THE ANALYST.

MAUMENÉ'S TEST FOR OILS.

By H. DROOP RICHMOND.

AFTER an extended trial of the various methods and forms of apparatus for performing the Maumené test which have been proposed from time to time, and after a study of the various reactions, so far as can be ascertained, which take place in that process, I have been led to devise the following method of procedure. By means of this modification, I feel assured that results more uniform and exact than those hitherto attainable can be secured:

Calorimeter.—A small, deep beaker is fitted inside a slightly larger one by means of a ring of cork; the outer beaker is placed in a tin cup padded with cotton-wool. The heat-capacity of this is estimated by placing 10 grammes of water inside, noting the temperature, and pouring in about 25 grammes of warm water (of known temperature), and noting the resulting temperature. The cooling in this calorimeter during a Maumené test does not amount to more than 1 per cent., and may be neglected.

Influence of Strength of Acid.—Between the limits, 92 per cent. to 100 per cent. H_2SO_4 , the rise of temperature is directly proportional to the strength of acid (difference not more than 1 per cent.).

Heat-Capacity of Mixture of Oil and Acid.—25 grammes of oil and 5 c.c. of sulphuric acid, when mixed, have a heat-capacity about equal to that of 20 grammes of water.

The total heat evolved per mean molecule may be calculated; this I call the "Relative Molecular Maumené" figure:

Let $x = \text{percentage of } H_2SO_4$ in acid (100 per cent. H_2SO_4 assumed as standard strength);

h = heat-capacity of calorimeter in grammes of water;

R = observed rise (25 grammes oil, 5 c.c. acid);

K = potash absorbed (19.5 per cent. KOH is assumed as a standard of comparison);

then R.M.M. = R ×
$$\frac{21.5}{x-78.5}$$
 × $\frac{20+h}{20}$ × $\frac{19.5}{K}$.

This action of the acid on the oil is probably in two stages—(i.) hydrolysis, (ii.) action on the unsaturated bonds.

I believe that a definite amount of the heat evolved is due to hydrolysis, and the remainder to the second stage. My experiments indicate that the R.M.M. of hydrolysis lies between 7° and 18°, and is probably about 12°, and that glycerides containing acid radicles with two unsaturated bonds evolve more than twice the amount of heat than those with one unsaturated bond.

Contribution to Butter Analysis—The Abbé-Zeiss Refractometer. J. Delaite. (Bull. de l'Assoc. belge des Chimistes, 1894, No. 5, pp. 145-152.)—According to Zeiss, the refractive index may be taken at any temperature, and the reading subsequently calculated to a standard temperature, e.g., 25°. But the author proves that this does not give concordant results with fats. Thus, the indices obtained from one butter, from readings taken at different temperatures and reduced to 25° C., were:

Temp. of Observation.	Refractive Index, calculated to 25° C.
45°	48.8
40°	49.3
35°	49.5
30°	49.65
25°	49.7

This inverse ratio of the refractive index to the temperature only occurs markedly in the case of pure normal butters. Abnormal and adulterated butters and margarines, though not giving constant results, show no order in their variation. Two margarines, for instance, gave the following results:

		Ter	nperatu:	res.			$_{ m In}$	dices ca	lculated	to 25° (J.	
	45°	40°	35°	30°	25°		45°	40°	35°	30°	25°	
Ref. Indices (1)					59.2				-, -	58·1 57·8		_
,, (2)	41	49.0	52.2	99	99.7	ļ	99.3	91.9	91.8	91.0	99.Z	

It is necessary, therefore, to select a definite temperature for taking the reading, and it is suggested that this should be 40° C. for fats and 25° for oils. By means of a current of warm water the temperature is brought to 45° or 50°, and is then allowed to fall very slowly to 40°. The error of reading introduced by the temperature of the whole instrument not being the same as that of the water, is thus avoided.

As to the value of the process, the author's opinion is that it cannot be relied upon to give alone even a preliminary decision as to the genuineness or otherwise of a butter. For although the average butter does not give a refractive index higher

than 44 at 40°, yet samples undoubtedly genuine often give higher figures. other hand, abnormal and adulterated butters have been known to give figures as low Moreover, rancidity appears to affect the result. A butter known to be adulterated was left for more than a year, and, when strongly rancid, gave a refractive index of 48.25 at 30°, while that of an average pure butter at that tempera-Hence, judged by the refractive index alone, this sample would have been declared genuine. The addition of cocoanut oil or palm oil to a butter also The refractive index of the former varies from 33.2 to 35.2 at 40°, lowers the index. that of the latter from 35.2 to 45.3. A mixture of a pure butter with refractive index of 44 at 40°, with 20 per cent. of cocoanut oil, showed an index of 42.2. A similar result was also obtained by adding horse-fat to butter. C. A. M.

THE ANALYST.

An Examination of the Twitchell Method for the Determination of Rosin in Soap. T. Evans and I. E. Black. (Amer. Chem. Journ., 1895, xvii., I., pp. 59-67.) —Twitchell's process,* although the most satisfactory known, has been proved to be far from exact, especially when the gravimetric method is employed, in which, as has been shown by Lewkowitch, a loss of from 10 to 15 per cent. is frequent. With the object of rendering it more accurate, the authors have made experiments on the same lines as Twitchell, preparing test-samples containing known quantities of rosin and fatty acids.

