Total Synthesis of (\pm) -Isoprosopinine B and (\pm) -Desoxoprosopinine

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The synthesis of the title compounds (10) and (16) in eleven and ten steps respectively from the imine (1) and 2-trimethylsilyloxycyclohexa-1,3-diene (2) *via* the azabicyclo-octanone (3) and the triol (5) illustrates a new general method for the preparation of the prosopis alkaloids.

Whereas numerous reports on the synthesis of all-*cis*-2,3,6-trisubstituted piperidines of the carpaine series have appeared over the past thirteen years,¹ fewer studies have been devoted to the synthesis of the 3,6-*cis*-2-*trans*-trisubstituted piperidines of the prosopis series.² We now describe a general synthesis involving the triol (5) which should serve as a common precursor to all the known naturally occurring prosopis piperidine alkaloids.

The synthesis of the triol (5) is shown in Scheme 1. The azabicyclo-octanone (3)^{\dagger} is prepared by cycloaddition of the imine (1)³ to 2-trimethylsilyloxycyclohexa-1,3-diene (2),⁴ followed by mild acid hydrolysis.⁵ The cycloaddition is

[†] All new compounds exhibited spectroscopic and analytical data consistent with the assigned structure. All compounds are racemic.



Scheme 1. Ts = p-MeC₆H₄SO₂. *Reagents:* i, C₆H₆, 5 °C to room temp., 3 h; ii, 0.005 M HCl-tetrahydrofuran(THF), room temp., 1 h (57%); iii, ACOOH, ACOH, NaOAc, 50 °C, 72 h (47%); iv, LiAlH₄, Et₂O, 0 °C, 1.5 h (80%). Note 1. The product (**3**) was accompanied by the *endo*-isomer (24%). Note 2. The product (**4**) was accompanied by the methylene-migrated lactone (4%).



Scheme 2. Ts = p-MeC₆H₄SO₂. *Reagents*: i, PhCHO, TsOH, C₆H₆, room temp., 12 h (60%); ii, pyridinium dichromate, 4 Å molecular sieve, CH₂Cl₂, room temp., 9 h (91%); iii, Ph₃P=CH[CH₂]_aC-(OLi)₂Bu,¹³ THF, 0 °C, 1 h (46%); iv, H₂, 5% Pd–C, EtOH; v, HOCH₂CH₂OH, TsOH, C₆H₆, reflux, 16 h; vi, Red-Al [sodium bis(2-methoxyethoxy)aluminium hydride], C₆H₆, reflux, 24 h; vii, 8 M HCl-MeOH, reflux, 16 h [66% from (8)].



Scheme 3. Ts = p-MeC₆H₄SO₂. *Reagents*: i, Me₂C(OMe)₂, TsOH, CH₂Cl₂, room temp., 16 h (77%); ii, pyridinium dichromate, 4 Å molecular sieve, CH₂Cl₂, room temp., 4 h (71%); iii, Ph₃P=CH[CH₂]₈Me, THF, 0 °C, 1 h (50%); iv, H₂, 5% Pd-C, EtOH (97%); v, Red-Al, C₆H₆, reflux, 24 h; vi, 8 M HCl-MeOH (38%).

rigorously regioselective and reasonably stereoselective. Baeyer–Villiger oxidation^{2b,8—10} of the bicyclic ketone (3) gives largely the bridgehead-migrated lactone (4)[†] which is reduced to the key triol (5).[†]

The conversion of the triol (5) into isoprosopinine B $(10)^{11,12}$ is outlined in Scheme 2. After protection of the triol as its benzylidene derivative (6),[†] the primary alcohol was oxidised to the aldehyde (7)[†] which underwent Wittig chain extension with the ylide derived from the reaction of n-butyl-lithium (3 equiv.) with 6-(triphenylphosphonio)hexanoic acid bromide¹³ to give (8), \dagger as a mixture of E and Z isomers. Hydrogenation [to give (9)], detosylation, and deprotection gave (10), which was identical (t.l.c. and n.m.r., i.r., and mass spectra) with one component of an authentic sample of an inseparable mixture of isoprosopinine B and its side-chain ketone isomer isoprosopinine A (11). The synthesis therefore establishes unambiguously the structure of isoprosopinine B. In order to obtain an exact comparison with a single compound derived from natural sources desoxoprosopinine (desoxyprosopine) (16)^{2c} was prepared as summarised in Scheme 3. The acetonide (12)[†] proved to be a superior protected form of the triol (5), being generally more stable, easier to handle, and less susceptible to hydrogenolysis than the corresponding benzylidene derivative (6). Elaboration of the aldehyde $(13)^{\dagger}$ through $(14)^{\dagger}$ (E/Z mixture) and $(15)^{\dagger}$ proceeded smoothly and yielded, after deprotection, compound (16) which was identical in all respects except optical rotation with an authentic sample.

In summary, a general synthetic route to prosopis alkaloids has been established. The method is the most efficient to have been reported both for any naturally occurring prosopis alkaloid and for desoxyprosopine (16).

[‡] The imine (1) adds in high yield to a wide variety of silyloxydienes,⁶ and appears in this respect to be superior to iminocarbamates.⁷

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