[1,3] Shifts of Potassium Alkoxides: the Bicyclo[3.2.0]heptene-Norbornene Transformation

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Summary exo-Bicyclo[3.2.0]hept-2-en-7-ol can be isomerized to exo-norbornenol on treatment with potassium hydride in tetrahydrofuran at room temperature.

Of all the [1,3] sigmatropic shifts of hydrocarbons¹ perhaps the best known is the classic case reported by Berson² $(1) \rightarrow (2)$. The reaction was shown to be concerted, with inversion of configuration of the migrating group. The recent report by Evans³ that oxy-Cope [3,3] sigmatropic rearrangements are strongly accelerated for potassium salts prompted us4 to investigate the effect of a potassium alkoxide substituent on a [1,3] sigmatropic⁵ shift. Although dramatic accelerations were observed for isomerizations of potassium salts such as $(3) \rightarrow (4)$, the mechanism of such reactions remains unclear. These formal [1,3] sigmatropic shifts can be considered 'reverse Grignard-type' reactions. Indeed in the case of $(3) \rightarrow (4)$ crossover products have been observed, although this and analogous reactions6 have been observed in protic solvents, hardly a suitable environment for an allyl anion. In the course of exploratory studies on the mechanism of such alkoxide accelerated shifts, we have prepared the exo-(5) and endo-alcohols (6) via the recently reported method of Fleming.⁷ Compound (5) isomerized (as the potassium salt) to a mixture of 67% (7), 8% (8), and 25% (5) (by g.l.c.) after 3 h at room temperature. Under the same conditions the endo-alcohol (6) was stable.† This result is consistent with a concerted mechanism.8 The exo-substituted compound isomerizes with (mainly) inversion at C-7 whereas the endo-substituted isomer (6) does not react, since the clockwise rotation about C-6-C-7 required by a concerted isomerization would force the alkoxide substituent into the cyclopentane ring. Under more vigorous conditions (refluxing tetrahydrofuran or in the presence of the

307 °C

ACO
H

(1)

(2)

Me
Ph

$$\frac{0 \text{ °C}}{\text{tetrahydroturan}}$$

(3)

(4)

 R^{1}

(5); R^{1} = H, R^{2} = OH
(6); R^{1} = OH, R^{2} = H

(8); R^{1} = OH, R^{2} = H

macrocyclic polyether 18-crown-6) compound (6) does isomerize yielding (7) and (8) in a 6:1 ratio. This again is consistent with the results of Berson who found that *endo*-substituted bicyclo[3.2.0]heptenes isomerize with retention. A concerted mechanism still may not be operating here since the thermodynamic ratio of (7):(8) is close to 6:1.

[†] The norbornenols (7) and (8) are stable under the reaction conditions.

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These results do show however that many well known thermal rearrangements may be accelerated by alkoxide substituents on the C-C bond undergoing cleavage in the reaction. Further examples of this process have been observed and will be reported elsewhere.

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⁴ S. R. Wilson, D. T. Mao, K. M. Jernberg, and S. T. Ezmirly, Tetrahedron Letters, 1977, 2559.

⁵ Throughout this discussion the term 'sigmatropic shift' is used to indicate the overall bonding change and not to imply a concerted mechanism (R. B. Woodward and R. Hoffmann, 'The Conservation of Orbital Symmetry,' Academic Press, New York, 1970).

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