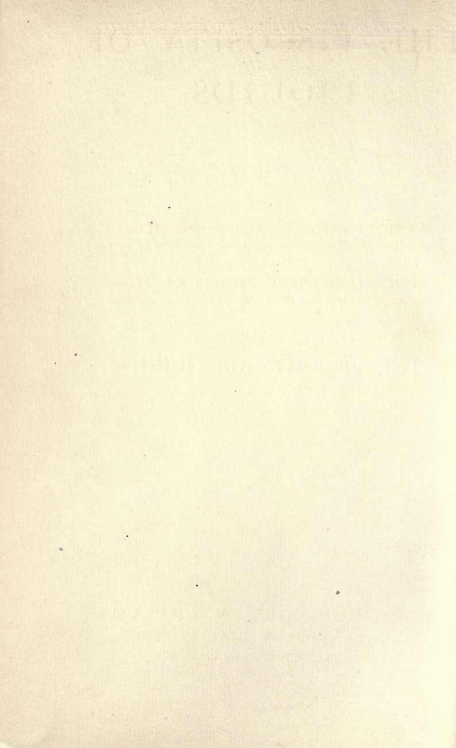


MONOGRAPHS	ON	INORGANIC A	ND	PHYSICAL	CHEMISTRY
En	TED	BY ALEXANDES	FI	NDLAY D.Sc	

THE VISCOSITY OF LIQUIDS



phone

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BY

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Monographs on Inorganic and Physical Chemistry

EDITED BY

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To those engaged in guiding the reading of advanced students of chemistry, the difficulty of obtaining adequately summarised accounts of the progress made in recent years, more especially along certain of the more actively pursued lines of advance, becomes ever more acutely felt. So great has now become the volume of chemical investigation, and so numerous the channels of its publication, that not only the Honours Student but also the worker desirous of undertaking Research in one or other department of his subject, feels it a growing difficulty to become au fait with the present state of the more important and more strenuously cultivated regions of his Science. To assist these two classes of students-those reading for an Honours Degree, and those undertaking Research—is the main aim of the present Series of Monographs.

In this Series of Monographs it is hoped to place before advanced students of Chemistry, accounts of certain sections of Inorganic and Physical Chemistry fuller and more extended in scope than can be obtained in ordinary text-books. Exhaustive treatment of the different subjects, however, so far as concerns work important in its time but now only of historical interest, will not be attempted; the chief attention will be given to recent investigations.

Arrangements have already been made to publish the following monographs, and should these prove themselves to be of value, others will be issued from time to time.

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PREFACE.

THE development of a physical property may follow two paths. Either the intrinsic importance of the property. may lead to investigation from a purely physical standpoint, or its application to the elucidation of chemical constitution may cause improvements to be sought for in method, degree of accuracy, and ease of application. the majority of physical properties utilized by the chemist, progress has gone on uniformly in each direction and a standardization of apparatus and technique has been arrived at, but in spite of a century of experimental work no very definite method has been set up for the measurement of viscosity, and no general agreement has been arrived at for the setting forth and interpretation of the results obtained. To a certain extent the inherent difficulty of understanding the strict physical meaning of viscosity is the cause of this lack of progress. The viscosity of a liquid is at least a dual phenomenon; there is first of all the mechanical friction of molecule against molecule, and this must depend on relative molecular surface and volume; then there is the resistance to deformation brought about by the mutual attraction of the molecules

It follows at once that viscosity must be largely a constitutive property, and perhaps it is this fact—that so little additivity is to be looked for—that has led investigators to turn their attention to properties where a mole-

cular value can readily be synthesized from atomic constants.

It should be borne in mind, however, that mere additivity is comparatively useless for the purpose of the chemist, who is searching for a physical confirmation of erratic chemical behaviour. When one considers, for example, such questions as the enol-keto tautomerism, the benzene problem, the effect of ring formation, and the contiguity of unsaturated groups, and the various physical methods which have been utilized for their elucidation, it will be obvious that it was the failure of the additive nature of such properties as refractive index, molecular volume, molecular magnetic rotation, and so on which turned out to be valuable. Viscosity stands out as an eminently constitutive property of the same order as optical rotatory power and the solution absorption of light. It is, therefore, of considerable importance to know how viscosity can be measured, and how the property can be utilized in the study of chemistry.

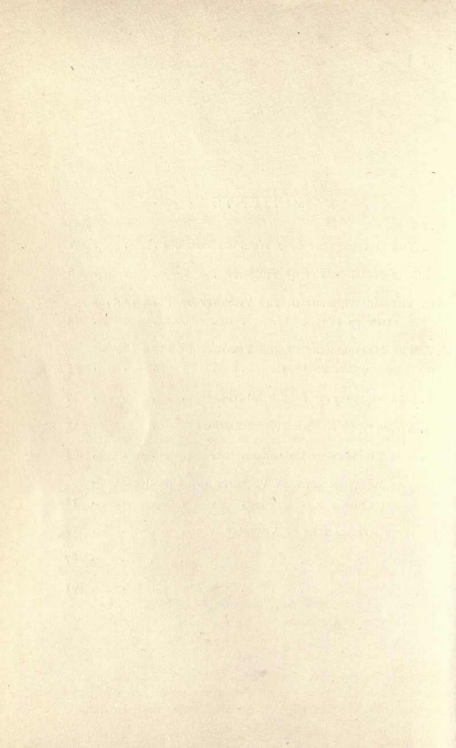
In the following pages will be given a brief description of some of the more important earlier researches which laid down the foundations of the subject, the development of a working formula, and the corrections which have been applied to it, the various apparatus which has been used at different times, and a series of chapters dealing with recent lines of work in which viscosity is playing an

important part.

The authors desire to thank the Councils of the Royal Society, the Chemical Society, and the Faraday Society for permission to reproduce diagrams which have appeared in their respective Transactions; they are indebted to Prof. Washburn for Fig. 2.

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CHAPTER I.

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THE DEVELOPMENT OF A WORKING FORMULA.

Poiseuille 1 and Girard 2 carried out the pioneer work on transpiration, or the flow of liquids through capillary tubes. The former was attracted to the problem from a physiological standpoint, and not only laid down definitely the law followed by transpiring liquids, but determined with consummate accuracy the viscosity of water through a considerable range of temperature. On the basis of his experiments Poiseuille deduced the formula

$$V = \frac{KD^4H}{L}$$

where V = volume transpired in c.cm.

L = length of capillary.

D = diameter of capillary in mm.

H, = pressure in mm. of mercury.

K = a factor which is constant for each liquid at a given temperature.

This empirical formula and its corrections have been substantiated mathematically by the labours of Stokes, Wiedemann, Hagenbach, Stefan, Couette, Wilberforce, Neumann, Jacobson, Knibbs, Boussinesq, 2 and Brillouin.

Making the assumptions that there is no slip at the surface of the capillary, that the liquid flows steadily and without eddies or turbulent motion, and that there is no kinetic energy of efflux, then the transpiration formula becomes for a liquid flowing under its own hydrostatic pressure

$$\eta = \frac{\pi}{8} \left[\frac{hgr^4}{l \times v} \cdot d \cdot t \right]$$

where η = the viscosity, h = the head of liquid, g = the gravitation constant, r = the radius of the capillary, d = the density of the liquid, t = the time of flow, l = the length of the capillary,

and v = the volume of the liquid which has transpired in the time t.

It is usual, however, to introduce a correction for loss of kinetic energy, viz., $-\frac{v}{8\pi l} \cdot \frac{d}{t}$ (the Couette-Finkener-Wilberforce correction).

Hence the corrected formula is $\eta = \frac{\pi}{8} \cdot \frac{hgr^4}{lv} dt - \frac{v}{8\pi l} \cdot \frac{d}{t}$.

For small pressures, high viscosities, and narrow capillaries uniformly blown out at the junctions, the correction tends to become vanishingly small, and indeed in many cases may be conveniently neglected.

This formula resolves itself into $\eta = A dt - B \frac{d}{t}$ where A and

B are the constants for any given apparatus, and t and d are the time of flow and the density respectively. It assumes that the area of the cross section of the capillary is uniform throughout, but if this is contracted or expanded, the correction will be correspondingly affected.

Knibbs, in who has thoroughly investigated the recorded experiments on the viscosity of water, concludes that the correct formula should be

$$\eta = \frac{\pi}{8} \frac{hgr^4}{lv} dt - 1.12 \frac{v}{8\pi l} \cdot \frac{d}{t}.$$

It will be noticed that the correction term has a slightly larger value than the Couette expression.

Couette also pointed out that the modification of the stream lines at the entrance to the capillary caused loss of pressure in overcoming friction, and estimated that this could be allowed for by adding to the observed length of the capillary an amount λ equal to thrice the diameter of the above, the formula thus becoming

$$\eta = \frac{\pi h g r^4 dt}{8v(l+\lambda)} - \frac{1.12 \ dv}{8\pi t \ (l+\lambda)}$$

Bingham and White ¹⁷ deal very fully with these various corrections and the reader is referred to their paper for a detailed account of the elucidation of a working formula for the determination of absolute values of viscosity.

Viscosity and Rate of Flow.—The important work of Grün-

eisen 14 leads to a method for standardizing a viscometer which shows deviations from the Poiseuille Law. This simple law states that $\eta = p \times t \times \text{const.}$, i.e. the same liquid flowing through the same capillary at different pressures, should give a constant value for pt. Grüneisen showed that this condition is by no means rigidly fulfilled by viscometers of the Ostwald type (vide infra) in which the liquid flows under its own hydrostatic pressure. The variation in the values of pt is due to the fact that above a certain limiting velocity the flow is not steady but turbulent, some of the potential energy being expended in forming eddies within the liquid. Hence the liquid is not forced down the capillary so quickly as it should be if Poiseuille's Law held good, and therefore the values of pt are greater than with a slower rate of flow. It follows then that the readings of a viscometer are only trustworthy when the times of flow are sufficiently great to ensure that the product pt remains constant over the whole range of observations. Grüneisen has tested several viscometers and has published the pt curves; he has also calculated a correction to be applied where pt is no longer constant. Evidently such a correction can be rendered unnecessary by reducing the diameter of the capillary, by shortening it or by reducing the hydrostatic pressure so as to lengthen the time of flow (Applebey, 15 Merton 91).

Bingham, 16 however, criticizes the application of Grüneisen's method and holds that the usual kinetic energy correction should be applied although it is obvious that a sufficiently long time of flow would cause $-B\frac{d}{t}$ to become vanishingly small.

The same author points out that near the entrance to the capillary the liquid is in accelerated motion for a short distance. In gaining this kinetic energy the pressure falls. The liquid at the exit end of the tube has the same velocity as at every other cross section, and its inertia carries it a considerable distance into the body of the liquid in the lower reservoir. Owing to the production of eddies this motion finally becomes disordered and disappears as heat. Bingham and White ¹⁷ measured the viscosity of water under the above conditions with a capillary which was afterwards cut into several pieces and they found that the kinetic energy corrections as demanded by the theory are directly proportional to the number of capillaries in series. The danger of

turbulent motion causing error has been dealt with by Osborne Reynolds ¹⁸ who has given an expression for this critical velocity above which eddy currents may be formed, viz. $\frac{700 \, \eta}{dr}$ cms. per second.

In connexion with the experimental determination of viscosity it should be noted that the flow of liquid is influenced in a very marked way by the driving pressure that is used. Bose and Rauert have made measurements at pressures from 0.005 to 2 kilograms per sq. cm., and find that whilst Poiseuille's Law holds for low pressures, very marked deviations are found when the pressure is increased, and in some instances the relative rates of flow are reversed, the more viscous of two liquids flowing more readily and becoming the less viscous at high pressure.

Viscosity and Temperature.—Poiseuille 1 found that his experimental results for the viscosity of water could be represented by the expression $\eta_t = \eta_o/(1 + at + \beta t^2)$ where a and β are constants.

O. E. Meyer ¹⁹ suggested the formula $\eta_t = \eta_o/(1 + at)$, but this was found to apply only over a limited temperature range.

Slotte ²⁰ proposed the equation $\eta_t = \frac{C}{a+t} - b$, where C, a and b are constants, but replaced this ²¹ by $\eta_t = \eta_o/(1 + \beta t)^n$ where β and n are constants.

Graetz ²² considered that the experimental data could be represented by the formula $\eta_t = \frac{A (\theta - t)}{t - t_1}$ in which $\theta =$ critical temperature, and $t_1 =$ a temperature below the melting point where the viscosity is infinite.

In their classical memoir Thorpe and Rodger,²³ after a critical examination of these formulæ, found that Slotte's second expression was the best, and they used it in the representation of the viscosity at various temperatures of no less than seventy liquids.

The expression given by Slotte, when expanded on the assumption that the temperature coefficient is small, simplifies to

$$\eta = \frac{A}{I + at + \beta t^2}$$

In the following table are quoted some of the results obtained by Thorpe and Rodger, which will illustrate the accuracy with which the formula of Slotte agrees with the experimental numbers.

Substance.			a.	β.	tº.	η× 10 ⁵ Calc.	$\eta \times 10^5$ Observed.	Difference.	
				27/4/2					
Pentane .			1000	.01044	'04230I	30°	212	212	0
Iso pentane			7.	.01088	'O41331	30°	204	204	0
Hexane .	. /			'0II22	'043337	60°	221	221	0
Iso hexane				.oiiio	.013200	60°	207	208	I
Heptane .				.01214	0,4004	90°	214	214	0
Iso heptane				.01100	.043863	80°	216	216	0
Octane .				.01394	.0,4926	120°	208	208	0
Allyl iodide		100		.01319	*0 ₄ 344I	100°	349	352	3
Allyl bromide				*01177	.042871	70°	315	316	I

40°

140°

130°

130°

1000

1100

1400

160°

90°

100°

50°

273

249

229

229

526

417

319

315

436

263

245

273

254

233

233

542

417

322

314

407

283

245

0

4

16

0

3

I

29

20

TABLE I .- TEMPERATURE COEFFICIENTS OF VISCOSITY.

·01111 ·0,2639

·01418 ·043923 ·01472 ·044578 ·02870 ·031695 ·01826 ·048537

01720 046941

'02109 '0₃1107

·02552 '032030

·03580 '0,2253

·01064 '043115

.045636

.01701

01418

Allyl chloride .

o-xylene .

m-xylene .

p-xylene . Formic acid

Acetic acid

Water

Acetone .

Propionic acid .

n-Butyric acid .

Allyl alcohol .

The values of α and β in the above table show that associated compounds such as water, the fatty acids and the alcohols have relatively large temperature coefficients of viscosity and that such substances, as might be expected, do not follow Slotte's expression with any great degree of accuracy. Acetic acid is an interesting case and apparently does not bear out the above statement; it must be remembered, however, that the association of this acid persists even in the state of vapour. It will be noticed, further, that in homologous series the numerical values of α and β increase with the molecular weight, but not regularly. Replacement of hydrogen by halogen raises the value of the coefficients. Fifty-five out of the seventy liquids gave differences between the observed and the calculated values of less than 5 per cent.

Brillouin 132 has reviewed the data of Thorpe and Rodger in order to elucidate the relationship between viscosity and temperature. He concludes that Slotte's formula is not the most advantageous and advocates the use of the hyperbolic formula of Graetz:-

$$\eta = A \frac{(\theta - t)}{(t - t_1)}$$

Graetz's formula was rejected by Thorpe and Rodger as it could not be applied to water or to alcohols, but Brillouin was able to show that it represents accurately the behaviour of a large range of non-polymerized liquids and breaks down only in the case of substances which exhibit abnormally large surface tensions at lower temperatures and which may be regarded as undergoing polymerization when allowed to cool. For such compounds a more complex formula is proposed, namely

$$\eta = A \sqrt{T} \cdot \frac{(t - t_1)^2 + C}{(t - t_0)^2 + C}$$

where T is absolute temperature and C and C' are constants.

Temperature of Comparison.—Although most investigators have measured the viscosity at some predetermined temperature, it is obvious that since the influence of temperature on the viscosity is different for different liquids, strictly comparable results can only be obtained when the compounds examined are at comparable temperatures. The boiling point suggests itself as the most convenient of such temperatures. It may be said at once that strictly additive relationships have not been established satisfactorily for viscosity at any one temperature although some progress has been made with its derived functions such as fluidity and the logarithm of the viscosity (p. 37).

A novel method of obtaining comparable results was devised by Thorpe and Rodger²³ who suggested that the temperatures where the viscosity-temperature curves attained equal slopes, i.e.

where $\frac{d\eta}{dt}$ was the same, should be chosen and that substances for

which the rate of change of viscosity was identical would probably be in a comparable physical condition. The rate of change of viscosity with temperature depends to a certain extent on the rate at which the dissociation of an associated compound is progressing, but it may be said with confidence that so far as viscosity is concerned the temperatures where the viscosity-temperature curves have equal slopes are corresponding temperatures for all substances.

By using this conception of temperatures of equal slope it has been found possible to reconstruct molecular values of viscosity which agree fairly satisfactorily with those observed. It was further shown that the ratio of the viscosities measured at two different temperatures of equal slope is constant. For thirty-three different liquids the ratio

$$\frac{\eta \text{ at slope o } \circ \circ_{4}987}{\eta \text{ at slope o } \circ_{4}323} = 2 \circ_{3} \text{ approx.}$$

was found to be practically the same, and hence it is clear that this method of comparison is independent of the value of the slope at which the comparison is made. On comparing the temperatures of equal slope in homologous series it is found, as shown by the following table, that an increment of CH₂ is accompanied by an increase in viscosity which is approximately constant.

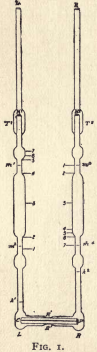
TABLE II .- TEMPERATURES AT WHICH THE SLOPE IS 0.04323.

Substance.		t.	Temperature Difference for CH ₂ .
Pentane Hexane Heptane Octane		- 5.4° 20.5° 41.1° 64.1°	25.9° 20.6° 23.0°
Acetone Diethyl ketone	:	17.8°	16.3°
Methyl ethyl ketone . Methyl propyl ketone	:	43°7° 56°5°	12.80
Methyl iodide Ethyl iodide Propyl iodide		42.9° 61.5° 83.6°	18.6° 22.1°
Ethyl bromide Propyl bromide		26·9° 54·7°	27.8°

CHAPTER II.

THE MEASUREMENT OF VISCOSITY.

Absolute Viscosity.—When absolute measurements of viscosity are desired it will be obvious from the formula given on page 2 that a knowledge of the driving pressure, the diameter and the length of the capillary, the volume of the liquid, the time of flow and the temperature must be obtained. The methods adopted by the earlier investigators such as Poiseuille, Graham 24, Rellstab 25, Pribram and Handl 26 are now of historical interest only. first comprehensive attempt to determine with a high degree of accuracy the various quantities mentioned above was that of Thorpe and Rodger²³. A diagram of the instrument used by them is given in Fig. 1. The capillary (diam. 0.008 cm.) and the



bulb (5 c.c.) were designed to give a minimum time of flow of 3 minutes which could be read to 0.2 second. This time of transpiration is equivalent to a maximum velocity of 66 cm./sec, which is well within that which causes turbulent flow (Osborne Reynolds 18). The dimensions of the capillary were determined in the usual way by weighing the mercury content and then the capillary was fused into position inside the thin glass sleeve connecting the bulbs A and B, every care being taken to avoid deformation during this process. A measured volume of liquid contained between the etched marks m₁ m₂, or m₄ m₃ was then forced through the capillary by an external pressure which could be measured by a suitable manometer. The movement of the liquid was observed by means of a telescope with cross wires, and the time of flow determined with the aid of a stop watch. The liquid was introduced into bulb A through the narrow end of the tube H2, and any small

excess was expelled by applying air pressure to the tube b until the levels K_1 and H_2 were obtained, the surplus running over into bulb T_2 . Alternate times of flow for bulbs A and B were observed and averaged. Extreme care was taken to avoid dust particles which would choke the capillary, and to this end each liquid was distilled in an all-glass apparatus until careful examination failed to detect any solid matter. Temperature control was achieved by means of a large thermostat and regulator.