In the volumetric method, too high results were obtained when the combining weight of the rosin was taken as 346, the number given by Twitchell. was attributed to incomplete etherification, but repeated titration proved that the combining weight of the sample was 328.9—a figure lower than any of those recorded by Twitchell or Lewkowitch. Taking this number as the factor, the results were much more accurate, as is shown in the following instances selected from a long table:

Percentage of Rosin.

Theory.	Combining Weight 346.	Difference.	Combining Weight 328.9.	Difference.
•	Found.		Found.	
40.13	42.19	+2.06	40.10	-0.03
40.13	41.74	+1.61	39.67	-0.46
40.13	42:16	+2.03	40:06	-0.07

By leaving the alcohol saturated with HCl, and before diluting with water, all night on the water-bath, the results could be considerably lowered, and still more so by adding H₂SO₄ before this treatment. The substitution of methyl alcohol for ethyl alcohol gave poorer results, and if left all night before evaporating the alcohol The results were also too low if the water surrounding the caused considerable loss. flask during etherification became too warm, or if the current of HCl were too rapid.

By using ethyl alcohol, and boiling out soon after etherization, concordant results were obtained, approximating closely to the theoretical.

Attempts were made to etherize rosin alone, but in each case the results were too low.

^{*} See Analyst, xvi., p. 169.

[†] Journ. Soc. Chem. Ind., xx., 1893.

Weight of Rosin.	Rosin calculated from Titration.	Difference.	Percentage loss.
$2.0968 \mathrm{\ grms}$.	2.054	0.0428	2.00
2.4723,	2.45	0.0223	0.90
2.035	2.016	0.019	0.93
2.03	2.016	0.014	0.69
2.115	2.091	0.024	1.14

With methyl alcohol the loss was still greater, being 6 or 7 per cent., a result probably due to decomposition.

As is shown in the first table, when the combining weight of a rosin is known the volumetric method is sufficiently accurate, but the combining weight varies greatly in different grades of rosins, ranging from 328.9 to 355.9. Accordingly, the authors made the attempt to separate the rosin in a given sample, determine its combining weight, and to use this number as the factor in the volumetric determination. A sample of resin (combining weight 328.9) was subjected to the gravimetric process, and the rosin acids dried at 105°, and titrated. The combining weights obtained were (1) 328.6, (2) 325.9.

Four similar experiments with a rosin with combining weight 351.4 gave the combining weights, 329, 323.9, 331.1, 329.4.

The explanation of these discrepancies was looked for in the loss which occurs in the gravimetric method. This loss was found to be in the petroleum ether layer from which the saponifiable rosin had been removed by caustic potash. A sample of rosin (combining weight 351·4) was treated as in the gravimetric method, and the residual petroleum ether, after removal of the rosin, evaporated to dryness, when a gum-like mass with a turpentine smell was left, corresponding to 9·71 per cent. of the original rosin. This residue was not saponifiable with alcoholic potash, and was therefore not an ether.

A mixture of fatty acids and rosin gave a loss of 5 per cent., which corresponded exactly with the percentage of unsaponifiable matter in the sample of rosin used.

Four other samples of rosin tested for unsaponifiable matter gave the following results:

Grade.	Unsapon. Matter per Cent.	Comb. Weight Original Rosin.	Calculated Combining Weight Saponifiable Rosin.
WW	7.34	339.9	315.0
\mathbf{WG}	5.00	328.9	312.4
N	9.00	351.4	319.7
N	8.21	347.5	319.0
${f M}$	7.61	339.8	314.0
		${f Mean} \qquad \dots$	316.0

The authors are investigating the subject further, and perfecting a method for the separation of the unsaponifiable rosin from the hydrocarbon oils used as soap-fillers, etc.

C. A. M.

The Determination of Sulphur in Volatile Organic Compounds. C. F. Maybury. (Amer. Chem. Journ., 1894, xvi. 544; through Chem. Zeit.)—The author has made a critical examination of various methods of determining sulphur, especially

in petroleum products, and has in particular examined the method of Carius and the process consisting in oxidizing the material to be analysed in a combustion tube in a stream of air or oxygen. The tube, which must be of refractory glass, is somewhat contracted in the middle; the stream of air or oxygen is let in through a piece of narrow tubing passing up to the point of contraction. The stopper which carries this narrow tube has a second perforation carrying a tube conveying carbonic acid, in order to drive the material to be analysed towards the forward end of the combustion tube. The products of combustion pass into a large U-tube containing 50 c.c. of caustic potash, of which each c.c. is equivalent to 1—5 milligrammes of sulphur, according to the content of the substance to be analysed. The liquid can be titrated in the U-tube itself, methyl orange being used as an indicator.

