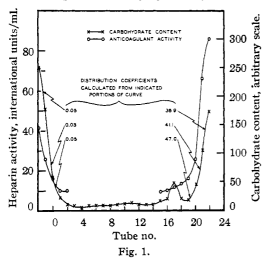
the application of the Craig² counter-current distribution technique several samples of sodium heparinate were distributed between amyl alcohol and an aqueous buffer at ρ H 6.5, using 2.5% laurylamine as a "carrier."³ After the distribution was completed, the material in the several solvent phases was recovered by shaking each with 0.5 Mdipotassium hydrogen phosphate.

Initially, peaks representing three fractions were located by means of anthrone⁴ (Curve A). Only the two larger fractions appear to have anticoagulant activity (Curve B). The approximate homogeneity of the two outer fractions is indicated by the relative constancy of the distribution coefficients as calculated² over several adjacent tubes. These coefficients are shown by the numbers superimposed on the graph of Fig. 1:



Separation of a 1-g. sample of sodium heparinate yielded, after removal of salts by dialysis, two major fractions. Preliminary data on these fractions are as follows (dry basis):

	Low coeff. fraction	High coeff. fraction
Ash, %	23.39	33.88
Potassium (calcd. from ash), $\%$	10.46	15.19
Nitrogen, %	3.33	2.93
Sulfur, %	8.42	13.33
S/N ratio	1.10	1.98
K/S ratio	1.02	0.93
Activity (intl. u./mg.)	$59 \neq 6$	215 ± 22

We wish to express our indebtedness to Dr. John Burke for the physiological assays and to Mr. Joseph Alicino for the microchemical analyses.

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(2) L. C. Craig, J. Biol. Chem., 155, 519 (1944); B. Williamson and L. C. Craig, *ibid.*, 168, 687 (1947).

(3) A. E. O'Keeffe, M. A. Dolliver and E. T. Stiller, THIS JOURNAL, in press.

(4) D. L. Morris, Science, 107, 254 (1948).

SYNTHESIS OF ISOPROPYLCYCLOPROPANE

Sir:

The preparation of ethylcyclopropane and isopropylcyclopropane from methylcyclopropyl ketone has been reported recently by Van Volkenburgh, Greenlee, Derfer and Boord.¹ Because of the publication of their paper, it is desirable to report that we have been investigating the syntheses of several alkylcyclopropanes from methylcyclopropyl ketone, the results of which are being withheld until a series of hydrocarbons is completed. Our method of hydrogenation of isopropenylcyclopropane to yield essentially pure isopropylcyclopropane may be of immediate interest, however, since Van Volkenburgh, *et al.*, were able to obtain a product of only 85 mole per cent. purity by their procedure.

Dimethylcyclopropylcarbinol and isopropenylcyclopropane were prepared in essentially the same manner as previously described.¹ Hydrogenation isopropenylcyclopropane $(n^{20}D \ 1.4256)$ at of 1500 to 2000 p.s.i. of hydrogen in the presence of a commercial barium-promoted copper chromite catalyst² at 100 to 130° was found to yield an extremely pure product with little or no ring cleavage. Fractionation of the hydrogenation product at 50-plate efficiency through a glass helix-packed column yielded distillate 80% of which had a refractive index of 1.3863-1.3864 (index range including forerun and residues was 1.3856 to 1.3864). Freezing curves were determined for consecutive cuts of this material and were found to vary from -113.30 to -113.17° . Physical constants of the purest cut, based on the freezing points, are given in the table with our constants for isopropenylcyclopropane:

·	Table I	
	lsopropenyl- cyclopropane	Isopropyl- cyclopropane
F. p., °C.	-102.34 m. p.	-113.17
B. p., °C. at 760 mm.	70.33	58.37
<i>n</i> ²⁰ D	1.42550	1.38639
d^{20} , g./ml.	0.75153	0.69829
Carbon, { Calcd. % { Found	87.7	85.6
% Found	87.6	85.6
Hydro- gen, % { Caled. Found	12.3	14.4
gen, $\% \setminus$ Found	12.2	14.4

From the freezing point of our material it appears that the calculation of purity and the "100 per cent. pure" freezing point of reference 1 for isopropylcyclopropane are in error. This is not surprising since the accuracy of such calculations decreases rapidly as purity decreases. The "100 per cent. pure" freezing point, calculated from our data by the geometrical method of Taylor and Rossini,⁸ is estimated to be $-113.07 \pm 0.05^{\circ}$. As-

⁽¹⁾ Van Volkenburgh, Greenlee, Derfer and Boord, THIS JOURNAL, 71, 172 (1949).

⁽²⁾ E. I. du Pont de Nemours, Ammonia Division, Wilmington, Delaware.

⁽³⁾ Taylor and Rossini, J. Research Natl. Bur. Standards, 32, 197 (1944).

Ν

suming that this is the true freezing point and that the Δt /mole per cent. impurity is 0.20° as given in reference 1, the purity of their sample is estimated to be 74% and ours is 99.5 mole per cent.

