

g mixture
ation of
question.
atchard's
olutions.
anselow's

activity co-
efficient
t available
ilities (γ_p)
ose for the
ously sup-
positions are

and R.

7.1

7.3

7.4

7.3

7.2

7.0

7.2

7.1

6.8

6.3

7.0 is used, γ_p

$\gamma_p = 0.6, 0.7$

and the "pro-
freezing point
content; the
are multi-
thermal data
y coefficient
s by Scat-
The agree-

1.0

-0.084

-4.6

ment is quite satisfactory. Similar data for solutions up to 0.1 M have been discussed by Scatchard.²²

The writer wishes to express his indebtedness and gratitude to the Elizabeth Thompson Science Fund for providing mechanical aid in the calculations.

Summary

A report is made of the determination of the freezing points of aqueous solutions of hydrochloric acid up to a concentration of 1.0 molal, making use of a platinum thermometer as a temperature measuring device.

The activity coefficients are calculated and compared with those derived from electromotive force measurements.

TUFTS COLLEGE, MASSACHUSETTS

(CONTRIBUTION FROM THE LABORATORY OF THE LINDE AIR PRODUCTS COMPANY)

SOME PHYSICAL PROPERTIES OF VINYL CHLORIDE

By L. I. DANA, J. N. BURDICK AND A. C. JENKINS

RECEIVED SEPTEMBER 16, 1927

PUBLISHED NOVEMBER 5, 1927

Although vinyl chloride (C_2H_3Cl) has been known as a chemical compound for a number of years, owing to the difficulties attending its preparation and purification, relatively little exact work has been done with it. Recent progress in chemical synthesis has made this material available in quantity commercially and therefore it has seemed worth while to review previous work on the subject in order to ascertain the present state of knowledge of the properties.

Beyond a few scattered values of the boiling point of vinyl chloride, very little has been published concerning its physical properties. Recently the desirability of utilizing vinyl chloride as a refrigerating fluid (boiling point -13.0°) has been suggested, and hence it has appeared desirable to conduct an experimental determination of its most important physical properties. In this investigation the measurements of the vapor-pressure curve, liquid densities and freezing point are reported; in addition a few of its other physical properties have been estimated from theory.

Description of Experiments

Vapor-Pressure Curve.—Commercial samples of vinyl chloride¹ were obtained in small steel cylinders. This liquid was purified by fractional distillation in a vacuum insulated, low-temperature Hempel column and was further subjected to a fractional distillation in a closed, low-

¹ Scatchard, *THIS JOURNAL*, 47, 641 (1925). See also LaMer, *Trans. Amer. Chem. Soc.*, 51, (1927), preprint No. 58; and Van Laar, *Z. energ. allgem. Chem.*, 139, 11521.

² Material obtained from the Carbide and Carbon Chemicals Corporation, New York City.

temperature fractionation apparatus, operated with a Langmuir pump in series for the purpose of removing the traces of moisture and all non-condensable gases. Finally, the samples were tested for boiling range and pressure drop in the vapor-pressure apparatus mentioned below. On distilling off over 95% of the sample at constant temperature, the pressure drop was found to be less than 2 mm. in 760 mm. of Hg, and on distilling off at constant pressure the temperature drop was less than 0.05°. It is probable that the purity was higher than 99.9%.

The vapor pressures were measured over the temperature range from -28 to 60°, corresponding to a pressure range of 40 to 760 mm. of Hg absolute or from 0.5 to 10 atm. absolute. Below and around one atmosphere a mercury manometer, with a vacuum over one arm, was employed for the pressure measurement. From one to five atmospheres a tall mercury manometer was used, while above this pressure a dead-weight piston gage served to measure the pressure. Various forms of containers for the sample were used. In general the experimental technique was similar to that used in the measurement of the vapor pressures of some paraffin hydrocarbons,² to which work reference is made for experimental details.

The boiling point at one atmosphere was determined a number of times and was found to be -13.9, ±0.1°. Biltz³ gives the boiling range as -13 to -15°, which would seem to be much too low temperatures and too wide a range to correspond with the pure material tested in this investigation. In the following table are given the values of the pressures and corresponding temperatures observed for the vapor pressures.