B. B.

The Determination of Free Sulphuric Acid in Leather. P. F. Jean. (Rev. Chim. Analyt., 1895, iii. 13; through Chem. Zeit.)—The process preferred by the author consists in extracting the dried and powdered leather in a Soxhlet apparatus with absolute alcohol containing a little caustic potash to fix the sulphuric acid. After thorough extraction, the alcohol is distilled off and the sulphuric acid determined in the residue as barium sulphate in the usual manner.

B. B.

Thioacetic Acid as a Substitute for Sulphuretted Hydrogen in Analysis. R. Schiff and N. Tarugi. (Gazz. Chim. ital., 1894, xxiv., vol. ii., 551; through Chem. Zeit.)—When a hydrochloric acid solution of the metals of the second group is treated with a slightly ammoniacal solution of ammonium thioacetate (30 per cent. strength) in the proportion of 1.5—2 c.c. for 1 gramme of the metals to be separated, and the mixture is warmed to 80-90° C., all the metals are precipitated as sulphides, although only a slight smell of sulphuretted hydrogen is perceptible. After cooling and filtering, the solution is found to be free from metals of the second group, and the analysis is completed in the ordinary manner.

Ammonium thioacetate is decomposed in the presence of hydrochloric acid with the formation of ammonium chloride, acetic acid, and sulphuretted hydrogen. Tin, bismuth, copper, platinum, and gold are only partly precipitated in the cold, but completely on keeping. Lead and mercury are precipitated in the cold as red chlor-sulphides, which on heating are converted into the ordinary black sulphides. With arseniates the reagent gives a slight white turbidity in the cold, but on heating the solution sudden and complete precipitation takes place. Cadmium sulphide dissolves in the warm hydrochloric acid, but again separates on cooling. Thioacetic acid reduces ferric salts and chromates.

The reagent is prepared by adding 10 per cent. ammonia solution in slight excess to thioacetic acid, and then diluting the mixture to treble the volume of the thioacetic acid used. The resulting liquid is yellow, and smells slightly of ammonium sulphide; it becomes turbid on heating, and needs to be freshly prepared every eight or ten days. The advantages which its use presents consist in avoidance of the smell of sulphuretted hydrogen, and of the need for a Kipp or similar apparatus; the rapidity and completeness of precipitation are also reasons for preferring this reagent.

B. B.

The Phospho-molybdic Acid Test as applied to Lard Analysis. George F. Tennille. (Jour. Amer. Chem. Soc., 1895, xvii. 1, pp. 33-41.)—The conclusion arrived at by Samelson (Analyst, xix., p. 251), that the phospho-molybdate test is of little analytical value, is here confirmed. It will certainly detect the addition of a large quantity of cotton-seed-oil; but if only a slight green coloration be obtained, adulteration cannot be considered as proved.

The author's experiments were made with the three grades of lard constituting the bulk of that sold to the New York refiners—viz.:

No. 1 Lard.—Steam-rendered, and consisting of gut fat, with occasionally a little trimmings, but no leaf.

Prime City Lard.—Rendered either by steam or in open kettles, and consisting of trimmings, head, foot, and back fat, and sometimes a little leaf.

Prime Western Lard. — Steam-rendered; from gut fat, with occasionally trimmings.

All the samples of No. 1 lard examined gave the green coloration, and this was also the case with a sample rendered in the laboratory from No. 1 stock. This was No. 10 in the table.

Nearly all the City lards gave negative results. Of fifty-seven undoubtedly pure samples, fifty gave no coloration, and seven gave a slight green tinge.

The Western steam lard often gave a slight green colour, which, however, was not so pronounced as that obtained with No. 1 lard.

The following table gives the particulars of the tests applied to ten samples of No. 1 lard:

					Cr	ystals fro	m	
						Ether		
Hübl Figure.	Dalican's Titre °C.	Rise of Temperature with H ₂ SO ₄ °C.	Becchi's Silver Test.	Milliau's Silver Test.	Phospho- Molybdic Acid Test.	Micro-	Free Fatty Acid, per Cent.	Spec. Grav. 98°C:15°C.
1. 54.5	41.4	30.5	Negative.	Slight discoloration.	Decided green. Blue if alkaline.	Lard only.		
2. 55.6	41.4	31.8	,,	,,	,,	,,		
3. 58.6	40.2	34.8	,,	,,	,,	,,	0.73	0.8595
4. 53.7	41.4	30.0	,,	,,	,,	,,		
5. 55.4	41.4	31.5	Slight dis- coloration.	,,	,,	,,		
6. 56.1	41.4	31.7	Negative.	,,	,,	,,		
7. 47.7	42.4	25.5	Slight dis- coloration.	"	: •	,,		
8. 51.7	41.8	31.5	Negative.	,,	,,	Indication of beef.	n	
9. 52.0	41.9	29 5	Slight dis- coloration.	,,	,,	,,		
10. 48.9) <u> </u>		_	,,	,,			

As leaf lard does not give a green colour with the reagent, the author thinks that the favourable reception given to the test on its first appearance was due to the experimenters using that lard.

