mediately rearrange to ocimene (VI).⁷ Other possible reactions of the intermediate and the stereochemical properties of the intermediate were not considered.

Application of the same concept to the pyrolysis of β -pinene (II) leads to the biradical (VII) which unlike V is asymmetric. Models show that hydrogen atoms from the isopropyl residue (positions (9) and (10)) may be transferred either to position (2) or (7). One would expect transfer to position (7) with the formation of an internal olefin (IX) to be favored over that to position (2) which leads to an exo-structure (VIII). Limonene (IX) so formed should be optically pure. Recombination of the biradical (VII) unlike that of V would regenerate an unracemized pinene. Bond fission between carbon atoms (3) and (4) would lead to myrcene (X).

Goldblatt and Palkin⁸ report the vapor phase pyrolysis of β -pinene to lead to myrcene (X) and *l*-IX of "singularly high rotation." The rotation of the isolated limonene was about 10% under the maximum possible.⁹ No VIII was reported. The small deficit in rotation of IX might originate from the presence of a little VIII.

Hunt and Hawkins¹⁰ report that the isomerization of β -pinene proceeds as a first order reaction unaffected by small quantities of hydroquinone. Dilution with *dl*-IV does not affect the rate constants. The reported activation energy is about 48 kcal.

Data which critically test the prediction that recovered β -pinene would be unracemized have not been reported.

The proposed mechanism provides a stereochemical correlation between l- β -pinene and llimonene.

(7) The formation of ocimene was later reported, F. O. Rice, U. S. Patent 2,190,369, February 13, 1940. No other investigator has reported ocimene.

(8) L. A. Goldblatt and S. Palkin, THIS JOURNAL, 63, 3517 (1941).

(9) Ref. 4, p. 152 gives 126.8°; ref. 8 reports 114.9°.
(10) H. G. Hunt and J. E. Hawkins, THIS JOURNAL, 72, 5618 (1950).

DEPARTMENT OF CHEMISTRY

NORTHWESTERN UNIVERSITY EVANSTON, ILLINOIS RECEIVED FEBRUARY 9, 1951

The Association of Tri-p-xenylmethyl

By Ting-li Chu and S. I. Weissman

Recent measurements of the magnetic susceptibility of solutions in toluene of tri-*p*-xenylmethyl¹ show that over the temperature range 0 to 60° and the concentration range 0.018 to 0.032 mole per liter the molar susceptibility is constant. Use of the conventional correction for diamagnetism leads to a constant degree of association of 0.20.

In connection with other work we have investigated the absorption spectra of solutions of tri-pxenylmethyl in toluene. We find that the Beer-Lambert law is obeyed over the concentration range 2×10^{-4} to 4.5×10^{-3} mole per liter, and that the molar extinction coefficient is independent of temperature over the range 10 to 40°. The shape of the absorption curves obtained by us is

(1) P. W. Selwood and R. M. Dobres, THIS JOURNAL, 78, 3860 (1950).

identical with those already given in the literature.^{2,3} The molar extinction coefficient at the peak at 6150 Å. is 1525. These results indicate that tri-p-xenylmethyl is not associated in toluene solution.

Our solutions were prepared on the vacuum bench by the method already described by Lewis, Lipkin and Magel.² A solution of a weighed sample of pure tri-p-xenylcarbinol4 in freshly distilled acetyl chloride was refluxed for 12 hours. The resulting solution was made up to a known volume and an aliquot pipetted into a Pyrex absorption cell. The cell was quickly attached to the vacuum line where all subsequent operations were carried out. The acetyl chloride was evaporated and the solid halide subjected to prolonged evacuation at about 80°. Any trace of acid remaining in the system was neutralized by introduction of triethylamine vapor. A measured volume of toluene and a trace of triethylamine (each dried over sodiumpotassium alloy) were distilled into the cell containing the halide, and silver amalgam was then introduced by a vacuum dumper. The cell was sealed off under vacuum, and reduction effected by several minutes shaking of the solution with the amalgam. The free radical solutions prepared in this way remained unchanged during several days storage in the dark at room temperature.

We may report, in addition, that α -naphthylphenyl-p-xenylmethyl also obeys Beer's law with a temperature-independent extinction coefficient. A mixture of this free radical with tri-p-xenylmethyl, however, does not obey Beer's law, nor is the absorption at fixed concentration independent of temperature. The two free radicals apparently associate in toluene solution.⁵ According to our data ΔF^0 for the association is about -4 kcal. per mole at 25°, while ΔH is about -12 kcal. per mole.

(2) G. N. Lewis, D. Lipkin and T. T. Magel, *ibid.*, 66, 1579 (1944).

(3) It has been demonstrated (ref. 2) that unless extreme precautions are taken, tri-*p*-xenylmethyl is converted to another free radical of unknown constitution. The pure tri-*p*-xenylmethyl is blue-green, the other free radical red. In a recent paper Dobres and Selwood (THIS JOURNAL, **72**, 5731 (1950)) describe tri-*p*-xenylmethyl as "deep burgundy." Apparently our blue-green substance and the one studied by Dobres and Selwood are different.

(4) This material was generously supplied by Professor Lipkin. It is a portion of the preparation used in the work of Lewis, Lipkin and Magel.

(5) A result pointing in the same direction has been obtained by C. S. Marvel and Chester M. Himel, THIS JOURNAL, **64**, 2227 (1942).

DEPARTMENT OF CHEMISTRY

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Rate of Oximation and Extent of Enolization of Alkyl Aryl Ketones¹

By Mary J. Craft,² Billy F. Landrum,² Edgar C. Suratt² and Chas. T. Lester³

We have previously reported the rate of oxime formation of four series of ketones.⁴ We have ex-

(1) This work was made possible through a contract with the Office of Naval Research.

 (2) Taken from respective Ph.D. Dissertations, Emory University, 1950.

(3) Responsible co-author.

(4) (a) E. C. Suratt, J. R. Proffitt, Jr., and C. T. Lester, THIS JOURNAL, 72, 1561 (1950); (b) M. J. Craft and C. T. Lester, *ibid.*, 73, 1127 (1951).