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The Solvolysis of 6-Methylenecyclodecyl Toluene-p-sulphonate

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Several recent reports², have been concerned with transannular bridging reactions of exomethylene ketones and toluene-p-sulphonates (Ia and Ib) to form adamantan-1-ol (II) and its derivatives. The reactions of (I) closely parallel an analogous ring closure of β -caryophyllene derivatives4 in that both systems can be regarded as involving a substituted cyclononylmethylene as the basic skeleton. We report results which further suggest that the reaction of medium-size⁵ cyclanes which incorporate an appropriately positioned exo-methylene unit and a (potential) positive centre results in formation of bicyclic systems with substitution at the bridgehead.

6-Methylenecyclodecanol (III), m.p. 40-41°, was prepared in 75% yield by reaction of methylenetriphenylphosphorane with 6-hydroxycyclodecanone in dimethyl sulphoxide at 65°. Reaction of (III) with toluene-p-sulphonyl chloride in pyridine at -10° gave the toluene-p-sulphonate ester (IV). The ester (IV) decomposed within one hour at room temperature (in a vacuum desiccator) to a mixture⁸ of components. Successful recrystallization was achieved using hexaneether at isopropyl alcohol-dry ice temperatures. The ester (IV) appears to have a double melting point, m.p. ca. 45° and 85°. Whether this phenomenon is due to a rearrangement9 or other causes cannot vet be explained.

Immediately after preparation and purification, (IV), in 75% acetone-water, containing an excess of calcium carbonate, at room temperature rapidly evolved carbon dioxide. The product, isolated in 90% yield, was shown (by alumina-column and thin-layer chromatography) to be a mixture of an alcohol (93%) and an olefin (7%). The alcohol, m.p. $55-56^{\circ}$, was cis-bicyclo[4,4,1]undecan-1-ol10 (V). The olefin fraction consisted of two, possibly three, components; the major one had an identical retention time with bicylo-[4,4,1]undec-1-ene¹⁰ (VI).

Under similar conditions, (IV) was solvolyzed

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⁸ Gas-liquid chromatography indicated the presence of at least six partially resolved components.

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 Cf., e.g., P. de Mayo, "The Mono and Sesquiterpenoids," Interscience Publishers, New York, 1956, pp. 292-296. ⁵ The medium-size cyclanes are generally defined as those containing eight to eleven members.

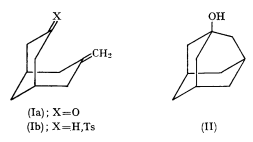
⁹ Cf. The rearrangement of trans-cyclodec-5-en-1-yl p-nitrobenzoate observed by H. L. Goering and W. D. Closson,

J. Amer. Chem. Soc., 1961, 83, 3511.

10 W. G. Dauben, T. L. Westman, and F. T. Bond, Abstracts, 141st Meeting, American Chemical Society, Washington, D.C., March 28, 1962, pp. 29-30; cf., also E. L. Eliel, "Stereochemistry of Carbon Compounds," McGraw-Hill Co., 1962, New York, p. 300.

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$$\begin{array}{cccc}
CH_2 & OH \\
OR & H \\
(III) R=H \\
(IV) R=OTs
\end{array}$$
(VI) (VI)

in glacial acetic acid or acetic acid-sodium acetate to a mixture of 74% (V), 19% (VI), and 7% of a mixture of two olefins. These latter two unidentified olefins had g.l.c. retention times different from those for cis- and trans-5-methylenecyclodecene. Again this reaction suggests a marked degree of transannular reaction. In contrast to cis- and trans-5-cyclodecen-1-yl p-nitrobenzoates,9 5-methylenecyclodec-1-yl p-nitrobenzoate was recovered unchanged after warming (85°) for 3 hours in a 75% dioxan-water solution.

The rapidity with which (IV) is solvolyzed and the high yield of bicyclic product further amplifies the fact that transannular participation of an exo-methylene, when appropriately positioned (or constrained), in solvolysis of medium-size cyclanes is apparently general.

Solvolytic rate measurements are being examined in order to answer questions about effects of ring-strain, solvent nucleophilicity and conformation.

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