C. A. M.

The Asphalt Question. S. F. Peckham. (Jour. Amer. Chem. Soc., 1895, xvii., No. 1., pp. 55-63.)—In a critical review of various papers that have been pub-

lished on the chemistry and analysis of asphalt, the author states his objection to the method employed by Richardson of determining the amount soluble in CS_2 , since that liquid does not dissolve all the bitumen. With Miss Linton's process,* in which the use of CS_2 is altogether discarded, he obtained very concordant results, agreeing to the third place of decimals. The petroleum ether used should be obtained from petroleum consisting of hydro-carbons of the paraffin series, and have a specific gravity of 0·7. D. Torrey's method, based on successive solutions of the asphalt in alcohol, does not offer any advantages over that proposed by Miss Linton.

C. A. M.

Rapid Estimation of Starch. P. L. Hibbard. (Jour. Amer. Chem. Soc., 1895, xvii. 1, pp. 64-68.)—Of the methods for determining starch in crude celluloses, none can claim absolute exactness. Lintner's process of heating in closed vessels, under high pressure, with water or a weak acid, gives good results when soluble convertible carbo-hydrates other than starch are absent; while Märker's process, which is the most satisfactory, has the same drawback, more or less of the various gums present being dissolved by the diastase. In the latter process the substance is boiled for a few minutes with water to gelatinize the starch, and after cooling to 60° diastase is added, and the liquid maintained at 65° for an hour. The solution is then filtered, the filtrate treated with HCl, and the converted dextrose estimated by Fehling's solution (see Analyst, xx., pp. 17-19).

A shorter and easier modification of this method is suggested by the author, the advantages of which over older processes are: (1) Prevention of clots by the addition of malt extract previous to heating; (2) prevention of lactic fermentation by rapid heating; (3) a minimum solution of bodies other than starch, owing to the short time of boiling.

The malt extract which is used instead of pure diastase is prepared by infusing the malt for several hours, with water containing 15 to 20 per cent. of alcohol, and filtering. The alcohol acts as a preservative, without interfering to any great extent with the diastatic power of the malt. The solution thus prepared will keep for a fortnight, even in warm weather.

In the estimation the finely-powdered substance is heated to boiling in a flask, together with 50 c.c. of water and one to two c.c. of malt infusion. (If the substance contain much fibrous material and but little starch, nothing is gained by adding the malt here, as the diastase is destroyed by the high temperature before it can act on the starch.) After boiling for a few minutes, the mixture is cooled to 50° or 60° C., and another two or three c.c. of malt extract added. It is then slowly heated for ten to fifteen minutes till boiling, again partially cooled, and tested with iodine solution. If any blue colour appear, the treatment with malt must be repeated. When all the starch is changed the liquid is cooled, made up to definite volume, and filtered through muslin or linen. An aliquot part, sufficient to contain 0.2 to 0.3 grammes of starch, is taken from the filtrate, and is boiled on a sand-bath, in a 100 c.c. flask, with 5 c.c. of HCl (30 per cent. gas), and sufficient water to make up about 60 c.c. No reflux

condenser need be used, and the conversion is complete after thirty minutes' actual boiling. When cool, the solution is neutralized with NaOH, and the dextrose estimated with Fehling's solution. Less than two hours is required for the determination. The following figures compare the results obtained by this method with those by the direct inversion method:

			Star	ch pe	r Cent.
Material.		Moisture.	Malt Method	i.	Acid direct.
Starch factory feed		10.0	 10.0		40.9
Impure moist starch		12.0	 81.3		$82 \cdot 3$
,, ,, ,,	• • •		 79.5		80:9
Good starch		12.8	 84.3		84.6
Purified dried starch	• .		 95.1		96.5
Corn, large white kernels	•••	11.0	 63.5		68.2
Wheat, hard red variety		10.0	 60.5		65.0
Bran from same wheat	•••	11.3	 28.4		$44 \cdot 4$
Middlings from same wheat	• • •	10.1	 53.8		60.0
Low flour ,, ,, ,,		10.2	 66.0		$67 \cdot 7$
Good flour ,, ,, ,,	• • •	11.0	 67.9		69.0
Wheat flour		10.0	 70.7		73.6
Wheat bran		10.1	 29.3		46.5
					C. A. M.

A New Method of Estimating Carbon in Iron. Leop. Schneider. (Stahl und Eisen, 1894, 1029; through Chem. Cent. Blatt.)—The author has found that the addition of powdered copper and lead to the iron causes the latter to burn much more readily in a current of oxygen. The lead is prepared by shaking the molten metal, and the copper by reduction of oxidized copper-wire in a current of hydrogen. In the estimation 3 grammes of the finely-divided steel are incorporated with 10 grammes of a mixture of three parts of lead-powder with one part of copper-powder, and introduced into the combustion-tube in a porcelain boat, below which is placed a little In front of the boat is placed 10 to 20 cm. of copper oxide. After the boat has been brought to a dull red heat, the current of oxygen is admitted, and the metallic mixture readily takes fire. The gases produced, after being dried by H₂SO₄, After the combustion the current of oxygen is continued for are led into soda. The whole operation requires three-quarters of an hour. another ten minutes.