TABLE I
VAPOR PRESSURES OF VINYL CHLORIDE

Temp., °C.	Press., mm. Hg.	Temp., °C.	Press., mm. Hg.	Temp., °C.	Press., mm. Hg.
-23.73	39.86	4.01	149.06	39.72	450.0
-23.02	51.30	5.53	158.2	46.80	543.4
-16.61	67.76	16.21	223.8	54.87	667.6
-13.61	76.75	25.72	302.7	60.34	758.6
- 8.32	94.60	33.53	378.9	60.34	760.3
- 1.57	122.48	39.72	449.2		

The experimental observations have been combined into the well-known Nernst⁴ equation of the form

$$\log p = a + b/T + 1.75 \log T + cT$$

in which the pressure p is expressed in atmospheres absolute and the temperature T in degrees centigrade absolute. The values for the constants are as follows:

$$a = 0.8420; b = -1150.9; c = -0.002415$$

¹ Dana, Jenkins, Burdick and Timm, *Refrigerating Eng.*, 12, 387 (1926).

² Biltz, *Ber.*, 35, 3525 (1902).

³ Nernst, "Theoretical Chemistry," Macmillan Co., New York, 1923, p. 815.

On the average by the formula

Liquid Density. The liquid density of the liquid densimeter and is similar to the Bureau of ammonia.⁵ The pycnometer was first pycnometer in coefficient of liquid was measured weight of liquid in the pycnometer determined. The m. lts. were pycnometer, constructed of After calibration meter carefully the sample and the pycnometer. Readings of stem were temperatures in a thermometer the temperature a platinum meter. By meter full a rected weight was found. Enough data able for comparison density. The coefficient of to 0.15% with the P. Corrected for the pycnometer. Nevertheless, however, since ever, the pycnometer

⁵ Cragg

air pump
re and all
for boiling
med below.
ature, the
Ig, and on
than 0.05°.

ange from
cm. of Hg
one atmos-
employed
res a tall
ad-weight
containers
was similar
ie paraffin
al details.
er of times
range as
tures and
in this in-
pressures
sures.

From,
cm. Hg
450.0
513.4
607.6
738.6
700.3

the well-

the tem-
constants

20).

p. 815.

On the average the agreement of the observed results with those computed by the formula is within the experimental error, 0.3%.

Liquid Densities.—The method employed for the determination of the liquid densities has been described in an earlier paper by the authors,² and is similar to that used by the Bureau of Standards on ammonia.³ Preliminary experiments were first made in a Pyrex pycnometer in which only the coefficient of expansion of the liquid was measured, since the weight of liquid vinyl chloride in the pycnometer was not ascertained. The final measurements were made with the pycnometer, shown in Fig. 1, constructed of Jena 10^{III} glass. After calibrating the pycnometer carefully with mercury, the sample was condensed in and the pycnometer sealed off. Readings of the liquid in the stem were observed at various temperatures from -13 to 60° in a thermo-regulated bath and the temperatures were read by a platinum resistance thermometer. By weighing the pycnometer full and empty, the corrected weight of vinyl chloride was found to be 5.9716 g. Enough data were then available for computing the absolute density. The results for the coefficient of expansion agreed to 0.1% with those obtained with the Pyrex pycnometer.

Correction has to be applied for the quantity of vapor in the vapor space above the liquid in the pycnometer. No experimental values for the vapor densities are available, however, so that these values were computed theoretically. Since, however, the maximum value for this correction is about 0.15% and an accuracy

² Craigie and Harper, Bureau of Standards, Scientific Paper, No. 430.

