

Bryostatin: a novel asymmetric synthesis of the C_{27} - C_{34} fragment starting from (*R*)-carvone as chiral template

J. De Brabander, B. A. Kulkarni, R. Garcia-Lopez and M. Vandewalle*

University of Gent, Department of Organic Chemistry, Laboratory for Organic Synthesis, Krijgslaan, 281 (S.4), B-9000 Gent, Belgium

Abstract: (R)-Carvone is a suitable chiral template for the synthesis of differently protected (4S,6R,7R)-trihydroxy-1-octyne derivatives, the C₂₇-C₃₄ fragment of bryostatins. Also other potentially interesting chiral building blocks are described. © 1997 Elsevier Science Ltd

The bryostatins 1 constitute a family of some 17 highly oxygenated marine macrolides based on a polyacetate-derived backbone (Scheme 1).¹ They exhibit exceptional antineoplastic activity against PS lymphocytic leukemia and ovarian carcinoma.² Next to the first completed total synthesis of bryostatin 7(1C) by Masamune *et al.*,³ other groups have described the synthesis of various fragments of the 20-membered ring lactone.^{4–6}





Previously we have described a synthesis of the $C_{17}-C_{27}$ fragment involving a $C_{27}-C_{34}$ acetylenic intermediate 2 obtained from D-isobutyllactate 3 as the chiral template.^{6a,b}

The synthesis of the differently protected eight carbon fragment 6, starting from L-threonine, has been described by Masamune *et al.*^{3b} Intermediate $C_{21}-C_{27}$ fragments has also been obtained by Roy *et al.*^{5a} from D-galactonolactone, by Evans *et al.*^{5b} via Sharpless epoxidation procedure and by Hale *et al.*^{5c} via a Sharpless asymmetric dihydroxylation–epoxidation sequence.

The present paper describes a new approach for the $C_{27}-C_{34}$ fragment based on (*R*)-carvone 7⁷ as the chiral template, a less obvious starting material for the synthesis of acyclic polyols. The described selective epoxidation of 7 led to 8.⁸ The organoselenium-mediated reductive opening⁹ of the epoxide gave the known¹⁰ alcohol 9 next to the 2-epimer in a 4:1 ratio; crystallization from EtOAc-hexane (5:95) led to pure 9. It has previously been obtained upon lithium liq. ammonia reduction¹⁰ of 8; the relative C-2, C-3 configuration was fully proven by ¹H NMR and is in accord with literature data.¹⁰

* Corresponding author.

Protection of the hydroxy group in 9 was best performed with *tert*-butyldimethylsilyl triflate as the use of TBSCl led to substantial elimination back to 7.

With 10 in hand we turned our attention to the oxidative removal of the isopropylidene substituent. Originally we had planned to perform a double Baeyer–Villiger oxidation on the corresponding ketone 11 to the desired lactone 14. However, in the beginning no conditions could be found for cleaving the exocyclic ketone; only oxidation of the cyclic keto function was observed leading to the lactone 13 in 63% yield. A viable route was found *via* ozonolysis of 10 in dry methanol to the methoxy-hydroperoxide which upon treatment with *p*-nitrobenzoyl chloride and *in situ* Criegee rearrangement¹² of the intermediate methoxy-peroxy ester afforded acetate 12. The reaction conditions are quite critical as traces of water have to be avoided during the whole process.^{13b} The mechanism is known^{12,13a} to proceed with retention of configuration; the structure of 12 was fully proven by ¹H NMR.¹¹ Baeyer–Villiger oxidation of 12 finally gave our key-intermediate 14,¹¹ a C₂₁–C₂₇ fragment with the correct stereogenic centers and a carboxyl function as the handle for further chain extention.

In order to avoid the rather critical Criegee rearrangement $(10\rightarrow 12)$ we decided to reinvestigate the alternative route to 14 based on the double Baeyer–Villiger oxidation of diketone 11. In the first experiments only oxidation of the cyclic ketone function, leading to 13, was observed after circa 24 h (*vide supra*). This indicates a much lower reactivity of the exocyclic ketone. Also more powerful reagents^{14b} did not lead to expected 14. Only upon performing the oxidation with 20 eq MCPBA for 6 days the desired product 14 was obtained in 70% yield. This observation deserves some comment. The Baeyer–Villiger oxidation of cyclohexylketones is normally an excellent process^{8,14a} and is substantiated with the formation of 24 from 23. In the transformation of 11 to 14, the carbonyl function in intermediate 13 is now exocyclic to a 7-membered ring; apparently this ring has a low migratory ability. This bears some parallel with reported problems on the cleavage of straight-chain ketones.^{14b} Furthermore to the best of our knowledge only one case of a Baeyer–Villiger oxidation of a cycloheptylketone has been reported.¹⁵

As can be deduced from Scheme 2 the synthesis of 14 via the double Baeyer–Villiger oxidation is the superior one and is furthermore easier to perform.

