Letter

## Preparation and reactions of 4-oxaspiro [2.3] hexanes<sup>†</sup>

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Received (in Gainesville, FL, USA) 13th December 2000, Accepted 14th February 2001 First published as an Advance Article on the web 12th April 2001

2-Methyleneoxetanes were converted in excellent yields to 4oxaspiro[2.3] hexanes under modified Simmons–Smith conditions. Treatment of the oxaspirohexanes with BF<sub>3</sub>·Et<sub>2</sub>O provided cyclopentanones, cyclobutanones or 4-methylenetetrahydrofurans, depending on the substituents.

We have recently developed the first general synthesis of 2methyleneoxetanes  $1^{1}$  and have found this class of strained heterocycles to display a diverse range of reactivities. We anticipated that the electron-rich exocyclic enol ethers of 1 would provide a useful handle for reactions with electrophiles. Indeed, 2-methyleneoxetanes were shown to undergo facile and efficient epoxidation with anhydrous dimethyldioxirane (DMDO) to give 1,5-dioxaspiro[3.2]hexanes 2.2 Moreover, 2 displayed a fascinating dichotomy in reactions with nucleophiles, yielding multifunctionalized ketones 3 with most neutral and anionic nucleophiles but resulting in 2,2-disubstituted oxetanes 4 in the presence of some Lewis acids (Scheme 1).<sup>3</sup> Another class of compounds that we anticipated would be accessible and of interesting reactivity was the 4oxaspiro[2.3]hexanes 5. In this communication, we report their preparation and describe our initial observations on their reactivity.

Previous preparations of compounds containing the 4oxaspiro[2.3]hexane framework are limited.<sup>4-9</sup> The one example directly related to our work involved the conversion of  $\beta$ -tosyloxyketone **6** to the  $\alpha$ -alkylidene oxetane **7**.<sup>7</sup> Reaction of **7** with dichlorocarbene afforded 4-oxaspiro[2.3]hexane **8**. Direct insertion of a methylene group would be more broadly useful and was, consequently, our goal in investigating the preparation of 4-oxaspiro[2.3]hexanes **5** from 2-methyleneoxetanes **1**.



† Electronic supplementary information (ESI) available: full experimental details and characterization data for all previously unreported compounds. See http://www.rsc.org/suppdata/nj/b0/b010095l/

2-Methyleneoxetanes  $1a-1c^{1}$  and  $1e^{10}$  were prepared as previously described by the reaction of the corresponding  $\beta$ lactones with dimethytitanocene. Compound 1f was prepared in the same manner from the known  $\beta$ -lactone 3,3-dimethyl-4, 4-diphenyloxetan-2-one,<sup>11</sup> and the  $\beta$ -lactone<sup>12</sup> required for 2methyleneoxetane 1d was prepared by benzylation of 9.<sup>1</sup>

Simmons and Smith disclosed the potential of iodomethylzinc iodide, generated *in situ* from the reaction of the zinccopper couple with diiodomethane, for the cyclopropanation of alkenes in 1958.<sup>13</sup> Since then, a variety of more convenient and reproducible procedures for preparing the Simmons-Smith reagent have been described.<sup>14,15</sup> 2-Methylenetetrahydrofurans have been cyclopropanated,<sup>16–18</sup> most successfully by the Furukawa procedure (diethylzinc and a dihalomethane).<sup>19</sup> Thus, we anticipated that this modification might be best for the cyclopropanation of 2-methyleneoxetanes. Nevertheless, we were concerned about the effect of Lewis acidic by-products on the reactants and/or products.

Initially, diiodomethane and diethylzinc in  $Et_2O$  at -5 °C were treated with 2-methylene-3-phenyloxetane (1a). The reaction was then allowed to warm to room temperature and stirred for 2 h. 4-Oxaspiro[2.3]hexane **5a** was isolated in low yield, and GC/MS showed numerous by-products. However, when the reaction temperature was maintained between -5 and 0 °C for 4 h, the starting material disappeared and only the desired product was produced; further warming resulted in numerous products. Quenching the reaction after 4 h at 0 °C, followed by purification, provided **5a** in 83% isolated yield. This protocol proved to be efficient for other 2-methyleneoxetanes, as is evident from the results in Table 1.

To our knowledge, there has been no examination of the behavior of 4-oxaspiro[2.3] hexanes in the presence of Lewis or protic acids. The corresponding 4-oxaspiro[2.2]pentanes 10 are known to undergo smooth rearrangements to cyclobutanones 12.<sup>20</sup> The reorganization is more rapid when R,  $R' \neq H$ , supporting the presumed intermediacy and subsequent rearrangement of a carbonium ion related to 11.21 Recently, another related system, a 5-oxaspiro[3.2]hexane 13 was reported to rearrange to cyclopentenone 14 when treated with LiI.<sup>22</sup> Also, 2-oxo-1-oxaspiro[3.2] hexanes 15 provided a complex mixture of products when treated with  $BF_3 \cdot Et_2O^{23}$ However, 15 did rearrange to lactones 16-18 when exposed to catalytic quantities of Cu(acac)<sub>2</sub>, suggesting a cyclopropyl cationic species of some sort that rearranges, based both on the catalyst and on the nature of R. Oxetanes (4-membered rings containing O) do not open as readily as oxiranes (3membered rings containing O). In light of this, if ring opening of 5 occurred in a fashion similar to that of 10, the resultant carbocation would not have the immediate pathway of a facile



DOI: 10.1039/b0100951

New J. Chem., 2001, 25, 673–675 673

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 Table 1
 Synthesis and reactions of 4-oxaspiro[2.3] hexanes

