Synthesis of a Biologically Active Analogue of Antibiotic A-32390A

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The total synthesis of a biologically active analogue of antibiotic A-32390A is described.

For some years we have been interested in the synthesis and biosynthesis of fungal metabolites containing the isonitrile functionality. Central to our syntheses of isonitrins A and B was the development of new methodology for the preparation of vinyl formamides from thiooximes. 1-3 Subsequent dehydration with trifluoromethane sulfonic anhydride afforded the corresponding vinyl isonitrile. In an effort to further define the scope and limitations of these reactions we have examined systems pertinent to A-32390A 1, an antibiotic isolated from a fungus of the genus *Pyranochaeta*⁵ (Fig. 1). Schöllkopf *et al.*

Scheme 1 Reagents and conditions: i, pToISCl (3 equiv.), propylene oxide (50 equiv.), 4 Å mol. sieves, CH_2Cl_2 , 2 h, room temp., 91%; ii, PPh₃ (3 equiv.), acetic formic anhydride (3 equiv.), propylene oxide (10 equiv.), CH_2Cl_2 , room temp; iii, DBU (5 equiv.), CH_2Cl_2 , 30 min, room temp., 70%; iv, DBU (5 equiv.), CH_2Cl_2 , 48 h, room temp., 40%; v, $Hg(OAc)_2$, (1 equiv.), Et_3N (5 equiv.), CH_2Cl_2 , 10 min, room temp., 80%; vi, $Hg(OAc)_2$ (1 equiv.), DBU (5 equiv.), CH_2Cl_2 , 15 min, room temp., 70%; vii, PPh₃ (3 equiv.), acetic formic anhydride (3 equiv.), propylene oxide (10 equiv.), CH_2Cl_2 , 24 h, room temp. then $Hg(OAc)_2$ (3 equiv.) then DBU (5.5 equiv.), 55%; viii, LiOH (1.1 equiv.), $THF-H_2O$ 4:1 then H_3O^+ , 85%

have synthesised 1 along with a number of structural variants of both the 'carbohydrate portion' and the vinyl isonitrile fragment and demonstrated their efficacy against various bacterial and fungal strains.^{6,7}

As no precedent existed for the rearrangement of α -carboxy thiooximes to vinyl formamides studies were first undertaken on simple systems. To this end thiooxime 2 was prepared from L-valine benzyl ester according to the method of Gordon.8 Exposure of 2 to the rearrangement conditions failed to deliver any of the desired vinyl formamide 3, however, two N-formylated products, the α -thioformamide 4 and the α-acetoxyformamide 5 were isolated in reasonable yield.† Preliminary experiments indicated that while the minor, α -acetoxy formamide 5, underwent facile elimination to the desired vinyl formamide upon treatment with 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), treatment of the α -thioformamide 4 under equivalent conditions gave rise to a sluggish reaction and only a modest (40%) yield of 3. Treatment of 4 with Hg(OAc)2-Et3N was found to lead smoothly to 5 in 86% yield. However, a combination of $Hg(OAc)_2$ and DBU in the same pot gave a 70% yield of the vinyl formamide 3 directly. Thus the desired transformation of 2 to 3 could be achieved by sequential treatment of the thiooxime 2 with PPh3, Hg(OAc)2 and DBU to afford the vinyl formamide 3 in 55% overall yield. Hydrolysis of 3 afforded the free acid 6, a key intermediate in Schöllkopf's synthesis of A-32390A 1,6 (Scheme 1). Dehydration of 3 proceeded uneventfully to deliver the isonitrile in good yield under previously described conditions.4,9