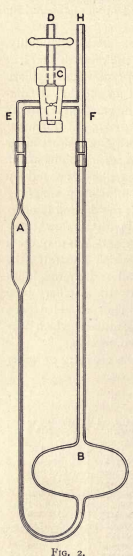
Bingham and White 17 in their recent work on fluidity, criticize Thorpe and Rodger's apparatus on the grounds that it is difficult to construct and to clean, that the constants of the capillary are determined before the viscometer is built up and that the diameter of the capillary is not quite accurately known. These authors recommend a capillary of approximately 0.025 cm. diameter and 10 cm. long which is ground into side tubes quite similar to those used by Thorpe and Rodger and possessing bulbs of the capacity 2.6 c.cm. The diameter of the capillary is found by the optical method for the ends and for the middle by weighing a mercury thread. The viscometer is kept at the desired temperature in a 50 litre bath which is provided with windows, stirring gear and thermo-regulation. Bingham and White quote a series of readings for the viscosity of water which illustrates the order of the experimental error in five careful and thorough investigations, all of which were made using the capillary tube method. Hosking's 27 apparatus differed from Thorpe and Rodger's and from Bingham's merely in possessing a capillary which was attached to the bulbs by means of a rubber connexion.

The following table gives the values of the viscosity of water obtained by different investigators.

Bingham and White. Temperature Thorpe and Poiseuille. Slotte. Sprung. Centigrade. Rodger. 00 0.01797 0.01776 0.01778 0.01808 0.01778 100 0.01300 0.01301 0'013025 0.01301 0'01314 20° 0.01008 0.01000 0.01003 0.01008 0.01012 30° 0.00803 0.00803 0.007908 0.00803 0.007975 40° 0.00653 0.00657 0.00657 0.006532 0.006563 50° 0.00553 0.00553 0.005475 0'005500 60° 0.00468 0.004735 0.00472 70° 80° 0.00408 0.00406 0.004075 0.00358 0'00356 0.003570 90° 0.003122 0'003143 0.00318 100° 0.00583 0.00282

TABLE III .- VISCOSITY OF WATER.

Relative Viscosity.—In the great majority of cases, however, it is not necessary to determine absolute viscosities directly and use may be made of any apparatus which has been previously calibrated with some liquid of known viscosity. The eminently simple and useful instrument designed by Ostwald ["Physico-Chemical Measurements," 1894, p. 163] has, either in its original or



in a slightly modified form, been largely used by workers in this subject. The conditions necessary for rendering it an accurate and trustworthy viscometer have formed the subject of recent discussion. Reference has already been made to the Applebey, investigations of Merton, Grüneisen, and Bingham with respect to the agreement of the values obtained with this type of apparatus with Poiseuille's Law; but there still remains a source of error due to the uncertainty which arises from the variation in the water constant of the instrument. There are four reasons for the variations in the water constant: (a) the presence of solid particles which block up the capillary, (b) the solvent action of water and cleaning liquids on the capillary, (c) the temperature hysteresis, (d) the contamination of the water by the dissolved glass. To surmount these difficulties Washburn and Williams 28 have devised an improved Ostwald viscometer constructed entirely of quartz (Fig. 2).

The dimensions were calculated according to Grüneisen's work so as to avoid errors due to deviations from Poiseuille's Law. The diameter of the capillary was 0.05 cm., wide enough to avoid choking by dust particles, and its length 19.5 cm. The volume of the upper bulb A was 9 c.c. and the large receiving bulb was 4.5 cm. high and 8.0 cm. wide. The mean head of liquid during its flow was 20 cm. At

the upper and lower ends of the capillary the change in diameter was made as gradual as possible so as to prevent eddy currents, whilst the constriction at the upper end of A was made symmetrical with the one at the lower end to eliminate any unequal surface tension effect when using liquids with surface tensions different from that of water. The viscometer was filled by means of a 65 c.c. pipette and the error in filling was shown to be quite negligible. The times of flow were determined by a recording chronograph operated by a motor and controlled by a standard clock, giving an error not exceeding 0.01 second. The temperature of the bath was kept constant within a few thousandths of a degree by using a large cylindrical silvered Dewar tube 50 × 12 cm. inside measurement provided with stirrer, heating coil, standard thermometer, and a Beckmann thermometer acting as a thermoscope. The liquid having been introduced, the threeway cock is attached as shown in the figure and the instrument is suspended in the bath from a metal stand and sunk until the horizontal connecting tube is immersed. The cock is then turned to connect D with the capillary arm of the viscometer and suction is applied till the bulb A is filled. The cock is then closed and turned to connect E with F. When the meniscus passes the upper mark of bulb A the observer taps the key, and repeats the operation when the meniscus passes the lower mark, the times of these two observations being recorded automatically on the chronograph tape.

Standardization of an Ostwald Viscometer.—To determine how closely a given instrument follows Poiseuille's Law, times of flow may be measured for the same liquid under different pressures or for different liquids under the same or different pressures.

sures, and since $\frac{\eta}{t \times p} = K$ where K depends only on the dimensions of the particular instrument, it follows that the $p \times t$

values should lie on a straight line.

Washburn and Williams (loc. cit.) first used the varying pressure method, connecting the viscometer with a large air reservoir and a manometer. The actual driving pressure in the viscometer is of course the sum of the manometric pressure and the head of liquid at mean time of flow, which was measured directly when half the time of flow had elapsed. The results of a series of readings are given in Table IV.

Total Pressure in mm. of Water.	Time of Flow in Seconds.	pt.	Deviation from Mea	
132.04	874'7	11550	0'04	
161.21	714.6	11542	0.03	
272.97	423.0	11547	0'02	
294'79	391.2	11541	0.03	

TABLE IV .- CALIBRATION OF A VISCOMETER.

It is obvious that for such pressures and times of flow, Poiseuille's Law is obeyed excellently, and a value for K can be obtained

since
$$K = \frac{\eta}{p \times t}$$
.

Seeing that quartz has so small a coefficient of expansion the dimensions of the instrument do not vary appreciably with temperature and accordingly the constant can equally well be determined by measuring the time of flow of water at varying temperatures under a pressure due to its own density. The formula then becomes

$$K = \frac{\eta}{t \times d}.$$

Three such measurements at o°, 25° and 50° gave

$$K_0 = 154.60 \times 10^{-7}$$

 $K_{25} = 154.72 \times 10^{-7}$
 $K_{50} = 154.72 \times 10^{-7}$
Mean = 154.68 × 10⁻⁷.

The apparatus just described have been devised for the extremely accurate determination of the viscosities of water or dilute solutions, but it by no means follows that this type of apparatus is essential or indeed desirable for all varieties of work where such a degree of accuracy is not aimed at, where the supply of material is very limited, or where special experimental difficulties To meet the special requirements of their investiare met with. gations, different workers have introduced a number of modifications of the Ostwald apparatus. Thus Findlay, 29 for investigating the viscosities of mixtures of volatile liquids near their boiling points, used an Ostwald viscometer, the two limbs of which were provided with bulbs containing some of the liquid under investigation. This device effectually stopped loss by evaporation. Faust, 30 to avoid choking the capillary with dust particles, allowed it to enter the side of a wide tube, so that solid matter might sink to the bottom and not get sucked up into the capillary. Thole 39 has described two Ostwald viscometers for use with organic liquids, where only small amounts of material are available. In these instruments horizontal capillaries, which enter the vertical limbs near the bottom, are made use of so as to avoid the entrance of solid particles. Heber Green ³¹ used a straight tube viscometer with two receiving bulbs, the one small for viscous liquids and the other large for more mobile fluids. The end of the tube dipped into a flask which was kept at a constant predetermined position with respect to the tube.

The Ostwald method has been used by Beck, 35 up to 250° C. for fused mercury salts, and up to 600° C. by Lorenz and Kalmus 36 . Goodwin and Mailey 37 used a platinum capillary, Woelter 38 measured the rate at which a platinum wire sinks in the liquid (fused silicate), whilst Arndt 34 also proposed to measure the viscosity of fused salts by observations on the time of sinking of a weight. An ingenious method due to Scarpa 133 and slightly modified by Farrow 134 consists in measuring the time t_1 which is occupied in drawing up, through a vertical capillary tube, sufficient liquid to fill a bulb at the top of this tube, and the time t_2 which this volume takes in flowing out from the bulb under the weight of the liquid. Given constant conditions the viscosity

of any liquid is proportional to the expression $\frac{t_1t_2}{t_1+t_2}$. The advantage of this method lies in the fact that no density determinations are necessary and that it is immaterial whether equal volumes of liquid are used in the different experiments.

Coulomb's method ³² of determining viscosity by the rate of damping of an oscillating suspended disc has not been used to any great extent owing to experimental difficulties and the need of a large volume of liquid, but Fawsitt ³³ has revived its use particularly with a view to measurements on fused salts, metals, and alloys.

For a full description of the chief methods of determining viscosity, reference should be made to the work of Arndt ⁴⁰.

CHAPTER III.

THE MEASUREMENT OF THE VISCOSITY OF PURE LIQUIDS PRIOR TO 1895.

ONCE a method for the measurement of viscosity had been established, attempts were made to find a connexion between this physical property and molecular weight, atomic grouping, and so The earliest of such investigations appears to be that of Rellstab 25 who examined some homologous series of acids, alcohols, esters, aldehydes and ketones, measuring the times of flow of equivalent weights of the substances at temperatures where their vapour pressures were equal. In brief Rellstab found that (I) the times of flow decrease as the temperature rises; (2) in an homologous series each increment of CH, increases the transpiration time; (3) increase of carbon diminishes whilst (4) unsaturation increases the time of flow; (5) increments of CHOH, H2 and O increase the viscosity; (6) metamers have different viscosities. Přibram and Handl 26, using the Poiseuille method, concluded that the substitution of halogen or NO, for hydrogen increased the viscosity; that isomeric esters have nearly the same viscosity, but that the normal ester is more viscous than the iso compound; that the alcohols have a higher viscosity than the corresponding aldehydes and ketones; and that in homologous series there is a fair proportionality between the increments of viscosity and of CH₂.

Similar results were obtained by Gartenmeister ⁴¹. Fatty alcohols and acids were shown, however, to possess remarkably high values and to give abnormal homologous relationships. Unsaturated compounds were found to be less viscous than the corresponding saturated ones.

The three series of investigations just mentioned agree in one respect, viz. their vagueness. No simple relationships are pointed out and no way is shown whereby molecular and atomic values of viscosity can be attained. In other words, the constitutive effects overshadowed the additive. These investigations, however,

cleared the way for the exhaustive researches of Thorpe and Rodger,²³ whose contribution to this subject surpasses any previous work in accuracy of measurement, wide scope of inquiry, ingenuity in manipulation and persistence in critical discussion of the experimental data. The apparatus which they used has already briefly been described.

Thorpe and Rodger determined the temperature-viscosity curves of about 70 exceptionally pure liquids, drawn from widely different families so as to include the effects of homology, substitution, isomerism and molecular complexity. They selected, as has already been mentioned (p. 6), the temperature of equal slope as that at which most truly comparable results should be obtained, and the results which they obtained lead to the following conclusions.

For unassociated liquids the viscosity increases regularly as an homologous series is ascended. An iso compound is generally less viscous than the normal isomeride. Allyl compounds are more viscous than iso propyl and less viscous than the normal propyl derivatives. Viscosity, indeed, is particularly susceptible to the arrangement of the atoms in the molecule. The substitution of chlorine, bromine and iodine for hydrogen causes a greater and greater increase in viscosity. Formic and acetic acids are highly anomalous in that they are more viscous than propionic acid. Associated liquids have a much higher viscosity temperature coefficient than unimolecular compounds have, whilst in most cases the initial members of a series show anomaly.

Recognizing that previous workers had not deduced a viscosity expression containing a molecular term, Thorpe and Rodger decided to compare not only viscosity but molecular viscosity and molecular viscosity work.

If Mv be the molecular volume of a liquid, that is the volume which contains for different substances an equal number of molecules, $(Mv)^{\frac{2}{3}}$ will represent an area over which an equal number of molecules is distributed; it may be called the molecular surface. The expression $\eta(Mv)^{\frac{2}{3}}$ will then represent the force in dynes required to keep a molecular surface in motion with unit velocity relative to another surface unit distance apart, and is called the molecular viscosity.

Since $(Mv)^{\frac{2}{3}}$ is the molecular surface $(Mv)^{\frac{1}{3}}$ will denote

molecular length, and will represent for different substances an equal number of molecules. Hence $\eta(Mv)^{\frac{2}{3}} \times (Mv)^{\frac{1}{3}}$ or $\eta(Mv)$ represents the *molecular viscosity work*, that is the work in ergs required to move a surface equal to the molecular surface through the molecular length with unit velocity.

In their endeavour to obtain evidence of additive relationships Thorpe and Rodger compared viscosity, molecular viscosity, and molecular viscosity work not only at the boiling-point but also at temperatures of equal slope, and they found that much better results were obtained in the latter cases. To illustrate the kind of results obtained at the boiling-point the following table may be given.

TABLE V.-VISCOSITY AT THE BOILING-POINT.

Subs	η×105.	Δ CH ₂ .			
Pentane		×		200	
Hexane				204	4
Heptane				199	-5
Octane				198	- I
Formic acid .				536	
Acetic acid .		37.		385	- 151
Propionic acid				319	- 66
Butyric acid .				309	-10
Methyl alcohol				329	
Ethyl alcohol .				441	112
Propyl alcohol.				463	22
-Butyl alcohol				405	-58

It was found that in ascending an homologous series the viscosity coefficients diminish, but for associated compounds the differences for an increment of CH_2 vary very irregularly. For corresponding alkyl chlorides, bromides, and iodides the increasing molecular weight caused an increase in the viscosity. The n-propyl compounds were uniformly more viscous than the corresponding allyl compounds, whilst iso-derivatives were not so viscous as the normal substances. The earlier members of the series showed the anomaly which is so characteristic whatever physical property is being studied, and in general the effects of mere molecular weight were often outbalanced by association and by the arrangement of the atoms in the molecule.

Better results were obtained by considering molecular viscosity

at the boiling-point, and although similar conclusions to the above were found to be justified, yet sufficient additivity was discovered to make possible the calculation of atomic values and the synthesis therefrom of a molecular value.

TABLE VI.—Atomic Viscosity Constants Derived from Molecular VISCOSITY AT THE BOILING-POINT.

Hydrogen .				80
Carbon .				- 98
Hydroxyl oxyge:	n			196
Ethereal oxygen				35
Carbonyl oxyger	1			248
Sulphur .				155
Chlorine .				284
Iso-union .				15
Double bond				113
Ring formation				610

TABLE VII.—CALCULATED AND OBSERVED MOLECULAR VISCOSITIES.

Substan	ice.		Observed Value.	Calculated.	Difference per cent.
Pentane			480	'470	2'I
Hexane			550	532	3.3
Heptane			593	594	I
Octane			646	656	1.2
Methyl alcohol.			401	421	: 5
Ethyl alcohol .			693	483	30.3
n-Propyl alcohol			869	545	37°3
n-Butyl alcohol			883	607	31.3
Ether			459		3.2
Propionic acid .			630	443 630	0
n-Butyric acid .			702	692	1.4

Very similar regularities were obtained when the values of molecular viscosity work were compared, and it was found possible to synthesise satisfactory molecular values for paraffins, monohalogen compounds, sulphides, ketones, oxides and acids, whilst unsaturated hydrocarbons, di- and poly-haloids, formic acid, benzene, water and the alcohols were quite anomalous.

Coming now to the cases when comparisons are made at temperatures of equal slope it is found that far more consistent results can be obtained. In the following table the values refer to the slope 0.04323.

TABLE VIII.—VISCOSITY AT TEMPERATURES OF EQUAL SLOPE.

A .- Homologous Compounds.

Substance.		η × 105.	Δ for CH ₂ .
Pentane		299	10
Hexane		318	19
Heptane	- 1	330	6
Octane		336	0
Isoprene		295	
Diallyl		304	9
Methyl iodide		399	-
Ethyl iodide		404	5
Propyl iodide		407	3
Ethyl bromide		368	
Propyl bromide		372	4
Ethylene bromide		455	
Propylene bromide		44I	- 14
Methyl sulphide		335	6 × 2
Ethyl sulphide		346	0 1 2
Acetone		329	7 V 0
Diethyl ketone		343	7 × 2
Methyl ethyl ketone		330	7.4
Methyl propyl ketone		344	14
Formic acid		373	2
Acetic acid		370	- 3
Propionic acid		390	-11
n-Butyric acid		379	-11
Acetic anhydride		378	I
Propionic anhydride		379	1
Benzene		330	24
Toluene		354	13
Ethyl benzene		367	13
Iso-propyl chloride		317	7.
Iso-butyl chloride		331	14
Methylene chloride	18 1	372	-
Ethylene chloride		377	5

B .- Normal and Iso Compounds.

Class.				Normal: η × 10 ⁵ .	Iso: η × 10 ⁵ .	Δ.
Pentanes				299	286	13
Hexanes				318	312	6
Heptanes				330	322	8
Propyl iodides .				407	390	17
Propyl bromides .				372	353	19
Propyl chlorides.				330	317	13

C .- Normal Propyl and Allyl Compounds.

	Class	•	198		n-Propyl: η × 105.	Allyl: η × 105.	Δ.
Hydrocarbon	s .				318	304 406	2 × 7
Iodides .					407	406	I
Bromides .					372	371	I
Chlorides .			Maria.		330	371 328	2

D.-Alcohols at Slope 0.04987.

(The curves did not admit of comparison at slope 0.04323.)

	Al	cohol.			η × 10 ⁵ .	Δ for CH ₂ .
Methyl Ethyl .	:				650 606	- 44
n Propyl					560	-46
n-Butyl Iso-propyl					575 490	15
Iso-butyl					525	35
Amyl . Trimethyl	carb	inol			474 461	00
Dimethyl e	thyl	carbi	nol		490	29

The following conclusions may be drawn from the above viscosities at temperatures of equal slope:—

- I. In homologous series there is an increase in viscosity corresponding with an increment of CH_2 , but the increase tends to diminish as the molecular weight increases. The alcohols, acids, and dichlorides are abnormal.
- 2. In general, increase of molecular weight causes an increase in viscosity in such cases as the alkyl chlorides, bromides, and iodides.
- 3. n-Propyl compounds have a higher viscosity than the allyl derivatives.
- 4. Iso compounds are less viscous than the corresponding normal substances.

Just as molecular viscosities at the boiling-point were found to show more regularities than the simple viscosity coefficients, so at temperatures of equal slope it is better to compare derived functions of viscosity.

TABLE IX.-Molecular Viscosities at Slope 0.04323.

