A general method for making bicyclic compounds with nitrogen at a bridgehead—application to the halichlorine spiro subunit†

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N-Protected β-amino aldehydes having the nitrogen in a ring are easily converted into Morita-Baylis-Hillman adducts; O-acetylation and N-deprotection result in spontaneous cyclization to bicyclic structures having nitrogen at a bridgehead.

A recent report from this laboratory described the preparation of spiro amine 1, which was made during studies on the total synthesis of the marine natural product halichlorine (2).² An obvious next step was to convert 1 into 3, and this has now been achieved through the development of a new and general method for making compounds that contain a bicyclic subunit with nitrogen at a bridgehead. The approach (Scheme 1) is based on sequential formation of Morita-Baylis-Hillman (MBH) alcohols $(4 \rightarrow 5)$ and intramolecular $S_N 2'$ displacement of the derived acetates $(6 \rightarrow 7 \rightarrow 8)$. In the present work we have made alcohols of type 5 by MBH³ condensation (and used only acrylates), but the same compounds should also be accessible by other^{3,4} methods.

While O-acetates of MBH alcohols are known to undergo intermolecular S_N2' displacement, 3,5 the intramolecular 6-endo pathway⁶ (Scheme 1, $7 \rightarrow 8$) requires that no competing

$$O$$
 Pg
 N
 RO_2C
 RO_2C

Scheme 1

stereoelectronic or reactivity factors intervene to direct cyclization onto the CO₂R group⁷ of 7 or to cause $O \rightarrow N$ acetyl transfer; in the event, the desired ring closure $(7 \rightarrow 8)$ occurs smoothly.

The lactam 9,1 an intermediate in the preparation of 1, was deprotected (Me₃SiBr, 84%, $9 \rightarrow 10$) and oxidized (Swern, 85%) to aldehyde 11. Wittig reaction with Ph₃P=CH(OMe) and hydrolysis (CSA, aqueous MeCN) of the intermediate enol ethers then gave the expected aldehyde (11 \rightarrow 12 \rightarrow 13, 74% overall). When aldehyde 138 was dissolved in methyl acrylate, condensation occurred on addition of DABCO and Sc(OTf)₃.9 Although the resulting alcohols (14a, more polar) and 14b (less polar) could be separated, it was more convenient to acetylate the mixture and separate the corresponding acetates 15a (37% from 13) and 15b (34%). When each of the acetates 15a and 15b was treated with Me₃OBF₄ and then with aqueous Na₂CO₃, the lactam ring was opened to amino esters 16a and 16b, respectively, and these cyclized in situ to afford the desired bis-ester 3 (77% for 15a and 72% for 15b). 10 Aldehyde 13 was recovered unchanged either after exposure to DABCO in CH2Cl2 for 3 days, or when the MBH condensation was worked up before completion. These observations show that epimerization by retro-Michael elimination and re-addition does not occur.

Scheme 2 Reagents and conditions: (i) Me₃SiBr, CH₂Cl₂, -10 °C, 2 h, 84%; (ii) Swern oxidation, 85%; (iii) MeOCH₂PPh₃Cl, t-BuOK, THF, 0 °C, 2 h; (iv) camphorsulfonic acid, MeCN-water, 4 h, 74% over two steps; (v) methyl acrylate, DABCO, Sc(OTf)3, 5 days; (vi) AcCl, pyridine, CH2Cl2, 0 °C, 1 h, 25 °C, 1 h, over two steps 37% of 15a, 34% of 15b; (vii) Me₃OBF₄, CH₂Cl₂, 1.5 h; (viii) 20% aqueous Na₂CO₃, MeCN, 2 h, 77% from 15a, 72% from 15b.

