A New P-Chiral Aminophosphine Ligand Containing a 2,2'-Coupled Pyrrolidine–Phospholane Ring System. Synthesis and Coordination Properties with Rhodium(ı) and Iridium(ı) Fragments

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A new approach to the synthesis of chiral P,N-ligands containing a stereogenic phosphorus atom is reported; the synthesis, characterization and coordination properties of (PR, 2R, 3R, 2'R, 3'S, 4'S) 2[2'(3',4'-di-tert-butoxy-1'-methyltetrahydropyrrolyl)]-3-methoxy-1-phenyltetrahydrophosphole (P*-N-1), which contains a stereogenic phosphorus in addition to five contiguous stereocentres in the ligand backbone; compound P*-N-1 represents the first example of a new class of chiral P,N-ligands with the heteroatoms incorporated in two 2,2'-coupled pyrrolidine–phospholane rings.

The design and development of chiral aminophosphine ligands is being actively pursued for applications in transition metal asymmetric catalysis. So far, however, very few P,N-ligands are known which contain a stereogenic phosphorus atom in addition to stereocentres in the ligand backbone. This report details our endeavours to design, synthesize and characterize a new class of optically pure P,N-ligands in which the phosphorus donor is a stereocentre and the heteroatoms are incorporated in two 2,2'-connected five-membered rings.

A structural arrangement of this type is expected to favour the formation of metal complexes in which the five-membered chelate ring is forced into a specific conformation, as is often the case for some widely used chiral diphosphine ligands, such as those adopting binaphthyl-like chelate rings.

We also report some preliminary results on the bonding capabilities of **P*-N-1** with selected organorhodium(I) and organoiridium(I) fragments, which are widely used in a variety of homogeneous catalytic reactions.

The synthesis of P*-N-1 was carried out as illustrated in Scheme 1. The isoxazolidine 4,³ obtained as the major isomer of the 1,3-dipolar cycloaddition of the nitrone 2⁴ with the phosphole 3,⁵ was treated with MeI in CH₂Cl₂ to give the isoxazolinium salt 5 in quantitative yield. The salt 5 was

OBu^t OBut Ph Mé _I-'n OBu^t OBut Ph Ph ButC Me HO Me MeO ίv 6 **OBu**t ButC

Scheme 1 Reagents and conditions: i, MeI; ii, MeCO₂H, Zn; iii, MeI, NaH; iv, HSiCl₃

reduced to the amino alcohol 6 with Zn–MeCO₂H,⁶ and after treatment with NaH in THF, was alkylated with MeI to afford the protected phosphine oxide 7. The key step of the synthesis of **P*-N-1** is thus the stereoselective reduction of 7. This reaction was carried out in refluxing benzene for 12 h with HSiCl₃ which, in the presence of NEt₃, is known to bring about the reduction of the P=O functionality with retention of configuration.^{2,7}† In this way, the optically pure aminophosphine ligand (PR, 2R, 3R, 2'R, 3'S, 4'S) 2[2'(3',4'-di-tert-butoxy-1'-methyltetrahydropyrrolyl)]-3-methoxy-1-phenyltetrahydrophosphole (**P*-N-1**)‡ was obtained in 55% yield (based on 4) as a single stereoisomer [³¹P{¹H} NMR (CDCl₃) δ –15.22 (s)] with a known absolute configuration at all six stereogenic centres. This methodology is suitable for the synthesis of a class of P,N-ligands bearing different substituents and in either enantiomeric form.

The ligand **P*-N-1** reacts in deaerated CH_2Cl_2 with an equivalent amount of $[Rh(CO)_2Cl]_2$, $[Rh(cod)Cl]_2$ or $[Ir-(cod)Cl]_2$ (cod = 1,5-cyclooctadiene) to give brown to orange microcrystals of the square-planar complexes (**P*-N-1**)Rh(CO)Cl **8**,‡ (η^1 -**P-P*-N-1**)Rh(cod)Cl **9**,‡ and (η^1 -**P-P*-N-1**)Ir(cod)Cl **10**,‡ respectively (Scheme 2). The ${}^{31}P{}^{1}H{}^{1}NMR$ spectra of **8** and **9** show the expected doublet multiplicity with coupling constants within the range for RhI square-planar complexes containing phosphorus donor ligands **8** [**8**, δ 76.51 (d,

Scheme 2

 $J_{\rm PRh}$ 171.2 Hz); **9**, δ 25.82 (d, $J_{\rm PRh}$ 146.2 Hz); **10**, δ 19.26 (s)]. The coordination of the nitrogen donor, demonstrated by the low-field shift of the N–Me signal, occurs only for the carbonyl complex **8**, which thus contains a bidentate **P*-N-1** ligand. The carbonyl ligand in **8** lies *trans* to the nitrogen atom as inferred from the 13 C{ 1 H} NMR spectrum, which shows the CO resonance as a doublet of doublets at δ 189.12 ($J_{\rm CRh}$ 70.9, $J_{\rm CP}$ 19.0 Hz). 10

Unlike 8, the two diene complexes 9 and 10 do not exhibit a bonding interaction of the pyrrolidine N atom with the metal as shown by the chemical shifts of both the N–Me group (9, δ 2.51; 10, δ 2.75) and the P nucleus (9, δ 29.49; 10, δ 19.26). Thus, a chloride, a bidentate cod ligand and a phosphorus atom complete the square-planar geometry about the metal centre in both 9 and 10. The coordination of the nitrogen donor atoms can readily be achieved by treatment of 9 and 10 in THF with a chloride scavenger such as NaBPh₄ in ethanol. As a result, the complexes [(P*-N-1)Rh(cod)]BPh₄ 11‡ and [(P*-N-1)Ir(cod)]BPh₄ 12‡ are quantitatively obtained, in which the presence of a chelating P,N-ligand ring is shown by the highfield shift of the phosphorus resonance (11, δ 58.50; 12, δ 54.14).8

The potential of P^* -N-1 and related ligands bearing different substituents on the pyrrolidine and phospholane rings in asymmetric catalysis is currently under investigation in our laboratories.

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Footnotes

 \dagger The retention of configuration at the phosphorus atom has been confirmed by reoxidation of the $P^*\text{-}N\text{-}1$ ligand to the starting product 7 by reaction

with hydrogen peroxide, which is known to proceed with complete retention of configuration (L. Horner, *Pure Appl. Chem.*, 1964, **9**, 225).

‡ Satisfactory spectroscopic and analytical data were obtained for all compounds.

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