Substance.			$\eta \ (Mv)^{\frac{2}{3}} \times 10^{4}.$	ΔCH ₂ .
Pentane			687	121
Hexane			818	131
Heptane			931	113
Octane			1035	104
Iso-pentane			663	136
Iso-hexane			799	
Iso-heptane			908	109
Isoprene			620	108
Methyl iodide.			728 638	
Ethyl iodide		1	778	140
n-Propyl iodide .			903	125
Iso-propyl iodide .			878	
Iso-butyl iodide .	1		1010	132
Ethyl bromide .			663	
n-Propyl bromide .			774	III
Iso-propyl bromide.		8 1	750	
Iso-butyl bromide .			877	127
Ethylene bromide .			973	05
Propylene bromide.			1068	95
Iso-propyl chloride.			644	116
Iso-butyl chloride .			760	110
Methylene chloride.			600	137
Ethylene chloride .			737	*3/
Methyl sulphide .	8.		578	2 × 117
Ethyl sulphide .			812	/
Acetone			572	2 X 107
Diethyl ketone .			785	
Methyl ethyl ketone			671	125
Methyl propyl ketone Formic acid			796	
Acetic acid			456	137
Propionic acid	1		593	
Butyric acid .			742 842	149
Acetic anhydride .			838	
Propionic anhydride		W.V	1037	2 × 99
Benzene			688	
l'oluene	N. M		821	133
Ethyl benzene .	100		939	0
-Xylene	1		954	118
n-Xylene			939	
-Xylene			923	

Careful examination of the above results shows that the increment of CH_2 in an homologous series causes a much more regular increase in viscosity than was apparent in the previous cases. With the exception of the fatty acids, some of the earlier members of the various series and the aromatic hydrocarbons, the differences are not very widely remote from the mean. It is unfortunate that Thorpe and Rodger did not investigate a few more members of each series so that this erratic behaviour of the

first one or two homologues could have been ignored. As it is the mean value for CH₂ comes to 120.

The value for the atomic viscosity of hydrogen can be obtained by subtracting the value for nCH_2 from that of C_n H_{2n+2} , where n is the number of carbon atoms in the molecule.

TABLE X .- ATOMIC VISCOSITY VALUE OF HYDROGEN.

Series.	n.	$\eta \ (Mv) \ \frac{2}{3} \times 10^4 \ \text{for} \ C_nH_{2n+2}.$	nCH ₂ Calc.	2H.
Paraffins	5 6	687 818	600 720	87 98
Paramns	7 8	931	840 960	91 75

Knowing the values for CH₂ and for H it is obviously possible to calculate those for the halogens in CH₃I, C₂H₅ Br, etc., for the oxygen in alcohols, carbonyl oxygen in ketones and so forth,

TABLE XI.—ATOMIC VISCOSITY VALUE OF CARBONYL OXYGEN.

Substance.		$\eta \ (Mv)^{\frac{2}{3}} \times 10^4$.	nCH ₂ .	=0.
Acetone Acetaldehyde Methyl ethyl ketone Methyl propyl ketone Diethyl ketone		572 448 671 796 785	360 240 480 600 600	212 208 191 196 185 mean 198

Similarly the other chief constants have been obtained.

TABLE XII.—VISCOSITY CONSTANTS AT TEMPERATURES OF EQUAL SLOPE.

	Slope 0'04323.	Slope 0.04987.
Hydrogen	44.5	86
Carbon	31.0	60
Sulphur	246	474
Hydroxyl oxygen	166	320
Carbonyl oxygen	198	382
Ether oxygen	58	II2
Chlorine in monochlorides	256	494
Chlorine in dichlorides .	244	470
Bromine in monobromides	372	717
Iodine in monoiodides .	499	962
Ring formation	244	465
Double bond	48	92
Iso union	-21	-40

It was now possible to calculate molecular values with a fair degree of accuracy from these atomic constants, and Thorpe and Rodger found that in the case of fifty compounds the average divergence of the observed from the calculated value was only about I per cent.

TABLE XIII.-Molecular Viscosities (x 104) AT Slope 0.04323

Substan	ce.		Observed.	Calculated.	Difference per cent.
Pentane			687	68g	- 0'3
Hexane			818	809	1.1
Heptane .			931	929	0.5
Methyl iodide			638	664	- 4.0
Ethyl iodide .			778	784	- o.8
Methyl sulphide			578	575	0.2
Diethyl ketone			785	798	- 1.6
Acetaldehyde .			448	438	2.2
Acetic acid .			593	604	- r·8
Ethyl ether .			635	627	1.3
Toluene			821	814	0.8
Formic acid .			456	484	- 6·I
Propionic acid			742	724	2.4
n-Butyric acid			842	844	- 0.3
Methyl alcohol			358	375	4.7
Water			206	255	- 23.8

TABLE XIV .- VALUES FOR MOLECULAR VISCOSITY WORK AT EQUAL SLOPE.

Substance.	$\eta \text{ (Mv)} \times 10^3.$	Δ CH ₂ .
Pentane	. 329	96
Hexane	. 415	86
Heptane	495	80
Octane	574	79
Iso-pentane	320	0
Iso-hexane	405	85
Iso-heptane	. 482	77
Methyl iodide	255	0.5
Ethyl iodide	341	86
Propyl iodide	. 425	84
Ethyl bromide	282	
Propyl bromide	. 353	71
Methyl sulphide	240	
Ethyl sulphide	. 393	2 × 77
Acetone	238	
Diethyl ketone	376	2 × 69
Methyl ethyl ketone	302	
Methyl propyl ketone	383	8r
Formic acid.	160	
Acetic acid	237	77
Propionic acid	325	87
<i>n</i> -Butyric acid	397	73
Benzene	314	
Toluene	396	82
Ethyl benzene	475	79

It is again noteworthy that the earlier members of a series and also associated compounds show considerable abnormality.

When the data are calculated in terms of molecular viscosity work, η (Mv), the relationships which are obtained closely resemble those for molecular viscosity.

The average value for CH2 was found to be 80. In the same way as before, atomic constants were calculated, and the following values obtained.

TABLE XV .- ATOMIC VISCOSITY WORK CONSTANTS.

		At Slope o'04323.	At Slope 0.04987.
Hydrogen		- 34	- 64
Carbon		148	278
Hydroxyl oxygen		100	188
Ethereal oxygen		43	73
Carbonyl oxygen		- 19	- 36
Sulphur		144	271
Chlorine in monochlorides.		89	167
Chlorine in dichlorides .		82	154
Bromine in monobromides		151	284
		218	410
Ring formation		- 369	- 694
Double bond		- 95	- 179
Iso-union		- 8	- 15

The above summary of Thorpe and Rodger's work shows that additive relationships become apparent only when proper temperatures are selected for making the comparison. Even then, compounds which are notably associated or in the case of which constitutional influences are at work, fall outside the scope of the additive law. The effects of the homologous increment, for example, gradually diminish as a series is ascended and when atoms or groups possessing much residual affinity are present in a molecule, considerable departure from the calculated values is observed. This is clearly shown in the case of the chlorine substitution derivatives of methane (Table XVI.).

TABLE XVI.-EFFECT OF SUBSTITUTION.

Substance.		Mol. Viscosity × 104 at Slope 0.04323.	Δ for Cl.
Methyl chloride . Methylene chloride Chloroform Carbon tetrachloride	: :	420 (calc.) 600 747 854	180 147 107

Each additional chlorine atom produces less and less effect on the molecular viscosity.

Reference has already been made to the anomalous behaviour of the alcohols; this is more clearly seen from the following table.

TABLE XVII.—THE ALCOHOLS.

Substance.	η × 105.	Δ.	η (Mv) $\frac{2}{3}$ × 104.	Δ.	$\eta (Mv) \times 10^3$.	Δ.
Water	650 606 560 575 490 525 461 490	 - 44 - 46 15 35	390 760 933 1041 1232 930 1137 1020 1190	362 173 108 191 107	105 260 367 449 570 405 529 480 527	155 107 82 121 124 47

It is obvious that there is no constant difference for the homologous increment in these cases and it is quite impossible to calculate a molecular value from the atomic constants previously given. This is by no means surprising, for the peculiar influence of association on viscosity has been pointed out by almost every worker in the subject. At the same time it must be emphasized again that Thorpe and Rodger had not sufficient material at their disposal to deal adequately with the question of atomic constants, and it is only recently that an attempt has been made to investigate a large number of homologous series each containing as many members as can conveniently be prepared.

CHAPTER IV.

THE MEASUREMENT OF THE VISCOSITY OF PURE LIQUIDS SUBSEQUENT TO 1895.

Temperature Co-efficients of Viscosity.—Although the work of Thorpe and Rodger did not go far enough to afford a final method of representing viscosity data, little has been done up to the present time to put the matter on a satisfactory basis. Brief reference may be made to the work of Mühlenbein ⁴² and of Wagner and Mühlenbein ⁴³ which confirmed previous work on the effects of substitution in the benzene series; to that of Beck ⁴⁴ who found that in homologous series there is a proportionality between viscosity and molecular weight, that *iso*-compounds are less viscous than normal isomers, and that the abnormally high viscosities of the alcohols and acids depends on the presence of potentially quadrivalent oxygen.

The next real advance is due to Bingham ⁴⁵ who pointed out that relationships of a more general and regular character are obtained when one considers not the viscosity but the reciprocal of the viscosity, or the *fluidity*. To the work of Bingham somewhat fuller consideration must be given.

Formulæ for Fluidity.—The fact that the mixture law is so rarely observed led Bingham to make use of the reciprocal of viscosity, or fluidity. Using the data of Thorpe and Rodger, Linebarger, and Dunstan, he found that mixtures which gave a sagged viscosity curve yielded a linear fluidity curve and naturally deduced that fluidity was an additive property. On applying this deduction to pure liquids it was found that the fluidity of unassociated compounds was very markedly proportional to the absolute temperature, and even for associated liquids this proportionality was found to hold good at sufficiently high temperatures.

The simple linear formula, however, was replaced by the expression

$$\phi = \frac{c}{\phi - at - b} \quad . \tag{Ia}$$

containing three constants a, b and c, and again by

$$t = A\phi - \frac{B}{\phi} + C \quad . \quad (1b)$$

These formulæ are really equivalent when

$$A = \frac{I}{a}$$
, $B = \frac{c}{a}$ and $C = -\frac{b}{a}$.

The above fluidity formulæ proved capable of representing a considerable amount of viscosity data with an ease and accuracy unequalled by any formula hitherto proposed, but an expression containing four constants was found necessary to reproduce the experimental values for water and hydroxylated compounds in general, viz.:—

$$t = A\phi + \frac{B}{\phi + D} + C \qquad . \tag{2}$$

Since a relationship may be reasonably expected to exist between fluidity and volume, van der Waals's equation

$$T = \frac{pv}{R} - \frac{pb}{R} + \frac{a}{Rv} - \frac{ab}{Rv^2}$$

might be recast in terms of fluidity as

$$T = a\phi - \beta + \frac{\gamma}{\phi} - \frac{\delta}{\phi^2}. \qquad (3)$$

where α , β , γ , δ are constants, and T represents the absolute temperature corresponding to a given fluidity ϕ . The applicability of this formula is shown by the following table.

TABLE XVIII.—FLUIDITY CALCULATED AND OBSERVED.

Absolute 7	emperature.	Percentage Difference
Observed.	Calculated.	Fercentage Difference
273°37°	273°37°	
274'13°	273'99°	- 0.02
276.09°	275°53°	-0.50
278.45°	277.62°	- 0.59
286.23°	285.80°	-0.25
303.42°	303°72°	-0.00
320.03°	320.28°	+0.07
373°00°	373.00°	_

^{*} The experimental data are from the work of Thorpe and Rodger.

B .- Octane.*

Absolute 7	l'emperature.	Possente se Differen		
Observed.	Calculated.	Percentage Difference		
273°25°	273°25°			
295'92°	295°33°	-0.10		
316.80°	316.88°	0.00		
339.46°	339°75°	0.08		
361.33°	361.25°	0.02		
395°07°	395°05°	_		

The above table shows that a formula based on van der Waals's equation reproduces fluidity data somewhat better than one which contains three constants only. It is, however, not so accurate as formula (2) which also contains four constants.

TABLE XIX .- AVERAGE PERCENTAGE DIFFERENCES FROM USING THE EQUATIONS (1b), (2), AND (3).

Substance.	2	Average per cent Difference from (1b).	Average per cent Difference from (2)	Average per cent Difference from (3)
Water		0°17 0°16 0°43 1°17	0.01 0.02 0.09	0.38 0.11 0.38

Additive Relations.—The additive nature of the relationships is seen more clearly when attempts are made to deduce atomic constants at some fixed value of the fluidity, where, according to Bingham, a truly "corresponding condition" is to be looked for.

In the following tables are given the values of the absolute temperature corresponding with a fluidity value of 300 in C.G.S. units ("Amer. Chem. Journ.," 1910, 43, 302). The absolute temperature values corresponding with the fluidity 200 have been calculated by Bingham and Miss Harrison ("Zeitschr. physikal. Chem.," 1909, 66, 1).

^{*} The experimental data are from the work of Thorpe and Rodger.

TABLE XX.—Temperature Value of a Methylene Grouping at Fluidity 300.*

Substance.	Absolute Temperature Observed.	Temperature Value CH ₂ .
Pentane	258·2°	30°5°
Hexane	288·7°	
Heptane	313.10	24'4°
Octane	337 ^{.8°}	24'7°
Iso-pentane	253.8°	29·2°
Iso-hexane	283.00	
Iso-heptane	305°3°	22'3°
Isoprene	. 250°0°	24'4°
Diallyl	. 274°4°	24 4
Methyl iodide	. 338°5°	20.80
Ethyl iodide	359°3°	200
Propyl iodide	382.5°	23°2°
Iso-propyl iodide	371.60	23°5°
Iso-butyl iodide	. 595°1°	43 3
Ethyl bromide	311.2°	29°0°
Propyl bromide	340°5°	29'0
Iso-propyl bromide	. 331.0°	26.80
Iso-butyl bromide	357.8°	20'0
Iso-propyl chloride	. 289.8°	20170
Iso-butyl chloride	. 322°5°	32.7°
Methyl propyl ether	265°5°	0000
Ethyl propyl ether	. 288·4°	22.0°
Dipropyl ether	314.20	25.8°
Methyl iso-butyl ether	. 284°9°	
Ethyl iso-butyl ether	304°5°	19.60

The mean value of a methylene grouping is 25.4°.

TABLE XXI.—TEMPERATURE VALUE OF THE ISO GROUPING AT FLUIDITY 300.

Substance.		Normal.	Iso.	Temperature Value		
Pentane Hexane Propyl iodide Propyl bromide Propyl chloride Butyric acid Methyl butyrate				258·2° 288·7° 313·1° 382·5° 340·5° 298·7° 426·5° 343·2°	253.8° 283.0° 305.3° 371.6° 331.0° 289.8° 416.7° 334.2°	4·4° 5·7° 7·8° 10·9° 9·5° 8·9° 9·8° 9·0°

The temperature of equal fluidity is, on the average, 8.2° lower in the case of the *iso*- than in the case of the normal compound.

^{*} The experimental data are from the work of Thorpe and Rodger.

TABLE XXII.-TEMPERATURE VALUE FOR HYDROGEN AT FLUIDITY 300.

Substance. n.		n.	Temperature Observed.	Calc. for $n \times CH_2$.	Temperature Value of 2H.	
Pentane	-		5	258·2° 288·7°	127.0°	131·2°
Hexane	70.		6		152.4°	136·3°
Heptane			7	313.1°	177.8° 203.2°	135°3°
Octane			8	337.8°	203.50	134.60

Hence the mean value for H is 67.8, and for carbon, 110.2.

TABLE XXIII .- TEMPERATURE VALUE FOR DOUBLE BOND AT FLUIDITY 300.

De	eriva	tive Su	bstand	e.	Normal Propyl.	Allyl.	Temperature Value of Double Bond.
Iodide					382·5° 340·5° 298·7°	380.2°	2.0°
Bromide Chloride				•	340.2	336.2°	4°0° 6.8°

Hence the temperatures of equal fluidity are, in the case of an allyl compound, about 4'3° lower than for the normal propyl compound, which contains two atoms of hydrogen more. Hence a double bond has a temperature value of 131.3°.

TABLE XXIV.—TEMPERATURE VALUE OF OXYGEN AT FLUIDITY 300.

		Absolute	Temperature.	T
Substance.		Observed.	Calculated for $C_nH_{2n} + 2$.	Temperature Value of - O
Methyl propyl ether . Ethyl propyl ether . Dipropyl ether . Methyl iso-butyl ether Ethyl iso-butyl ether	: : :	265'5° 288'4° 314'2° 284'9° 304'5°	237·2° 262·6° 288·0° 254·4° 279·8°	28·3° 25·8° 26·2° 30·5° 24·7°

The average value for $-0 - is 27^{\circ}1^{\circ}$.

TABLE XXV .- VALUE OF THE BENZENE RING AT FLUIDITY 300.

			Absolute	Temperature.	Towns and town Males
Substance.			Observed.	Calculated for $C_nH_{2n} - 6$.	Temperature Value of the Benzene Ring.
Benzene			348·1° 347·5°	139.5°	208·6° 182·6°
Ethyl benzene			362·1° 377·5°	190.30	171.8° 176.2°
m-Xylene.			355.5° 357.4°	190.30	164.3°

The minimum value 164.2 was chosen to avoid error due to "unlike association".

In Table XXVI are grouped together for comparison the constants at fluidities 200 and 300.

TABLE XXVI.—CONSTANTS AT FLUIDITY 200 AND FLUIDITY 300.

Atom or Grouping.							$\phi = 200.$	$\phi = 300.$
Carbon Hydrogen Oxygen Iso-union							- 95·7 59·2 24·2 - 7·6	-110°2 67°8 27°1 - 8°2
Double bon Benzene rin							114.4	131.3

A few illustrations may be given showing the application of these values in constructing a "molecular temperature" at fluidity 300.

Where, however, a compound is associated, Bingham considers that the ratio between the observed and calculated results (as in the case of methyl butyrate) should give the association factor since the value of the constants are additive. Such a list of association factors is given in Table XXVII.

TABLE XXVII.—Association Factors at Fluidity 300.

		Absolute T	emperature.	Acceptation Produce
Substance.		Observed.	Calculated.	Association Factor.
Water		358·5°	162.7°	2°20
Dimethyl ketone .		289.5°	207'5°	1.23
Diethyl ketone		326.5°	285'4°	1'14
Diethyl ketone Methyl ethyl ketone .		315.6°	260.00	1.31
Methyl propyl ketone		330.0°	285'4°	1.17
Acetic acid		407'9°	236·3°	1.73
Propionic acid		408.2°	261.7°	1.22
n-Butyric acid		426.5°	287'I°	1'48
Acetic anhydride .		388·1°	300.00	1.25
Propionic anhydride.		402.40	360°7°	1.11
Benzene		348·1°	303°7°	1'14
Toluene		347.5°	329'I°	1.00
Ethyl benzene		362.1°	354°5°	1.03
Methyl alcohol Ethyl alcohol		336°9°	188.10	1.79
Ethyl alcohol		371.2°	213.2°	1.74
Methyl formate		297.5°	236·3°	1.36
Ethyl formate		311.40	261.7°	1.10
n-Propyl formate .		333'9°	287·1°	1.19
Methyl acetate		306.0°	261.7°	1.12
		320.8°	287·1°	1.15
n-Propyl acetate .		343°0°	312.2°	I.II

Interesting as is the above method of calculating the degree of association of a liquid, still too great stress ought not, in the present condition of our knowledge, to be laid on the actual values so obtained, for it must be borne in mind that the differences between the first and second columns are due not only to association but to want of sufficient data for calculating accurately the atomic "constants" and also to constitutional effects, such as mutual influence of groupings in the molecule, symmetry and so forth.