C. A. M.

On the Use of Litmus and Methyl Orange as Indicators in Volumetric Analysis. G. Lunge. (Zcit. für angewandte Chemie, 1894, 24, pp. 733-738.)— The author admits that litmus, when prepared and used according to Reinitzer's directions (Analyst, xix. 255), is more sensitive than methyl orange, but denies that the superiority is as great as claimed. With 250 c.c. of distilled water, and using normal or decinormal acid, it is not eight times, but only twice, as sensitive. Since Reinitzer only gave comparative results when the indicators were used with distilled water, the author has made experiments to determine their merits when used with 50 c.c. of a $\frac{1}{5}$ normal soda solution. With normal acid practically identical results are obtained, but methyl orange is preferable on account of its speed and the

precautions to be observed in the use of litmus. With semi-normal acid the change of colour is more difficult to observe in the case of methyl orange, but a practised observer can be sure to a drop; and when liquids containing carbonic acid have to be frequently titrated, the gain of time is a compensating advantage. It is only with decinormal acid that litmus is undoubtedly superior, and Reinitzer's method of titration must be observed.

C. A. M.

The Determination of Potash in Manures. W. E. Garrigues. (Jour. Amer. Chem. Soc., 1895, xvii., No. 1, pp. 47-51.)—The process recommended depends on (1) conversion of all the potash present and removal of ammonia and organic matter by ignition with $\rm H_2SO_4$; (2) precipitation of the $\rm H_2SO_4$ with $\rm BaCl_2$; (3) precipitation of the excess of barium and other earthy bases with $\rm Na_2CO_3$; (4) evaporation of the filtrate with $\rm PtCl_4$.

In the estimation, 10 grammes of the sample are ignited in a platinum basin with concentrated H_2SO_4 , finishing with a blast. The residue is powdered in a mortar with a little hot water, washed into a 500 c.c. flask, and boiled for half an hour with 250 c.c. of water. A 10 per cent solution of $BaCl_2$ is then introduced, 5 c.c. at a time, until no more precipitate forms. One drop of phenol-pthalein solution is then added, and sodium carbonate until a faint pink tinge is observed. The liquid is boiled, cooled, made up to the mark, and filtered. To 50 c.c. of the filtrate (=1 gramme of the sample) HCl is added to slight acid reaction, then PtCl₄, and the liquid evaporated to dryness on the water-bath. The precipitate is washed with alcohol (specific gravity 0.848), dried at 100°, and weighed. For potassium salts ignition with H_2SO_4 is omitted.

The author lays stress on the following points: (1) Cautious addition of the H_2SO_4 ; (2) fine pulverization of the ignited mass in order to prevent calcium sulphate mechanically enclosing potassium sulphate; (3) avoidance of an excess of Na_2CO_3 , which might cause decomposition of the precipitated $BaSO_4$.

From the table of results given, this method, which, in common with other methods, does not account for the entire potash present in samples of known composition, appears to be superior to the Lindo-Gladding process.

C. A. M.

The Addition of Calcium Chloride to the Solution of a Fertilizer in the Determination of Potash. Rudolf de Roode. (Jour. Amer. Chem. Soc., 1895, xvii., No. 1, p. 46.)—In order to allow platinum basins to be used in the Lindo-Gladding method, Huston recommended the addition of sufficient CaCl₂ to form calcium phosphate with the phosphoric acid present. The author finds that, in order to make this effective, it is necessary to filter off the precipitate formed by the CaCl₂ and NH₃, and then to add ammonium oxalate to the filtrate. As this involves two filtrations, the suggestion does not seem to be an improvement on the present alternative method.

C. A. M.

Viscometrical Examination of Butter. C. Killing. (Zeit. für angewandte Chemie, 1894, pp. 643-645, and pp. 739, 740.)—The apparatus used by the author

consists of a wide glass cylinder, closed at the bottom by an indiarubber cork, through which passes a short tube, having its top ground to receive a sort of pipette. This holds about 50 c.c., and has an arrangement in its body to allow of the introduction of a small thermometer. The upper tube of the pipette, which passes through the top cork of the cylinder, is closed by a tap, and the pipette has three marks—one below the body and two above, the latter being placed about 1 cm. apart. The cork closing the top of the cylinder is made to remove in two halves, and a second thermometer is passed through one of these. The whole apparatus is fixed in a clamp, and a beaker is placed below to receive the fat.

The standard of viscosity is the time a definite volume of water at 20° C. takes to run out. The viscosity of butter is taken at 40° C., and it is essential that both thermometers should be compared with a standard thermometer, and a correction made if different.

In using the apparatus the top cork of the cylinder, with the thermometer, is removed and the pipette withdrawn. About 60 c.c. of the clear melted fat is cooled to 40.5° , the pipette thermometer inserted in its place, and the butter sucked to the upper of the two top marks. The tap is then closed, and the pipette placed in the cylinder; the latter is filled to the top with water at 42° , and the cork covers replaced. When both thermometers stand at 40° , the tap is slightly opened and the fat allowed to run slowly down to the lower of the upper marks. The tap is then opened wide, and the fat runs out into the beaker until it reaches the lowest mark, when the tap is closed and the time noted. The mean of two or three determinations, which should not vary more than $\frac{1}{4}$ to $\frac{1}{2}$ a second, is taken, and this compared with water as 100, gives the viscosity number.

