

Stereochemistry of Theaspirone and the Blumenols

By GEORGE WEISS, MASATO KOREEDA, and KOJI NAKANISHI*

(Department of Chemistry, Columbia University, New York, New York 10027)

Summary The stereochemistry of theaspirone (9), blumenol A (5), and blumenol B (8) has been established; a stereospecific preparation of theaspirone is also described.

In connection with studies¹ on (+)-abscisic acid (ABA), the plant growth regulator, to which the absolute configuration (7) has been assigned,^{1,2} we have carried out experiments which elucidate the full stereochemistry of blumenol A (5)³ and blumenol B (8).³ The absolute configuration of (-)-theaspirone,⁴⁻⁶ an important component of tea aroma, has also been established as (9). In addition the effect of γ -substituents on the c.d. of conjugated enones is discussed.

The racemic bis-enone (1) [prepared by t-butyl chromate oxidation⁷ of (\pm)- α -ionone] was reduced with sodium borohydride in ethanol at 0° (quantitative yield), and the resulting diastereoisomeric mixture of 6,9-diols was separated into (2) and (3) by high speed liquid chromatography (l.c.).

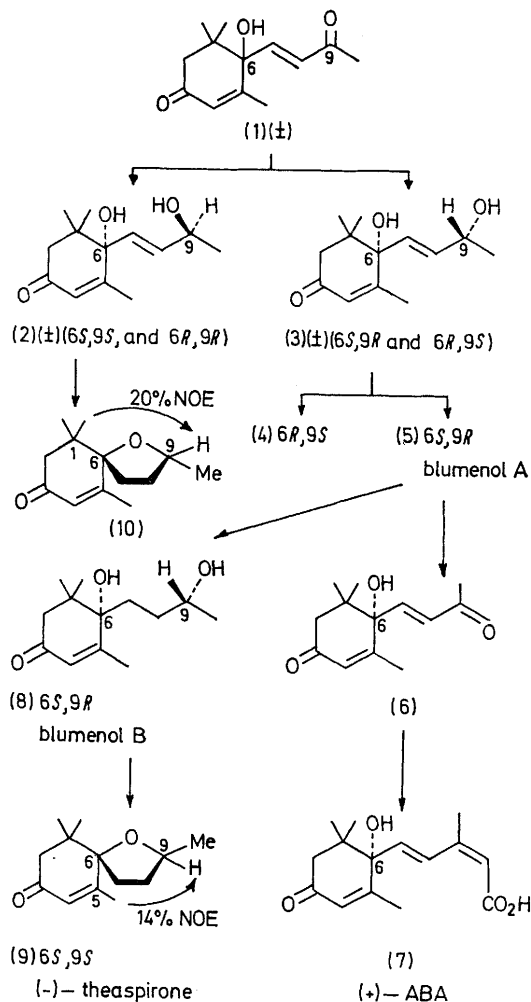
Comparisons of i.r. spectra (KBr) of the two diastereoisomers, (\pm)-(2) (m.p. 116–118°) and (\pm)-(3) (m.p. 112–114°) with that of blumenol A[†] clearly showed that the less polar (by l.c.) (3) corresponded in relative configurations at C-6 and C-9 to the natural product.

The racemic diol (3) was resolved by preparation of the diastereoisomeric (+)- α -methoxy- α -trifluoromethylphenyl-acetyl (MTPA) esters⁸ followed by l.c. Basic hydrolysis (quantitative yield) of the separated MTPA esters thus gave the two enantiomers diol (4) [(enantiomer of structure (3)), c.d. (MeOH) $\Delta\epsilon_{241} - 8.7$, $\Delta\epsilon_{317} + 0.39$, and diol (5) [(same as structure (3)), c.d. (MeOH) $\Delta\epsilon_{242} + 9.6$, $\Delta\epsilon_{317} - 0.44$. The c.d. data of (5) corresponded to those of blumenol A, and hence the two are identical.†

Diol (5) was oxidized with Jones reagent to the bis-enone (6), the optical properties of which corresponded to the bis-enone [$\Delta\epsilon_{208} - 30.2/\Delta\epsilon_{242} + 38.4$ (exciton-splitting), $\Delta\epsilon_{320} - 2.28$ (in methanol)] that had earlier¹ been converted into natural (+)-ABA (7). This defines the C(6) configuration of diol (5) (blumenol A) as S.⁹

Blumenol B (8) was obtained by hydrogenation³ of blumenol A with platinum oxide-ethyl acetate (90%). Interestingly the sign of the main π, π^* c.d. Cotton effect

(CE) of blumenol B (8), $\Delta\epsilon_{217} + 11.4$, $\Delta\epsilon_{250} - 8.2$, and $\Delta\epsilon_{325} + 1.01$, was opposite to that of blumenol A (5). It has been



† We thank Dr. D. H. S. Horn, C.S.I.R.O., Melbourne, for this information.

shown¹⁰ that the helicity between an enone moiety and a γ -hydroxy-group is in agreement with the sign of the longest π, π^* enone CE. This indeed is the case with the 250 nm CE of blumenol B (8). However, the fact that it is reversed in blumenol A (5) (242 nm) suggests that the effect of the homoconjugated 7-ene outweighs that of the γ -hydroxy-group. ‡

Mesylation of (8) at C(9) followed by refluxing in pyridine-benzene gave the spiro-ether (9) [75% yield from (8); stereospecific inversion at C-9], c.d. (MeOH) $\Delta\epsilon_{221} +7.0$, $\Delta\epsilon_{252} -7.0$, and $\Delta\epsilon_{317} +0.90$. Irradiation (n.m.r.) of the 5-Me group in (9) exhibited a 14% intramolecular nuclear Overhauser effect (NOE) on 9(H). As the C(6) configuration is known (6S), this establishes the configuration at C(9) in (9) (9S), and also that at C(9) in blumenol B (8) (9R) and

blumenol A (5) (9R). Since natural (-)-theaspiron shows NOE (10%)⁵ and optical properties⁶ essentially the same as those of (9), (-)-theaspiron can be fully represented by structure (9).

In a similar reaction sequence (\pm)-(2) was converted into "trans-theaspiron"⁵ (10), which showed a 20% NOE at 9(H) upon irradiation of the 1-Me group and thus corroborated the C(9) configuration in theaspiron (9). The fact that the C(6) configurations of natural products, (+)-ABA (7), blumenol A (5), blumenol B (8), and (-)-theaspiron (9), are identical is of biogenetic significance.§

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‡ Molecular models show that the helicity is independent of the configuration of the cyclohexenone ring.

§ The same configuration at C(6) for blumenols A and B has been established by Galbraith and Horn. We thank Dr. Horn for this information prior to publication.

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