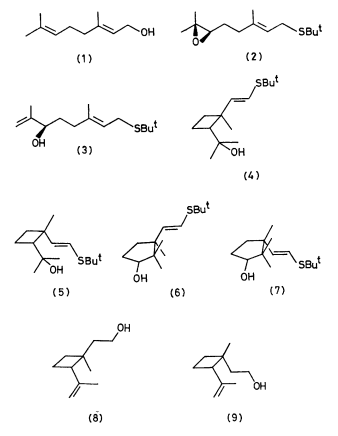
Transformation of the Geraniol Skeleton into the Fragranol and Grandisol Skeletons

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Summary Racemic 6,7-epoxygeranyl t-butyl sulphide (2) was lithiated [butyl-lithium-1,2-bis(dimethylamino)ethane] at C(1) to give the (1E)-epoxyallyl-lithium (10) which cyclised to give the alcoholates of the (racemic) cyclobutyl carbinols (4) and (5) (fragranol and grandisol skeletons) and of a single (racemic) cyclopentanol, (6) or (7).

EXTREME cyclisations which involve displacement (exo- and endo-cyclic displacements) are of current interest.^{1,2} I report cyclisations of a geraniol (1) derivative to give racemic fragranol (8)[†] and grandisol (9)[†] derivatives,³ via displacement of an epoxide O-atom by a stabilized allyl anion generated by abstraction of a proton. This principle, previously used by Itô⁴ to cyclise farnesol (\rightarrow 10-membered ring) and geranylgeraniol (\rightarrow 14-membered ring) derivatives, is a useful anionic counterpart of the 'corresponding'[‡] biological and biomimetic cyclisations. Stork's⁵ epoxynitrile cyclisations are related.⁶

Geranyl and neryl chloride, as a mixture, were transformed into the corresponding t-butyl sulphides which were epoxidized at the C(6)-C(7) double bond⁷ and racemic 6,7-epoxygeranyl t-butyl sulphide (2)[†] was isolated by g.l.c. The sulphide (2) was added at once to excess of butyllithium-1,2-bis(dimethylamino)ethane (5 equiv. of each) in tetrahydrofuran (THF)-hexane [ca. -75 °C, 0.08-0.09 M initial concentration of (2) in THF-hexane (ca. 2.4:1 v/v)] and the resulting yellow solution was kept at ca. -75 °C for 3 h and then at ca. 5 °C for 2 h. Hydrolysis, work up, and distillation afforded a mixture (g.l.c.) of two (racemic) cyclobutyl carbinols, (4)[†] (ca. 43%, fragranol skeleton) and

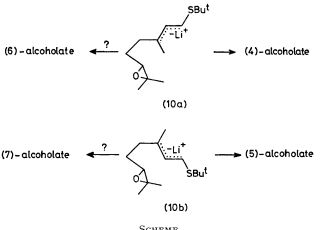


[†] One enantiomer is depicted, which, in the case of (9), is (+)-grandisol.

‡ Cyclisation via addition of a cation, generated by opening of an epoxide, to a double bond.

(5)† (ca. 29%, grandisol skeleton), a single (racemic) cyclopentanol, (6)† or (7)† [ca. 25%, <ca. 1% of the isomer, (7) or (6), present], and the (racemic) open-chain alcohol $(3)^{\dagger}$ (ca. 3%), in ca. 64% combined yield.

At ca. -75 to ca. -25 °C under otherwise the same conditions, (2) was deprotonated at C(1) (ca. 75% conversion within 10 min at ca. -25 °C) and the cyclisations of the resulting anion were slow, quenching with D₂O giving $[1-^{2}H]-(2)$. The geometry $(1E)^{2}$ of the anion $(10)^{\dagger}$ (Scheme),



Scheme

follows from that of the derived cycles. Apart from the stereochemistry with respect to the rings, structures (4), (5), and (6) or (7) [and (8)] were deduced from the spectra [1H n.m.r. spectrum (CDCl_3) of (4): δ 1.09, 1.20, 1.38 (s, 3H each), 1.33 (s, 9H), 1.5–2.4 (m, 6H), and 6.04 (AB, δ_{AB} 0.06 p.p.m., J = 15 Hz; (5): $\delta = 1.04$, 1.19, 1.29 (s, 3H each),

1.36 (s, 9H), 1.5–2.4 (m, 6H), and 6.28 (AB, δ_{AB} 0.27 p.p.m., J 16 Hz); (6) or (7): δ 0.84, 0.87, 1.07 (s, 3H each), 1.34 (s, 9H), 1·4-2·4 (m, 5H), 3·8-4·0 (br. m, 1H), and 5·98 (AB, δ_{AB} very small]. That (4) is trans- and (5) is cis-substituted at the ring (as drawn) was shown by ¹H n.m.r. spectrometry using Eu(fod)₃ complexation. The analogous distinction between (6) or (7) has not yet been made.

I interpret the results as follows (see Scheme). In the epoxyallyl-lithium (10), intramolecular displacement of the epoxide O-atom at C(6) or C(7), by the distal C(1) (to give a trans-cyclohexene and -cycloheptene, respectively) is not geometrically feasable, but such displacement is feasible by the proximate C(3), and is more favourable than an intermolecular displacement. Exocyclic displacement^{1,2} by C(3) at C(6) within folded conformers of type $(10a)^{\dagger}$ and (10b)[†] gives the lithium alcoholates of (4) and (5), respectively, and endocyclic displacement^{1,2} by C(3) at C(7) within these types of conformers would give the lithium alcoholates of (6) and (7), respectively. Molecular models show that the displacement leading to (7), although sterically quite hindered, is sterically more favourable, and structure (7) is therefore more likely.

The sulphide function stabilizes the allyl-lithium (10) and facilitates its selective formation from (2) and butyllithium; attack at C(8) to give the alcoholate of (3) is minimal. The t-butyl group was introduced to protect the sulphur atom during epoxidation. Synthetically, these particular choices lead to an impasse since the cyclisations are neither regio- nor stereo-selective and since the tbutylthio function in (4) and (5) turns out to block liberation of the latent aldehyde functions, but they provide a well characterized model for further, systematic studies, which are under way.

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⁶ For a further related study, see W. C. Still, Tetrahedron Letters, 1976, 2115.

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