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Asymmetric Codimerization of Cyclohexa=1,3=diene with Ethylene Catalysed by Chiral Aminophosphine-nickel Complexes

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Summary The asymmetric codimerization of cyclohexa-1,3-diene with ethylene is catalysed by the Ni(cycloocta-1,5-diene)₂-AlEt₂Cl-L system where L are chiral aminophosphines; optimization of this catalytic system $(-70 \text{ °C}; \text{ L/Ni} = 10)$ gives $(+)$ - (S) -3-vinylcyclohexene with 73.5% optical purity in 87% chemical yield when $(-)$ - (R) -N-methyl-N-(1-phenylethylamino)diphenylphosphine is used.

IN recent years much interest has been shown in the synthesis of optically active substances catalysed by transition metal complexes containing chiral ligands.^{1,2} In many instances, catalytic processes to form C-H bonds with simultaneous creation of chirality give almost quantitative optical yields.³ For the formation of C-C bonds, however, such reactions have been much less successful. Important reactions of this type are the codimerizations of 1,3-dienes with ethylene in dichloromethane catalysed by Wilke's $(\pi - C_3H_5NiX)_2 - AlX_3 - P^*R^1R^2R^3$ system.⁴ Utilization of $(-)$ -dimenthylphenylphosphine afforded increasing optical yields of $(-)$ - (S) -3-vinylcyclo-octene with decreasing temperatures **(23.5%,** *0* "C; **53%,** *-75* "C) when a **1/2-5/1-2** molar ratio was employed. The highest enantiomeric excess (e.e.) **(70%)** was attained at *0* "C using an Ni/P ratio of **1/3-8,5** in which case, however, the observed chemical yields were low.^{3b}

We have previously shown^{6,7} that active species prepared from the bis(cyclo-octa-1,5-diene)nickel-diethylaluminium chloride-aminophosphine system were able to catalyse the above-mentioned condensation with good rates at low temperatures; we now report that chiral aminophosphines are effective in creating chirality in the catalysed synthesis of pure 3-vinylcyclohexene in almost quantitative yields.

The optically active aminophosphines **(3)** $([\alpha]_p^{20} - 58.2^\circ;$ **c** 1.1, CHCl₃) and **(4)** $([\alpha]_D^{20} - 69.9^\circ; c \ 1.7$, CHCl₃) were synthesized, following a standard procedure, from PPh_2Cl and $(+)$ - (R) -phenylethylamine and the $(+)$ - (R) -N-methyl derivative, respectively.^{8,9} The diaminodiphosphine (5) $([\alpha]_D^{20}$ -21.7°; *c* 1.04, CH_2Cl_2 ; 70 °C) was synthesized

| | | TABLE 1. Effect of the nature of the chiral ligand on the optical yields. ⁸ | | |
|--|---------------------------|--|---|---|
| Ligand | Conversion/ $\frac{9}{6}$ | Selectivity/ $\%$ ^c | $\lceil \alpha_{\rm D} \rceil$ ^d | Optical yield/ $\frac{9}{6}$ ^e |
| | $98 - 5$ | 99 | $22 - 1$ | 13 |
| $\begin{array}{c} (3) \\ (4) \\ (5) \end{array}$ | 99.5 | $98 - 5$ | 45.9 | 27 |
| | 97.51 | 97 | $11-2$ | $6-6$ |

All codimerizations were run in a bomb (300 ml) with **144** mmol of cyclohexa-1,3-diene under an initial ethylene pressure of After hydrolysis, pure hydrocarbon **(2)**

letermined by g.l.c. \circ 99—97% of **(2)**,
 *c*d after dilution in CHCl₃. \circ Calculated **12** atm, 0.36 mmol of Ni(1,5-cod)₂, 0.36 mmol of (4) and 1 ml of AlEt₂Cl for 6 h at 0 °C. was isolated by vacuum distillation. **b** Conversion based on the amount of (1) used was 1-3 % of *cis-* and **trans-3-ethylidenecyclohexenes.** on the basis of the value for the optically pure enantiometric 3-vinylcyclohexene (see text). ^b Conversion based on the amount of (1) used was determined by g.l.c. ^c 99-
clohexenes. ^dAll rotations (in degrees) were measured after dilution in CHCl₃.
Ily pure enantiometric 3-vinylcyclohexene (see text). ¹50

TABLE *2.* Asymmetric codimerizations with **(4).** Effect of temperature."

| T /°C ^b | Conversion/ $\frac{6}{6}$ | Selectivity/ $\%$ | Optical yield/ $\frac{9}{6}$ |
|----------------------|---------------------------|-------------------|------------------------------|
| | $99 - 5$ | 99 | 27 _d |
| -20 | 98 | 99 | 32 |
| -30 | 96.5 | 99 | 34 |
| -40 | 95 | 99 | 36.5 |
| -50 | 92.5 | 97.5 | $40-5$ |
| -70 | 88.5 | 97 | 48 |