In the author's apparatus, distilled water at 20° C. ran out in 80.33 seconds, and supposing a butter to take 3 minutes 42 seconds, this would give as the viscosity number

$$\frac{222.00 \times 100}{80.33} = 276.3.$$

The following tables give the results obtained with different samples of margarines and butters:

A.—MARGARINES.

		Description	n.			Time rui Min.	nning out.	Viscosity Number.	Remarks.
1.	Dutch m	argarine				4	12.50	314.3	
		margarine,	M.D.	Quality	I.	4	13.66	315.8)	From the same
3.	,,	,,		,,	Π.	4	11.66	313.3	factory.
								1	All from factory.
4.	,,	,,	М.В.	,,	1.	4	11.66	313.3	No. 4 contained
5.	,,	,,		,,	II.	4	13.83	315.9 {	earth-nut oil;
6.	,,	,,		,,	III.	4	11.83		Nos. 5 & 6 cotton-
									seed & sesame oil.
7.	Rotterda	m margarin	e			4	12.16	313.9°	•
8.	German	margarine,	M.G.	,,	I.	4	12.38	314.2)	From same
9.	,,	,,		,,	II.	4	15.00	317:4)	factory.
10.	,,	,, I	M.H.	,,	I.	4	14.00	316·2)	•
11.	,,	,,		,,	II.	4	11.83	313∙5∫	,,,
		Mean		• • •	• • •	4	12.77	$314 \cdot 7$	

B.—Butters. Time running out. Description. Viscosity Number. 1. Westphalian 3 min. 42.00 sec. 276.3 . . . 2. Rhine 3 41.90276.2... . . . 3. 3 280.545.33. ,, . . . 4. Dutch 276.53 42.16... . . . Oldenburg 45.163 280.3٠.. ,, Westphalian 3 43.42278.17. Holstein 3 46.00281.3. 8. Rhine 3 44.13279.0... . . . 3 min. 43.76 sec. 278.5 Mean

Taking the mean numbers, the amount of margarine in an adulterated butter may be calculated from the formula:

$$x = (v - 278.5) \frac{100}{314.7 - 278.5}$$
, or $x = 2.76(v - 278.5)$, where $v =$ viscosity number.

The products used in the manufacture of margarine gave the following viscosity numbers:

German oleo-mar	garine	•••	•••	• • •	339.2
American ,,	•	•••			332.7
Earth-nut oil	•••		•••		296.3
Sesame oil		•••	•••		273.9
Cotton-seed oil	•••		•••	•••	258.9

A mixture of 75.6 parts of American oleo-margarine with 24.4 parts of cotton-seed oil gave the mean viscosity number of margarine 314.7. A mixture of 50 per cent. of each gave a viscosity number of 295.8, but the product was so fluid that it would have been unsaleable. It thus appears that it is impossible to make a butter from cotton-seed oil and oleo-margarine with the right viscosity number—viz., 278.5.

The one oil from which an artificial butter with a low viscosity number can be prepared is cocoa-nut oil, which gives 223·1, but according to the author this is no longer employed in the manufacture of margarine, since it has been found impossible to get rid of the taste.

Mixtures of butter and margarine with the respective viscosity numbers of 278.8 and 315.6 showed a proportionate increase in the numbers:

Butter Fat.		Margarine Fat.	Viscosity Number Found.	Corresponding to Margarine %.
95.17	+	4.83	280.20	3.81
75.53	+	24.47	287.56	23.83
55.70	+	44.30	295.50	45.42
42.45	+	57.55	300.16	58.09
25.20	+	74.80	306.09	74.23

The smallest amount of added margarine that can be detected will depend to a large extent on the amount of variation of butters among themselves, and this seems to be considerable. Butter from the milk of a cow fed on beetroot only gave a viscosity number of 270.76, whilst another cow fed exclusively on distillers' refuse produced a butter with viscosity number 278.23. Nevertheless, the author claims for this process superiority over the Reichert-Meissl method.

C. A. M.

The Rapid and Accurate Analysis of Bone-black. W. D. Horne. (Journ. Amer. Chem. Soc., 1895, xvii., 1, pp. 51-55.)—In new char the amount of moisture is of importance, being frequently limited by contract to 3 per cent. It is determined by heating 2 grammes for two hours in an air-bath at 140° C.

Calcium carbonate may be conveniently estimated in Lunge's nitrometer, using mercury. The flask containing the sample and the small tube of HCl is connected with the three-way cock of the nitrometer, and the liberated CO₂ measured with the usual corrections for temperature and pressure. The char in the flask is previously moistened with 3 c.c. of saturated solution of mercuric chloride, which absorbs any H₂S formed.

The carbon, iron, calcium sulphate, and insoluble ash are determined on one portion. This is prepared by first removing all particles of iron with a magnet, and then grinding to an almost impalpable powder. Two grammes of this are boiled in a beaker with 20 c.c. HCl for thirty minutes, and the beaker then filled up with boiling water. The residue settling down is washed several times by decantation, and then transferred to a weighed platinum Gooch's crucible with asbestos plug. It is then washed with 80 per cent. alcohol, followed by 90 per cent. alcohol, and finally with ether, after which it is dried in the oven at 100° and weighed. This gives the carbon and the ash, and the loss of weight on burning over a blast when deducted from this gives the carbon alone.

