# Conformational isomerism in 3,5,8-trioxabicyclo[5.1.0]octane and its diastereomeric 4-methyl derivatives. A combined IR, X-ray and ab initio study 

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#### Abstract

IR spectroscopy and ab initio calculations have been applied to the title compounds. Bicyclic acetals were found to exist as mixture of chair and twist-boat conformations, a parent epoxide being in ternary equilibrium. X-ray data on 4 -methyl (exo) derivative display a twist-boat form. © 2006 Elsevier B.V. All rights reserved.


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## 1. Introduction

Investigations of 1,3-dioxanes with substituent at $C^{5}$ (polar especially) show a number of interesting trends [1]. 3,5-Dioxabicyclo[5.1.0]octanes with exo- and endo chairlike conformations relate closely to above 1,3-dioxanes if one assumes that three-membered cycles in these molecules are equivalent to substituents at $\mathrm{C}^{5}$ in the stereochemical sence (axial $\equiv$ endo and equatorial $\equiv$ exo). The noticeable feature of bicyclic acetals is the real propensity to exist in both chair conformations and a twist-boat form in addition [2-9]. It is worth noting that seven-membered acetals with a planar fragment have been inspected thoroughly and untraditional chair-twist-boat $(\mathrm{C} \rightleftharpoons \mathrm{TB})$ equilibrium has been clarified [10].

[^0]

a


$\mathrm{C}_{\mathrm{exo}}$


TB


Cer $_{\text {e }} \quad \mathrm{R}=\mathrm{H}$, Alkyl, Aryl $\quad$ TB

One can expect that bicyclic acetals with $\mathrm{X}=\mathrm{C}, \mathrm{O}, \mathrm{S}, \mathrm{N}$ may serve as suitable models for the investigation of polar and steric interactions between acetal moiety and three-membered rings. Encouraged by available stereochemical information implying both a broad range of


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