[†] Electronic supplementary information (ESI) available: characterization data for compounds 3, 21, 26, 32 and 38. See http://www.rsc.org/suppdata/ cc/b4/b413481h/

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Scheme 3 Reagents and conditions: (i) References 10–12; (ii) methyl acrylate, DABCO, 3 days, 39% (19a), 34% (19b); (iii) AcCl, pyridine, CH₂Cl₂, 1 h, 89% (20a), 85% (20b); (iv) CF₃CO₂H, CH₂Cl₂, 1 h; aq Na₂CO₃, MeCN, 1 h, 77%; (v) (a) (Boc)₂O, Et₃N, CH₂Cl₂, 0 °C, 1 h, room temperature, 12 h, 87%; (b) Dess-Martin periodinane, CH₂Cl₂, 1.5 h, 88%; (vi) as in (ii), 46% (less polar), 38% (more polar); (vii) as in (iii), 94% for **24a**, 92% for **24b**; (viii) as in (iv), 77%; (ix) (a) NaN₃, aq H₂SO₄, 0 °C, 16 h, 74%; (b) LiAlH₄, dioxane, reflux, 24 h, 80%; (x) (a) (Boc)₂O, EtOAc, 16 h, 98%; (b) Swern, 2 h, 87%; (xi) as in (ii), 5 days, 39% (less polar), 47% (more polar); (xii) as in (iii), 92% for 30a, 94% for 30b; (xiii) as in (iv) 90% (more polar), 87% (less polar); (xiv) (a) NaN₃, aq H₂SO₄, 0 °C, 16 h, 80%; (b) LiAlH₄, THF, reflux, 24 h, 71%; (xv) (a) as in (x), 83%; (b) as in (x), 80%; (xvi) as in (ix) 49% (more polar 36a), 45% (less polar 36b); (xvii) as in (iii), 78% for 37a,79% for 37b; (xviii) as in (iv), 84% for 37a, 93% for 37b.

The approach of Scheme 2 appears to be general, and we have applied it to several other cases (Scheme 3).

L-Proline (17) was converted by literature methods^{11–13} into aldehyde 18, which underwent condensation with methyl acrylate, affording a separable mixture of 19a (more polar, 39%) and 19b (less polar, 34%). Acetylation produced the corresponding acetates 20a (89%) and 20b (85%). Finally, exposure of a mixture of both acetates to CF₃CO₂H resulted in N-deprotection, at which point, treatment with aqueous Na₂CO₃ caused spontaneous cyclization to 21 (77% yield). HPLC analysis [Chiracel OD-H, 1% EtOHhexane] showed the material to have an enantiomeric purity of 99.8%, indicating that little, if any, racemization occurs in the synthetic sequence.

In another series of experiments, commercial (2-hydroxyethyl)piperidine (22) was converted by N-protection (Boc₂O, 87%) and Dess-Martin oxidation (88%) into aldehyde 23, which underwent efficient MBH condensation [23 → 24a (more polar, 38%) and 24b (less polar, 46%)]. Once again, acetylation (94% for 24a, 92% for 24b), N-deprotection (CF₃CO₂H), and treatment with aqueous Na₂CO₃ resulted in ring closure, giving 26 (77% from a mixture of both acetates).

We also investigated two other ring sizes for the starting amine. Keto ester 27 was converted by Schmidt reaction and LiAlH₄ reduction into 28,14 which was protected on nitrogen [Boc2O, 98%], oxidized (Swern, 87%), and subjected to the MBH condensation [29 \rightarrow 30a (more polar, 47%) and 30b (less polar, 39%]. Acetylation (92% for **31a**, 94% for **31b**) and *N*-deprotection (CF₃CO₂H) and basification gave 32 (90% from 31a, 87% from 31b). Similarly, keto ester 33 was converted into amino alcohol 34

and then into aldehyde 35. Our standard sequence (35 \rightarrow 36a,b, \rightarrow 37a,b, \rightarrow 38) then proceeded in the expected way.

In summary, synthetic work related to halichlorine has led to the development of a method for generating bicyclic amines with nitrogen at a bridgehead. The process occurs with preservation of stereochemistry α to the nitrogen.

All new compounds were fully characterized by spectroscopic methods, including high resolution mass spectrometry. We thank NSERC for financial support and C. Boucher [Boehringer Ingelheim (Canada)] for ee measurements. M.Y. holds a Province of Alberta Graduate Fellowship.

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