The Relation between Fluidity and Vapour Pressure.—The fact that the fluidities of saturated aliphatic hydrocarbons and of ethers are for each class practically identical at the boiling-points of the substances suggests the question whether this relation holds for other temperatures of equal vapour pressure. Using the vapour pressure determinations of Young and his collaborators and also independent observations of his own, Bingham (loc. cit.) finds that the fluidity-vapour pressure curves for ethers and hydrocarbons are entirely similar and indicate that at a sufficiently high temperature, fluidity is a linear function of the vapour pressure. It follows then for ethers and hydrocarbons that at temperatures corresponding to any given fluidity the vapour pressures of these compounds are equal, and hence when liquids of a given class have equal fluidities they are in a corresponding condition.

Viscosity Relations.—Although it is fairly evident that no method of obtaining additive relationships can expect to be trustworthy till a larger mass of data is available, yet attention was drawn by Dunstan and Thole 46 to the fact that a fairly exact

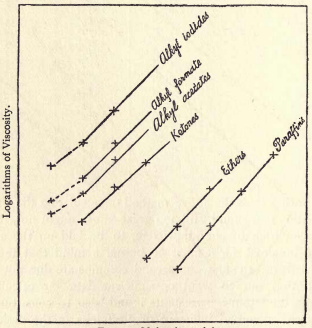


Fig. 3.-Molecular weights.

linear relationship exists between molecular weights and logarithms of viscosity in several homologous series. In one of these series, the methyl alkyl carbinols, all the eight members, ranging from *iso*-propyl alcohol to methyl-nonyl-carbinol were found to give log. viscosities which lie on a straight line. In the other series,

the iso-propyl alkyl carbinols, the agreement is scarcely so good, but a glance at the figure will show that a straight line adequately represents the relationship between molecular weight and the logarithm of the viscosity, in the case, at least, of the higher members of a series. A long series of dibasic esters from the

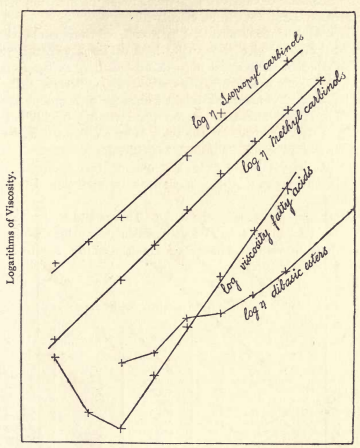


Fig. 4.-Molecular weight.

oxalic to the sebacic term shows little regularity until the first three members are passed, since conjugation effects are pronounced in these cases, but from glutaric ester onwards the linearity of the curve becomes marked. Gartenmeister ⁴⁷ measured the viscosities of the first nine members of the fatty acids at 20°, and on plotting log. viscosity against molecular

weight a linear curve is afforded from the third member onwards. Adding to these cases, all of which embody a homologous series of some length, the measurements of Thorpe and Rodger ²³ for the first four or five members of some ten homologous series, the same linear relationship is found to obtain and hence additive values of log. viscosity for the more important elements and radicles can be calculated.

It should be mentioned that hydroxylated compounds, which almost invariably give abnormally high values for viscosity, are not suitable for the purpose of such a calculation, as is apparent when the slope of the curves is considered. Whereas in Fig. 8 the log.-viscosity-molecular-weight curves for such groups as the paraffins, ethers, esters, and ketones are parallel, it will be seen in Fig. 4 that the curve for the acids is by no means parallel to that for the carbinols. Further, as with almost every physical property, the results for the first one or two members of a homologous series tend to be anomalous, and should not be included.

In the following tables the data of Thorpe and Rodger at 20° have been taken as a basis for the calculated values, while the logarithms of viscosity have been multiplied by 10⁵ to avoid barred characteristics.

TABLE	XXVIII.	-METHYL	ALKYL	CARBINOLS	AT	25°.
-------	---------	---------	-------	-----------	----	------

		Viscosity.	Log η × 105.	Δ CH ₂ .
Iso-propyl alcohol . Methyl n-propyl carbinol Methyl n-butyl carbinol Methyl n-amyl carbinol Methyl n-hexyl carbinol Methyl n-heytyl carbinol Methyl n-octyl carbinol Methyl n-nonyl carbinol Methyl n-nonyl carbinol		0°0206 0°0309 0°0398 0°0506 0°0655 0°0828 0°1010	3'3139 3'4901 3'6001 3'7037 3'8162 3'9180 4'0043 4'0892	0.0881 0.1100 0.1036 0.1125 0.1018 0.0863 0.0849

		Viscosity.	Log η × 105
Methyl iso-propyl carbinol		0.0321	3.2449
Ethyl iso-propyl carbinol.		0°0403	3.6050
n-Propyl iso-propyl carbinol		0.0474	3.6761
n-Butyl iso-propyl carbinol		0.0200	3.8505
n-Amyl iso-propyl carbinol		0.0729	3.8629
n-Hexyl iso-propyl carbinol		0.1018	4'0077
n-Octyl iso-propyl carbinol		0.1431	4°1557

A marked abnormality in the fourth member disturbs the regularity of this series and interferes with it as a means of obtaining CH₂ values.

TABLE XXX.-Esters of the Oxalic Series at 25°.

				Viscosity.	Log η × 105.	Δ CH ₂ ,
Diethyl	loxalate			0.01721	3*2433	
,,,	malonate			0.01872	3.2730	
"	succinate			0.02400	3.3802	
"	glutarate			0.02480	3'3945	
19	adipate	•		0.02768	3'4425	
,,	pimelate			0.03280	3.2160	0.0899
11	suberate			0.04036	3.6029	0.0033
"	sebacate			0.02023	3.7023	0 0994

This series shows anomaly up to the glutaric member.

TABLE XXXI.-FATTY ACIDS AT 20°.

1.334				12	Viscosity.	Log η × 105.	Δ CH ₂ .
Formic :	acid		-		0.01804	3.2562	a sufficient
Acetic	22				0.01535	3.0006	
Propionic		-			0'01107	3.0441	
Butyric	,,				0.01208	3.5036	0.1202
Valeric	22				0.02236	3'3495	0.1459
Hexoic	99				0.03501	3.2023	0.1528
Heptoic	22				0.04356	3.6301	0.1338
Octoic	22				0.05749	3.7596	0.1202
Nonoic	22				0.08310	3'9201	0.1602

TABLE XXXII .- THE VALUE FOR CH2.

Substance.	η.	Δ	$\log \eta \times 10^5$ for CH ₂ .
Hexane	. 0'00320		
Heptane	. 0'00411	0,100	0.100
Octane	. 0.00538	0'117	paraffins
Iso-hexane	. 0.00300	0.103	
Iso-heptane	. 0.00370		
Ethyl iodide	. 0.00583	0'102	0'102
Propyl ,,	. 0.00737		alkyl iodides
Iso-propyl iodide	. 0.00600	0.101	
Iso-butyl ,,	. 0.00870	ATT I I	
Methyl propyl ether	. 0.002515	- day -	
Ethyl " "	. 0.003175	0.101	
Propyl ,,	. 0'00420		
Ethyl ,,	. 0.002345	0.139	0.108
Propyl "	. 0.00420		ethers
Methyl propyl ,,	. 0.002515	O.III	
" iso-butyl "	. 0.003065		
Ethyl ,, ,,	. 0.003785	0.003	
Ethyl acetate	. 0.00449		
Propyl ,,	. 0.00281	0,115	0'107
Methyl propionate	. 0'00454		esters
" butyrate	0.00575	0.103)	
Acetone	. 0'003225	6	
Methyl propyl ketone	. 0.00201	0.000	0.106
" ethyl "	. 0.00423	12	ketones
Acetone	. 0.00201	0.114	
Propyl alcohol	. 0.0226	63	
Butyl ,,		0,119	0.116
Iso-butyl ,,		2	alcohols
Iso-amyl ,,	. 0.0200	0.112	
Methyl sulphide	. 0.00203	1	0.001
Ethyl ,,	. 0.00455	0.001	sulphides
	133		

Value at 20° of log $\eta \times 10^5$ for the Chief Groups.

Hydrogen.—Subtracting the CH_2 value from $\log \eta \times 10^5$ of the paraffins:—

```
From hexane . . . . 0.931 , heptane . . . 0.932 , 0.932 , octane . . . 0.938 Mean for H = 0.934.
```

Alcoholic Hydroxyl.—Subtracting the value for CH_2 and H from the log $\eta \times 10^5$ of the higher alcohols:—

```
From propyl alcohol . . . 2'097 Mean for OH = 2'102.
```

Ethereal Oxygen.—Subtracting the value for the alkyl radicles from the $\log \eta \times 10^5$ of the ethers:—

```
From methyl propyl ether . . . 0°105

,, ethyl ,, ,, . . . 0°099

,, propyl ether . . . 0°13

,, ethyl ,, . . . 0°074

Mean for O = 0°098.
```

Carbethoxyl.—Subtracting the value for the alkyl radicles from $\log \eta \times 10^5$ of the higher fatty esters:—

Carbon.—Subtracting the value for hydrogen from that of CH₂:—

$$C = -1.761.$$

Carbonyl (ketonic).—Subtracting the value of the alkyl radicles from $\log \eta \times 10^5$ of the ketone:—

```
From acetone . . . . 0'427

,, methyl ethyl ketone . . 0'437

,, propyl ,, . . 0'404

,, diethyl ketone . . 0'372

Mean for CO = 0'407.
```

Iso-Union.—Subtracting the value for the normal compounds from those of the *iso-*isomeride:—

```
From iso-hexane – hexane . . . – 0.028 , iso-heptane – heptane . . . – 0.035 , iso-propyl bromide – propyl bromide – 0.030 , iso-propyl iodide – propyl iodide. – 0.030 = – 0.030.
```

Double Bond.—Subtracting the value for the saturated compound from those of the unsaturated body and two H values:—

```
From allyl iodide + 2H - propyl iodide . 1.862

,, ,, bromide + 2H - propyl bromide 1.850

,, ,, chloride + 2H - propyl chloride 1.840

,, \frac{1}{2} [diallyl + 4H - hexane] . 1.835
```

Collected Values at 20°.

CH ₂		0.102	O (ethereal)			0.008
н		0.934	CO ₂ Et .		•	1.573
OH (alcoholic)		2.103	C			- 1.761
CO (ketonic) .		0.407	Iso-union			- 0.030
Double bond .		1.847				

Owing to the additive character of the logarithms of viscosity, it becomes possible to calculate, with fair accuracy, the viscosity of a compound from the values of the atomic and group constants just given, as the following examples indicate:—

(I) Ethyl Diethylacetoacetate. $\eta_{20}^{\circ} = 0.0344$ (Gartenmeister). CH₃·CO·CEt₂·CO₂Et:
6 CH₂ = 0.642
H = 0.934
CO = 0.407
CO₂Et = 1.573
3.556 = log of 0.0360 × 10⁵.

 $2.578 = \log \text{ of } 0.0378 \times 10^5$

- (3) Iso-Amylene. $\eta_{20}^{\circ} = 0.00212$ (Thorpe and Rodger) (CH₃)₂CH·CH: CH₂: 5CH₂ = 0.535 = 1.847 2.382 iso = -0.030 $2.352 = \log of 0.0225 \times 10^{5}$
- (4) Ethyl Acetoacetate. $\eta_{20}^{\circ} = 0.0168$ (Gartenmeister).

Ketonic.

$$CH_3 \cdot CO \cdot CH_2 \cdot CO_2Et$$
 $2CH_2 = 0^{\circ}214$
 $H = 0^{\circ}934$
 $CO = 0^{\circ}407$
 $CO_2Et = 1^{\circ}573$
 $CO_2Et = 1^{\circ}573$

Enolic.

$$CH_2 \cdot C(OH) : CH \cdot CO_2Et$$
 $2CH_2 = 0 \cdot 214$
 $OH = 2 \cdot 102$
 $CO_2Et = 1 \cdot 573$
 $= 1 \cdot 847$
 $C = -1 \cdot 761$
 $3 \cdot 975 = \log of 0 \cdot 0944$

Therefore, percentage of enolic form = 4.2.

CHAPTER V.

THE VISCOSITY OF LIQUID MIXTURES.

THE viscosity of binary liquid mixtures has been a subject of frequent investigation but up to the present no formula has been deduced which will satisfactorily reproduce the experimental facts. No linear formula can be utilized since in every curve there is a divergence, greater or less, from that demanded by the mixture law. In the majority of cases a sagging takes place in the viscosity concentration curve, and since it is hardly conceivable that two substances should be entirely indifferent towards each other, this, in general, slight departure from linearity may be accounted for by an equally slight mutual action. Further, it should be noted that the divergences become less with elevation of the temperature, a fact which again leads to the assumption that the exercise of some residual affinity leading to the formation of loosely aggregated complexes may be postulated.

Historical.—Brief reference may be first made to the earlier work in this direction.

Poiseuille,¹ who appears to have been the first investigator to attack the problem of the viscosity of liquid mixtures, found that the ethyl alcohol-water pair gave a maximum viscosity at about the composition represented by $C_2H_6O.3H_2O$. Graham ²⁴ discovered that aqueous solutions of many common acids and alcohols give similar maxima at points corresponding with simple molecular concentration, e.g. $H_2SO_4.H_2O$; $CH_3COOH.H_2O$; $HCl.12H_2O$.

Wijkander ⁴⁸ followed this up with an investigation of the viscosities of binary mixtures of acetic acid, water, aniline, benzene, ether chloroform, carbon disulphide and ethyl alcohol. Such of the above pairs as mixed without appreciable change gave more or less sagged curves, ether-chloroform and ether-carbon disulphide afforded inflected curves, but the aqueous solu-

tions of acetic acid showed, at various temperatures, a distinct drift in the position of the maximum point. This behaviour was observed also by Noack 49 in connexion with aqueous ethyl alcoholic solutions, by Traube 50 and by Pagliani and Battelli 51 for aqueous alcohols and acids. Linebarger 52 investigated binary mixtures of ethyl acetate, carbon bisulphide, carbon tetrachloride, chloroform, nitrobenzene, ethyl iodide, toluene, ethyl benzoate and ether. In nearly all these cases the observed viscosity was less than that calculated from the mixture law but for the liquid pairs benzene-toluene and toluene-carbon disulphide, a very slight sag indeed was perceptible. Thorpe and Rodger 53 extended their work on the viscosity of pure liquids to that of liquid mixtures, viz. benzene—carbon tetrachloride; methyl iodide-carbon disulphide and ether-chloroform. In none of these cases was the mixture law observed, the third pair giving an inflected curve, which suggests possible combination between the components. Varenne and Godefroy 54 found a series of discontinuities on the viscosity-concentration curve of ethyl alcohol-water and considered these as evidence for the existence of several hydrates in solution. Wagner and Mühlenbein 55 quoted several cases of the occurrence of maximum and minimum points. Benzonitrile, nitromethane, and allyl iso-thiocarbimide depress the viscosity of ethyl alcohol, whilst nitrobenzene lowers that of iso-butyl alcohol; o-nitrotoluene in ethyl alcohol gives an inflected curve and pnitrotoluene in the same solvent shows a minimum point. Dunstan ⁵⁶ investigated a considerable number of binary mixtures and concluded that viscosity concentration curves of binary mixtures could be classified as follows:-

Type I.—Those which obey approximately the mixture law, being concave to the axis of percentage composition; e.g. ethyl acetate-benzene.

Type 2.—Those which exhibit definite maxima, sometimes corresponding with points of simple molecular composition. Nearly all experimental work in this class has been done on mixtures between the compounds of which chemical action might well be postulated, and the occurrence of a maximum may be regarded as pointing unmistakably to the formation of molecular complexes of a more or less stable kind.

Type 3.—Those which display a minimum point. Seeing that the connexion between viscosity and molecular

volume is so close (cf. p. 73) these three types of mixture may be defined respectively as being composed of (1) mutually indifferent components, (2) components which undergo chemical combination to a greater or less extent, (3) associated components which bring about mutual dissociation.

Findlay ⁵⁷ determined several viscosity concentration curves at the boiling-points of the individual mixtures and emphasized the fact that no simple formula could be obtained which would express the connexion between viscosity and composition. By plotting values calculated from the fluidity formula

$$\frac{\mathrm{I}}{\eta} = \frac{v_1}{\eta_1} + \frac{v_2}{\eta_2}$$

where η_1 and η_2 refer to the viscosities of the constituents at the temperature of the experiment, and v_1 v_2 are the volumes contained in I c.cm. of the mixture, it was found in most cases that the observed values are smaller than those calculated by the mixture formula. The maximum deviations from the calculated values are regarded by Findlay as important, since they give an indication of the magnitude of the interaction, of whatever nature it may be, which produces divergence from the ordinary law of mixtures. A similar point of view has just been reached by Denison ⁶⁸. Bingham ⁵⁹ using the data of previous workers showed that mixtures of Type I gave nearly linear fluidity $\left(\frac{I}{n}\right)$

curves, and concluded therefrom that fluidities are additive.

Mixture Formulæ.—The failure of the linear formula

$$\eta = v_1 \eta_1 + v_2 \eta_2$$

where η_1 and η_2 are the viscosities of the two components and $v_1 \, v_2$ the percentage volume concentrations has already been indicated.

The logarithmic formula of Arrhenius 60

$$\eta = \eta_1^{\upsilon_1} \cdot \eta_2^{\upsilon_2}$$
 or $\log \eta = \upsilon_1 \log \eta_1 + \upsilon_2 \log \eta_2$

holds fairly well for mixtures of liquids when one component is present to the extent of 90 per cent or more, but is not valid throughout the whole range of mixtures. Lees ⁶¹, after an examination of the various formulæ, found that the fluidity formula,

$$\frac{\mathrm{I}}{\eta} = \frac{v_1}{\eta_1} + \frac{v_2}{\eta_2}$$

is also unsatisfactory and that exact agreement can only be obtained by the introduction of a supplementary constant depending on the particular mixture considered. Jones, Bingham and McMaster ⁶² considered that fluidity and not viscosity is the characteristic additive property, but the calculations of Lees show that this is not the case if the concentrations are expressed as volume percentages, nor is it when they are expressed as weight percentages as urged by Drucker and Kassel. ⁶³ Kendall ⁶⁴ in view of these failures proposes a modified Arrhenius formula

$$\log \eta = \eta_1 \log \eta_1 + \eta_2 \log \eta_2$$

where η_1 η_2 are the molecular percentage concentrations of the two components in the mixture. The following illustrations show the applicability of the formula.

TABLE XXXIII.—CARBON TETRACHLORIDE AND BENZENE, 60° C. (THORPE AND RODGER'S DATA).

Mols. of C ₆ H ₆ per cent.	Observed.	Calculated.	Percentage Difference.
0.00	0.00582		
36.10	0.00203	0.00204	0.0
60°42	0.00456	0.00457	0.0
80.41	0.00422	0'00423	0.0
100.00	0.00331		

TABLE XXXIV.—Ethyl Acetoacetate and Pyridine at 25°. (Dunstan and Stubbs' Data.)

Mols. of Ester per cent.	Observed.	Calculated.	Percentage Difference
0.00	0.008802		
11.14	0.009297	0'00934	0
22'30	0.009318	0.00992	0
38.76	0.01000	0.01082	0
59.88	0.01233	0.01512	- I
100.00	0.01208		

Kendall also tried a molecular fluidity formula, but it gave no results superior to those cited above. These facts militate against the view which recently has been persistently set forth, that fluidity and not viscosity is the essential additive property, and they show that neither fluidity nor viscosity is the true additive

property but the logarithms of these quantities (since obviously $\frac{1}{\eta}$ may be substituted for η without affecting the result).

