Observation of a stable *cis***-diphosphine solvate rhodium dihydride derived from PHANEPHOS**

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Received (in Cambridge, UK) 12th March 2001, Accepted 24th May 2001 First published as an Advance Article on the web 27th June 2001

The methanol solvate rhodium(PHANEPHOS) forms a stable dihydride which has been characterised in solution by NMR as a pair of equilibrating diastereomers.

Early experiments designed to elicit the mechanism of asymmetric homogeneous hydrogenation provided a contrast between chelate diphosphine and bisphosphine rhodium complexes. The solvate 1a showed no tendency to react with



ambient hydrogen, whilst the solvate 2 formed a characterised dihydride;¹ the difference was attributed to a requirement for trans-diphosphine geometry in the stable solvate with H correspondingly trans to solvent oxygen. This observation has generally been sustained until recently. Aside from reversible ortho-para dihydrogen equilibration by complexes 1a and 1b,² and the likely mediation of a *cis*-dihydride in the formation of the dimeric species 3,³ no further progress had been made prior to the work of Gridney, Imamoto and coworkers.⁴ They demonstrated that the corresponding solvate 4 from a simple Pchiral alkylphosphine ligand formed significant quantities of the *cis*-dihydride 5 (20% at -95 °C and ambient pressure), with two diastereomers formed in a ratio of 10:1.5 Further, this intermediate reacted with the catalytic substrate 6a to form a Rh alkylhydride,6 which then underwent reductive elimination at -50 °C to give the hydrogenated product. Taken together with labelling studies, the results are compatible with path A in Scheme 1, in contrast to the more generally accepted sequence **B**.7

We recently demonstrated the presence of an agostic dihydride intermediate 7 in the hydrogenation cycle of compound **6a** by [PHANEPHOS]Rh⁺, employing *para*-enriched hydrogen and the precursor complex **8** (or the NBD analogue) to identify the transient at -10 to -30 °C by ¹H NMR.⁸ When hydrogenation is complete and the substrate exhausted a second species can be observed, however. By carrying out the hydrogenation of the catalyst precursor in the absence of



Scheme 1 The possible paths for addition of dihydrogen to a dehydroamino ester; path A: H_2 addition prior to substrate (dihydride route). Path **B**: substrate addition prior to H_2 addition (unsaturate route).



substrate the same intermediate is seen, optimally at -40 °C. The δ and J values are entirely consistent with a *cis*-dihydride structure **9**, with one hydride *trans* to phosphorus (δ *ca*. -11) and one *trans* to one of the two solvent oxygens (δ *ca*. -20). Both the intensity and persistence of the signals indicate that it is a relatively robust species. There are two diastereomers **9a** and **9b** in 2:1 ratio, and their NMR spectra have been fully assigned using PHIP++ [Fig. 1(a)].⁹ The chemical shifts are very different from the previous case⁴ where the observed major diastereomer resonates at δ -7.7 and -23.0.

When hydrogenation is carried out under conventional NMR conditions at -80 °C, the same dihydride **9** may be observed, along with small amounts of other Rh hydride resonances not seen in the PHIP spectrum. It is stable up to -40 °C [Fig. 1(b)], and a rough estimate based on integration of the high-field ¹H NMR signals against the CH₂-region of the ligand indicates that 45% of species **9** is formed at equilibrium, making it more accessible than the previously observed case,^{4,5} and to higher

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Fig. 1 (a) The PHIP ¹H NMR spectrum (CD₃OD, 200 MHz) of dihydrides **9a** and **9b** taken after parahydrogen (98% enriched) passage through a solution of complex **8** in CD₃OD at -40° C. Minor diastereomer: $\delta - 10.87$ ($J_{\rm HP}$ 170, 24, $J_{\rm HRh}$ 14, $J_{\rm HH}$ -9.0 Hz), -21.77, ($J_{\rm HP}$ 32, 12, $J_{\rm HRh}$ 21 Hz); major diastereomer : $\delta - 11.46$ ($J_{\rm HP}$ 171, 28, $J_{\rm HRh}$ 15.5, 15, $J_{\rm HH}$ -7.5 Hz), -20.91, ($J_{\rm HP}$ 29, 16, $J_{\rm HRh}$ 22.5 Hz). (b) The ¹H NMR spectrum (CD₃OD, 500 MHz) of dihydrides **9a** and **9b** formed in the hydrogenation of complex **8** at -80 °C, taken at -40 °C, with comparable J and δ values.

temperatures. This may be attributed to the high level of electron donation ensuing from the [2.2]paracyclophane backbone,¹⁰ together with the large bite angle of PHANEPHOS,¹¹ which will favour the dihydride at equilibrium. The two diastereomers are in equilibrium by an unselective mechanism, as indicated by a selective homodecoupling experiment.¹²

When the solution containing complex $\hat{9}$ is held at -80 °C and a solution of compound **6a** in MeOH added, rapid formation of the agostic dihydride **7a** occurs. The signals at $\delta -2$ and -19 are broad at that temperature, and at -70 °C they decay over time without formation of any further observable intermediates. The absence of a 'classical' alkylhydride **10** indicates that **7a** is the only accessible intermediate on the hydrogenation pathway. Further, it must be formed directly from an assumed dihydride precursor rather than by reinsertion of rhodium into the β -CH of **10** after formation of the latter, since the latter pathway would vitiate the earlier PHIP experiment by uncoupling the H–H spins.

In the earlier publication of Gridnev, Imamoto and coworkers,⁴ it was suggested that path **A** could be a viable alternative to the accepted reaction mechanism of path **B** (Scheme 1). We



Fig. 2 The PHIP ¹H NMR spectrum of the hydrogenation of reactant **6b** in CD₃OD in the presence of Rh complex **8** at -27 °C, after 40 5 s pulses of parahydrogen (98%). Spectra taken earlier in the sequence after 16 pulses show only traces of complex **9**.

observed that when the PHIP experiment was carried out with **6b** as substrate at the lower temperature of -27 °C, the solvate dihydride **9a**, **b** could be observed in significant amount, but only late in the reaction sequence when the substrate concentration was depleted (Fig. 2). This opens up the possibility that path **A** may contribute to catalytic turnover in the PHANEPHOS case. Earlier INEPT experiments demonstrated that the agostic intermediate **7a** is in reversible equilibrium with the solvate complex and substrate.⁷ This makes the discrimination between the two pathways quite subtle. Given that both species **7** and **9** are observed in the same experiment under turnover conditions, the result is accessible in principle and a challenge for further work.

We thank Philip Pye and Kai Rossen (Merck, Rahway) for a generous gift of PHANEPHOS, and Johnson-Matthey for the loan of RhCl₃. R. G. thanks BASF AG and Studienstiftung des Deutschen Volkes for a Fellowship. JMB is very pleased to acknowledge an unrestricted grant from Merck, Inc. H. H. and J. B. thank the Deutsche Forschungsgemeinschaft for financial support.

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