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## Intramolecular Acylations of γ-Benzoyloxy Phosphine Oxides: Synthesis of Optically Active Cyclopropyl Ketones

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Abstract: Intramolecular acylation of  $\gamma$ -benzoyloxy phosphine oxides with LDA in the presence of Me<sub>3</sub>SiCl gives silyl ethers with high stereoselectivity: treatment of these silyl ethers with t-BuOK gives optically active di- and trisubstituted cyclopropyl ketones in good yield.

We have previously described a synthesis of racemic cyclopropyl ketones *trans*-3. Intramolecular acylation of diphenylphosphinoyl esters 1 generated hydroxy ketones 2 which were cleanly converted into cyclopropyl ketones *trans*-3 upon treatment with potassium *tert*-butoxide.<sup>1</sup> This route was far from ideal: hydroxy ketones 2 exist as a mixture of open chain and closed hemiketal forms and decompose in solution to dihydrofurans 4. However, carrying out the acylation in the presence of an internal Me<sub>3</sub>SiCl trap solved these problems since treatment of benzoate esters *anti*-5 with Me<sub>3</sub>SiCl and then LDA gave single diastereoisomers of silyl ethers  $6.^2$ 



We now report syntheses of optically active silyl ethers and their direct conversion into optically active cyclopropyl ketones (with two or three ring substituents), an approach which combines the chemical stability of silyl ethers such as **6** with the synthetic utility of hydroxyketones such as **2**. Compounds containing cyclopropane rings show considerable promise as antiviral agents<sup>3</sup> and conformationally restricted analogues of natural amino acids.<sup>4</sup>

1,2 Diols 7a (76% ee) and 7b (86% ee), synthesised by Sharpless asymmetric dihydroxylation,<sup>5</sup> were converted into optically active hydroxy phosphine oxides 9 and 10 using a two step sequence. Vinyl phosphine oxides 8 were obtained by DBU elimination<sup>6</sup> of the cyclic sulfites<sup>7</sup> derived from 1,2 diols 7. Addition of LiAlH<sub>4</sub> or Me<sub>3</sub>CuLi<sub>2</sub> gave hydroxy phosphine oxides 9<sup>8</sup> and 10. Phosphine oxide 10 was produced with good *anti* selectivity<sup>9</sup> (82:18) as a mixture easily separable by HPLC.



Benzoylation of hydroxy phosphine oxides 9 and 10, followed by intramolecular acylation using the LDA/Me<sub>3</sub>SiCl internal quench procedure gave good yields of silyl ethers 13 and 14 with excellent stereoselectivity (see Table). In particular, intramolecular acylation of benzoates 11 with only one chiral centre ( $\gamma$  to phosphorus) gave good yields of silyl ethers 13 as single diastereoisomers (entries 1 and 2), the sense of induction being determined by a 500 MHz NOESY analysis of silyl ether 13a. We interpret these results in terms of the configurational instability of lithiated phosphine oxides:<sup>10</sup> lithiation of benzoates 11 is followed by equilibration to the lithium derivative whose true structure is 16. Acylation then leads to the observed diastereoisomer.<sup>2</sup>



Table: Intramolecular acylation of benzoates 11 and 12								
	Alcohol 9 or 10			Benzoate 11 or 12		Silyl ethers 13-14 or 15		
Entry		R	R'		Yield (%)		Yield (%)	dias.ratio
1	9a	n-Bu	Н	11a	94	13a	55	>98:2
2	9b	Ph	Н	11b	94	13b	88	>98:2
3	anti-10	n-Bu	Me	anti-12	86	14a	75	95:5
4	syn-10	n-Bu	Me	syn-12	85	14b 15	64 20 <sup>a</sup>	>98:2 87:13

<sup>a</sup> Decomposed to the dihydrofuran on standing.

The successful intramolecular acylation of benzoates *syn*- and *anti*-10 also suggests that the chiral centre  $\gamma$  to phosphorus is controlling the stereoselectivity of the reaction: both *syn*- and *anti*-10 generated silyl ethers 14 with the same relative stereochemistry between the chiral centres  $\alpha$  and  $\gamma$  to phosphorus (entries 3-4).<sup>11</sup> Thus, in intermediate 17, it is relatively unimportant whether the methyl group sits in the axial position **B** (benzoate *syn*-12) or in the equatorial position **A** (benzoate *anti*-12). This is, perhaps, not surprising since the axial position **B** does not suffer from 1,3 diaxial interactions.

