## **Chelation Control in the Hydroformylation of Terminal Alkenes**

## **W. Roy Jackson, Patrick Perlmutter," and Guem-Hee Suh**

*Department of Chemistry, Monash University, Clayton, Victoria, Australia 3168* 

Hydroformylation of phosphine-bearing terminal alkenes can lead to a complete reversal of regioselection to that observed for simpler alkenes.

Over the last few decades the hydroformylation of alkenes (the **'0x0'** reaction) has found important applications in industrial syntheses.<sup>1,2</sup> The emphasis in these processes has been on selective production of *linear* products **(2)** and **(4)**  (Scheme 1), which has been achieved through the appropriate choice of metal and ligands.3



From a synthetic standpoint it is the *branched* isomers **(3)**  and *(5)* which appear to be the more interesting and useful. **As**  typical reaction conditions involve a rhodium(0) catalyst associated with a phosphine ligand, inclusion of a phosphine in the alkene framework might lead to a more selective reaction. Adjustment of the distance of the phosphine from the alkene might then provide a means for fine-tuning selectivity (Scheme 2).

In order to test these ideas we have prepared a series of alkenylphosphines **(6)** and **(ll).?** Each **of** these was then subjected to the same set of hydroformylation conditions  $\{ \text{except} \}$  where otherwise indicated in Table 1, {except where otherwise indicated in Table 1,

t **(6a)** and **(6b)** were prepared by reaction of the appropriate Grignard reagent with chlorodiphenylphosphine (ref. **4). (6c)** and **(6d)** were prepared by reaction of lithium diphenylphosphine with the appropriate bromoalkene (ref. *5).* **(ll),** the phosphine oxide of **(6b)** was prepared by either coupling chlorodiphenylphosphine oxide with the appropriate Grignard reagent (ref. **4)** or oxidation of **(6b)** (air, or  $H<sub>2</sub>O<sub>2</sub>$  in acetone).

Table 1. Results of the hydroformylation of various **diphenylphosphinylalk-l-enes.** 

	Alkene	o isano oi proguer				
Entry		(7)	(8)	(9)	(10)	% Yield <sup>b</sup>
	$(6a)^c$			30	20	68
2	(6b)			100		86
3	$(6b)^d$			75	25	85s
4	$(6c)^c$	$21 - 26$	$2 - 9$	$27 - 52$		64—95
5	$(6d)^c$	60	32			96
6	$(6d)$ e	50	27	13	8	54
	(11)	36	64			55f
8	$(1; R = Bu^n)$	25(3)	75(2)			71

 $\%$  Ratio of producta

**<sup>a</sup>**Product ratios determined from the 300 MHz 1H n.m.r. spectra of the crude product in each case, for at least two runs. *b* Yield after distillation. All crude yields were close to quantitative, however significant decomposition accompanied distillation in each case. <sup>c</sup> The remainder of product was the alkane. <sup>d</sup> 50 equiv. of PPh<sub>3</sub> added. *e* No PPh<sub>3</sub> added. f After recrystallisation from ethyl acetate-light petroleum; crude yield 94%. **g** Crude yield.







alkene : PPh<sub>3</sub> : [Rh(OAc)<sub>2</sub>]<sub>2</sub>, (200 : 4 : 1), H<sub>2</sub>/CO (1 : 1, 400) psi), EtOAc, 100°C, 5 h}. Consistent with earlier studies of rhodium-catalysed hydroformylations no significant amounts of aldol by-products were formed.6 The results, for the reactions summarised in Scheme 3, are collected in Table 1. $\ddagger$ 

Several trends emerge from these results. First, in a number of cases, reduction of the product aldehydes **(7)** and **(8)** had occurred yielding the corresponding alcohols **(9)** and **(10).** A linear correlation was observed between the number of intervening methylenes and the oxidation state of the products. For short chains,  $n = 1$  or 2, only alcohols were formed,



whereas for the longer chain,  $n = 3$ , a mixture was produced. The longest chain,  $n = 4$ , yielded only aldehydes (Table 1, entry *5).* 

Secondly, in all cases selection for the branched isomers was better than that for a hydrocarbon analogue, hex-l-ene (entry **8).** Most notably, **4-diphenylphosphinylbut-l-ene (6b)** yielded only *one* product **(9b)** and in high yield **(86%)** (Scheme **4).**  This represents the first example of the total reversal of regioselection in hydroformylation reactions. Although its homologue, **(6c),** produced a mixture of aldehydes and alcohols, the regioselection was again good. In this case alone, however, the yields and product distribution proved difficult to reproduce (Table 1, entry **4).** 

Thirdly, variation in the amount of  $PPh<sub>3</sub>$  present influenced the product distribution. Thus, addition of a large excess of PPh3 led to a reduction in regioselection *(cf.* entries 2 and 3). This is to be expected if intramolecular chelation is controlling the reaction. Alternatively, the absence of  $PPh_3$  resulted in the appearance of some alcohols in the products without significant change in the regioselection *(cf.* entries 5 and **6).** 

Finally, hydroformylation of **(ll),** the phosphine oxide of **(Sb),** was also carried out (entry **7).** Only aldehydes were produced in this case and in a ratio which is only slightly different to that for hex-l-ene (entry 8).

In conclusion, we have demonstrated that terminal alkenes bearing a phosphine group undergo the hydroformylation reaction with high regioselection. This selection is the reverse to that for simple hydrocarbon alkenes. Secondly, we have shown that a simple correlation exists between the oxidation state of the products and the number of intervening methylenes in the phosphinylalkene. These observations appear to reflect the influence of a cyclic transition state, the result of *intramolecular* chelation (Scheme 2), on the nature of the products. These ideas may have potential application to more complex alkenes as well as to other reaction types.§

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 $\ddagger$  All new compounds gave satisfactory spectroscopic and elemental analyses.

<sup>§</sup> While this manuscript was in preparation a report of the application of a chelation-controlled hydroformylation reaction to the preparation of (+)-phyllanthocin appeared: **S.** D. Burke and J. E. Cobb, *Tetrahedron Lett.,* 1986, 27, 4237.