

A Rearrangement of $\beta\beta$ -Disubstituted Mannich Bases

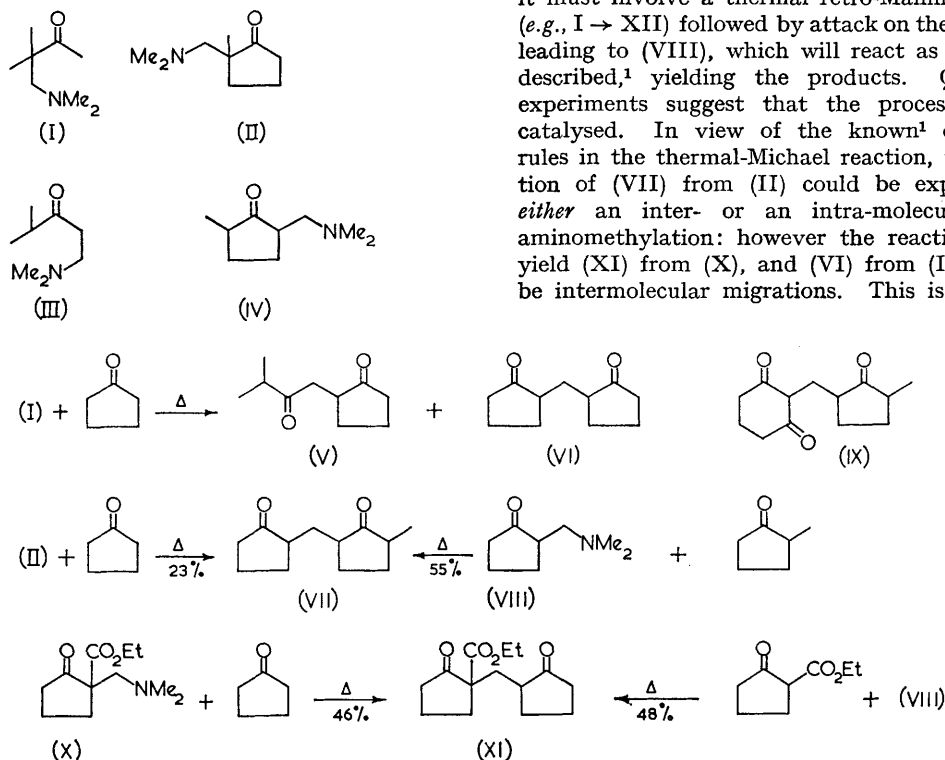
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OUR interest in Mannich bases of ketones as enone-equivalents in the *thermal*-Michael reaction¹ has currently compelled us to investigate and verify their structures in certain cases where there has been disagreement.² For example, it can be shown by n.m.r. studies that the Mannich bases prepared from isopropyl methyl ketone and 2-methylcyclopentanone are formulated correctly as (I)^{2,3} and (II)^{2,4}. Nevertheless, we find that in

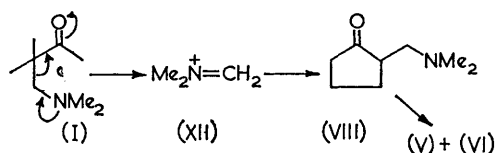
the thermal-Michael reaction these bases behave as if they had the alternate structures (III) and (IV). Thus the condensation of (I) with cyclopentanone affords (V) in 45% yield, and the reaction of (II) with cyclopentanone affords (VII), the product also obtained by the thermal condensation of 2-methylcyclopentanone with (VIII). Under similar conditions (II) reacts with dihydroresorcinol to give a 60% yield of (IX), and

(X) reacts with cyclopentanone giving (XI). The structure of (XI) is corroborated by independent synthesis from (VIII) and 2-ethoxycarbonylcyclopentanone under thermal conditions, or (in 60% yield) by the classical Michael condensation of the latter with 2-methylenecyclopentanone. In each of these reactions, the structure of the product has been established by n.m.r. or u.v. spectroscopy and analysis.



In seeking to explain these findings, it is important to recognise that crude Mannich bases often contain small amounts of isomers,² e.g., (III) and (IV), which could account for the above reaction products. However this explanation cannot account for the yields we have obtained, and it is therefore necessary to postulate a rearrangement. The intervention of a rearrangement is convincingly demonstrated by the formation of (XI) from (X) and by our observation that

with our experimental practice of using a three-fold excess of substrate.



(Received, September 14th, 1966; Com. 691.)

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