The most expedient route to an acetylenic C_{27} - C_{34} precursor would involve DIBAH reduction of 14 to the lactol with concomitant deprotection of the 23-hydroxy group followed by *in situ* treatment with diazomethylphosphonate.¹⁶ Unfortunately the reduction step led to substantial decomposition of the β -hydroxy aldehyde.

This forced us to first protect the hydroxy function. Base mediated methanolysis of the acetate in 14 caused decomposition. On the other hand, enzyme catalyzed hydrolysis afforded in high yield alcohol 15, which was then transformed to 16. Reduction of 16 to the lactol 17 (in equilibrium with the corresponding aldehyde) followed by treatment of this crude mixture with dimethyl (diazomethyl) phosphonate¹⁵ afforded alkyne 4.

An alternative route to the differently protected $C_{27}-C_{34}$ fragment 6 involves as intermediate the heptyltetrol 18, obtained by reduction of 14. Selective protection of the α -diol unit and of the 3-hydroxy group led to primary alcohol 21. Oxidation to the aldehyde 22 and formation of the alkyne function, employing the Seyfert reagent,¹⁵ finally afforded the target molecule 6.

The use of the fragments 2 and 6 (5) for bryostatin synthesis has already been documented.^{3b,6a,b} These and other described intermediates derived from (R)-carvone (or (S)-carvone for the enantiomeric series) could be of interest for the synthesis of other natural products possessing a polyol structure.

Acknowledgements

We thank the "NFWO" and the "Ministerie voor Wetenschapsbeleid" for financial assistance to the laboratory.

Bryostatin



(a) H₂O₂, NaOH, MeOH, -10 °C, 3 h; (b) Ph₂Se₂, NaBH₄, EtOH, HOAc, 0 °C, 15 min; (c) TBSOTf, CH₂Cl₂, 2,6-lutidine, 0 °C, 20 min; (d) KIO₄, OsO₄, THF-H₂O (1:1), 12 h; (e) O₃, CH₂Cl₂, MeOH, -78 °C, 40 min; then dry PhH and evaporation; then CH₂Cl₂, py, p-NO₂C₆H₄COCl, 0 °C, 1 h, Δ 15 h; (f) MCPBA (10 eq), CH₂Cl₂, r.t., 2 d; then Me₂S; (g) MCPBA (20 eq), CH₂Cl₂, r.t., 6 d; then Me₂S; (h) PLE (EC 3.1.1.1), phosphate buffer pH 7, Me₂CO, 35 °C; (i) MEMCl, DIPEA, CH₂Cl₂, r.t., 16 h; (j) DIBAH, CH₂Cl₂, -78 °C, 1 h; (k) (MeO)₂P(O)CHN₂, t-BuOK, THF, -78 to -30 °C, 12 h; (l) LiBH₄, THF, r.t., 6 h; then Amberlyst A-15, MeOH-THF, 1 h; (m) (i) (MeO)₂CMe₂, THF, PPTS, r.t., 2 h; (ii) MeOH, PPTS, r.t., 2 h; (ii) TBAF, THF, r.t., 4 h; (p) SO₃.py, Et₃N, DMSO, -10 °C, 4 h.

Scheme 2.

References

- 1. Petit, G.R.; Gao, F.; Sengupta, J.M.; Coll, J.C.; Herald, C.L.; Doubek, D.L.; Schmidt, J.M.; Van Camp, J.R.; Rudloe, J.J.; Nieman, R.A. *Tetrahedron* **1991**, *47*, 3601, and references cited therein.
- 2. Petit, G.R.; Day, J.F.; Hartwell, J.L.; Wood, H.B. Nature 1970, 227, 962.
- (a) Blanchette, M.A.; Malamas, M.S.; Nantz, M.H.; Roberts, J.C.; Somfai, P.; Whritenour, D.C.; Masamune, S.; Kageyama, M.; Tamura, T. J. Org. Chem. 1989, 54, 2817. (b) Masamune, S. Pure Appl. Chem. 1988, 60, 1587. (c) Kageyama, M.; Tamura, T.; Nantz, M.H.; Roberts, J.C.; Somfai, P.; Whritenour, D.C.; Masamune, S. J. Am. Chem. Soc. 1990, 112, 7407.
- 4. Norcross, M.H.; Paterson, I. Chem. Rev. 1995, 95, 2041 and references cited therein.
- (a) Roy, R.; Rey, A.W.; Charron, M.; Molino, R. J. Chem. Soc., Chem. Commun. 1989, 1308. (b) Evans, D.A.; Gauchet-Prunet, J.A.; Carreira, E.M.; Charette, A.B. J. Org. Chem. 1991, 56, 741.
 (c) Hale, K.J.; Lennon, S.A.; Manaviarar, S.; Javaid, M.H.; Hobbs, C.J. Tetrahedron Lett., 1995, 36, 1359.