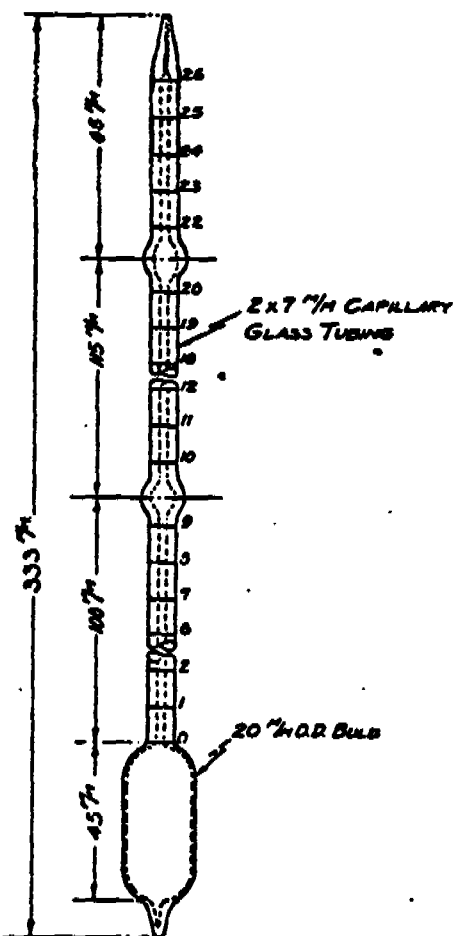


Fig. 1.—Pycnometer for liquid densities.

of 0.1% is claimed for the results, it is evident that an accuracy of 50% for the vapor density correction is sufficient. It is believed that the vapor density can be computed with much higher accuracy and as a matter of interest this has been done and the results are given later on.

RESISTANCE THERMOMETER

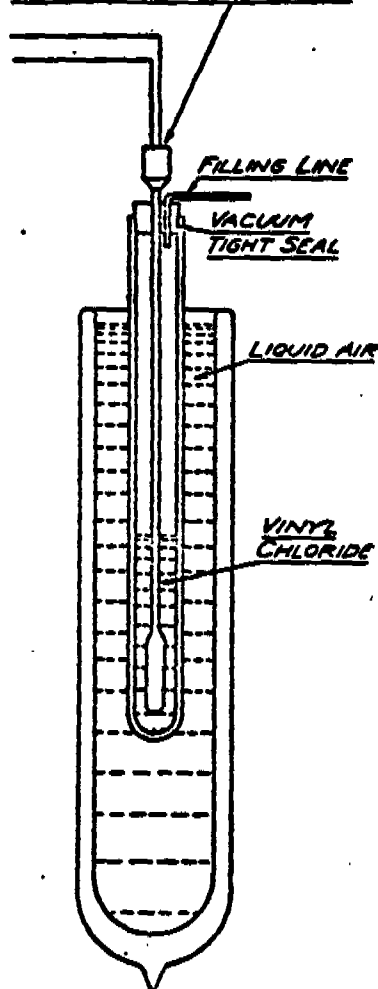


Fig. 2.—Freezing-point apparatus.

ously been calibrated at the freezing point of mercury, the sublimation point of carbon dioxide and the boiling point of oxygen.² The period of freezing lasted about seven minutes and the freezing temperature was

The liquid densities of vinyl chloride are shown in Table II.

It is believed that the results are good to 0.1%.

An empirical formula representing the results from -13 to 60° is as follows

$$d = 0.9471 - 0.001746t - 0.00000324t^2$$

in which d is the liquid density in grams per cc. and t is the temperature in degrees centigrade. The data agree with the formula to better than 0.1%.

Freezing Point.—Since relatively large quantities of vinyl chloride, purified by fractional distillation for the vapor-pressure measurements, had been prepared, it was decided to determine the freezing point by direct immersion of the thermometer in a large sample. The apparatus is shown in Fig. 2. Throughout the measurements the sample was handled in a closed system to keep out moisture. Liquid air was employed as the cooling medium. With a single tube the rate of cooling was too rapid; hence a double-walled tube was used. This also had the effect of producing a uniform temperature gradient in the freezing-point sample. Preliminary determinations with a calibrated, propane filled, low temperature thermometer resulted in the value -150° . The final and most accurate determination was made with a platinum resistance thermometer which had previously been calibrated at the freezing point of mercury, the sublimation point of carbon dioxide and the boiling point of oxygen.²

constant to within 0.1° at it is safe to claim an accu

Temp., °C.	Liq. Li
-12.06	
1.32	
13.49	
23.11	

Estimated

Since the vapor densities, it appeared so that the values might be the vapor-pressure curve with the aid of Clape

Unfortunately the curve so that the law of Clape but van Laar⁴ has been used by means of which the curve serves in other relations. By these means while the estimated

In the following table vapor, estimated by

ESTIMATED SPECIFIC

Temp., °C.
-30
-20
-10
0
+10

To compute the latent heats were used and the results graphically.