Viscosity Formulæ for Solutions of Solids in Liquids.— The formulæ which have been suggested to express the viscosity of solutions are, in general, similar to those for binary mixtures of liquids and are similarly unsatisfactory (Kendall, loc. cit.). Thus the linear formula

 $\eta = (I + Ax) \eta_0$ where η_o is the viscosity of the pure solvent, x the volume of solute per unit volume of solution and A a constant, holds only for very dilute solutions and the same is true of

Arrhenius' logarithmic expression $\eta = A^z \eta_o$ or $\log \frac{\eta}{\eta_o} = x \log A$.

Green ⁶⁵ found in the case of sucrose solutions that none of the current formulæ was valid over a wide range of concentration and put forward the equation

 $\eta = A^{\stackrel{v}{\widehat{w}}} \eta_o$ where v is the volume of sucrose and w that of water in unit volume of solution. Kendall again suggested a logarithmic expression,

$$\log \frac{\eta}{\eta_o} = x \log A$$

which is the Arrhenius formula modified by substituting *molecular* concentration of solute in a fixed weight of solvent. Two illustrations of the application of this expression may be given.

TABLE XXXV.—The Octyl Hydrogen Phthalates (r, d and l) in Benzene at 25°. (Dunstan and Thole's Data.)

Mols. Solute in 100 Mols. Solvent.	Observed.	Calculated.
0*00	0.006030	
7. 1.805	0.006762	0.006787
7. 2.597	0.007180	0.007126
7. 3'363	0'007520	0.007520
r. 5°097	0'008432	0.008424
r. 5.228	0.008604	0.008498
r. 6.490	0.000100	0.000330
d. 1.010	0.006844	0.006832
d. 3°478	0'007575	0.007575
d. 3.642	0.002631	0.007657
d. 6.996	0.009225	0.009244
l. 6.928	0.000418	0.000499

Mols. of Sucrose per 1000 σrm. of Water. A.

0* 0.008953 0.4382 0.013083 2.38

0.0666

1.618

2.440

3.216

5'440

TABLE XXXVI.—Sucrose in Water at 25° C. (Green's Data.)

0.03102

0.03802

0.07973

0'2072

1.028

2.42

2.44

2.45

2.44

2.41

The constancy of the value of A shows that the formula holds satisfactorily throughout the whole range of concentration even when the viscosity of the solution is more than one hundred times greater than that of water.

Quite recently Dolezalek and Schulze, ¹²⁷ investigating the system ether-chloroform, have shown that the equilibrium $C_4H_{10}O + CHCl_3 = C_4H_{10}O$. CHCl₃ is established and that the percentage amount of the compound can be calculated by the application of the law of mass action.

The viscosities of the various mixtures of ether and chloroform which vary very considerably from those calculated by the mixture law are shown to agree excellently with those calculated on the supposition that each mixture contains an amount of the compound determined by the above considerations.

Viscosity Maxima and their Interpretation.—Having dealt with the viscosity concentration curves which belong to Type I, it is now necessary to discuss the very important cases in which maxima appear.

Based on the view that associated liquids in general, and hydroxylated liquids in particular have a relatively high viscosity coefficient, the conclusion may be drawn that a maximum point in a viscosity-concentration curve means further association proceeding in the direction of complex formation, whilst the existence of a minimum point tends to the opposite view, namely, that some dissociation has resulted.

The simple view taken by Graham ²⁴ that the maximum point definitely indicated the existence of a compound, and, moreover, established its composition, has been generally criticized as being far too explicit. It seems probable, however, that in some cases

this view is sound, as will be seen later. Where, however, the amount of chemical affinity between the components is not sufficient entirely to bring about combination, the position of the maximum will vary within limits depending on temperature, the viscosity of the constituent liquids, and the degree of dissociation of the compound.

Washburn, 66 in a valuable summary of the theory of hydration, dealing with the deviations of property-composition curves from those demanded by the law of mixtures, agrees that striking irregularities exist, which may be quite invariant with temperature, such as the expansion-coefficient of aqueous solutions of nitric acid and sulphuric acid, which clearly indicate the existence of the hydrates HNO₃, H₂O; H₂SO₄, H₂O. Such irregularities are not found for mixtures the components of which show no chemical affinity for each other. However, Washburn sums up his review by stating that the deviations of a physical property from the mixture law cannot give any conclusive evidence regarding the existence of a hydrate (or complex). This statement is directly contrary to the views which have been held by a great number of investigators in this field (see Trans. Chem. Soc., 1909, 95, 1556).

Tsakalotos 67 has investigated maximum points on viscosityconcentration curves, and holds the opinion that the study of the viscosity curves in conjunction with the freezing-point curves proves undoubtedly that the viscosity maxima must be attributed to molecular compounds in a state of partial dissociation. The discrepancy between the compositions of the mixtures of maximum viscosity and maximum freezing-point for aniline-m-cresol, p-toluidine-m-cresol must be attributed to the difference existing between the viscosity of the two components and the partial dissociation of the molecular aggregate. He imagines the mixture as containing two different components, namely, (1) the inactive mixture of the original constituents, and (2) the complex anilinephenol. Mixture (1) gives a linear mixture-law curve. The effect of (2) alone would be a maximum at 50 mols. per cent. The resultant of these curves has a maximum displaced towards the component of greater viscosity.

Faust ³⁰ points out that it is very probable, although not absolutely proved, that in those cases where a maximum viscosity occurs a chemical compound is produced. Thus there is always

such a maximum when the components are an acid and a base. In all cases where there is an evolution of heat on mixing, the compounds will dissociate with rising temperature, and so the viscosity curve will reproduce the dissociation by showing a smaller and smaller deviation, and at high temperatures will approach linearity. It is, however, untrustworthy to deduce from the position of the maximum the chemical composition of the compound, since the position frequently alters with temperature, and depends on the relative viscosities of the two components. On the other hand, when the viscosity curve is sagged, it must be interpreted as a result of dissociation (compare Dunstan ⁵⁶). Faust also noticed that maxima on viscosity curves correspond with minima on vapour-pressure curves, a fact which is compatible with the formation of a molecular compound.

Denison 58 considers that the formation of a maximum does imply molecular aggregation, but prefers to determine the nature of the complex by the maximum deviation from the mixture law rather than from the position of the maximum itself. It should be mentioned that other observers, notably Findlay 57 have emphasized the fact that the greatest deviation should be considered, and not the actual maximum. Denison assumes that where complex formation is possible, there will be a solution of the complex in a mixture of the unaltered components. This mixture of complex and unchanged components will tend to form a maximum at some point along the curve, depending on its relative viscosity. It follows as a corollary that if most or all of the components associate in this way, the maximum point must agree with the composition of the complex. As Baker 68 has indicated, however, the complex may be of less viscosity than one or other of the components (assumed to be very associated), in which case a sagged curve or a minimum might be obtained. Baker, in fact, draws the conclusion that in mixtures of alcohol-ether, used for dissolving nitrocellulose, there is a complex existing, but the shape of the curve does not in any way indicate it.

On surveying these various opinions, it will be evident that they conform to the belief that a maximum viscosity does indicate some molecular aggregation, but the mass of evidence is decidedly against any predication that the position of the maximum point may indicate the composition of the compound. It will be obvious that this position can be influenced by a variety of causes. Suppose two liquids, A and B, the association factors of which are n and m, tend on mixing to form the compound AB. Then

$$A_n + B_m \rightleftharpoons (n-x) A + (m-x) B + xAB$$
,

where x is the amount of AB present.

Four cases present themselves:-

Case 1.—x is very small, and A_n and B_m are stable, and have little mutual action. A linear or slightly sagged curve will result; for example, phenol—a-naphthylamine.

Case 2.—x is fairly large, there is considerable chemical affinity, and the complexes A_n and B_m are mutually fairly stable. Then if the viscosity of the complex AB is less than that of one or other of the components, there will be a sagged curve; for example, phenol-acetone, lactic acid-water.

Case 3.—x is fairly large, there is considerable chemical affinity, and the viscosity of the complex is greater than that of A_n and B_m , which are mutually somewhat unstable. Then there will be a maximum point, but its position will depend on temperature and the relative viscosities of A_n , B_m , and AB; for example, alcohol-water.

Case 4.—x is very large, and chemical action very considerable; the complexes A_n and B_m are dissociated. Then there will be a definite maximum corresponding with the actual molecular concentration of A and B in AB; for example, sulphuric acid-water.

It has been found that in the large majority of cases a maximum in the melting-point curve means a maximum in the viscosity curve as will be evident from the following table.

TABLE XXXVII.—LIQUID MIXTURES WHICH SHOW MAXIMUM POINTS.

Mixture.	Maximum.	Observer.	Fusion Curve.
Water – methyl alcohol	Approximately	Getman, Dunstan	
Water - ethyl alcohol	MeOH, 3H ₂ O Approximately	Poiseuille, Graham,	
	EtOH, 3H2O	Dunstan, Traube	
Water - propyl	Approximately	Dunstan	
alcohol Water – acetic acid	PrOH, 2H ₂ O CH ₃ ·CO ₂ H, H ₂ O	Graham Dungtan	No maximum, Kre-
water - acetic acid		Wijkander	mann, "Sitzungs- ber. K. Akad. Wien," 1907, 116,
			795
Water - propionic	Approximately	Tsakalotos	
acid Water – pyridine	C ₂ H ₅ ·CO ₂ H, H ₂ O Approximately	Hartley, Thomas and	
water - pyriame	2C ₅ H ₅ N, 5H ₂ O		THE RESERVE
	Approximately 30	Dunstan and Faust	
	mols. per cent		
Water minharia	pyridine	Graham, Knietsch,70	M-
Water - sulphuric acid	H_2SO_4 , H_2O	Dunstan, Kremann	
		and Ehrlich	Knietsch, Kremann
Sulphuric acid - sul-	H ₂ SO ₄ , SO ₃	Dunstan and Wilson	Knietsch, maximum
phur trioxide	A	Tanlalatas	M
Aniline - m-cresol	Approximately 2 m- cresol: 1 aniline	1 sakalotos	Maximum, Kremann, "Sitzungsber. K.
	0.0001.1.4.4.4.4.4.4		Akad. Wien," 1905,
			112, 1201.
o-Toluidine - m-	Approximately 4 m-		Kremann, loc. cit.
cresol Acetone – chloroform	Approximately	Tsakalotos	
rectone - emototorm	chloroform : I	2 Sakaiotos	
	acetone at o°		
The selection of the se	and 3 chloroform	Faust	
THE RESERVE N	: 1 acetone at 13° No maximum at 19°	Fanct	ile il oropelaliti
Pyridine - acetic acid	Approximately 4	Tsakalotos	
	acetic acid: 1		
	pyridine	E.	
di-el autura (in	Approximately 4 acetic acid: 1		
THE RESERVE OF THE PERSON NAMED IN COLUMN TWO IS NOT THE PERSON NAMED IN COLUMN TO THE PERSON NA	pyridine acid. I		The same of the same of
	Very slight shift of	Faust	
	maximum be-		
Pyridine - butyric	tween o° and 99° Approximately 4	Tsakalotos	
acid butylic	butyric acid: 1		
Aniline - acetic acid	Approximately 2	Faust, Thole, Dun-	Carlo Later In
	acetic acid: I		
Acetic anhydride -	aniline at 18°—60° Approximately 2	Faust	
water	water: I anhy-		
	dride, invariant		
	between oo and		
Iso-butyric acid -	73°	Tsakalotos	
water	Approximately 3 water: 1 acid	1 Sanaiutus	
		MALES METERS	

TABLE XXXVII .- LIQUID MIXTURES WHICH SHOW MAXIMUM POINTS-(Cont.).

Mixture.	Maximum.	Observer.	Fusion Curve.
n-Butyric acid –	Approximately 2 water: 1 acid	Tsakalotos	
Triethylamine - water		Tsakalotos	
Nicotine-water		Tsakalotos	
Allyl thiocarbimide – piperidine	Exactly 1 amine: 1 thiocarbimide	Kurnakov and Schemtschushni 71	
Phenyl thiocarbimide - diethylamine	Exactly 1 amine: 1 thiocarbimide	"	
Allyl thiocarbimide – methylaniline	Exactly 1 amine: 1 thiocarbimide	"	
Ethyl thiocarbimide – piperidine		"	
Aniline - o-chloro- phenol	Nearly I amine: I phenol	Thole, Dunstan and Mussell	
Aniline - m-chloro- phenol	Approximately 1 amine: 1 phenol	,,	
Aniline - p-chloro- phenol	20 per cent aniline	37	
Phenyl hydrazine - phenol	i base: i phenol exactly	,,	Ciusa and Bernardi "Gazzetta," 49,
			I, 159, 1910, maximum at 1 base: 1 phenol
Phenyl hydrazine – o-chlorophenol	r base : r phenol exactly	,,	
	I: I nearly	Kurnakov and Efremov 126	
Chloral – alcohol	I: I nearly	"	
p-Cresol – aniline	Indefinite	Thole, Dunstan and Mussell	Maximum at 1 amine: 1 phenol, Philip, "Trans., Chem. Soc.," 1903,183, 814
Phenol - p-toluidine	Approximately I amine: I phenol	,,	,, 1903,103, 814
α-Naphthylamine –	Indefinite	,,	,,
Aniline – phenol	Indefinite	'n	Maximum at 1 amine : 1 phenol, Schreine- makers, "Zeits. Physikal. Chem.," 1899, 31, 581

It will be noticed from the above table that in some cases the evidence of the fusion curves is closely paralleled by that of the viscosity curves, for example, for sulphuric acid-water, o-chlorophenol-aniline, phenol-phenylhydrazine, aniline-phenol, o-chlorophenol-phenylhydrazine. In the majority of maxima, however, there is a distinct shift from the point of equimolecular concentration. One must therefore incline to the view that the appear-

ance of a maximum on a viscosity concentration curve does indicate the existence of chemical combination between the components, but the position depends on several independent factors, namely, the actual viscosity of the components (which is intimately connected with their degree of association), and the degree of dissociation of the compound itself at the temperature of observation. It by no means follows that the absence of a maximum point precludes complex formation, as, indeed, has

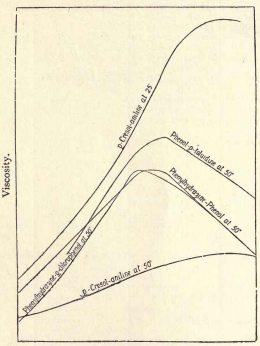


Fig. 5.—Percentage Composition.

been pointed out repeatedly; for obviously the viscosity of one highly associated component may be greater than that of the complex (see also Baker, loc. cit.). Comparison of the viscosity with the fusion curves and with the vapour pressure curves is valuable, for, as Faust has indicated, there is a close connexion between vapour pressure and viscosity, since whatever forces operate in complex building would similarly depress the vapour pressure.

A few curves where viscosity maxima are to be observed are shown in Fig. 5.

A very striking confirmation of the validity of the above views has been obtained by Kurnakov and Schemtschushni.⁷¹ These authors have investigated at various temperatures the viscosity concentration curves of irreversible binary systems where liquid and completely miscible compounds are formed. The viscosity isotherms consist of two separate branches convex to the composition axis intersecting at an angle (increasing in acuteness as the temperature is lowered), and this maximum point occurs at exactly equimolecular concentration.

CHAPTER VI.

VISCOSITY OF ELECTROLYTIC SOLUTIONS.

THE viscosity of salt solutions naturally attracted the attention of the earlier workers, and so far back as 1873 Hübner ⁷⁸ determined the viscosities of a series of solutions of alkaline haloids of equal density and found that some of the salts diminished the viscosity of water, whilst Sprung ⁷⁴ discovered that his experimental material fell into two classes:—

(1) KCl, KBr, KI, KNO $_3$, KClO $_3$, NH $_4$ Cl, NH $_4$ Br, NH $_4$ NO $_3$; (2) K $_2$ SO $_4$, NaCl, NaBr, NaI, NaNO $_3$, NaClO, Na $_2$ SO $_4$, (NH $_4$) $_2$ SO $_4$, BaCl $_2$, SrCl $_2$, CaCl $_2$, LiCl, MgSO $_4$.

At low temperatures the salts in the first group depress the viscosity of water and at high temperatures they increase it, whilst salts of the second group always increase the viscosity of the solvent. Slotte, Arrhenius, Wagner, Ranken and Taylor Tand Getman kneed also experimented in this direction. The last mentioned considers that the negative viscosity shown by salts of the first class is due to the specific effect of the potassium and ammonium ion, since the undissociated molecule (and the anions) bring about an increase in viscosity. This view is shared by Jones and Veazey who state that cations of large ionic volume diminish the viscosity of water.

It is obvious that one must consider not only the amounts of the ions and the undissociated salt present but also the degree of hydration of each of these components and further the effect of the salt and of its ions on the molecular complexity of the solvent. Further it must be remembered that all these quantities vary with temperature and concentration. It is not surprising, therefore, that little progress has been made in the direction of finding a quantitative expression which will represent the viscosity of a salt solution.

Viscosity and Conductivity.—As early as 1856 Wiedemann 80 drew attention to the fact that viscosity and conductivity are

closely related and showed that $\frac{K\eta}{m} = \text{constant}$, where K is the conductivity of the solution whose concentration is m, and similar empirical relationships have been pointed out by Grotian, ⁸¹ Grossmann, ⁸² Arrhenius ⁸³ and Kohlrausch. ⁸⁴

To illustrate the nature of this connexion the following results obtained by Getman 78 may be quoted.

TABLE XXXVIII .- EQUIVALENT CONDUCTIVITY AND VISCOSITY AT 18°.

Salt.	η.	Λ.	η Δ.
NaCl	0.0119	74.4	*863
NaBr	0.0112	70'5	.811
NaNO,	0.0114	66.0	.752
KC1	0.0102	98.2	1,031
KBr	0.0103	103'7	1.060
KI	0.0101	103.4	1.041
KNO,	0.0104	80.4	.839
NH Cl	0 0103	97.0	.999
NH ₄ Br	0.0101	102.0	1.030
NHI	0.0008	104.0	1.019
NHANO,	0.0101	88.0	.901

It is evident that some other factor besides those given is required to establish the relation between viscosity and conductivity.

It has been shown by Bousfield 85 that the application of Stokes' theory respecting the limiting velocity of a small sphere moving under a constant force in a viscous medium is of great value in studying the various causes which may affect the mobility of an ion. The limiting velocity of a sphere of radius r, moving in a medium of viscosity η , under a constant force p is proportional to $\frac{p}{r \cdot \eta}$. Applying this to the case of an ion

moving with a spherical solvent atmosphere around it, the mobility of the ion will depend on the viscosity of the solution and on the size of the envelope.

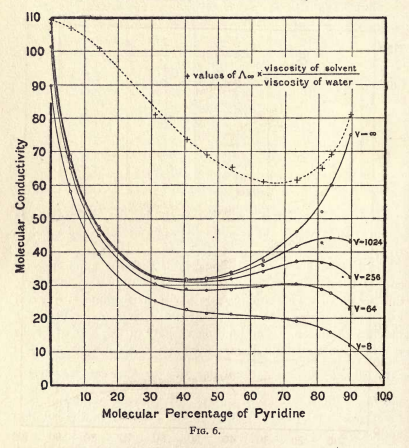
Hartley, Thomas and Applebey ⁸⁶ employed Stokes' theory in connexion with the conductivity of lithium nitrate dissolved in various aqueous pyridine solutions (Figs. 6 and 7). Assuming that U_A and U_C are the absolute velocities of anion and cation at infinite dilution under unit gradient, r_a and r_c the radii of the ions

$$U_A = C \frac{I}{r^a \eta}$$
 and $U_C = C \frac{I}{r^c \eta}$

and since $\Lambda_{\infty} = Q (U_A + U_C)$, Q being the charge liberated by I gram equivalent of a univalent ion,

$$\Lambda_{\infty} = \frac{Qc}{\eta} \left(\frac{I}{r_a} + \frac{I}{r_o} \right).$$

It follows then that any variation in $\Lambda_{\infty} \eta$ indicates a



change in the size of the solvent envelope attached to the ions. The following table shows that as the amount of pyridine is increased the ion atmospheres increase till the solution contains about 67 mols. per cent of pyridine and then steadily become less.