^aIdentical experimental conditions to those described in Table **1** (except for temperature). **b If1** "C. *C* Based on **(1). d** At **0** °C, cyclo-octa-1,3-diene was converted into $(+)$ - (R) -3-vinylcyclo-octene by the catalytic system Ni^o-AlEt_aCl- $(+)$ - (4) $(1/4/1)$ with the surprisingly low optical yield of **9%.**

similarly in excellent yields $(80-90\%)$ from the $(-)$ - (R) - $1,2$ -diaminopropane.^{8,9} In contrast with many other chelating phosphines, all enantiomers of the amino phosphines **(3)' (4),** and *(5)* are readily available, so both configurations of 3-vinylcyclohexene are accessible.

Thus, it appears (Table 1) that the asymmetric aminophosphine **(4)** is the most efficient for inducing chirality in the catalysed cyclohexa-1,3-diene-ethylene codimerization and this reaction occurs almost without isomerisation, as expected.6

Tables 2 and 3 show clearly that the optical yields are greatly improved without alteration of the chemical yields by carrying out the catalysis at low temperature or in the presence of an excess of **(4).** This is in contrast With the results of Bogdanovic who observed deactivation of the catalyst when an excess of phosphine was used. 3^b Optimized conditions $(-70 \degree C, P/Ni = 10)$ furnished the best optical yield (73.5%) with the formation of 87% of $(+)$ - (S) -3-vinylcyclohexene $([\alpha]_D^{25} + 125.8^\circ; c \ 3.2$, CHCl₃).

The optical yields were determined in two ways. (a) From application of Brewster's method,¹⁰ which can be used as a more general form of Mill's empirical rules¹¹ for

alkylcyclohexenes; a calculated value of $+170^{\circ}$ for the specific rotation was predicted for $(+)$ - (S) -3-vinylcyclohexene. (b) Stereoselective hydroboration of the most optically active $(+)$ -3-vinylcyclohexene sample $([\alpha]_D^{2\delta})$ $+125.8^{\circ}$; c 3.2, CHCl₃) with di-isopentylborane¹² led to $(+)$ -(S)-3-(2-hydroxyethyl)cyclohexene **(7)** $([\alpha]_D^{25} + 62.8^{\circ};$ c 1.5, CHCl₃). This compound was quantitatively converted into the corresponding diastereomeric mixture of (+ **)-(R)-a-methoxy-a-trifluoromethylphenylacetic** acid esters by Mosher's procedure.¹³ Eu(dpm)₃-induced shift of the methoxy signal¹⁴ permits the determination of enantiomeric purity (73 \pm 1%) by simple integration of MeO peak areas.[†] There is an excellent correlation between the highest optical yields $(74 \text{ vs. } 73\%)$ calculated from these two independent methods.

To prove its *(S)* absolute configuration, **(2)** $([\alpha]_D^{25} + 44^\circ;$ c 1, CHCl₃) was selectively reduced¹⁵ by homogeneous hydrogenation using Vaska's catalyst¹⁶ to $(+)$ -3-ethylcyclohexene $([\alpha]_D^{25} + 12.6^\circ; c \, 1$, CHCl₃) known to have the (R) absolute configuration.¹⁷

a Based on **(1)** ; lower when calculations are based on ethylene (formation of butenes). **b** Remaining 1-13 % made **up** of 3-ethylidenecyclohexenes.

 \dagger This result [73.5% e.e. for $(+)$ -(S)-(2)] was confirmed also because the $[\alpha]_{20}^{25}$ of pure $(+)$ -alcohol (7) was found independently to be + 85.4°. This latter value was determined by initial resolution of cyclohex-2-enylacetic acid (6) by the best resolving agent, brucine, Subsequent reduction of (6) by LiAlH₄ gave $(+)$ - (S) alcohol (7) The use of quinine, phenyl which afforded the $(+)$ - (S) acid (6) $[(\alpha_{B}^{25}]$ + 65° ; *c* 2.66, CHCl₃). $((\alpha)^{26}_{\text{D}}+71.7^{\circ}; c \cdot 1.02, \text{CHCl}_3)$ with 84% e.e. (found by the lanthanoid-induced shift n.m.r. technique). ethylamine, and cinchonine was less successful in our hands.

consists mainly in two points. (i) The most optically chirality on a C_6 ring, are produced catalytically in nearly active aminophosphine ligands are easily prepared and quantitative yields. both enantiomers of 3-vinylcyclohexene are available. (ii) The side-reactions such as olefin isomerisation are limited so that the optically active products, which could *(Received,* **19th** *May* **1980;** *Cow.* **543.)**

We can conclude that the virtue of the present study be used for further synthetic application in producing

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