A Facile and Enantiospecific Synthesis of 2(S)- and 2(R)-[1'(S)-Azido-2-phenylethyl]oxirane

Arun K. Ghosh,* Sean P. McKee, Hee Yoon Lee and Wayne J. Thompson

Department of Medicinal Chemistry, Merck Sharp & Dohme Research Laboratories, West Point, Pennsylvania 19486, USA

The title azidoalkyl oxiranes were synthesized efficiently in an enantiomerically pure form utilizing readily available diethyl p-tartrate.

Protected aminoalkyl epoxides have been utilized extensively in the synthesis of potent and selective inhibitors of HIV-1 protease¹ and other aspartic proteases.² Accordingly, several stereoselective syntheses of variously protected aminoalkyl epoxides have appeared in the literature.³ The majority of previous syntheses, however, have shortcomings with regard to variation of substituents and stereochemical control as well as optical purity due to the use of rather sensitive α -amino aldehydes as the intermediates. In conjunction with our studies on the design and synthesis of HIV-1 protease inhibitors, we were interested in the synthesis of the previously unknown 2(S)- and 2(R)-[1'(S)-azido-2-phenylethyl]oxiranes (1 and 2) in optically pure form. In this paper, we report an efficient and enantiospecific route to azido oxiranes 1 and 2 starting from optically pure and commercially available diethyl D-tartrate (Scheme 1). The strategy is quite efficient and potentially can provide a convenient access to other azidoalkyl oxiranes in enantiomerically pure form.

The known⁴ benzylidene acetal 4 was prepared conveniently in multigram quantity by reaction of diethyl p-tartrate with benzaldehyde and triethyl orthoformate in the presence of a catalytic amount of toluene-p-sulfonic acid (84% after distillation, b.p. 146 °C/0.2 mmHg). Reductive cleavage of the acetal 4 with a mixture of lithium aluminium hydride (LAH) and aluminum chloride following Seebach's procedure5 afforded the D-threitol derivative ($[\alpha]_D^{23} - 16.4, c \, 0.2, EtOH$) in 97% isolated yield (Scheme 2). Reaction of the resulting threitol derivative with acetone in the presence of toluene-psulfonic acid provided the desired isopropylidene derivative 5 exclusively in 91% yield after silica gel chromatography $([\alpha]_D^{23} + 16.8, c \ 1.8, CHCl_3)$.⁶ Protected threitol **5** was then converted to the desired epoxide 6 in the following two step sequence: (i) catalytic hydrogenation of 5 with Pearlman's catalyst in absolute ethanol under atmospheric pressure to afford the diol and (ii) treatment of the resulting diol with triphenylphosphine and diethyl azodicarboxylate to result in epoxide 6 in 77% yield after distillation (b.p. 30°C/0.8 mmHg).7

Introduction of the alkyl side chain was readily achieved by regiospecific opening of the epoxide ring of **6** with a carbon nucleophile. Thus, treatment of epoxide **6** with phenylmagnesium bromide (2.2 equiv.) in the presence of a catalytic amount of copper(1) cyanide (10 mol%) at -40 to 0 °C for 3 h furnished the alcohol **7** (93% yield) exclusively. Mitsunobu azidation⁸ of alcohol **7** with triphenylphosphine (1.2 equiv.), diethyl azodicarboxylate (1.2 equiv.) and diphenylphosphoryl azide (1.2 equiv.) in tetrahydrofuran (THF) (-10 to 23 °C for 12 h) afforded a mixture (3:1) of azide and the corresponding elimination product, which, without separation, was subjected to 40% aqueous acetic acid at 90 °C for 2 h to provide pure azido diol **8** in 64% yield after silica gel chromatography.



Scheme I

Azido diol **8** was then converted efficiently to the desired oxiranes **1** and **2** in the following two- and three-step sequences (see Scheme 2). Thus, reaction of **8** with 2-acetoxy-isobutyryl chloride (1.5 equiv.) in chloroform at 23 °C for 4 h followed by exposure of the resulting chloroacetate derivative to sodium methoxide (5 equiv.) in THF (23 °C for 6 h) resulted in oxirane **1** ($[\alpha]_D^{23} + 12.9, c$ 1.15, CHCl₃)⁹ after silica gel chromatography (72% yield). On the other hand, selective benzoylation of the primary alcohol of **8** with benzoyl chloride (1.1 equiv.) in pyridine at -10 to 23 °C for 12 h and subsequent mesylation with mesyl chloride (1.5 equiv.) in pyridine (-10 to 23 °C for 12 h) provided the corresponding mesylate. Alkaline hydrolysis of the benzoate with sodium methoxide in THF furnished the oxirane **2** ($[\alpha]_D^{23} + 20.1, c$ 1.29, CHCl₃)[†] in 66% yield starting from azido diol **8**.