The acid filtrate is nearly neutralized with ammonia, and ammonium acetate added till the solution turns yellow, then a few drops in excess, and the liquid heated below 70° C. until the iron and aluminium phosphates are completely precipitated. The precipitates washed free from chlorides are dissolved in a 6-oz. flask by dilute H_2SO_4 , 10 c.c. of sodium sulphite added and the liquid boiled until the iron is completely reduced and the SO_2 expelled, after which it is cooled and titrated with permanganate. If the aluminum is to be determined, the above precipitate may be treated with 150 c.c. of ammonium molybdate solution and filtered from the phosphoric acid. The iron and aluminum in the filtrate are then precipitated by ammonia, the precipitate filtered off, dissolved in HCl, and reprecipitated by ammonia, collected, ignited, and weighed. The two may then be separated by any of the known methods.

The filtrate from the iron and alumina precipitation which contains the CaSO₄ is acidified with HCl, the sulphuric acid precipitated by BaCl₂ and the BaSO₄ calculated to CaSO₄.

Calcium sulphide is estimated by evaporating 5 grammes of char nearly to dryness with 20 c.c. $\mathrm{HNO_3}$, and after adding 20 c.c. of HCl again evaporating low to expel the $\mathrm{HNO_3}$. The residue is taken up in HCl, and the sulphuric acid precipitated in an aliquot part of the filtered solution. From the $\mathrm{BaSO_4}$ is deducted that found above as $\mathrm{CaSO_4}$.

The physical conditions of the char should be thoroughly examined. The grist is estimated by shaking gently 100 grammes on sieves of known mesh and weighing what passes through.

The density is determined, both loose and packed, by introducing the char into a weighed 50 c.c. or 100 c.c. flask, making up to the mark, and weighing. From this the density and the weight per cubic foot is calculated.

The porosity is determined by introducing the char little by little into the weighed flask filled with distilled water. The flask is tapped to pack the char, and when filled up to the mark is heated on the water-bath to expel air. After cooling, the supernatant water is removed, and the flask and its contents weighed. The increase of weight over that of the flask packed with char gives the amount of contained water.

To determine the durability of a char under continued friction, the author has devised the following test: 25 grammes of the char are shaken ten times backwards and forwards in a sieve with circular holes one-fiftieth of an inch in diameter, and the dust which passes through weighed. Dust and char are then shaken together in a cylinder of turned iron two inches deep and four inches in diameter with six glazed porcelain marbles. After shaking 200 times the marbles are removed, the char sifted as before, and the increased weight of dust calculated to per cent. of the char used. Fairly constant results were obtained from different parts of the same sample. Thus a new char yielded 1.72, 1.46, and 1.76 per cent. of dust. Two grades made by one firm gave 2.16 and 2.16 per cent. in one case, and 2.86 and 2.87 per cent. in the other. Char which had been in use ten months gave 0.92 and 0.94 per cent., which showed that its wearing condition had improved by use, the softer parts having been separated.

C. A. M.

On the Examination of Linseed Oil and Boiled Linseed Oil. Dr. Hugo Amsel. (Zcit. für angewand. Chemic, 1895, Heft. iii., pp. 73-78.)—With the exception of the Köttstorfer number, none of the usual tests applied to an oil (c.g., the specific gravity, iodine-number, elaidin reaction, etc.), is of much value in the case of linseed oil, since they give results varying so widely in different specimens of the pure oil.

The method proposed by Filsinger (*Chem. Zeit.*, 1894, 1005 and 1867), in which the sample is mixed with chloroform and alcohol, and examined with the polariscope, is useless when mineral oil, which is optically inactive, has been used to adulterate the linseed oil. Moreover, in the case of boiled oils the solution is often so dark as to prevent an accurate observation being made.

As the result of an experience of many years, the author declares that adulteration may be detected with certainty by determining:

- 1. The capacity of the oil to dry when smeared on glass.
- The saponification number, and the behaviour of the alcoholic solution on adding water.

And, according to the results from the above,

- 3. The mineral matter.
- 4. The amount soluble in alcohol.

Pure linseed oil without "driers" ought to dry quite hard in three days at most when spread in a thin layer on glass. With the addition of 5 per cent. driers it should not require longer than twenty-four hours, which should also be the time taken by boiled linseed oil. The addition of only 5 per cent. of rape oil prevents the layer from becoming hard, even after eight days.

In testing an oil the Köttstorfer number is first taken, about two grammes being saponified in the usual way with alcoholic potash. Since mineral oils are not thus saponifiable, and the saponification number of rosin oils very low, the result will give

an approximate idea as to the character of the oil. The average saponification number (i.e., the KHO required per 1000 of oil) of linseed oil is 188, while that for boiled linseed oil varies from 190 to 195.