- 6. (a) De Brabander, J.; Vandewalle, M. Synlett 1994, 231. (b) De Brabander, J.; Vandewalle, M. Synthesis 1994, 8, 855.
- 7. Purchased from Aldrich (98% e.e.); the %d.e. of all derived compounds was checked by anal. HPLC.
- 8. Baggiolini, E.G.; Iacobelli, J.A.; Hennessy, B.M.; Batcho, A.D.; Sereno, J.F.; Uskocovic, M.R. J. Org. Chem. 1986, 51, 3098.
- 9. Miyashita, M.; Suzuki, T.; Yoshikoshi, A. Tetrahedron Lett. 1987, 28, 4293.
- 10. Mc Chesney, J.D.; Bloum, T.J.F. J. Org. Chem. 1985, 50, 3473.
- 11. Selected analytical data. **12**: ¹H NMR (500 MHz, CDCl₃): δ 1.06 (3H, d, J=6.7 Hz), 1.84 (1H, ddd, J=12.9, 10.6, 2.1 Hz), 2.04 (3H, s), 2.37 (1H, dddd, J=12.9, 4.5, 4.5, 2.0 Hz), 2.40 (1H, ddd, J=13.8, 10.9, 1.1 Hz), 2.46 (1H, ddq, J=6.7, 2.5, 1.1 Hz), 2.83 (1H, ddd, J=13.8, 5.3, 2.0 Hz), 4.19 (1H, ddd, J=4.5, 2.5, 2.1 Hz), 5.32 (1H, dddd, J=10.9, 10.6, 5.3, 4.5 Hz) ppm. **14**: ¹H NMR (500 MHz, CDCl₃): δ 1.39 (3H, d, J=6.7 Hz), 1.8 (1H, ddd, J=2.9, 10, 11.2 Hz), 2.05 (3H, s), 2.28 (1H, dd, J=4.7, 13.5 Hz), 2.91 (2H, d, J=7.7 Hz), 3.92 (1H, dd, J=2.9, 5.4 Hz), 4.48 (1H, q, J=6.7 Hz), 5.35 (1H, dddd, J=5.0, 5.0, 10.3, 15.4 Hz) ppm. [α]D²⁰ values in CHCl₃ or otherwise stated for: **4**; +24.8 (c=1.2), **6**; +71.7 (c=1.0), **8**; +1.7 (c=2.0), **9**; -18.6 (c=1.7), **10**; -27.7 (c=2.0), **12**; -14.3 (c=1.7), **14**; +6.0 (c=1.0), **15**; +14.5 (c=1.0), **16**; +29.7 (c=1.0), **18**; +27.6 (c=2.9, MeOH), **19**; +16.9 (c=1.1), **20**; +17.3 (c=1.1), **21**; +25.5 (c=0.8), **22**; +27.6 (c=1.4), **23**; -25.7 (c=1.6), **24**; -15.6 (c=1.8).
- 12. Schreiber, S.L.; Liew, W.F. Tetrahedron Lett. 1983, 48, 2226.
- 13. (a) Okamura, W.H.; Aurrecoechea, J.M.; Gibbs, R.A.; Norman, A.W. J. Org. Chem. 1989, 54, 4072. (b) We thank Professor Okamura for sending us details on the experimental conditions.
- 14. (a) Krow, G.R. Org. Reactions, John Wiley & Sons Inc. 1993, 43 p 251. (b) Ibid. p 260.
- 15. Momose, T.; Muraoka, O. Chem. Pharm. Bull. 1982, 26, 2589. Also oxidation of 3-acetylbicyclo[1.2.3]octane with MCPBA took 8 d.
- (a) Seyferth, D.; Marmor, R.S.; Hilbert, P. J. Org. Chem. 1971, 36, 1379. (b) Gilbert, J.C.; Weerasooriya, U. J. Org. Chem. 1979, 44, 4997. (c) Gilbert, J.C.; Weerasooriya, U. J. Org. Chem. 1982, 47, 1837.

(Received in UK 21 March 1997)