TABLE XXXIX.—Effect of Pyridine on Size of Solvent Atmosphere.

Mols. per cent Pyridine.	Λ_{∞} .	η.	Λ_{∞} . $\frac{\eta \text{ (Solvent).}}{\eta \text{ (Water).}}$
0,00	57*10	0.01748	57.1
14'47	19.05	0.04735	50.7
40.64	13'40	0'04990	37.6
54'35	16.25	0.03625	33.I
73.76	27.5	0.02022	32.0
83.98	39.0	0.01642	36.0

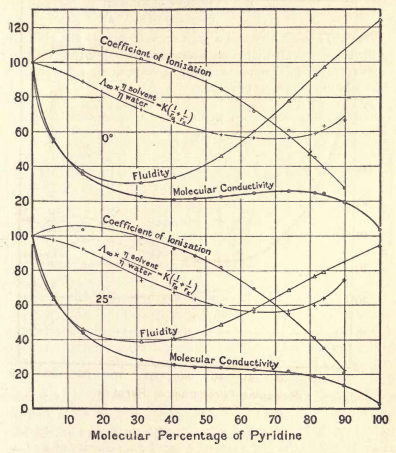


Fig. 7.

The ordinary equation used in calculating coefficients of ionization, namely, $a=\frac{A_v}{A_{\infty}}$

neglects changes in the mobilities of the ions at different concentrations due to changes in the viscosity of the solution and to variations in the size of the solvent atmosphere attached to them. Hartley and his co-workers point out (loc. cit.) that on the assumption that the mobility of the ion varies inversely with the viscosity of the medium (see Bousfield and Lowry ⁸⁷)

$$a = \frac{\Lambda_*}{\Lambda_{\infty}} \times \frac{\text{viscosity of solution}}{\text{viscosity of solvent}}$$

Heber Green,⁸⁸ however, using sucrose to increase the viscosity of the solvent, finds that Λ does not vary directly as the fluidity, but as some function more nearly approximating to ϕ [§] as is shown in the following table.

TABLE XL.—Conductivities of Mixtures of Sucrose and Lithium Extrapolated to the Ratio $\frac{LiCl}{Sucrose} = \frac{N}{n} = 0$.

Mols. of Sucrose N.	Λ × 103.	$\Lambda/\phi \times 10^3$.	$\Lambda/\phi^{0.7} \times 10^3$.
0	114.0		
0.02	108.8	1.012	4.135
0°2	95.8	1.025	4.000
0.6	68.7	1.122	3.862
1.0	46.6	1.508	3.802
1°4 1°8	29'5	1.288	3.812
1.8	16.8	2'099	3.020

Green expresses doubt not only as to the applicability of Stokes' theory in a case where the moving body is of the same order of dimensions as the particles of the medium but also with reference to the need for assuming the existence of an aqueous envelope around each ion. He considers that the calculation of the ionization coefficient can be made in the following way.

Comparing the conductivity Λ of a pure lithium chloride solution with that Λ_f of a solution infinitely dilute with respect to lithium chloride but containing enough sucrose to give it the same fluidity as the first solution, then the coefficient of ionization will be

$$x = \frac{\Lambda}{\Lambda_f}$$

TABLE XLI.—Ionization Coefficients for Solutions of Mixtures of Lithium Chloride and Sucrose. Ratio $\frac{Mols.\ of\ LiCl}{Mols.\ of\ Sucrose} = 5$ °o.

f.	$\Lambda \times 10$	$x=\Lambda/\Lambda f.$
111.2		1,000
111.50	1.070	0.944
110.30	1.019	0'904
109'45	0.0000	0.885
89.13	0.8045	0.7795
44.84	0°3875	0.410
18.21	0.10102	0.649
5.131	0.0722	0.578

These figures for x are not to be regarded as true ionization factors since the ionic mobilities in solutions of the same fluidity will vary according as the viscosity is mainly due to lithium salt, sucrose, or other added substances (see also Washburn ⁹² who developed a similar formula $\Lambda = K\phi^m$, m being not far removed from unity).

This conclusion becomes additionally interesting from the fact that similar considerations apply to the relationship between the fluidity and ionic mobility of aqueous solutions when these are modified continuously by changing the temperature instead of by the addition of a foreign substance. From the work of Noyes and his colleagues, Johnston ¹²⁸ has deduced the following values for a series of ions from the relationship $\mu_{\infty} = K^n \phi^n$.

n	к. o·887	Na. 0.97	NH ₄ . 0.891	Ag. 0.949	⅓Ba. 0°986	½Ca. 1°008
log K	+0.024	-0.581	+0.045	-0.143	-0.313	-0.286
μ_{∞} for o°	40.4	26	40°2	32.9	33	30
	n	CI. o·88	NO ₃ . 0.807	C ₂ H ₃ O ₂ . 1.008	‡FeC ₆ N ₆ . 0°929	
	log K μ _∞ for o°	+0.074	+0.104	-0°455	+0°139	

It will be noticed that the more mobile of the ions are those that show the greatest deviation from the linear law, the index n being less than I in each case. The less mobile ions give an index very near to unity with an almost exact proportionality between mobility and fluidity.

The subject was developed still further by Applebey 89 who continued the work of Grüneisen and extended it considerably. Grüneisen had shown that the viscosity concentration curves for all dilute salt solutions exhibit a remarkable change of curvature

in the sense that the first portions of salt added to water have a greater effect in increasing or a less effect in diminishing the viscosity than subsequent additions. This effect is most clearly shown when one plots $\frac{\text{relative } \text{viscosity} - \text{I}}{\text{mol, concentration}} \text{ against cube root of concentration (Fig. 8)}.$

No such effect as this has ever been observed with non-electrolytes, and it is not surprising therefore that Grüneisen en-

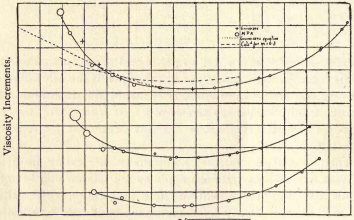


Fig. 8.—10 × 3/Concentration.

deavoured to connect the phenomenon with ionization. The following formula was found to reproduce the observations with a fair degree of accuracy:—

$$\frac{\frac{\eta}{\eta_o} - I}{m} = Aa + B(I - a) + Cm$$

where $\eta | \eta_o =$ the relative viscosity; m = molecular concentration; a = degree of ionization and A, B, C are constants.

A series of results for lithium nitrate obtained by Applebey is given on opposite page.

Applebey found that the results for the very dilute solutions departed somewhat from those calculated by Grüneisen's formula and this divergence is apparent in figure 8, where the calculated values are shown by a dotted line. Now Grüneisen's equation contains the assumption that the effects of the ion and of undissociated salt respectively are distinct and separable and that each is in all cases proportional to the concentration of the component

Concentration (m).	10m1/3	<u>η</u> .	$\frac{\eta/\eta_0-1}{m}$.
0.0174	2.29	1.0026	0.140
0.0567	3.84	1.0062	0.110
0'1071	4.75	1.0152	0'1165
0.3643	7'142	1.0402	0.1111
o·8666	9.533	1.0080	0.1131
1.316	10.00	1.1567	0.1101
3.8541	15.678	1.74075	0'1922
4.578	16.60	2.0577	0.5310
5.8.0	-8100	010077	212.6

TABLE XLII.—VISCOSITY OF LITHIUM NITRATE SOLUTIONS.

considered. This can be true only if the process of solution is merely a mechanical mixing, and if the view, generally accepted now, that the act of solution interferes with the molecular complexity of the solvent is to be rejected.

On the assumption that the viscosity is proportional to the mean molecular volume and that the different molecular species present in a salt solution are: (a) simple water molecules, H_2O ; (b) associated water molecules assumed to be triple, $(H_2O)_3$; (c) ions, hydrated to an unknown extent, $\text{Li.}xH_2O$, $\text{NO'}_3.yH_2O$; (d) undissociated molecules, possibly combined with water, LiNO_3 ; (e) salt complexes in strong solutions, $(\text{LiNO}_3)_n$, Applebey deduces the formula

$$\frac{\eta}{\eta_o} = \frac{c'_1 + c'_3}{c_1 + c_3 + ms (I + a)}$$

where c_1 is the concentration of single H_2O molecules, c_3 the concentrations of triple molecules in the solution, c_1 and c_3 the respective concentrations in pure water, m the weight normality, s the density, and a the degree of ionization.

Using Kohlrausch and Mailey's 90 conductivity determinations to find α from the relation $\alpha = \frac{\Lambda}{\Lambda_{\infty}} \times \frac{\eta}{\eta_{o}}$, it may be shown that an approximate value of the degree of hydration of lithium nitrate is determinable.

Unfortunately, however, Applebey's formula cannot at present be submitted to a quantitative test; but there is no doubt that a formula of this kind will have to be taken into account when sufficiently accurate determinations of the variables involved are available. In an investigation of the viscosity of caesium nitrate solutions Merton,⁹¹ following the procedure of Applebey, found that at all temperatures the viscosity of these solutions is less than that of water, and confirmed Grüneisen's discovery anent the change of curvature at the dilute end of the viscosity-concentration curve. Qualitatively, viscosity changes follow the classification of the elements in the periodic system and so, seeing that the caesium ion in aqueous solution is the smallest of its group, it possesses the largest ionic mobility.

On surveying the behaviour of aqueous electrolytic solutions outlined above it will be obvious that the whole problem is complicated by the anomalous behaviour of water, and that simple relationships might be expected from an organic solvent. This aspect of the question has been successfully attacked by Walden 93 who has used such salts as tetramethyl ammonium iodide, and sodium iodide dissolved in a variety of organic liquids. In all cases, no matter what the solvent may be, it appears that at infinite dilution the relationship

$$\Lambda_{\infty} \eta_{\infty} = \text{constant}$$

is very nearly correct. The following numbers, obtained by Walden, may be quoted in support of this.

TABLE XLIII.—Solutions of Tetra Methyl Ammonium Iodide at o° C. and 25° C.

Solvent.	lina.	$\Lambda_{\infty} \eta_{\infty}$ at 0° C.	Λ m n m at 25° C
Acetone	7	0.403	0.411
Propionitrile		0.918	0.682
Nitromethane		0.772	0'743
Methyl alcohol		0.461	0.210
Ethyl iso-thiocyanate.		0.694	0.655
Methyl iso-thiocyanate		0.707	0.690
Acetyl acetone		0.656	0.640
Sym, ethyl sulphite .		0.655	
Acetic anhydride		0.683	0.654
Epichlorhydrin		0.674	0.688
Ethyl alcohol		0.662	0.648
Benzonitrile		0.679	0.706
Furfurol		0.744	0.745
Diethyl sulphate .		0.686	0.688
Dimethyl sulphate .		0.756	0.757
			- 757
		0'700	•700

TABLE XLIV .- Solutions of Tetra Propyl Ammonium Iodide at 25° C.

Solve		Λ_{∞} η_{∞} .		
Acetone				0.632
Propionitrile .				0.620
Ethylene chloride				0.612
Epichlorhydrin .				0.911
Benzonitrile .				0.636
Acetoacetic ester				0.019
Nitrobenzene .				0.637
Benzyl cyanide	0.			0.614
Iso-butyl alcohol				0.616
m-Chloraniline .				0.635
			140	0.624

TABLE XLV.—Solutions of Potassium Iodide at Various Temperatures.

	Solven	t.		t.	Λ_{∞} η_{∞} .
Methyl alcoho "" "Acetone ." Acetonitrile Pyridine . ""				25° 0° 18° 25° 0° 25° 0° 25° 0° 25°	0*647 0*630 0*616 0*636 0*652 0*647 0*676 0*693 0*630 0*639

These numbers also show that the relationship Λ_{∞} $\eta_{\infty} = K$ holds good at different temperatures, e.g. the value of K for tetramethyl ammonium iodide is 700 both at 0°C and 25°C.

We may therefore write $\eta_{\infty}^{t} \Lambda_{\infty}^{t} = \eta_{\infty}^{t} \Lambda_{\infty}^{t} = K$

or
$$\frac{\eta^t_{\infty}}{\Lambda^t_{\infty}} = \frac{\Lambda^{t'}_{\infty}}{\eta^{t'}_{\infty}}$$
.

Results for water, formamide and lactonitrile are uniformly high, and so are their association coefficients and dielectric constants. It is probable, therefore, that a knowledge of these last two factors and the way in which they vary will be necessary before the behaviour of aqueous solutions is rendered intelligible.

To illustrate the result of considering dielectric constant and association factor, the following table may be consulted (Walden⁹⁴).

TABLE XLVI.-KCl IN WATER.

	$t = 0^{\circ}$.	$t=50^{\circ}$.	t = 100°.
η_{∞}	81.4	232·5	414
	0.0178	0·00548	0.00283
	1.449	1·271	1.173
	2.52	2·24	1.98
	0.575	0·568	0.592
Dielectric constant e	87°33	70°0	56.2
	0°0418	0°0407	0.0411

The validity of Walden's rule has been questioned by Dutoit and Duperthius 129 who, after making measurements of viscosity from 0° to 40° or 80° found that the magnitude of the quantity η_{∞} Λ_{∞} was affected by change of temperature even when all the other conditions were kept constant. Thus, for solutions of sodium iodide in pyridine the product varied from 0.573 at 0° to 0.559 at 80° and in *iso*-amyl alcohol to an even larger extent, from 0.374 to 0.225. Schmidt and Jones 180° also call attention to similar discrepancies in the case of potassium iodide dissolved in methyl alcohol, ethylene glycol and glycerol.

CHAPTER VII.

THE VISCOSITY OF COLLOIDS.

Historical.—In previous chapters viscosity has been dealt with only as applied to homogeneous liquids, i.e. to one phase systems. It is obvious that an entirely different state of affairs will hold for two phase or colloidal systems. Considering first of all the case of the suspensoid where particles of extremely small size are suspended in a homogeneous medium, it has been found that the presence of the suspended particles affects but slightly the viscosity of the dispersion medium. Bingham 118 for instance, working with suspensions of china clay, graphite, and infusorial earth, concludes that the decrease in fluidity is directly proportional to the volume concentration of the suspended solid, i.e. it is very small. Much work in this field has been carried out in connexion with the so-called critical solution phenomena which manifest themselves when solutions of partially miscible liquids become heterogeneous on alteration of temperature. For instance Friedländer 112 found a marked increase in the viscosity of iso-butyric acid-water mixtures as the solution was cooled to the temperature where opalescence became evident and the critical solution point approached. extended his work to aqueous solutions of phenol and the ternary system benzene-water-acetic acid, and concluded that the increase in viscosity was due to the formation of drops. Bose 111 and his co-workers have also found that abnormally high viscosities are to be found at the critical solution temperatures. and consider that this may be due to the rolling of drops in the capillary of the viscometer. They also discovered that similar abnormalities occur in the case of anisotropic liquids and regarded these "liquid crystals" as emulsions of very long life. It was pointed out that the formation of orderly oriented "swarms" of the crystalline liquid molecules would cause an increase in viscosity and Bose proposed to test this by determin-

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ing the viscosity under the conditions for turbulent flow, which might conceivably break up the "swarms". Something of this sort apparently happened, for the abnormalities certainly decreased as the transpiration velocity increased.

With regard to true colloidal solutions (emulsoids) Griffiths ¹¹⁰ found that gelatine sols containing sodium chloride had a conductivity greater than that of a gelatine-free solution but a viscosity enormously greater, whilst Garrett ¹⁰⁹ determined the viscosity of colloidal solutions by means of an oscillating disc and discovered that the logarithmic decrements of the amplitude of vibration increased, two facts which are explicable on the assumption that the emulsoids possess a definite structure.

Von Schroeder 108 used the viscosity method for studying the behaviour of gelatine solutions and found that the viscosity of a given solution at a given temperature was not constant but depended largely on the previous history of the system. The observed phenomena were accounted for by the overlapping of two processes:—

- (1) An irreversible change, probably a chemical reaction, between the gelatine and water.
 - (2) The reversible change hydrosol = hydrogel.

The speed of the first reaction was measured by heating a gelatine solution to 100° C. for varying periods. The rate of change of viscosity is expressed by

 $\frac{d\eta}{dt} = K(\eta - \eta_i)$ where η_i is the final value of the viscosity.

Von Schroeder calls this the "saponification reaction" since it is analogous to the hydrolysis of cane sugar.

Acids and bases accelerate the change, whilst salts are without influence. With regard to the reversible reaction a fall in temperature favours the gel formation but the change takes place slowly. Rise of temperature favours the sol formation but is accompanied by hydrolysis of the gelatine, so that if the solution is heated till the sol is completely "saponified" then on cooling no gel is formed and the viscosity remains constant.

The Viscometer as Colloidoscope. ¹—In a general introduction to the subject of colloids and their viscosity Wo. Ostwald ¹⁰⁷ draws particular attention to the enormous qualitative and quan-

¹ The material in this section has been largely taken from the recent "General Discussion on Colloids and their Viscosity," "Trans. Faraday Society," 1913, 9, 34.

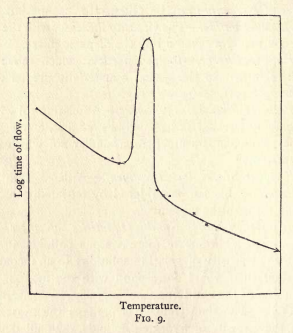
titative variability of the viscosity of substances in the colloidal state. Whereas the viscosity of a molecular solution is defined completely by the concentration and the temperature, that of a colloid presents an entirely different problem. In the simplest case, viz. a suspensoid colloid in which the disperse phase still retains the properties of the solid state, one further variable must be considered, the size of the particles or degree of dispersity.

A greater number of variables defines the viscosity of the emulsoid colloids, where not only concentration, temperature and degree of dispensity must be taken into account but also

- (1) solvate formation.—The viscosity increases with the amount of dispersion medium taken up by the disperse phase.
- (2) Electric Charge.—Ionized particles which move in the electric field impart to the system a materially greater viscosity than uncharged particles.
- (3) Previous Thermal Treatment.—A gelatine sol which has been repeatedly warmed and cooled has, at first, a lower viscosity at the same temperature than a similar sol which has not been so treated.
- (4) Previous Mechanical Treatment.—A decrease in the viscosity of gelatine sols can be produced by repeatedly forcing the same liquid through the viscometer.
- (5) Inoculation with Small Quantities of more Viscous Colloids.—Small additions of more viscous colloids after some time raise the viscosity of gelatine solutions to an incomparably higher extent than would correspond with the increase in concentration due to the addition.
- (6) Time.—The time factor has perhaps the most striking effect on the viscosity of hydrated colloids. In all determinations of the viscosity of such colloids different values are obtained according to the age of the sol, and according to the velocity with which the temperature is being altered. The ageviscosity curve appears generally to be S shaped.
- (7) Addition of Electrolytes and Non-electrolytes.—Most varied effects are produced by relatively small additions of substances.

It is particularly remarkable that relatively enormous changes of viscosity occur within very narrow limits of temperature and concentration. A gelatine sol may, by merely altering these variables or even by mere ageing, pass through all values of viscosity from that of water to that of a solid. Soap sols can attain the viscosities of solids at concentrations less than I per cent.