The latent heats corresponding states, carbon chloride.⁵ At the boiling while at 30° they are

⁴ Van Laar, "Z. Phys. Chem." 1901.
⁵ Holst, "Compt. Rend. Acad. Sci. Paris," 1901.

constant to within 0.1° at -150.7° . Considering the purity of the sample it is safe to claim an accuracy of 0.1° for this freezing point.

TABLE II
LIQUID DENSITIES OF VINYL CHLORIDE

Temp., $^\circ\text{C}$	Liquid density, g./cc.	Temp., $^\circ\text{C}$	Liquid density, g./cc.
-12.06	0.9692	39.57	0.8733
1.32	.9443	48.30	.8555
18.49	.9223	59.91	.8310
28.11	.8955		

Estimated Vapor Densities and Latent Heats

Since the vapor densities were required for correction terms to the liquid densities, it appeared worth while to estimate them as accurately as possible so that the values might be of use elsewhere. At the same time, since the vapor-pressure curve was measured, the latent heats could be estimated with the aid of Clapeyron's equation.

Unfortunately the critical constants of vinyl chloride are not known so that the law of corresponding states could not be employed directly; but van Laar⁶ has described methods based on empirical relationships by means of which the critical constants may be computed, which in turn serve in other relationships from which the specific volumes can be ascertained. By these methods the estimated critical pressure is 52.2 atm. while the estimated critical temperature is 142° .

In the following table the estimated specific volumes of the saturated vapor, estimated by the method of van Laar, are given.

TABLE III
ESTIMATED SPECIFIC VOLUMES OF SATURATED VAPOR OF VINYL CHLORIDE

Temp., $^\circ\text{C}$	Spec. vol., cc./g.	Temp., $^\circ\text{C}$	Spec. vol., cc./g.
-30	635	20	105.4
-20	418	30	79.7
-10	284	40	60.3
0	199	50	46.3
+10	143.3	60	36.2

To compute the latent heats the above values for the specific volumes were used and the slopes of the vapor-pressure curve were determined graphically.

The latent heats were also computed by means of the law of corresponding states, comparing with the values given by Holst⁷ for methyl chloride. At the boiling point the values by the two methods agree to 1% while at 50° they agree to 5%.

⁶ Van Laar, "Die Zustandsgleichung," Leopold Voss, Leipzig, 1924, pp. 186, 21.

⁷ Holst, *Comm. Phys. Lab. Univ. Leiden*, No. 144c.

TABLE IV
ESTIMATED LATENT HEATS OF VINYL CHLORIDE

Temp., °C.	Range of vapor pressure mm. of Hg./°C.	Latent heat, cal./g.	
		By Chappuis's equation	By corresponding states
-20	25.4	85.4	85.7
-10	35.4	81.1	81.1
0	47.1	81.2	81.0
+10	61.8	79.8	81.7
20	79.7	77.7	80.2
30	100.2	76.2	78.8
40	124	73.2	76.6
50	151	70.1	74.4

Summary

The vapor pressure of vinyl chloride was measured from -28 to 60°. The normal boiling point was found to be -13.9, $\pm 0.1^\circ$.

The liquid densities were measured over the temperature range -13 to 60°.

The freezing point was found to be -150.7, $\pm 0.1^\circ$.

Estimated values of the vapor densities and latent heats were computed from theoretical relationships.

BUFFALO, NEW YORK

NOTE

Calculation of Heats of Combustion.—Kharasch and Sher¹ have applied the electronic conception of valence to the calculation of heats of combustion of organic compounds. They find that the energy change which accompanies the displacement of one valence electron from the position which it occupies in methane, a carbon chain or the benzene nucleus to the position after combustion in carbon dioxide is 26.05 kg. cal. per mole. Thus the heat of combustion of a hydrocarbon is given by the product $26.05 \times N$, where N is the number of electrons displaced.² Electrons associated with bonds other than C—C and C—H are accompanied by an additional energy change during combustion. To the term $26.05 N$, Kharasch and Sher add 26 kg. cal. for each aliphatic ether group, 13 kg. cal. for each ethylenic linkage, primary alcohol, ester or aromatic ether group and 6.5 kg. cal. for each secondary alcohol or ketone group. No correction is added for tertiary alcohol, phenol or acid groups. In most cases this method gives good results, but for aromatic compounds the calculated values are uniformly higher than the experimental values and the

¹ Kharasch and Sher, *J. Phys. Chem.*, 29, 625 (1925).

