The carbonate fractions are seen to contain on the order of 0.3-0.35 part per million of protactinium, which is mainly bound with the acid-insoluble silicious portion of this fraction. The ionium, present to some 15-20 parts per million, is found in the readily soluble portion of the Fraction. Information furnished about the carbonate fraction is that it is 215 parts by weight for 580 parts by weight of the crude ore—*i. e.*, 37%. For pitchblende which is 65% by weight U_3O_8 , this indicates that on the order of two-thirds the original protactinium content of the ore was dissolved in the sulfuric-nitric acid treatment, and precipitated by the carbonate treatment, with more than three-fourths of the ionium content following the same path. The apparent high level of ionium in the sulfate residues is in part at least a reflection of the fact that much of the soluble portion of the ore has been removed, giving a bulk concentration. For example, removal of the uranium alone gives a concentration of about threefold. It is interesting to note that the radioactivity surviving to the ether residues is practically entirely ionium (together with a certain amount of uranium). It is difficult to assign a concentration factor for this material, but it must be rather large due to the previous treatments.

On the basis of the results obtained, a number of milligrams of protactinium and approximately one-half gram of ionium (with three times as much Th²³²) have been obtained in pure state, from a carbonate fraction.^{9,10,11} The yields obtained for the protactinium correspond to 50-60% of the material indicated by preliminary analysis. No attempt was made to obtain complete recovery of ionium.

(9) K. Kraus and Q. Van Winkle, "PPR," Vol. 17B, No. 6.2.

(10) R. Larson, L. I. Katzin and E. Hausman, *ibid.*, Vol. 17B, No. 6.3.

(11) R. C. Thompson, Q. Van Winkle and J. G. Malm, *ibid.*, Vol. **17B**, No. 6.4.

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The Melting Point of Lead Bromide

By L. M. KNOWLES

The literature references to lead bromide may be divided into two general groups which report the melting point at approximately $370^{\circ 1}$ and 495° ,² respectively. Recent investigations in this Laboratory, which are summarized below, reveal that the cause of this discrepancy was the failure of many investigators to procure pure lead bromide. We have determined the melting point of lead bromide to be $370.0 \pm 0.2^{\circ}$ for various rates of heating and cooling; in contact with platinum, quartz, Vycor, Pyrex and magnesium oxide crucibles; and under atmospheres of air and nitrogen, and *in vacuo*. Each sample, tested chemically and spectroscopically both

(1) Ramsay and Eumorfopoulos, Phil. Mag., 41, 360 (1896); Herrmann, Z. anorg. allgem. Chem., 71, 287 (1911); C. Sandonnini, Gass. chim. ital., 41, II, 145 (1911); Lorenz and Fox, Z. physik. Chem., 63, 109 (1908); Tubandt and Eggert, Z. anorg. Chem., 110, 219 (1920); Mönkemeyer, Neues Jahrb. Mineral. Geol., 22, (B. B.) 1 (1906); Calingaert, Lamb and Meyer, THIS JOURNAL, 71, 3709 (1949).

 Czepinski, Z. anorg. allgem. Chem., 19, 208 (1899); Goodwin and Kalmus, Phys. Rev., 28, 1 (1909); Helfstein, Z. anorg. Chem., 23, 255 (1900); Weber, *ibid.*, 21, 305 (1899); Ehrhardt. Ann. Physik, 24, 215 (1885); Camelley, J. Chem. Soc., 33, 278 (1878). before and after each thermal run, contained at least 99.7% lead bromide. Temperature measurements with iron-constantan and platinumplatinum 10% rhodium thermocouples, gave a consistent melting point of $370.0 \pm 0.2^{\circ}$. In no case was a value near 495° obtained.

The literature references to lead bromide are wholly inadequate in reporting methods of preparation or analyses. Consequently, it is impossible to determine the true composition of the material reported to melt near 495°. However, recent unpublished data from our laboratory on the lead bromide-lead oxide system show a persistent transformation temperature at 497° for compositions of 35-55 mole % lead oxide. It is this oxybromide transition temperature which the authors reporting the higher value probably measured.

The accepted correlation of the heat capacities and heat of fusion of lead bromide is the work of Kelley.³ Unfortunately Kelley's calculations are based on the physical properties of a lead bromide sample reported to melt at 488°. Consequently, the accepted specific heat equations and other thermodynamic data apply to the basic salt rather than to pure lead bromide.

(3) Kelley, U. S. Bureau of Mines Bull., 371, 31 (1934); 383, 61 (1935); 476, 95 (1949); Quill, "The Chemistry and Metallurgy of Miscellaneous Materials, Thermodynamics," McGraw-Hill Book Co., Inc., New York, N. Y., 1950; Natl. Bur. Standards, Selected Values of Chemical Thermodynamic Properties, Table I: 27-3, Sept. 30, 1948; Table II: 27-2, Sept. 80, 1948.

ETHYL CORPORATION DETROIT, MICHIGAN

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The Interaction of Diacyl Peroxides with Diazomethane

By JOHN E. LEFFLER¹

Diacyl peroxides might be expected to react with diazomethane either by analogy with acid anhydrides² or by analogy with sources of free radicals like hexaphenylethane.³ However, no reaction was found between benzoyl peroxide and diazomethane in ether solution, the peroxide being recovered unchanged. This result may be due to a lesser degree of positive charge on the carbonyl group of the peroxide. The resonance form shown below would be of lower energy in the peroxide than in the acid anhydride, since in the anhydride the positive charge would be adjacent to the positive end of a carbonyl group.



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(3) Schlenk, Ann., 394, 183 (1912).

⁽²⁾ Bradley and Robinson, THIS JOURNAL. 52, 1558 (1930).