The infrared spectrum of the sample of isopropylcyclopropane used to determine the physical constants gave the absorption maxima in Table II.

TABLE II							
λ, microns		3.40	4.92	5.26	6.77	6.96	7.21
Transmission,	%	5	72	74	6	19	11
λ, microns		7.30	7.45	7.58	7.74	8.21	8.57
Transmission,	%	11	58	39	14	33	18
λ, microns		8.84	9.05	9.56	9.78	9.94	10.46
Transmission,	%	40	45	15	10	17	14
λ, microns			10.82	11.01	11.39	12.18	13.48
Transmission,	%		35	22	12	13	46
NATIONAL ADVISORY COMMITTEE FOR Aeronautics V. A. Slabby							
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RECEIVED FEBRUARY 17, 1949

POLAROGRAPHIC DETERMINATION OF THE MOLECULAR WEIGHT OF SERUM ALBUMIN BY ITS EFFECT ON THE DIFFUSION CURRENT OF METHYL ORANGE

Sir:

The purpose of this note is to describe the great difference in the effect of gelatin and serum albumin on the diffusion current of 10^{-4} M methyl orange in a 0.01 N phosphate buffer of pH of 7.2, or in very dilute potassium chloride solutions and to show that the complex formation between serum albumin and methyl orange can be studied quantitatively by the polarographic method.

The polarographic behavior of methyl orange in water in the presence of various supporting electrolytes without proteins shows several complicated features which are the subject of further work. Under the conditions described below in the presence of proteins these complications were eliminated.

Å small amount of gelatin (0.01%) reduces i_d by 10%, but larger amounts (up to 1%) exert hardly any further effect. The addition of serum albumin causes a continuous reduction of i_d until the protein concentration is about 0.5%. These observations can be interpreted in agreement with the results of Klotz,¹ et al., who found that serum albumin combines, but that gelatin does not combine with methyl orange. Polarographically we found that at a concentration of about 0.5% serum albumin practically all the methyl orange $(10^{-4} M)$ has combined with the protein. Application of the Ilkovič equation gave a value of the diffusion coefficient of the complex of 9.05 \times 10⁻⁷ cm.² sec.⁻¹ at 25°. Using this value in the Stokes-Einstein equation yields a value of 66,000 for the molecular weight, which is in good agreement with reported

(1) I. M. Klotz, F. M. Walker and R. B. Pivan, THIS JOURNAL, 68, 1486 (1946); Klotz, *ibid.*, 68, 2299 (1946); Klotz, Triwush and Walker, *ibid.*, 70, 2935 (1948); Klotz and Urguhart, J. Phys. Coll. Chem., 53, 100 (1949). values of molecular weight (68,000 to 70,000) of serum albumin. The decrease of i_d of methyl orange by serum albumin at various concentrations can be made use of in the calculation of the binding constant.

The effect of serum albumin on the diffusion current of $10^{-4} M$ methyl orange in 0.01 to 0.05 N potassium hydroxide solutions is negligibly small. This again is in agreement with the results of Klotz,¹ who found that serum albumin does not combine with the dye at these high ρ H values.

More extensive polarographic studies on the effect of proteins in the native and denatured state on various dyes and other compounds are planned.

The authors acknowledge a grant from the Public Health Service, which enabled them to carry out this work.

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RECEIVED MARCH 7, 1949

INHIBITING EFFECT OF NITRIC OXIDE ON THE THERMAL DECOMPOSITION OF DIETHYL KETONE Sir:

Previous studies on the thermal decomposition of acetone¹ and methyl ethyl ketone² established the fact that the rates are uninhibited by the addition of nitric oxide. Instead, small amounts of nitric oxide, probably due to its reaction with the ketene intermediate, cause marked catalysis and this effect increases with the amount of inhibitor added. Propylene, on the other hand, reduces the rates of both reactions to a limiting value, indicating that free radicals are indeed present.

In our present investigations we find that small amounts of nitric oxide reduce the rate of decomposition of diethyl ketone to a definite, limiting value which is readily reproducible. The figure, where ρ is the rate with a given amount of inhibitor, ρ_0 , that with none, and ρ_{∞} , the limiting rate, shows that the amount of inhibition is essentially independent of ketone pressure, indicating that the radical producing reaction must be of the first order. No catalytic effect was observed up to 20 mm. partial pressure of nitric oxide. The mean chain length is 2.0, the same value as determined for the thermal decomposition of methyl ethyl ketone with propylene. Preliminary experiments reveal that the decomposition of methyl propyl and methyl isopropyl ketones are uninhibited by nitric oxide.

The inhibiting effect of nitric oxide on diethyl ketone is in striking contrast to its effect on other ketones investigated so far. One would logically expect the primary decomposition reactions to be basically similar, and certainly methyl ethyl and diethyl ketones should initially produce the same kind of chain propagating radicals. This surpris-

(1) Smith and Hinshelwood, Proc. Roy. Soc. (London), **A183**, 33 (1944).

(2) Waring and Mutter, THIS JOURNAL, 70, 4073 (1948).