² Each bond corresponds to two electrons. In oxygen compounds the two oxygen bonds are disregarded. Thus N is 8 for methane, 30 for benzene, 58 for phenyl benzoate, etc.

deviations become greater necessitates a correction for this correction follows:

The electrons associated (μ electrons) will be one the other twelve (λ electrons) 26.05 kg. cal. per mole; the compound being burned at constant pressure of is given by the equation group + 19.5 for each primary alcohol or ester + 3.2 for each phenol ketone group.

Of the 57 aromatic compounds whose heats of combustion new method gives results the same in 7 cases. mental and calculated shows per cent. difference for 171 aromatic compounds.

DIFFERENCE BETWEEN C.

Aromatic compounds containing C and H, or C, H and

Saturated hydrocarbons
Ethylenic hydrocarbons
Phenols
Ethers
Aldehydes
Ketones
Acids
Esters

Total

CONTRIBUTION FROM THE
DEPARTMENT OF CHEMISTRY
THE PENNSYLVANIA STATE
UNIVERSITY, PENNSYLVANIA
RECEIVED MARCH 11, 1927
PUBLISHED NOVEMBER 9, 1927

³ See Pauling, *THIS*

⁴ λ and μ are the number of electrons in the benzene nucleus contains anthracene; $\lambda = 40$, $\mu =$

deviations become greater as the number of benzene nuclei increases. This necessitates a correction for each benzene nucleus. The method proposed for this correction follows.

The electrons associated with three of the bonds of the benzene nucleus (μ electrons) will be assumed to differ in energy relations from those of the other twelve (λ electrons).¹ There is an energy change of 24.85 and 26.05 kg. cal. per mole, respectively, for each μ and λ electron present in the compound being burned. The heat of combustion in kg. cal. per mole at constant pressure of compounds containing C and H, or C, H and O, is given by the equation⁴ $Q_p = 26.05\lambda + 24.85\mu + 22.7$ for each ether group + 19.5 for each aldehyde group + 13 for each ethylenic linkage, primary alcohol or ester group + 6.5 for each secondary alcohol group + 3.2 for each phenol or tertiary alcohol group + 0.0 for each acid or ketone group.

Of the 57 aromatic compounds containing C and H, or C, H and O, whose heats of combustion were calculated by Kharasch and Sher, the new method gives results closer to the experimental values in 37 cases and the same in 7 cases. The average per cent. difference between experimental and calculated values is lowered from 0.52 to 0.32%. Table I shows per cent. difference between calculated and experimental values for 171 aromatic compounds of different types.

TABLE I

DIFFERENCE BETWEEN CALCULATED AND EXPERIMENTAL HEATS OF COMBUSTION

Aromatic compounds containing C and H, or C, H and O	No. of calculations	Difference between calculated and experimental values	
		Total %	Average %
Saturated hydrocarbons	26	8.4	0.32
Ethylenic hydrocarbons	11	2.7	0.25
Phenols	24	10.7	0.44
Ethers	23	8.0	0.36
Aldehydes	12	7.3	0.61
Ketones	15	5.5	0.57
Acids	33	13.3	0.40
Esters	28	8.3	0.30
Total	171	64.2	0.35

CONTRIBUTION FROM THE
DEPARTMENT OF CHEMISTRY,
THE PENNSYLVANIA STATE COLLEGE,
STATE COLLEGE, PENNSYLVANIA

RECEIVED MARCH 11, 1927
PUBLISHED NOVEMBER 5, 1927

J. H. SEYMOUR

¹ See Pauling, *THIS JOURNAL*, 48, 1132 (1926).

⁴ λ and μ are the numbers of λ and μ electrons present. In aliphatic compounds there are no μ electrons and λ is the same as N in Kharasch and Sher's equation. Each benzene nucleus contains 6 μ electrons. Thus $\lambda = 38$ for hexane; $\lambda = 48$, $\mu = 18$ for anthracene; $\lambda = 46$, $\mu = 12$ for phenyl benzoate, etc.