The drying test is next made, and this will prove whether mineral oils be present. If present, the thin layer on the glass will not be hard after three days.

Another important characteristic of a pure linseed oil is that its alcoholic soap solution remains perfectly clear on adding water, whilst an oil adulterated with mineral oil, rosin, or resinated metallic oxides becomes milky. The following instances, selected from a long table, show this:

TABLE I.

	Saponifica- tion No.	Alcoholic Soap Solution + Water.	Calculated Addition from the Saponification No. per cent.
1. Linseed oil	188	Clear	
2. Boiled linseed oil	196	,,	
3. Blue rosin oil	13.5	Milky	
4. American mineral oil	2	,,	
7. Scotch mineral oil	9.6	"	
10. Linseed oil + 5 per cent.			
mineral oil	177	,,	5.9
11. Linseed oil + 10 per cent.			
mineral oil	171.4	,,	9
16. Linseed oil + 20 per cent.	150		
rosin oil	156	"	18
22. Boiled linseed oil + about 2.5			1
per cent. resinated metal-	107		41 40
lic oxides	187	,,	About 3

The saponification number of No. 22 was so nearly normal that had the alcoholic soap solution remained clear it would readily have been passed as genuine.

TABLE II.—LINSEED OILS.

	Saponification No.	Alcoholic Soap Solution on addition of Water.	Iodine No.	Drying Capacity on Glass +5 per cent Driers.
1.	183	Clear	180	Hard in 24 hours.
2.	188	,,	180	,,
3.	173	Milky		Viscous after 24 hours.
4.	179	,,	112	After 5 days, as at beginning.
5.	189	Clear	180	Hard in 24 hours.
6.	180	,,		,, ,,
7.	174	Milky		Viscous after 24 hours.
8.	165	,,	157	Viscous after 48 hours.
9.	190	Clear	183.5	Hard in 24 hours.
10.	191	,,	167	After 5 days, as at beginning.
11.	190	,,	-	Hard in 24 hours.
12.	189	,,		Hard in 24 hours.

The foregoing tests showed that Nos. 1, 2, 5, 6, 9, 11, and 12 were pure linseed oils. Nos. 3 and 7 were adulterated with rosin oil; No. 8. with mineral oil; and Nos. 4 and 10 with rape oil.

THE ANALYST.

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	Saponifica- No.	Alcoholic Soap Solution + Water.	Iodine No.	Drying Capacity.	Soluble in Alcohol, per cent.	Mineral Matter, per cent.
1.	117	Milky		After 3 days viscous		
2.	141	,, ,	153	Dry after 2 days.		
3.	188	,,		,, ,,		
4.	193	Clear		Hard in 24 hours.		
5.	138	Milky	128	Not quite hard after 2 days		
6.	153	,,		Dry after 2 days	1	
7	195	Clear	181	Hard in 24 hours		Ì
8.	172	Milky		Dry after 2 days	26.3	
9.	199	Clear	174	Hard in 24 hours		
10.	92	Milky	148	Dry in 36 hours	50	
11.	198	Clear	183	Hard in 24 hours.		
12.	191	,,		,, ,,	24	1
13.	197	,,	189	Hard in 24 hours.		
14.	189	Milky	165	Dry after 2 days.	32	
15.	193	Clear	169	Hard after 24 hours.	19.5	
16.	174	Milky	155	Still viscous after 3 days	19	0.21
17.	171.8) ,,	146	,, ,, ,,	18	0.21
18.	175	,,	161	Dry after 2 days	25	0.23
19.	191	Clear	125	After 7 days as at first	8	
20.	125	Milky		Dry after 2 days	37.8	0.4
21.	190	Clear		Hard in 24 hours		
22.	193	,,		Hard in 18 hours		
23.	161.7	Milky		Viscous after 2 days	i	
24.	133	,,		Dry after 2 days		
25.	183	,,	_	,, ,,	1	1

Of the above, Nos. 4, 7, 9, 11, 12, 13, 15, 21, and 22 were pure boiled linseed oil. No. 19 contained 50 to 60 per cent. of rape oil. Nos. 16 and 17 contained mineral oil. Nos. 1, 2, 3, 5, 6, 8, 10, 14, 18, 20, 23, 24, and 25 contained rosin, rosin oil, or resinated metallic oxides.

TABLE IV.—LINSEED OILS, THICKENED BY HEAT AND OXIDATION.

	Saponification No.	Alcoholic Soap Solution + Water.	Iodine No.	Per cent. Soluble in 98 per cent. Alcohol.
1.	195	Clear	94.4	18.5
2.	194	22	104	17.7
3.	209	,,	_	9.2
4.	192	,,		6
5.	111	Turbid		49

Of these, No. 5 was adulterated with about 40 per cent. of rosin, while the others were pure linseed oils.

In the case of thickened oils, the determination of the amount soluble in alcohol is of value. About 1 gramme of the oil is mixed with 20 c.c. of alcohol, allowed to stand at the ordinary temperature for twenty-four hours, with constant shaking, and then filtered. For linseed oils the mean amount is about 25 per cent., while thickened oils always yield under 20 per cent.

C. A. M.