Scheme 2 Reagents and conditions: i, LAH, AlCl₃, Et₂O–CH₂Cl₂; ii, MeC₆H₄-*p*-SO₃H, acetone, 23 °C; iii, H₂, Pd(OH)₂; iv, Ph₃P, EtO₂CN=NCO₂Et, benzene; v, PhMgBr, CuCN, THF, -40 to 0 °C; vi, (PhO)₂P(O)N₃, Ph₃P, EtO₂CN=NCO₂Et, THF, -10 to 23°C; vii, aq. AcOH (40%), 90 °C; viii, MeCO₂C(Me)₂COCl, CHCl₃, 23 °C; ix, NaOMe, THF, 23 °C; x, C₆H₅COCl, pyridine, -10 to 23 °C; vi, MeSO₂Cl, pyridine, -10 to 23 °C

[†] All new compounds gave satisfactory spectroscopic and analytical results. For example, oxirane 1: $([α]_D^{23} + 12.9, c \ 1.15, CHCl_3)$; ¹H NMR (300 MHz): δ 7.4–7.2 (m, 5H), 3.6 (m, 1H), 3.1 (m, 1H), 2.95 (dd, 1H, *J* 4.6, 13.9 Hz), 2.8 (m, 3H). Oxirane 2: $([α]_D^{23} + 20.1, c \ 1.29, CHCl_3)$; ¹H NMR (300 MHz): δ 7.4–7.2 (m, 5H), 3.4 (dd, 1H, *J* 7.3, 13.1 Hz), 3.1 (m, 1H), 2.9 (dd, 2H, *J* 3.3, 7.5 Hz), 2.75 (dd, 1H, *J* 4.2, 4.6 Hz), 2.51 (dd, 1H, *J* 2.6, 4.8 Hz).

In conclusion, this methodology offers an efficient and enantiospecific route to titled azido oxiranes and also holds considerable promise as a general route to other structurally related oxiranes. Further studies and application of this sequence in the synthesis of potent HIV-1 protease inhibitors are currently in progress.

The authors thank Professor Samuel Danishefsky for helpful discussions and acknowledge the encouragement and support of Dr Joel R. Huff and Dr Paul S. Anderson.

Received, 20th November 1991; Com. 1/05896G

References

I. S. Sigal, J. R. Huff, P. L. Darke, J. P. Vacca, S. D. Young, S. J. deSolms, W. J. Thompson, T. A. Lyle, S. L. Graham and A. K. Ghosh, Eur. Pat. 0337714, 1988; N. A. Roberts, J. A. Martin, D. Kinchington, A. V. Broadhurst, C. J. Craig, I. B. Duncan, S. A. Galpin, B. K. Handa, J. Kay, A. Krohn, R. W. Lambert, J. H. Merrett, J. S. Mills, K.-E. B. Parkes, S. Redshaw, A. J. Ritchie, D. L. Taylor, G. J. Thomas and P. J. Machin, *Science*, 1990, 248, 358; D. H. Rich, C.-Q. Sun, J. V. N. Vara Prasad, A. Pathiaseril, M. V. Toth, G. R. Marshall, M. Clare, R. A. Mueller and K. Houseman, *J. Med. Chem.*, 1991, 34, 1225; T. A. Lyle, C. M. Wiscount, J. P. Guare, W. J. Thompson, P. S. Anderson, P. L.

Darke, J. A. Zugay, E. A. Emini, W. A. Schleif, J. C. Quintero, R. A. F. Dixon, I. S. Sigal and J. R. Huff, *J. Med. Chem.*, 1991, **34**, 1228 and references cited therein.

- 2 W. J. Greenlee, J. Med. Res. Rev., 1990, 10, 173 and references cited therein.
- 3 B. E. Evans, K. E. Rittle, C. F. Homnick, J. P. Springer, J. Hirshfield and D. F. Veber, J. Org. Chem., 1985, 50, 4615; J. R. Luly, J. F. Dellaria, J. J. Plattner, J. L. Soderquist and N. Yi, J. Org. Chem., 1987, 52, 1487.
- 4 R. M. Wenger, *Helv. Chim. Acta*, 1983, **66**, 2308; D. W. Curtis, D. A. Laidler, J. F. Stoddart and G. H. Jones, *J. Chem. Soc.*, *Chem. Commun.*, 1975, 833.
- 5 D. Seebach and E. Hungerbuhler, *Modern Synthetic Methods*, ed.
 R. Scheffold, Salle and Sauerlander, Berlin, 1980, 152.
- 6 M. L. Lewbart and J. J. Schneider, *J. Org. Chem.*, 1969, 34, 3505;
 S. Valverde, B. Herradon and M. Martin-Lomas, *Tetrahedron Lett.*, 1985, 26, 3731.
- 7 E. Abushanab, P. Vemishetti, R. W. Leiby, H. K. Singh, A. B. Mikkilineni, D. C.-J. Wu, R. Saibaba and R. P. Panzica, J. Org. Chem., 1988, 53, 2598.
- 8 B. Lal, B. N. Pramanik, M. S. Manhas and A. K. Bose, *Tetrahedron Lett.*, 1977, 23, 1977; T. Schiori, K. Ninomiya and S. Yamada, J. Am. Chem. Soc., 1972, 94, 6203; O. Mitsunobu, *Synthesis*, 1981, 1.
- 9 J. P. H. Verheyden and J. G. Moffatt, J. Org. Chem., 1972, 37, 2289; S. Greenberg and J. G. Moffat, J. Am. Chem. Soc., 1973, 95, 4016.