Palladium-catalysed Enantioselective Grignard Cross Coupling with Use of a new

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A new **ferrocenylaminophosphine** ligand, cyclopentadienyl(7-dimethylamino-1 **-diphenylphosphino-4,5,6,7** tetrahydroindeny1)iron (ptfa), and its palladium dichloride complex have been synthesised, structurally characterised by X-ray analysis of the palladium complex, and shown to be effective in the enantioselective Grignard cross coupling reaction of phenethylmagnesium chloride and vinyl bromide giving 3-phenylbut-1-ene with 79% enantiomeric excess.

Enantiomerically pure ferrocenylphosphines have proved to be excellent chiral ligands for various types of transition metal catalysts.¹ Like 2-(1-N, N-dimethylaminoethyl)-1-diphenylphosphino-ferrocene, ppfa,2 almost all such ferrocenylaminophosphine ligands are based on **1-N,N-dimethylaminoethyl**ferrocene as the enantiomerically pure starting material.' Typically, transition metal complexes of such ferrocenylaminophosphines catalyse hydrogenations, hydrosilations, and aldol-type or other carbon-carbon bond forming reactions, in many cases in high chemical and optical yields.¹ Still challenging however remains enantioselective cross-coupling as exemplified by the reaction of phenethylmagnesium chloride and vinyl bromide giving 3-phenylbut-1-ene as the product. Enantioselectivites are in general moderate for these reactions, except when either very air-sensitive chiral amino acid derivatives like t-Leuphos4 are used or when a large

Ferrocenylaminophosphine Ligand

excess of zinc halide is added to the reaction mixture, as in the case of ppfa5 or similar ferrocene-based ligands *.6*

We here report the synthesis of a new enantiomerically pure ferrocenyl aminophosphine ligand, based on a homoannularly disubstituted ferrocenylamine, its palladium dichloride complex and its use for palladium-catalysed Grignard cross coupling reactions. The enantiomerically pure ferrocenylaminophosphine, **cyclopentadienyl(7-dimethylamino-1-diphenylphosphino-4,5,6,7-tetrahydroindenyl)iron** (ptfa) **2** was prepared by lithiation of the easily accessible $(-)$ -ferrocene derivative^{7.8} 1, and subsequent treatment with chlorodiby an
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and the ferrocentrical experiment catalysed Grignar
reactions. The enantiomerically pure ferroce
hine, cyclopentadieny (7-dimethylamino-1-di

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Ph \nCH-MgCl + H_2C = CH \nCH \nM_2
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M_3
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M_4
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Scheme 1
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Ph \nCH-CH
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M_4
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Scheme 2 *Reagents and conditons:* i, BuLi-hexane **(1.4** equiv.), diethyl ether, room temp., **2** h; ii, CIPPhz **(2** equiv.), reflux, **3** h **(77% 2,20%** *2a);* iii, PdC12(MeCN)2 **(1** equiv.), benzene, room temp. **(95%)**

phenylphosphine. The air-stable homosubsituted main product 2^{\dagger} was isolated in 77% yield $\{[\alpha]_D^{20} +233$ (c 0.456, CHC13), m.p. 186°C). **As** a byproduct, the heterosubstituted aminophosphine **2a** was isolated in 20% yield $\{[\alpha]_D^{20} -3.3$ (c 0.394, CHCl₃), orange oil}. The fact that monolithiation partly takes place at the unsubstituted cyclopentadienyl ring strongly reflects the influence of the *endo*-orientation of the dimethylamino group of **1.** Finally, **2** was treated with $PdCl₂(MeCN)₂$ in benzene to give the palladium dichloride complex 3 in 95% yield $\{[\alpha]_D^{20} -154$ (c 0.173, CH₂Cl₂), m.p. 205-210 °C }. The molecular structure of (\pm) -3 was determined by X-ray crystallography \ddagger and is shown in Fig. 1. From

j. *NMR spectral data* of ptfa **2:** 1H NMR (CDC13, Me4Si, **400.1** MHz) 6 **1.42** (m, **lH), 1.83** (m, **2H), 1.98** (m, **lH), 2.21 (s, 6H), 2.31** (dd, **J -16.0, 6.5 Hz, lH), 2.68** (ddd, **J -16.0,** 6.5, **12.4** *Hz,* **lH), 2.83** (m, lH), **3.21** (m, **1H); 4.07** (s, 5H), **4.17** (m, **lH), 7.15-7.40** (m, **10H);** l3C{lH) NMR (CDCl3 **100.6** MHz) 6 **21.0, 24.0, 24.5, 60.8** (d, **Jc-p 20.0** Hz), **67.3,68.2** (d, **Jc-p 4.0** Hz), **71.1,71.7** (d, **Jc-p 8.9 Hz), 86.4, 94.1, 127.3, 127.8 (d, J_{C-P} 6.6 Hz), 128.1, 132.2 (d, J_{C-P} 20.1 Hz), 134.5** (d, **Jc-p 21.3** Hz), **140.2** (d, **Jc-p 16.2** Hz), **140.2** (d, **Jc-p 12.6** Hz); **31P{lH}** NMR (CDCl3, **85%** H3P04, **162.0 MHz)** 6 **-16.4.**

 \pm *Crystal structure analysis of* (\pm)-3: data collected at 90 K; space group $P2_1/n$, $Z = 4$ for $C_{28}H_{30}Cl_2FeNPPd$, $M_r = 644.65$. $a =$ **15.996(16),** $b = 9.987(9)$, $c = 16.129(14)$ Å, $\beta = 95.82(7)$ °, $V =$ 2563.4(1.2) \mathring{A}^3 , $D_c = 1.67$ g cm⁻³. Intensity data were collected for two octants with $5.5 \le 20.55$ °, 6588 observed, 5749 unique, and 4005 significant $[F_{obs} > 4\sigma(F)]$ structure factors. Structure refined with anisotropic atomic displacement parameters for all non-hydrogen atoms, \dot{H} atoms at calculated positions, $R = 0.0432$ (unit weights) for **307** parameters and **4005** observations. A final difference Fourier synthesis showed features up to $0.9(1)$ e A^{-3} in the vicinity of the metal atoms. Atomic coordinates, bond lengths and angles, and thermal parameters have been deposited at the Cambridge Crystallographic Data Centre. See Notice to Authors, Issue No. **1.**

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Fig. **1** Molecular structure of 3

the known absolute configuration of **17** the configuration of **(-)-3 is** (R) at the benzylic carbon C-7 and (1S,7aR,3aR) for the ferrocene unit.

The air-stable and very easy to handle palladium dichloride complex **3** was found to catalyse efficiently enantioselective Grignard cross-coupling reactions. For example, in the presence of 0.4 mol% of $(-)$ -3, phenethylmagnesium chloride (2 equiv.) reacts with vinyl bromide (1 equiv.) in diethyl ether $(0^{\circ}C; 20 h)$ to give (R) -3-phenylbut-1-ene in 95% yield (isolated), $\{[\alpha]_{D}^{22} - 4.67 \text{ (neat)}$, 79.3% enantiomeric excess (e.e)}. Thus, the enantioselectivity of **3** closely resembles that of t-Leuphos (83% e.e.), which is the highest asymmetric induction *so* far described for that particular Grignard cross coupling reaction. Further examples of such cross couplings are being studied, and results will be published in a full paper.

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