We sought to apply the modified methodology to a total synthesis of 7, a biologically active analogue of A-32390A 1.6 Thus 7 can be disconnected to the erythritol fragment and two identical vinyl isonitrile portions. The stereochemistry of the central portion suggested that it could be prepared from a suitable L-tartrate ester while the vinyl isonitriles would be prepared as described above. Thus diethyl L-tartrate was protected as its tert-butyldimethylsilyl (TBDMS) ether¹⁰ and reduced with LiEt₃BH¹¹ to afford 10. At this point we elected to attach the amino acid residues to the central portion prior to establishing the vinyl formamide in order to probe the thiooxime to the vinyl formamide rearrangement in a more demanding environment. The resulting primary diol 10 was coupled to L-N-Z-valine using 1,3-dicyclohexylcarbodiimide (DCC)-4-dimethylaminopyridine (DMAP)¹² then deprotected by hydrogenolysis to afford the free diamine 11 in essentially quantitative yield over the two steps. Treatment of 11 with 6 equiv. of p-TolSCl in the presence of propylene oxide8 smoothly converted 11 to the thiooxime 12 as a mixture of geometric isomers. After treatment of 12 with acetic formic anhydride-PPh₃ then Hg(OAc)₂ the ¹H NMR spectrum of crude product indicated that the desired α -acetoxy compound had indeed formed but silica gel chromatography led to complete decomposition, thus we effected an in situ elimination to deliver vinyl formamide 13. Unfortunately this very polar compound co-eluted with the triphenylphospine oxide. This practical problem was resolved by use of polymer

Scheme 2 Reagents and conditions: i, TBDMSCI (2.4 equiv.), inidazole (5 equiv.), dimethylformamide (DMF), 14 h, 35 °C, quant.; ii, LiEt₃BH (6 equiv.), THF, 30 min, 0 °C, 75%; iii, L-N-Z-valine (2.1 equiv.), DCC (2.1 equiv.), DMAP (0.2 equiv.), CH₂Cl₂, 24 h, room temp., quant.; iv, H₂, 10% Pd/C, EtOAc, quant.; v, p-TolSCl (6 equiv.), propylene oxide (100 equiv.), 4 Å mol. sieves, CH₂Cl₂, 3 h, room temp., 85%; vi, polymer supported PPh3 (6 equiv.), acetic formic anhydride (6 equiv.), propylene oxide (20 equiv.), CH₂Cl₂, 24 h, room temp., then Hg(OAc)₂ (2 equiv.) then DBU (5 equiv.), room temp., 68%; vii, Tf₂O (2 equiv.), Pri₂EtN (12 equiv.), CH₂Cl₂, 30 min, -78 °C, 82%

Scheme 3 Reagents and conditions: i, 90% formic acid, 4 h, room temp., quant.; ii, TMSCN, DMF then Tf₂O (2 equiv.), Pri₂EtN (12 equiv.), CH₂Cl₂, 30 min, -78 °C, 79%; iii, methanolic citric acid, 15 min, room temp., 85%

supported triphenylphosphine^{13,14} which could be removed by filtration to afford 13 as a crystalline solid, (Scheme 2).

Vinyl formamide 13 was dehydrated to the isonitrile 14 in 82% yield upon treatment with 2 equiv. of Tf_2O in the presence of $Pr^i{}_2EtN$ (12 equiv.).^{4,9} Attempted deprotection with a variety of fluoride sources led to complete decomposition presumably via hydrolysis of the ester linkages by residual water in the fluoride source. Thus we were forced to exchange protecting groups at the vinyl formamide stage. The TBDMS groups were quantitatively removed by stirring with 90% formic acid at room temp. for 4 h. The resulting crystalline diol-formamide 15 has been prepared previously by Schöllkopf⁶ and as such represents a formal total synthesis. In order to effect dehydration to an isonitrile Schöllkopf employed prior protection of the hydroxy groups as formate esters. However, they experienced significant decomposition of the target molecule during the final base catalysed deprotection.6 In an effort to overcome this problem the diol was blocked with trimethylsilyl (TMS) groups¹⁵ and the crude product was dehydrated to afford 16 in 79% yield over two steps. The TMS groups were successfully removed in 85% yield by a brief exposure to methanolic citric acid, 16 to afford 7,‡ (Scheme 3).

In summary, the methodology for the preparation of vinyl isonitriles from thiooximes has been extended to α -carboxy systems and applied to the total synthesis of a biologically active analogue of the natural product, Antibiotic A-32390A

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Footnotes

- † A discussion of the mechanism will appear in a forthcoming full
- ‡ Compound 7 had identical chemical and physical data to that reported previously.

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