Viscosity as a Methodic Principle.—If it is desired to follow quantitatively the alterations of dispersity or of solvate formation, such as coagulation, peptonization, gelatinization and so on, any physico-chemical property of the system may be used as an indicator of such changes. In practice that property will be chosen which shows the largest possible variation with small



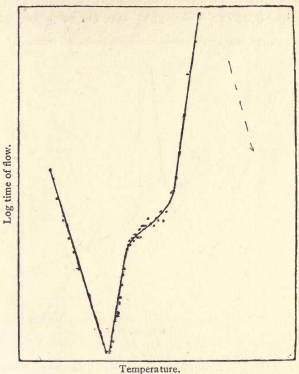
changes in the colloidal condition and which lends itself to convenient measurement. These requirements are eminently fulfilled by viscosity.

As an illustration of the application of the viscosity method may be quoted the change undergone by a dilute albumen sol on heating.

Fig. 9 shows the viscosity changes, with temperatures as abscissæ and logarithms of times of flow as ordinates. Between 50° and 57° the viscosity decreases in the normal manner. At 57.5°, shortly before the appearance of turbidity, a large increase in viscosity takes place which at 60° gives place to an equally

steep decrease. After that the viscosity curve of the coagulated albumen proceeds practically in the original direction as if nothing had happened.

Another interesting case is the viscosimetric behaviour of a starch suspension during heating, a process which leads to the well-known gelatinization (Fig. 10). At first the viscosity falls in the normal way and then extremely rapidly at 57° a sharp

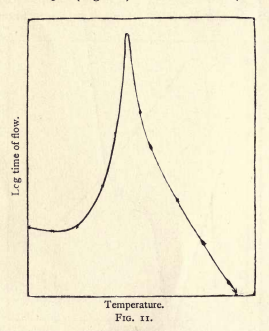


Temperature. Fig. 10.

break occurs and the curve rises sharply to 65°, then bends away and again rises to 95°.

Viscosity and Continuity.—Modern colloidal chemistry has led to the discovery that continuous transitions are possible between coarse suspensions, colloidal solutions, and systems of molecular dispersity. A number of transitional systems on the border of colloidal and molecular dispersity may be mentioned which show pronounced anomalies of viscosity. First may be instanced the critical mixtures of liquids, e.g. isobutyric acid and

water at certain temperatures and concentrations (Fig. 11). Then the percentage increase in viscosity of a critical water isobutyric acid mixture containing 38.6 per cent of acid amounts to 34.3 per cent per degree. Other systems exhibiting entirely analogous anomalies of viscosity are, for instance, liquid sulphur above 130° and the opalescent liquid crystals; and in these cases very great temperature coefficients of viscosity are met with. On cooling molten sulphur from 400° to 200° the viscosity rises from 150 to 50,000 and falls to 8 at 150° (Fig. 12). Wo. Ostwald (loc. cit.) justly



points out that the remarkable similarity between the cases just mentioned and purely colloidal phenomena such as the gelatinization of starch can only be explained by the theory that colloids, critical mixtures, crystalline liquids, melted sulphur, etc., are all disperse systems, i.e. systems whose properties are periodically discontinuous in space, this period being extraordinarily small. In other words this theory applies not only to coarsely heterogeneous systems but also to purely molecular systems.

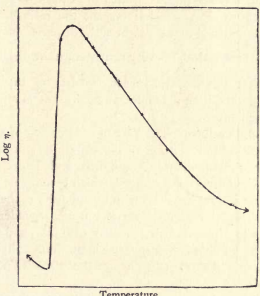
Hatschek's Work.—Apparently the only successful attempt to find a mathematical expression for the viscosity of a two phases system is due to Hatschek¹¹⁵ who deduced the equation

$$\eta_1 = \eta \left(\mathbf{I} + 4.5 f \right)$$

where η_1 is the viscosity of the system, η that of the liquid continuous phase and

f is the ratio $\frac{\text{total volume of disperse phase}}{\text{total volume of system}}$.

Einstein 106 obtained a similar expression, viz. $\eta_1 = \eta (\mathbf{I} + f)$. These formulæ state that the viscosity of a system of deformable particles suspended in a liquid is independent of the size of the particles and is a linear function of the volume of the disperse phase only. Whilst Hatschek's formula has been found to hold



Temperature. Fig. 12.

good in the majority of cases for low concentrations and for fairly coarse particles, discrepancies become apparent (I) at higher concentrations, and (2) with ultra-microscopic particles. Hatschek draws attention to the fact that the capillary viscometer, with its high rate of shear, may be responsible for the want of agreement, and further that nothing is known about the shape and density of ultra-microscopic particles, quantities which may conceivably be important in this connexion.

When, however, Couette's apparatus was used permitting a variable rate of shear the results obtained show a closer agree-

ment with theory. The same author considers that the volume of the adsorption envelope must be considered as well as that of the disperse phase itself, and calculates, using figures for sulphur sols, that the thickness of this layer is 0.87 $\mu\mu$.

In the case of emulsoids Hatschek ¹⁰⁵, assuming that the velocity of displacement of juxtaposed layers exceeds a certain critical value, shows that the viscosity of an emulsoid η' is given

by the equation $\eta' = \frac{\eta \sqrt[3]{A}}{\sqrt[3]{A'} - 1}$ where η is the viscosity of the dis-

persive medium and A is the ratio of the volume of the emulsoid to that of the disperse phase. When the values of A are compared with those of A' which represent the ratio between the volume of the emulsoid to the weight of the disperse phase present, it is found that the ratio $\frac{A}{A'}$ is very nearly constant, leading to the

conclusion that the disperse phase consists of the dissolved substance together with a definite and constant amount of the dispersive medium.

Other Applications.—The rate of coagulation of aluminium hydroxide sols by the addition of electrolyte has been followed by viscosity measurement. Freundlich and Ishizaka ¹⁰⁴ find that the maximum viscosity of a sol, which is completely precipitated by an electrolyte increases with the quantity of the colloid in the sol. The maximum viscosity is further dependent on the nature of the precipitant, being smaller with NO₃ and CNS ions than with the sulphion and the succinate ion. Shaking and other mechanical disturbances change the viscosity of the concentrated sols.

The Nature of Aqueous Soap Solutions.—Soap solutions have a high electrical conductivity and fall into the class of electrolytic colloids, to which congo red, casein and many other compounds belong, and which are characterized by containing both colloidal and electrolytic constituents. McBain, Miss Cornish and Bowden 116 have shown that when electrolytes such as caustic soda are added to aqueous solutions of sodium palmitate the viscosity falls at first, reaches a minimum and then increases rapidly as the electrolyte concentration is still further increased. The initial diminution of viscosity may be due to the abstraction of water from the disperse phase, with a consequent diminution

in the bulk of the latter; the rapid increase in viscosity above a certain electrolyte concentration is due probably to the formation of colloidal aggregates, the size of which may increase with increasing concentration. Results similar to the above have been obtained by Goldschmidt and Weissmann 114 with solutions of soft soap, and these authors consider that the addition of electrolytes with a common ion in dilute solution diminishes the hydrolysis and degree of dissociation of the alkali salts of the fatty acids, whereas in higher electrolyte concentration the soap is largely present as a hydrophile colloid.

The Chemistry of Nitro-Cellulose.—Schwarz 103 determined the viscosity of solutions of nitro-cellulose in camphor-alcohol, ether-alcohol and in acetone. The viscosity increases very rapidly with concentration and the solutions exhibit the phenomena of ageing. The ageing is accompanied by a very considerable increase in the viscosity which is more pronounced in the case of the more concentrated solutions. Baker 102 in a similar investigation showed that the nitro-cellulose solutions follow the law $\eta = \eta_a (I + ac)$ when a and k are constants dependent on the nature of solvent and solute.

The Valuation of Rubber.—Schidrowitz 186 has shown that an intimate connexion exists between the mechanical properties of rubber and the viscosity of its solutions. Fol 101 pursuing the same line of work finds that violent shaking during the preparation of the solution affects the viscosity considerably. The results are affected by lapse of time, between making the solutions and carrying out the measurements, and by temperature.

CHAPTER VIII.

THE RELATION BETWEEN VISCOSITY AND CHEMICAL CONSTITUTION.

It has been pointed out in Chap. III. that the earlier workers experienced very considerable difficulty in establishing the additive nature of viscosity on account of the apparently erratic results afforded by certain of the substances examined. Particularly abnormal are the aliphatic alcohols and acids and, indeed, all hydroxylic compounds; but many other constitutive factors such as ring formation, branched chains, unsaturation (simple or conjugated), etc., exert a marked disturbing influence on the viscosity of a liquid.

So sensitive indeed is viscosity to constitutive changes that special care must be taken to eliminate as far as possible the influence of these when calculating atomic or group values, and the figures so obtained can only be used for calculating the viscosities of "normal" liquids.

The influence of the hydroxyl group in raising viscosity was first recognized by Gartenmeister and its action is no doubt intimately connected with the potential quadrivalence of the oxygen atoms inducing association. The effect it produces will be evident from the following figures.

TABLE XLVII .- VISCOSITY AND ASSOCIATION.

Substance. Ethane	N.			٧.	Viscosity (25° C.).
Ethyl alcohol					0.0100
Ethylene glycol					0°1733

Its influence again is very well illustrated by the steady increase in the viscosity of freshly distilled ethyl acetoacetate owing to the gradual enolization of the ketonic form. 96

Since, however, the great exaltation of viscosity produced by the hydroxyl group is at any rate mainly due to association induced by the latent affinities of the oxygen atom we should expect the introduction of another unsaturated group into the molecule to result in the partial neutralization of the residual affinity of the oxygen atom owing to mutual action of the two unsaturated centres. This would result in a reduction of the association and consequently to a diminution of the viscosity.

Moreover the diminution of association should depend on the degree of unsaturation of this second group and also on the degree of its proximity to the hydroxyl group.

The correctness of these views is clearly borne out by the numerical values obtained for a number of phenols and their derivatives. 96

TABLE XLVIII .- VISCOSITY OF THE PHENOLS.

Substance	e.				Viscosity.	$\frac{\eta \times \text{Mol. Vol.}}{10^6}$
Phenol	-		(19)		0*0404	453
Anisole					0.00741	66.6
Phenetole					0.00822	63.8
p-Chlorophenol .					0.0603	590
m-Chlorophenol .					0'0472	459
o-Chlorophenol .					0°0225	212
p-Cresol					0.0261	527
m-Cresol					0.0206	475
o-Cresol					0.0321	333
m-Tolyl methyl ether					0.00875	68.5
o-Tolyl methyl ether					0.00849	66.7
b-Tolyl methyl ether					0.00809	62.8
-Nitrophenol					0'0234	199
Ethyl salicylate .		1.		5.5	0.0172	118
Benzyl alcohol .					0.0301	286
Benzyl methyl ether					0.0103	81.1

It will be noticed that the value of the expression

 $\frac{\eta \times \text{Mol. Vol.}}{10^6}$ has been used in the above table as a means of comparison of the liquids employed and as an indication of the presence and the extent of association.

Dunstan ⁹⁷ pointed out that in a number of series the members of which are similar in constitution and generally recognized as unassociated, the value of $\frac{\eta \times \text{Mol. Vol.}}{10^6}$ only varies

between comparatively small limits. The average value for the lower alkyl chlorides is 37, for the bromides 51, for the iodides 68, for the ketones 43, and for the ethers 25. The values for

benzene, anisole, phenetole, and the three tolyl methyl ethers all lie between 60 and 70. On the other hand association enhances the value to a most marked extent, the numbers for water, ethyl alcohol and glycol being respectively 494, 193 and 2750, and the expression affords a fairly good test for molecular aggregation. Too much weight, however, must not be attached to the values afforded by this expression since it is undoubtedly only a first approximation, being influenced considerably by many other factors such as molecular symmetry, conjugation of unsaturated groups and linkings and ring formation.

Influence of Symmetry.—The figures in Table XLVIII illustrate a further constitutive influence on viscosity by the difference in the viscosities of the isomeric and presumably unassociated tolyl methyl ethers and benzyl methyl ether, the more compact molecules of the former exhibiting a distinctly lower viscosity. That this is an example of a very general rule is clearly shown by comparison of the viscosities of the following normal and *iso* compounds.

TABLE XLIX.—Effect of Symmetry on Viscosity.

Substance.	Viscosity (o° C.).	Substance.	Viscosity (o° C.).
n-Pentane Propyl iodide Methyl butyrate Butyric acid Butylamine	010000	Iso-pentane	0°00273 0°00878 0°00672 0°01885

Influence of Unsaturation.—Until the last twenty years the term unsaturation was almost exclusively employed to denote the presence of an ethylenic or acetylenic union. With the development of modern views on "residual affinity" and "latent valency," however, the scope of the term unsaturation has been very greatly widened. Accordingly, in view especially of the additional complications introduced by "conjugation" it will be advisable to consider separately the various effects produced by the manifestations of what is still a very vaguely understood phenomenon.

Dealing first with the influence of a simple ethylenic and acetylenic linkage one cannot draw any definite conclusions from the numerical data available. It may be stated that as a general rule the presence of such groupings raises the viscosity above that of the corresponding saturated compound. The viscosity of the acetylenic compound, however, is sometimes greater and sometimes less than that of its ethylenic analogue. The following results are typical. 98

TABLE L.-Effect of Unsaturation on Viscosity.

Substance.	Formula.	Viscosity (25° C.).
Propyl ethyl ether Allyl ethyl ether Propargyl ethyl ether Phenyl ethane Phenyl ethylene Phenyl acetylene	$\begin{array}{c} . & CH_3 \cdot CH_2 \cdot CH_2OEt \\ . & CH_2 = CH \cdot CH_2 \cdot OEt \\ . & CH \equiv C \cdot CH_3OEt \\ . & C_6H_5 \cdot CH_2 \cdot CH_3 \\ . & C_6H_5CH : CH_2 \\ . & C_6H_5C \in CH \end{array}$	0.00339 0.00345 0.00527 0.00607 0.01109 0.00883

Very much more striking, however, are the effects produced by the alternation of double with single linkages (the "conjugated double bonds" of Thiele).

The abnormalities produced by the presence of such conjugated double bonds have been recognized in many physical properties but in none more markedly than in viscosity, which undergoes remarkable exaltation in conjugated compounds.

The following results 99 illustrate the effect on viscosity of conjugation of double linkage.

TABLE LI.-EFFECT OF UNSATURATION ON VISCOSITY.

Substance.	Formula.	Viscosity.
Acetone	. CH ₃ . CO. CH ₃	0.00302
Mesityl oxide .	$\begin{array}{c} CH_3 \\ CH_3 \end{array} C = CH \cdot CO \cdot CH_3 \end{array}$	0.00876
Phorone	$\begin{array}{c c} CH_3 & C = CH \cdot CO \cdot CH = C & CH_3 \\ CH_3 & CH_3 & CH_3 \end{array}$	0.0130
Eugenol	HO CH ₂ . CH: CH ₂	0.0693
Iso-eugenol	CH ₂ O CH: CH. CH ₃	0.2672
Safrol	-CH ₂ .CH:CH ₂	0.0229
Iso-safrol	-CH: CH. CH ₃	0.0398

It is interesting to compare the effect of conjugation on other physical properties.

TABLE LII.—Effect of Conjugation of Double Bonds on Physical Properties.

Substance.	Molecular	Refr	raction.	Optical Anomaly.	Mol. Volume.	Viscosity.	Boiling Point.
Acetone .	observed calc		16·05 16·04	+ 0.01	73.8	0*00305	56
Mesityl oxide	observed calc	:	30.13	+ 0.74	114.6	0.00876	130
Phorone .	observed calc	:	45°39 42°73	+ 2.66	153.9	0.0130	195
Eugenol .	observed calc		48.1	+ 0.2	_	0.0693	247
Iso-eugenol .	observed calc	:	49°3 47°6	+ 1.7	T -	0.2672	260
Safrol	observed calc		45°9 45°8	+ 0.1	_	0.0330	232
Iso-safrol .	observed calc		47.6 45.8	+ 1.8	_	0.0308	249

It might well be expected that conjugation of groups possessing notable residual affinity would produce a very similar effect in enhancing the viscosity of a substance. This is well illustrated by the following table which gives the viscosity (in amylacetate solution at equivalent concentration) of a number of derivatives of acetone.

TABLE LIII.—Effect of Conjugation of Unsaturated Groups on Viscosity.

Substance.	Formula.	Viscosity at Mol. Wt. per cent Concentration.
Acetone	$\begin{array}{c} \text{CH}_3.\text{CO}.\text{CH}_3 \\ \text{CH}_3\text{CO}.\text{C}_6\text{H}_5 \\ \text{C}_6\text{H}_5\text{COC}_6\text{H}_5 \\ \text{C}_6\text{H}_5\text{CH}_2.\text{CH}_2.\text{COCH}_3 \\ \text{C}_6\text{H}_5\text{CH}_2.\text{CH}_2.\text{COC}_6\text{H}_5 \end{array}$	0.00752 0.00808 0.00892 0.00830 0.00877

By combining the effects due to the conjugation of unsaturated bonds and the conjugation of unsaturated groups we obtain very great exaltation of viscosity. The degree may be gauged if it is remembered that the solutions in Table LIV have half the concentration of those in Table LIII.

TABLE LIV.—Effect of Conjugation of Double Bonds and Unsaturated Groups on Viscosity.

Substance.	Formula.	Viscosity at Mol. Wt. per cent.
Benzylidene acetone . Benzyl acetophenone Benzylidene acetophenone Cinnamylidene acetophenone Dibenzylidene acetone .	$\begin{array}{c} C_6H_5\cdot CH_2\cdot CH_2\cdot CO\cdot CH_3\\ C_6H_5CH:CH\cdot COCH_3\\ C_6H_5CH:CH\cdot COC_6H_5\\ C_6H_5CH:CH\cdot CO\cdot C_6H_5\\ C_6H_5CH:CH\cdot CH\cdot CH\cdot CH\cdot COC_6H_5\\ C_6H_5CH:CH\cdot CO\cdot CH:CH\cdot CC_6H_5\\ C_6H_5CH:CH\cdot CO\cdot CH:CH\cdot CC_6H_5\\ C_6H_5CH:CH\cdot CH\cdot CO\cdot CH:CH\cdot CC_6H_5\\ C_6H_5CH:CH\cdot CH\cdot CH\cdot CH\cdot CO\cdot CH:CH\cdot CH\cdot CH\cdot CC_6H_5\\ \end{array}$	o'00811 o'00833 o'00850 o'00888 o'00893 o'00934

The validity of the conclusions drawn from the above data regarding the influence of symmetry and of unsaturated linkages and groupings on viscosity have been tested by means of a large number of aliphatic and aromatic amines and their derivatives 100.

TABLE LV .- VISCOSITY OF ALIPHATIC AMINES.

Substance.			Viscosity.	η × 10 ⁶ Mol. Vol.	
n-Propylamine.			0'00353	42.0	
Allylamine .			0.00372	50.0	
n-Butylamine .			0.00081	70.8	
Iso-butylamine.			0'00553	56.8	
Diethylamine .			0.00362	36°2	
Di iso-butylamine			0.00682	40.0	
Triethylamine .			0.00363	26.1	

TABLE LVI.-VISCOSITY OF AROMATIC AMINES.

