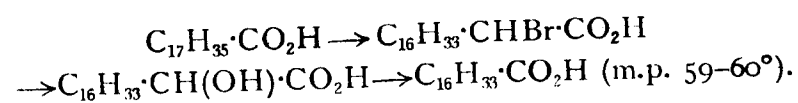
Fraction No.	Palmitic	Stearic	<i>n</i> -Eicosanic	Lignoceric
I	5.1	3.8	0.8	...
II	5.1	5.1	1.5	...
III	3.3	4.9	3.8	...
IV	4.9	8.4	1.2	...
V	1.6	2.7	15.8	...
VI	...	3.5	13.6	...
VII	...	1.4	9.0	...
VIII	...	0.3	9.7	...
IX	...	...	10.5	1.3
X	...	...	8.4	5.0
XI	...	...	3.5	1.7
XII	...	...	...	1.1
Total in grams ...	20.0	30.1	77.8	9.1

PREPARATION OF PURE *n*-EICOSANIC AND *n*-HEPTADECOIC ACIDS.

A specimen of the first named acid was prepared from stearic acid by the following series of reactions, adopting the methods given by Adam and Dyer (*J.C.S.*, 1925, 127, 70) and Levene and Taylor (*J. Biol. Chem.*, 1924, 59, 916):—



n-Heptadecoic acid was prepared also from stearic acid by the usual method of degrading higher fatty acids as outlined below (Levene and West, *J. Biol. Chem.*, 1913, 16, 477):—



The unsaponifiable matter yields about 30 per cent. of a sterol melting at 115–120°, which is most probably a mixture, as its alcoholic solution deposits two distinct types of crystals (short prismatic needles, and hexagonal plates) as seen under the microscope.

SUMMARY.

The seed-kernels of *Sapindus trifolius* fruits contain 44·7 per cent. of a non-drying fatty oil, the analytical constants of which have been determined.

The oil is composed of the glycerides of palmitic (5·6 per cent.), stearic (8·5 per cent.), *n*-eicosanic (21·9 per cent.), lignoceric (2·5 per cent.) and oleic (61·5 per cent.) acids.

The oil is a rich source of *n*-eicosanic acid.

Department of Organic Chemistry,
Indian Institute of Science,
Bangalore.

[Accepted, 9-8-29.]

PRINTED AND PUBLISHED BY GEORGE KENNETH AT THE DIOCESAN PRESS, POST BOX 455, MADRAS—1928 C8986

- Part VI.—The Fermentation of Toddy and an Account of the Micro-Organisms producing it. By M. Damodaran. Price, As. 12.
- Part VII.—I. Estimation of Iodine in Soils. By Roland V. Norris and D. A. Rama Rao. II. Carbohydrate Changes during the Ripening of Plantains. By S. Ranganathan (Junior). Price, As. 12.
- Part VIII.—I. The Oxidation of Sulphur in Suspensions of Activated Sludge and its Influence on the Solubilisation of Mineral Phosphates. By C. V. Ramaswami Ayyar, T. S. S. Perumal and Roland V. Norris. II. Studies in the Proteins of Indian Foodstuffs. Part I. The Proteins of Ragi (*Eleusine coracana*): Eleusina, the Alcohol-Soluble Protein. By Nuggehalli Narayana and Roland V. Norris. Price, Re. 1.
- Part IX.—I. Contributions to the Study of Spike Disease of Sandal (*Santalum album*, Linn.). Part II. Analysis of Leaves from Healthy and Spiked Trees. By A. V. Varadaraja Iyengar. II. Contributions to the Study of Spike Disease of Sandal (*Santalum album*, Linn.). Part III. Physico-Chemical Study of the Leaf-Sap. By A. V. Varadaraja Iyengar. Price, Re. 1.
- Part X.—I. Studies on Soil Protozoa. Part I. Protozoan Fauna of Some Mysore Soils. By H. S. Madhava Rao. II. Studies on Soil Protozoa. Part II. The Function of Mitochondria in Some Soil Protozoa. By H. S. Madhava Rao. Price, Re. 1.
- Part XI.—I. Studies in Enzyme Action. Part I. Amylase from Chulam (*Sorghum vulgare*). By V. N. Patwardhan and Roland V. Norris. II. Studies in Enzyme Action. Part II. The Nature of Amylase. By D. Narayana-murti and Roland V. Norris. Price, Rs. 1-8.
- Part XII.—A Bio-Chemical Study of some Soil Fungi with special reference to Ammonia Production. By A. K. Thakur and Roland V. Norris. Price, Re. 1.
- Part XIII.—The Atomic Weight of Antimony from Different Sources. By K. R. Krishnaswami. Price, Re. 1.
- Part XIV.—Oil from the Seeds of *Adenanthera Pavonina*. A Source of Lignoceric Acid. By S. M. Mudbidri, P. Ramaswami Ayyar and H. E. Watson. Price, As. 8.
- Part XV.—I. Constituents of some Indian Essential Oils. Part XXII. The Essential Oil from the Flower Heads of *Cymbopogon Coloratus*, Stapf. By P. P. Pillay, B. Sanjiva Rao and J. L. Simonsen. Part XXIII. The Essential Oil from the Fruits of *Piper Cubeba*, Linn. By B. Sanjiva Rao, V. P. Shintre and J. L. Simonsen. Part XXIV. The Essential Oil from the Rhizomes of *Curcuma Zedoaria*, Roscoe. By B. Sanjiva Rao, V. P. Shintre and J. L. Simonsen. II. The Constituents of Indian Turpentine from *Pinus Longifolia*, Roxb. Part IV. By P. Parameswaran Pillay and J. L. Simonsen. Price, Rs. 1-4.
- Part XVI.—I. α -isoPropylglutaconic Acid. By K. V. Hariharan, K. N. Menon and J. L. Simonsen. II. Derivatives of Methyl 2: 2-Dimethylcyclo-Pentan-3-one-1-Carboxylate. By C. S. Gibson, K. V. Hariharan and J. L. Simonsen. Price, As. 12.
- Part XVII.—I. Thiophthalic Acids. Part I. By Gopāl Chandra Chakravarti. II. Organic Cyclic Polysulphides: Condensation of Ethylene Mercaptan with Di- and Trichloroacetic Acids. By Gopāl Chandra Chakravarti and Jogendra Mohan Saha. Price, As. 8.
- Part XVIII.—Bis-Semidine Inversion in Aromatic Dihydrazo-Compounds. By Praphulla Chandra Guha and Hirendra Kumar Banerjee. Price, As. 8.