Treatment of silvl ethers 13a-b and 14a-b with potassium *tert*-butoxide in *tert*-butanol initiated a remarkable cascade of events which led to the formation of cyclopropyl ketones 20a-b, 22 and 23. We propose that desilvlation was followed by ring opening to generate alkoxy ketones 18 which rearranged by phosphinoyl transfer to enolates 19. Cyclisation of these enolates led to the formation of the cyclopropyl ketones. In the synthesis of cyclopropyl ketone 22, each chiral centre is controlled by a different factor: centre (a) is controlled by the inversion of a displacement reaction  $(21\rightarrow 22)$ , centre (b) is already present in benzoate *anti*-12 and centre (c) is controlled by which face of the enolate reacts.



In particular, reaction of silyl ethers 13a-b with potassium *tert*-butoxide in *tert*-butanol generated cyclopropyl ketones 20a (*trans:cis* >95:5) and 20b (*trans:cis* >95:5).<sup>12</sup> The cyclisation of enolates 19 is stereoselective because the substituents prefer to be *trans* in the transition state. On standing in acidic solution (CDCl<sub>3</sub>) 20b gave a thermodynamic 67:33 *trans:cis* ratio. No equilibration occurred in basic solution and all other ratios quoted in this letter are kinetically controlled.

Our route is particularly well suited to the synthesis of optically active cyclopropyl ketones with a chiral centre at each corner of the three-membered ring: compound 22 was obtained as a 94:6 mixture of diastereoisomers in 84% yield. This is a result of the highly stereoselective cyclisation of enolate *syn-21* in which both alkyl groups are *trans* to the forming phenyl ketone (figure 24). This favourable cyclisation is not available to enolate *anti-21*, and therefore reaction of silyl ether 14b gave a 61:27:11 mixture of cyclopropyl ketones *trans-23*, *cis-23* and *ent-22* in a poor 55% yield. Presumably, a S<sub>N</sub>1 mechanism is competitive with stereospecific S<sub>N</sub>2 inversion.

Optically active di- and trisubstituted cyclopropyl esters and amides are most usually synthesised by the metal-catalysed addition of carbene equivalents to alkenes in the presence of chiral ligands<sup>13</sup>, but the diastereoselectivities are poor unless very bulky diazoacetates are used<sup>14</sup> or the reaction is intramolecular.<sup>15</sup> Several asymmetric syntheses of cyclopropanes using homochiral nucleophiles as chiral auxiliaries are known. In particular, Johnson<sup>16</sup> has synthesised ketone **20b** by adding the anion of a chiral sulfoximine to an unsaturated ketone and Hanessian<sup>17</sup> has synthesised tri- and tetrasubstituted cyclopropyl carbonyl compounds by adding the anion of a *trans*-chloroallyl phosphonamide to unsaturated ketones, esters and amides.

Our route combines the advantages of these two approaches: our cyclisation precursors are synthesised by a catalytic asymmetric method, the Sharpless dihydroxylation reaction, and the intermediacy of anionic intermediates in the ring-closure enables our reactions to be highly diastereoselective in favourable cases. The method is particularly useful for the synthesis of di- and trisubstituted cyclopropyl ketones with one or two alkyl groups *trans* to the acyl substituent.

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## **References and Notes**

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- 9. The *anti* selectivity of these additions was established by 500 MHz NMR analysis of silyl ether 14 (R = Ph; R' = Me) and by coupling constant correlations (Footnote 11).
- 10. We have recently demonstrated that lithiated phosphine oxides are not configurationally stable in THF at -78 °C: O'Brien, P.; Warren, S. *Tetrahedron Lett.*, **1995**, *36*, 8473-8476.
- 11. For silyl ethers 13 and 14, we have noticed the following trends in proton NMR coupling constants:



- Typical <sup>3</sup>J<sub>HH</sub> coupling constant values for adjacent protons in cyclopropyl rings: <sup>3</sup>J<sub>HH</sub>(trans)=4-9.5 Hz and <sup>3</sup>J<sub>HH</sub>(cis)=7-13 Hz: Williams, D.H.; Fleming, I. Spectroscopic Methods in Organic Chemistry, McGraw-Hill, London, 5th edition, 1995, page 164.
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