Substance.	Viscosity (55°).	Viscosity (130°).	η ₅₅ × 10 ⁶ Mol. Vol.	η ₁₃₀ × 10 ⁶ Mol. Vol.
Aniline Methyl aniline Dimethyl aniline Allyl aniline Benzyl aniline Diphenylamine Triphenylamine Benzylidene aniline Acetanilide Methyl acetanilide (at 120° (Benzylamine Dibenzylamine Tribenzylamine Methyl diphenylamine		0°00555 — 0°00506 0°0120 0°0104 0°0640 — 0°0190 0°00818 0°00442 0°00812	182 96°9 64°7 101 306 266 — 229 —	56·5 — 34·2 64·5 61·1 262 — 142 53·6 37·0 39·5 69·9 34·2

From the above figures which are drawn from a much larger number quoted in the original papers ample confirmation is obtained of the recognized influence of symmetry and conjugation of unsaturated nuclei.

The effect of replacing the hydrogen atoms of the amino group by alkyl radicles is steadily to diminish the viscosity whether the primary amine be aliphatic or aromatic. The effect of symmetry of the alkyl group in the isomeric normal, and isobutylamines has already been pointed out.

The results obtained for the aromatic amines, however, are particularly interesting in illustrating the effect of accumulating unsaturated nuclei. Whereas methylaniline and dimethylaniline are much less viscous than aniline, substances such as phenylaniline, benzylaniline, benzylidene aniline, acetanilide and especially triphenylamine have notably high viscosities. Exactly parallel are benzylamine, dibenzylamine and tribenzylamine. It is evident that here the exaltation due to accumulation of unsaturated nuclei more than counteracts the depression which might be expected owing to increased symmetry. That this latter influence is, however, still perceptible even in the highly conjugated compounds is evident when one compares diphenylamine and acetanilide with methyldiphenylamine and methylacetanilide.

The effect of introducing various substituent groups into the phenyl group of aniline and its derivatives has been studied by Thole (loc. cit.), the results in general following those previously described in connexion with the substituted phenols.

CHAPTER IX.

SOME APPLICATIONS OF VISCOSITY.

THE value of a physical property in the eyes of the average chemist depends largely on its applicability to problems which arise in the various branches of chemistry and which sometimes cannot be definitely solved by purely chemical methods.

Mention need only be made of the application of refractivity and absorption spectra to the classical controversies concerning the constitutions of ethyl acetoacetate and of isatin and of the work of Patterson, Lowry and others on the connexion between rotatory power and velocity of chemical reaction. Since viscosity is so highly a constitutive property it may and has been applied to a number of problems arising in physical, inorganic and organic chemistry and has proved particularly useful in certain cases such as in determining the constitution of the acetaldehyde phenylhydrazones where chemical evidence is not available.

Measurement of Rate of Reaction.—Since the viscosities of different substances are as a rule widely different, it is evidently possible to follow the course of a reaction by determining the viscosity of the reacting materials at definite intervals of time. It is necessary, of course, that the reaction should proceed to a small extent only during each viscosity observation though in some cases it is possible to examine more rapid reactions by removing samples of the reacting mixture at definite times, stopping the reaction by diluting, for example, with a cold solvent and then measuring the viscosity of the resulting solution.

The first worker in this direction appears to be von Schroeder, who in 1903 found that the viscosity of a hot aqueous solution of gelatine slowly fell owing to hydrolysis of the gelatine. He found that $\frac{d\eta}{dt} = K(\eta_1 - \eta_2)$ indicating a monomolecular reaction.

Dunstan and Mussell ¹¹⁸ in 1911 applied the method to the following reactions: Freshly distilled acetoacetic ester (ketonic) → equilibrium mixture; aniline + ammonium thiocyanate → phenyl thiocarbamide; ammonium cyanate → carbamide; ammonium thiocyanate → thiocarbamide; acetic anhydride + water → acetic acid; benzoyl chloride + water → benzoic acid.

Two typical sets of figures will give an indication of the results obtained.

TABLE LVII.—Change of Ammonium Cyanate into Carbamide (in Aqueous Solution at 25° C.).

Time in Minutes from liddle of First Observation.	Time of Flow.	a-x.	$\int_{t}^{1} \log_{10} \frac{a}{a-x}.$
0	4*29"	36.0	
50	4'30" 4'32"	35.0	0'00024
120	4.32"	33.0	0,00031
238	4°34″	30°4	0.00031
247	4'35"	30.0	0'00032
290	4'36"	29.0	0'00032
350	4'38"	26.7	0.00037
560	4'45"	20.0	0'00045
after 3 days	5'5"		_

It may be concluded, therefore, that this reaction is not unimolecular, a result in agreement with those of Walker, Hambly, Kay and Wood.

TABLE LVIII.—Hydrolysis of Benzoyl Chloride by Water (in Acetone Solution).

Time in Minutes.	$\frac{1}{t}\log_{10}\frac{a}{a-x}.$
11.2	
14.5	0.0002
22.5	0.0105
44'5	0'0095
53.0	0.0096
80.0	0.0103
140.0	STATE OF THE REAL PROPERTY.

The values in the last column agree sufficiently well to indicate a reaction of the first order.

Determination of Transition Points.—If saturated solutions of a substance at a series of temperatures are prepared and the viscosities of these solutions plotted against the temperatures, a

smooth curve is in general obtained. If, however, the chemical composition of the substance changes at some point within the temperature range chosen the viscosity of the solution will naturally change and the temperature of transition will be indicated by a singular point on the viscosity-temperature curve. Hence viscosity measurements can be added to the already well-known dilatometric and solubility methods for determining transition points, and the results obtained by the various methods agree well.

The curves obtained on examining the classical cases of sodium sulphate and sodium carbonate ¹¹⁹ indicate clearly the transition of Na₂SO₄. 10 H₂O to Na₂SO₄ and of Na₂CO₃. 10 H₂O to Na₂CO₃. 7 H₂O and Na₂CO₃. H₂O.

Detection of Association and Conjugation.—The effect of these molecular influences on viscosity have been dealt with previously (pp. 72-75), and it need merely be repeated that both produce a very marked exaltation in the viscosity coefficient.

The Existence of Racemic Compounds in the Liquid State.—This problem has been the source of much controversy, and many methods have been suggested to decide whether a racemate on fusion or solution is converted into a mixture of the antipodes or whether it retains its individuality.

The bulk of the evidence is very unsatisfactory, usually because dilute solutions of the racemate have been investigated and this condition would tend to promote dissociation of the racemate.

Thus comparisons of affinity constants, molecular weights by the cryoscopic method and absorption spectra of d-, l- and racemic tartaric acids in dilute solution show no indication of the presence of the latter substance. On the other hand measurements of colour intensity, molecular weight, density, and absorption spectrum in concentrated solutions show in many cases undoubted signs of the existence of a racemic compound in solution.

Viscosity was first applied to the question by Beck ¹²⁰ who examined fused mixtures of dextro and lævo camphoroxime. He found that the viscosities of the antipodes of the inactive mixture were identical. The same results were obtained with the oximes of the carvones showing that in neither of these cases is any racemic compound produced. These results might be expected from the shape of the fusion point curves determined

by Adriani for similar mixtures, and have been confirmed by subsequent viscosity measurements of the fused and also of the dissolved components.¹²¹ The same investigator also carried out series of measurements with fused dimethyl-d-tartrate and racemate and with the acetyl derivatives of these esters.

TABLE LIX.—Viscosity of Dimethyl-d-Tartrates—Dimethyl Racemate Mixtures.

Per cent of Dextro-ester.	$\eta_{85^{\circ}\text{C.}}(\eta_{\text{H}_2\text{O}}^{25^{\circ}\text{C.}}=1).$
100	14.023
95 80	14.251
80	13.971
50	14.580
0	14.464

TABLE LX.—VISCOSITY OF ACETYL DIMETHYL-D-TARTRATE—ACETYL DIMETHYL RACEMATE.

Percentage of Dextro-ester.	$\eta_{104^{\circ} \text{ C.}} (\eta_{H_2O}^{25^{\circ}\text{C.}} = 1)$
o	6*764
10	6°764 6°666
40	6.015
100	6·912 6·933

Beck considered that in both series the viscosity values remain sufficiently constant to show complete dissociation of the racemic ester. Thole, 121 however, has pointed out that Bruni's cryoscopic measurements in ethylene dibromide solution undoubtedly demonstrate the possibility of actual existence of the dissolved racemic esters and that in Table LX the viscosity values with one exception seem rather to point to the inactive ester having a distinctly different viscosity from the active compound.

He examined several pairs of esters and considers that distinct indications are afforded of the existence of liquid racemates. Evidence of this is afforded by the following table.

TABLE LXI.-VISCOSITY OF RACEMATES.

Ester.		Temperature.	Viscosity.
Methyl d-tartrate		85° 85°	0.133
Methyl racemate Ethyl d-tartrate		85°	0°130
Ethyl racemate	1	25° 25° 85° 85°	1,360
l-Methyl l-mandelate .		85°	0.0632
l-Methyl dl-mandelate .		85°	0.0623

The free tartaric and mandelic acids were also investigated, in the latter case a number of different solvents being used in order to find what dissociating influence these had.

The viscosity-concentration curves for aqueous solutions of active and racemic tartaric acids show that the amount of undissociated racemic acid increases with increasing concentration.¹²²

Stewart had previously found that the absorption spectra curves of similar solutions of dextro and racemic tartaric acid are at low concentrations identical but at higher concentrations (about 14 per cent) show a distinct divergence.

In dissociating solvents such as water and pyridine no difference is to be found between active and inactive mandelic acids, but in a comparatively non-dissociating solvent such as amyl acetate curves similar to those of the tartaric acids are obtained.

The viscosimetric examination of a number of active and inactive alcohols and other substances has shown, however, that the free existence of a liquid or dissolved racemic compound is very exceptional.

The Viscosity of Geometrical Isomerides.—The first attempt to correlate geometrical isomerism with viscosity was made by Lauenstein ¹²³ who, working with the sodium salts of certain dibasic ethylenic acids, could find no regularities, sodium maleate being less viscous than sodium fumarate, whilst with the methyl derivatives of these acids the reverse holds. These results are, however, complicated by the different degrees of ionization of the salts.

A much more complete investigation has been carried out by Thole ¹²⁴ who examined the ethylenic acids and the esters, and also a number of oximes and phenylhydrazones, both in the dissolved and in the fused conditions.

The results obtained indicate that the viscosity and other

physical properties depend on the relative positions in the molecule of the two groups which exhibit most residual affinity, and a most convenient uniformity is gained by describing those compounds in which these groups approach each other as "adjacent" compounds and their isomerides as "opposed". Thus benzanti aldoxime is a typical "adjacent" compound and fumaric acid an "opposed" compound.

TABLE LXII .- VISCOSITIES OF GEOMETRICAL ISOMERIDES.

Substance.	Configuration.	Solvent.	Viscosity.
Maleic acid	adjacent	methyl alcohol	0.005802
Fumaric acid	opposed		0.002884
Citraconic acid	adjacent	" "	0'005908
Mesaconic acid	opposed	" "	0.002902
Oleic acid	adjacent	amyl acetate	0'009549
Elaidic acid	opposed	" "	0'009740
α-Bromo cinnamic acid		" "	0.008212
α-Bromo-allo-cinnamic acid	adjacent	17 77	0.008674
Ethyl maleate	,,		0.03016
" fumarate	opposed		0.02455
,, citraconate	adjacent	_	0'03251
" mesaconate	opposed	_	0.05399
" oleate	adjacent	_	0.08948
" elaidate	opposed	_	0.08175
" oleate	adjacent	amyl acetate	0.009260
_,, elaidate	opposed	" "	0.000144
Benzaldehyde-α-phenyl hydrazone .	17	" "	0.008782
Benzaldehyde-\beta-phenyl hydrazone .	adjacent	,, ,,	0.008030
Acetaldehyde-a-phenyl hydrazone .	West - Line	-	0.008323
Acetaldehyde-\beta-phenyl hydrazone .		-	0.008348
Camphor quinone-a-phenyl hydrazone	opposed?	chloroform	0.006632
Camphor quinone-\(\beta\)-phenyl hydrazone	adjacent?	99	0.006334
a-Benzaldoxime	"	amyl acetate	0.008344
8-Benzaldoxime	opposed	" "	0.008433
z-Anisaldoxime	adjacent	" "	0.008640
8-Anisaldoxime	opposed	" "	0.008722
z-Piperonaloxime	adjacent	"	0.008840
3-Piperonaloxime	opposed	,, ,,	0.000100
z-Benziloxime	adjacent	" "	0.000300
3-Benziloxime	opposed	,, ,,	0'009402

Where a solvent was used the viscosities quoted are those of solutions of concentration $\frac{\text{mol. wt.}}{40}$ per cent of solute.

Summing up these results this worker concludes that the viscosities of the isomers depend not only on the relative positions of the unsaturated groups but also on the degree of residual affinity they possess. Where these radicles possess small residual affinity the opposed compound has a lower viscosity than the adjacent

isomeride. This result is indeed to be expected since in the latter a tendency to potential ring formation will exist, a condition which is well known to enhance viscosity values.

On the other hand, when the groups possess a considerable degree of residual affinity as in the dibasic ethylenic acids, the exaltation of viscosity due to potential ring formation is masked by the depression of viscosity due to intramolecular neutralization of the residual affinities and consequent inhibition of molecular association, the result being that the adjacent compound has the lower viscosity. It is interesting to note in this connexion that barium fumarate crystallizes with three molecules of water while barium maleate, in which the residual affinities of the carboxyl groups are more nearly mutually satisfied, combines with only one molecule of water.

The value of these conclusions shows itself when they are applied to the hitherto unproved and in one case unknown constitutions of certain geometrically isomeric phenylhydrazones.

Thus in the case of the two camphorquinone phenylhydrazones

$$C_8H_{14}$$
 C_8H_{14}
 C_8H

it is highly probable that the form I in which the carbonyl and NHPh groups are adjacent will have the lower viscosity. Measurement showed that the lower melting, more soluble isomer had the lower viscosity and this form therefore will have the adjacent constitution I. By analogy with other camphor derivatives, though without any direct evidence, this same configuration was assigned by Forster and Zimmerli 125. The configurations of the benzaldehyde phenylhydrazones have been proved by chemical means and the adjacent form possesses the higher viscosity, showing that the effect due to potential ring formation is greater than that due to inhibition of association. This is due to the small residual affinity of the phenyl group as compared with that of the carbonyl group in the case just considered.

With the data now available one can approach the difficult problem of the configuration of the two acetaldehyde phenylhydrazones. If ordinary acetaldehyde phenylhydrazone is crystallized from alcohol containing a trace of acid it separates as the so-called β form melting at 56°. In the presence of a trace of alkali, however, the α form melting at 98° is obtained.* These changes take place so readily that the determination of the configuration by chemical methods is almost impossible, but measurement of the viscosities of solutions of the isomerides shows that the α form has the lower viscosity. Now since the methyl group undoubtedly possesses less residual affinity than the phenyl residue in the benzaldehyde phenylhydrazones the isomeride of lower viscosity should be the opposed compound. The following configurations may therefore be assigned to these compounds.

^{*} It has been suggested that the differences are due to dimorphism and not to geometrical isomerism. The different viscosities of the solutions effectually dispose of this argument.

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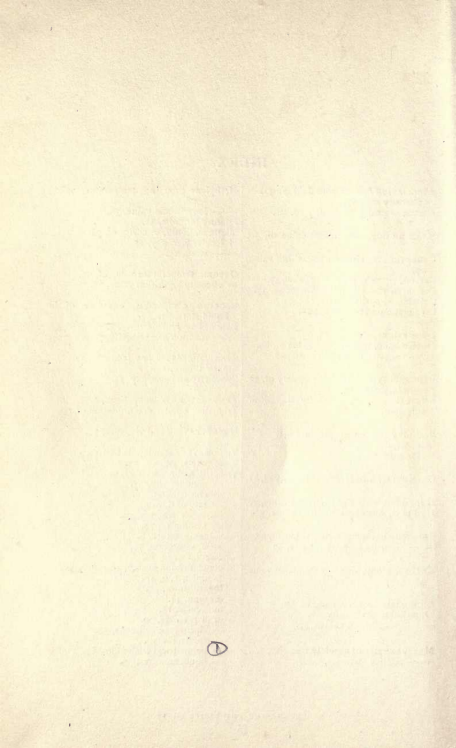
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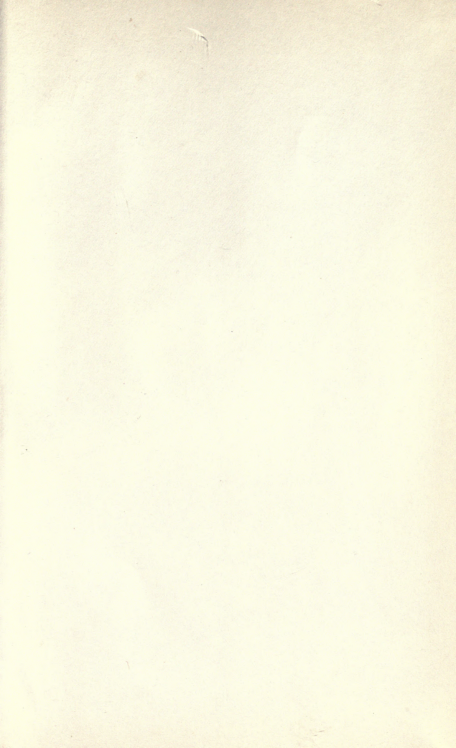
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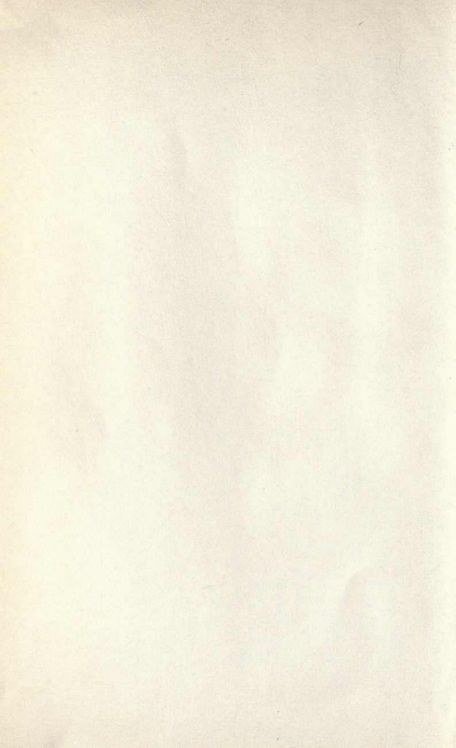
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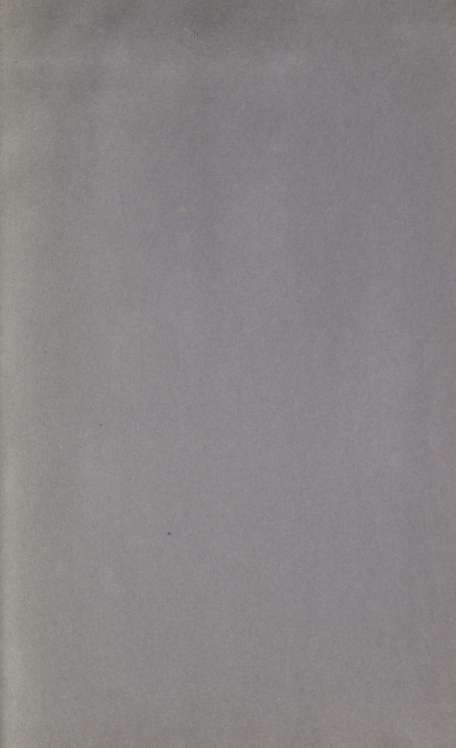
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