CONFORMATIONAL EFFECTS IN THE OPENING OF
CIS/TRANS 1,4-DIALKYL SUBSTITUTED 1,2-
CYCLOHEXENE EPOXIDES WITH ACETIC ACID

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In our continuing studies on the reactions of
terpane oxides² we wish to report the following cases
of stereospecific epoxide openings in the 2-menthene
series.

The peracid epoxidation of limonene (I), menthene-1
(II) and menthene-3 (III) leads, in each case, to
approximately a 1:1 mixture of the corresponding cis
and trans epoxides³ as would be expected from the
recent findings of Rickborn and Lwo³d. Although
the cis/trans epoxide mixtures are separable by dis­
tillation only with difficulty, they may be separated
conveniently on a 15 ft. 15% Carbowax 20M on chromo­
sorb W OLPC column⁴. In each of the three cases cited,
reaction of the cis epoxide (IVA,b,c) with a sodium

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acetate buffered solution of acetic acid leads exclusively to a single product containing trans tertiary acetoxy and secondary hydroxyl groups and trans 1,4-alkyl groups (VII a, b, c), as would be expected by an extension of the Fürst-Plattner rule to monocyclic systems, assuming that R₂ "fixes" the conformation of the cyclohexane ring system. Similarly, reaction of the trans epoxides (Va, b, c) leads predominantly to epoxide opening resulting from cleavage of the "equatorial" C-O
linkage in the transition state (VIII) to afford the
diaxial products (IXa, b, c) with secondary acetoxy and
tertiary hydroxyl groups. The proof of structure in
each case follows from pyrolysis of the acetate to
yield known products$^{2,3a}$. For example, (±)-cis-3-
menthene epoxide (IVc) affords a hydroxycetate (VIIc)
which pyrolyzes at 390° to a mixture of (±)-trans-3-
menthene-5-ol (X)$^7a$ and (±)-menthone-isomenthone (XI) in

\[
\text{VIIc} \xrightarrow{390^\circ} \begin{array}{c}
\text{XII} \\
\text{X} \\
\text{XIII} \\
\text{XIV}
\end{array}
\]

\[
\text{IXc} \xrightarrow{450^\circ} \begin{array}{c}
\text{XII} \\
\text{XV}
\end{array}
\]

a product ratio, 7:3. The (±)-trans-3-menthene epoxide
affords a crystalline (±)-hydroxycetate (IXc), m.p.
86.5-89.5°, which pyrolyzed at 450° to (±)-trans-3-
menthene-4-ol\textsuperscript{7b} (XII). In each case, the infrared spectra of the pyrolysis products were superposable on those of authentic samples\textsuperscript{7a,b}. Further structure proof of (X) came from hydrogenation (PtO\textsubscript{2} - ethyl acetate) to a mixture of \(88\%\) (\(\pm\))-isomenthol (XIII) and \(12\%\) (\(\pm\))-neomenthol (XIV). Hydrogenation of (XII) afforded trans-p-menthan-4-ol (XV).

The high degree of stereospecificity in these epoxide openings leads us to propose the following:

1. The appreciable amount of abnormal, "diquatorial", opening observed with nucleophilic reagents on monocyclic \(\text{cis-1,4-dialkyl cyclohexene epoxides (vide infra)}\) does not necessarily arise from reaction of the less stable conformational isomer\textsuperscript{8} (as is observed in conformationally mobile cyclohexene epoxide systems\textsuperscript{9}) but rather, perhaps, from the inate difficulty of nucleophilic displacements at tertiary centers\textsuperscript{10}.

2. The 4-alkyl substituent determines the conformation in which the \(\text{cis-}\) and \(\text{trans}\) \(1,4\)-alkyl substituted cyclohexene-1,2-epoxides react with acidic reagents (i.e., H\text{C\textsubscript{2}}\text{CO}) via a normal, "diazial", opening (Fürst-Plattner Rule\textsuperscript{5}).
Thus the lithium aluminum hydride (LAH) reduction of (+)-cis-1,2-limonene epoxide (IVA)\textsuperscript{3a,11a} affords a product ratio of 15\% cis-\(\beta\)-terpineol (XVI) via "effective diequatorial opening" and 85\% (+)-neodihydrocarveol (XVII) via "normal diaxial opening". With other nucleophilic reagents, i.e., dimethylamine, the percentage of "diequatorial" opening is even greater (30-40\%)\textsuperscript{11b}.

There are two ways in which such diequatorial openings may occur; (a) through diaxial opening of the less stable epoxide conformer with subsequent inversion (process "A") or (b) via diaxial opening of a "twist" transition state (process "B"). Inasmuch as there should be a considerable steric (shielding) effect exhibited by the "axial" 4-alkyl group on the incoming nucleophile in process "A", in addition to the already energetically unfavorable epoxide conformation, we tend to favor
process "B".

In the opening of the cis-dialkyl epoxide (VIa) with acetic acid, although $R_1$ is isopropyl and $R_2$ is methyl, only a single product is obtained (governed by diaxial opening with $R_2$ equatorial). Therefore, even though the 1,4-dialkyl substituted cyclohexene epoxide has been postulated to assume a "nearly chair" conformation in the reaction transition state, the 1-alkyl substituent plays little or no part in determining the conformational preference of the transition state. It is probable, therefore, that the transition states for the reactions of the cis/trans-1,4-dialkyl cyclohexene epoxides as illustrated above (VI and VIII) are a bit exaggerated, since to develop a true "chair" conformation the C-O bond must effectively be broken (or at the least very stretched). Transition states such as (VIId) and (VIIIId) might be better representations. In such transition states, as the protonated oxirane starts to assume the status of an axial hydroxyl there is still effective overlap between the p-orbital of oxygen and the vacant p-orbital of the "incipient carbonium ion". Since $R_1$ in such a transition state for the cis epoxide (VI) can be neither axial nor equatorial it should not have any effect on conformational preference (as indeed is the case). Such overlap would be less favorable for a
di-equatorial opening (which is not observed for the cis epoxides of type IV). The small percentage (ca. 10%) of an "effective di-equatorial opening," observed for the trans epoxides (Va and Vb) may occur by either a process of type "A" or "B" (vide supra). Although the results presented do not distinguish between these two mechanisms it is again feasible that process "B" may function inasmuch as the normal diaxial opening of a "twist" conformer of (V) would be enhanced by formation of an incipient tertiary positive carbon center.

In any event, the stereospecific opening of 4-alkyl substituted cyclohexene epoxides, coupled with the convenient pyrolysis of the resultant hydroxyacetates affords an excellent method for the preparation of specific allylic alcohols.

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REFERENCES


(4). J. C. Leffingwell, unpublished results; Ref. 3a, footnote 5.


(6). In the case of (Va,b) ca. 10% of the abnormal product arising from an effective "diequatorial" opening was observed; see Ref. 3a.


(9). For examples see Ref. 3a,c; W. F. Newhall, J. Org. Chem., 24, 1637 (1959); 29, 185 (1964); H. Kuczynski and A. Zabza, Roczniki Chem., 37, 733 (1963).


(13). The term 1-alkyl and 4-alkyl substituent refers to the systematic naming of cyclohexane systems in general. Therefore, in each of the cases cited here, the alkyl substituent substituted on the epoxide bearing carbon is the 1-alkyl substituent. The fact that the isopropyl group in the 3-menthene epoxides is on the 4-carbon atom according to nomenclature in the p-menthane systems thus should not confuse the reader in the interpretation of the general effect.