- Part XIX.—I. Contributions to the Study of Spike-Disease of Sandal (*Santalum Album*, Linn.). Part IV. Chemical Composition of Healthy and Spiked Sandal Stems. By D. A. Rama Rao and M. Sreenivasaya. II. Contributions to the Study of Spike Disease of Sandal (*Santalum Album*, Linn.). Part V. Transmission of Spike by Budding. By M. Sreenivasaya and G. Gopalaswami Naidu. Price, Re. 1.

VOLUME XII (A)—1929.

- Part I.—Phototropic Compounds of Mercury. By Bh. S. V. Raghava Rao and H. E. Watson. Price, Re. 1.
- Part II.—Photoelectric Emission from Phototropic Mercury Compounds. By Bh. S. V. Raghava Rao and H. E. Watson. Price, As. 12.
- Part III.—Attempts to Synthesise *Ortho*-Thiolphenylhydrazine. By Praphulla Chandra Guha and Tejendra Nath Ghosh. Price, As. 6.
- Part IV.—I. Characterisation of very small quantities of Proteins by Van Slyke's Method. By Nuggihalli Narayana and Mothnahalli Sreenivasaya. II. The Determination of Pyruvic Acid. By Basettihalli Hanumantha Rao Krishna and Mothnahalli Sreenivasaya. Price, As. 12.
- Part V.—I. Studies on Soil *Actinomyces*. Part I. Introduction. By V. Subrahmanyam and Roland V. Norris. II. Studies on Soil *Actinomyces*. Part II. Their Mode of Occurrence in the Soil. By V. Subrahmanyam. Price, Re. 1.
- Part VI.—I. A Study of the Symbiotic Fungus from the Mysore Lac Insect. By M. Sreenivasaya and S. Mahdihassan. II. The Golgi Apparatus of Free-Living Protozoa. By H. S. Madhava Rao. Price, As. 12.
- Part VII.—I. Formation of Heterocyclic Compounds from Diethylxanthic Formic Ester. By Praphulla Chandra Guha and Devendra Nath Datta. II. Ring Closure of Hydrazo-Monothiodicarbonamides with Acetic Anhydride: Formation of Iminothiobiazolones and Iminothioltriazoles. By Praphulla Chandra Guha and Tarani Kanta Chakraborty. Price, Rs. 1-4.
- Part VIII.—I. Studies in Enzyme Action. Part III. Amylase from *Camba* (*Pennisetum typhoides*). By D. Narayanamurti, C. V. Ramaswami Ayyar and Roland V. Norris. II. Studies in Enzyme Action. Part IV. Tyrosinase I. By D. Narayanamurti and C. V. Ramaswami Ayyar. Price, Rs. 1-8.
- Part IX.—The Activated Sludge Process of Sewage Treatment. Report on the Working of the Plant at the Indian Institute of Science, Bangalore. By N. Swaminathan. Price, Rs. 1-4.
- Part X.—Contributions to the Study of Spike-Disease of Sandal (*Santalum Album*, Linn.). Part VI. Nitrogen Metabolism in Healthy and Spiked Sandal Leaves. By N. Narasimhamurthy and M. Sreenivasaya. Price, As. 12.
- Part XI.—Lengthened *Ortho*-Di-Derivatives of Benzene and their Ring-Closure: Formation of Polymembered Heterocyclic Compounds from Substituted Phenylene-Dicarbamides. By Tejendra Nath Ghosh and Praphulla Chandra Guha. Price, As. 12.
- Part XII.—Oil from the Seeds of *Sapindus Trifoliat* (Linn.). By D. R. Paranjpe and P. Ramaswami Ayyar. Price, As. 6.