	$\begin{array}{c} R \sqsubseteq O \\ \Box O \\ 1a-f \end{array} \xrightarrow{CH_2 I_2, \ Zr} \\ Et_2 O, \ O \xrightarrow{O} \\ \end{array}$	$\begin{array}{c} \text{HEt}_2 \\ \text{C, 4 h} \\ \textbf{5a-f} \end{array} \xrightarrow{\text{BF}_3 \circ \text{OEt}_2 (5 \text{ ec})} \\ \begin{array}{c} \text{BF}_3 \circ \text{OEt}_2 (5 \text{ ec}) \\ \text{CH}_2 \text{CH}_2 \text{CH}_2 \end{array}$	uiv.) → 19–24	
Oxaspirohexane	Isolated yield (%)	$BF_3 \cdot Et_2O$ conditions	Product	Isolated yield (%)
Ph O 5a	83	35 °C, 17 h	Ph0 19	35
Ph O 5b	87	35 °C, 17 h	Ph O	5
Ph 5c	90	70 °C, 13 h	=	7
Ph Ph 5d	82	35 °C, 35 h	Ph Ph 22	16
Ph 5e	81	0°C, 2 h		48
Ph + O Ph 5f	98	0 °C, 2 h	Ph Ph	42

1,2-shift to an oxygen-stabilized carbonium ion. Thus, the fate of **5** is not easily predictable.

Our prediction that the 4-oxaspiro[2.3]hexanes 5 might not react as readily as the corresponding epoxides 10 was confirmed. When 5a was treated with LiI and heated at reflux in THF for 24 h, the starting material was recovered. Conversely, oxaspirohexanes 5a-f underwent rearrangements when treated with  $BF_3 \cdot Et_2O$ . However, the requirements for reaction and the outcome of the rearrangements varied significantly.

Cyclobutanones 19, 20 and 23 (Table 1) were isolated from oxaspirohexanes 5a, 5b and 5e, respectively. For compounds 5a and 5e, the indicated products were the only ones of significance in the crude reaction mixture, based on GC/MS. The rearrangement of 5b was not as clean; however, cyclobutanone 20 was the major product in GC/MS traces. Oxaspirohexanes 5c and 5d gave a number of isomeric products, based on GC/MS. The predominant product in each case was the 4-methylenetetrahydrofuran (21 and 22). We were unable to separate 21 from one of its structural isomers.<sup>12</sup> Based on <sup>1</sup>H and <sup>13</sup>C NMR, it appears likely that this isomer is cyclobutanone **25**. Oxaspirohexane **5f** provided cyclopentanone **24** as the only product of significance seen in GC/MS traces.

Initially, the divergent outcomes of the rearrangements were somewhat surprising. On further consideration, the results can be rationalized as shown in Scheme 2. Considering a generalized oxaspiro [2.3] hexane complexed to  $BF_3 \cdot Et_2O$  (26), initial ring opening could occur in two directions. Cleavage at O4–C5 would provide carbocation 27. A 1,3-bond migration involving the cyclopropane would then provide an oxygenstabilized carbonium ion 28 that results in a cyclopentanone. Alternatively, 27 could undergo consecutive 1,2-shifts, ultimately leading to oxygen-stabilized cations 30 or 32 (via 29 or 31, depending on the number of 1,2-shifts). If initial bond rupture occurred at O4-C3, the resulting cyclopropyl cation 33 would be expected to reorganize, and one possible product would be methylenetetrahydrofuran 34. There are, obviously, other possible fates for the cations shown, and this could explain the low isolated yields. That the only cyclopentanone observed resulted from substrate 5f suggests that only highly stabilized carbocations rearrange to the cyclopentanones. Even sub-





strate 5e, which would be expected to provide a relatively stable benzylic cation (see 27), undergoes multiple 1,2-shifts, resulting in 23. It is likely that in the formation of 19 and 20, oxaspirohexanes 5a and 5b undergo phenyl migration concurrently with ring opening (equivalent to direct conversion of 26 to 29). With 5a the carbonium ion corresponding to 29 appears to be converted to the product in a reasonably efficient fashion. However, this is not the case with 5b. The presence of other isomers was indicated by GC/MS, although we were unable to isolate any other clean, clearly identifiable products. It seems likely that both directions of initial rupture operate with 5c and 5d. Although 4methylenetetrahydrodrofurans 21 and 22 were the products present in the largest yield, our observation of 25 in the rearrangement of 5c confirms that initial O4-C5 cleavage also occurred.

The importance of the nature of the substituents in controlling reaction outcomes is further illustrated by the temperatures required for the reactions. Those substrates containing a phenyl at C5 provided products under the mildest conditions. The compounds lacking substitution at C5 required an elevated temperature for reaction. There was no improvement in results at an elevated temperature for the other oxaspirohexanes. Because of the involvement of multiple steps in going from reactant to product and the availability of alternative pathways, it is not surprising that the rearrangements of oxaspirohexanes occur with less efficiency and greater divergence in outcomes than the reorganizations of the corresponding oxaspiropentanes (see **10** to **12**).

In summary, we have demonstrated that oxaspirohexanes can be efficiently prepared from the corresponding 2-methyleneoxetanes using the Furukawa modification of the Simmons–Smith reaction. Preliminary studies of the reactivity of the oxaspirohexanes have shown that their behavior on exposure to  $BF_3 \cdot Et_2O$  is governed by the nature of the substituents on the oxetane ring. Further studies are underway to see if the efficiency of any of these reactions can be improved by employing alternative Lewis acids or if the outcome can be more effectively controlled.

## Acknowledgement

A. R. H. thanks the NSF for a CAREER award.

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