

化学的不斉増殖法の新展開

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CENTENARY LECTURE Chemical Multiplication of Chirality Science and Applications

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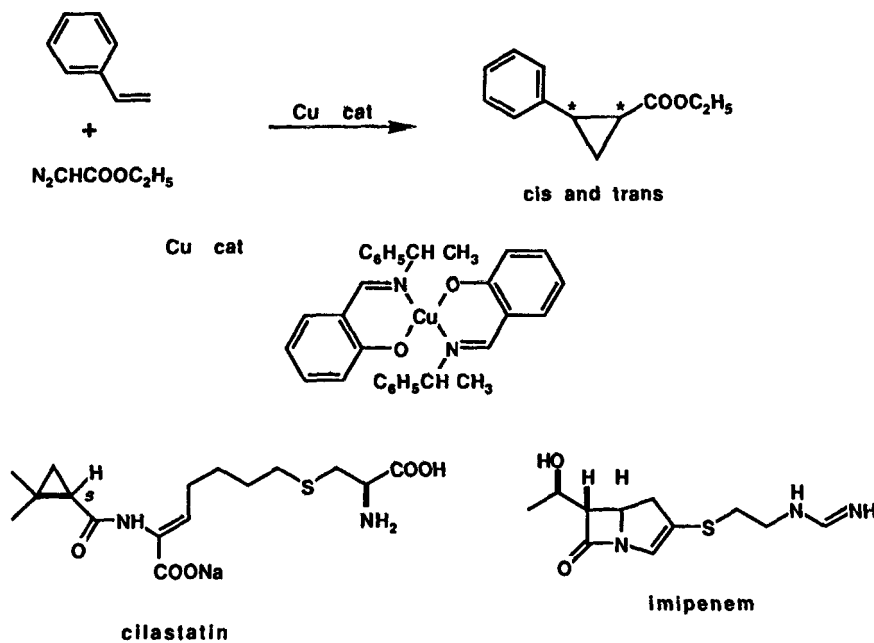
1 Introduction

Chirality plays a central role in science and technology. A wide range of significant physical, chemical, and biological functions are generated through precise molecular recognition which requires strict matching of chirality. For a long time, access to highly enantiomerically pure compounds, at least in a practical sense, was thought to be Nature's monopoly and has indeed been accomplished by biological or biochemical transformations. Efficient creation of optically active organic molecules from prochiral compounds by chemical means, though it is challenging, has remained difficult, and only optical resolution and structural modification of naturally occurring chiral substances have provided complements in this respect. However, assiduous efforts made by synthetic organic chemists in the last two decades are converting the chemists' dream into reality. In order to maximize synthetic efficiency, multiplication of chirality, namely stereoselective production of a large quantity of a chiral target compound utilizing a catalytic amount of chiral source, is obviously desirable. Enantioselective catalysis using chiral metal complexes among various possibilities provides one of the most general flexible methods for this purpose.¹ Metallic elements possess a variety of catalytic activities, and permutation of organic ligands or auxiliaries directing the steric course of the reaction is practically unlimited. Accordingly, in principle, one can generate any dynamic properties at will through molecular architecture using accumulated chemical knowledge. To this end, creation of a single highly reactive catalytic species possessing excellent chiral recognition ability is required. Besides the choice of central metals, therefore, molecular design of the chiral modifiers is a particularly significant task. The efficient ligands must be endowed with a suitable functionality, an appropriate element of symmetry, substituents capable of differentiating space, either sterically or electronically, skeletal rigidity or flexibility (depending on the nature of the reaction), etc.—all of which contribute to accomplish highly enantioselective catalyses.²

¹ For the present state of this subject, see R. Noyori and M. Kitamura, in *Modern Synthetic Methods*, 1989, ed. R. Scheffold, Springer Verlag, Berlin, p. 115.

² R. Noyori and H. Takaya, *Chem. Scr.* 1985, 25, 83.

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Scheme 1

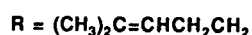
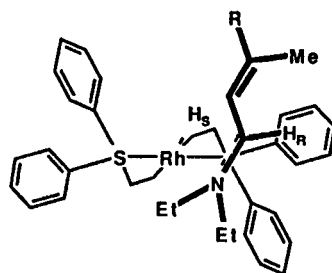
2 Discovery and Opportunities

To our best knowledge the first example of asymmetric synthesis from prochiral compounds *catalysed* by homogeneous chiral metal complexes appeared in the literature in 1966.³ A chiral Schiff base-Cu^{II} complex was found to catalyse decomposition of ethyl diazoacetate in styrene to give *cis* and *trans* 2-phenylcyclopropanecarboxylates in <10% ee proving the existence of a reactive Cu carbenoid placed in a chiral environment. The intermediary carbenoid was also trapped by racemic 2-phenyloxetane leading to optically active furan derivatives. Later extensive systematic screening of the chiral Schiff bases resulted in a dramatic improvement of the optical yield of the cyclopropanation allowing asymmetric synthesis of chrysanthemic acid derivatives in up to 94% ee.⁴ This chemistry has been successfully applied to industrial synthesis of (*S*)-2,2-dimethylcyclopropanecarboxylic acid a component of cilastatin which serves as an excellent inhibitor of dehydropeptidase I increasing *in vivo* stability of antibiotic imipenem (Sumitomo Chemical Co Japan and Merck Sharp & Dohme Co USA) (Scheme 1).

Among other asymmetric catalyses working in industry at this moment perhaps the largest is a process involved in the synthesis of (-)-menthol

³ (a) H. Nozaki, S. Moriuti, H. Takaya, and R. Noyori, *Tetrahedron Lett.* 1966, 5239. (b) H. Nozaki, H. Takaya, S. Moriuti, and R. Noyori, *Tetrahedron* 1968, 24, 3655.

⁴ T. Aratani, *Pure Appl Chem.* 1985, 57, 1839.



(1)

(Takasaga International Co Japan) The key step is the Rh–BINAP⁵ catalysed enantioselective isomerization of diethylgeranylamine to citronellal diethylenamine proceeding in 96–99% optical yield⁶ The optical purity of the synthetic citronellol is much higher than that of the natural product *ca* 80%. The technical refinement has led to an innovative catalytic process working on up to a 7 ton scale Here use of atropisomeric BINAP ligand has played a key role in the successful asymmetric catalysis The fully aromatic diphosphine having an axial element of chirality was first prepared by optical resolution of the racemate through an optically active amine–Pd^{II} complex^{5a,b} but is now obtainable more conveniently by resolution of its dioxide, BINAPO with camphorsulphonic acid or *O* dibenzoyltartaric acid followed by reduction with trichlorosilane⁷ A number of BINAP analogues can be prepared in such a way (Scheme 2)

Olefinic double bonds are known to shift *via* a metal hydride addition–elimination mechanism or a π allylmetal hydride pathway However the allylamine to enamine isomerization was revealed to occur *via* a new nitrogen triggered mechanism (Scheme 3)⁸ The nitrogen coordinated allylamine–Rh⁺ complex causes four centred hydride elimination from C(1) to generate a transient iminium–RhH complex Delivery of the hydrogen from Rh to C(3) gives the enamine η^2 and then η^3 complexes The latter having an aza allyl structure serves as the chain carrier in the catalytic cycle The overall 1,3 hydrogen shift in the geranylamine occurs in a suprafacial manner from its *s trans* type conformer as proved by the deuterium labelling experiments The cationic Rh–BINAP complexes differentiate efficiently between pro *R* and pro *S* hydrogens at C(1) through interaction with the adjacent nitrogen atom (Scheme 4) A transition

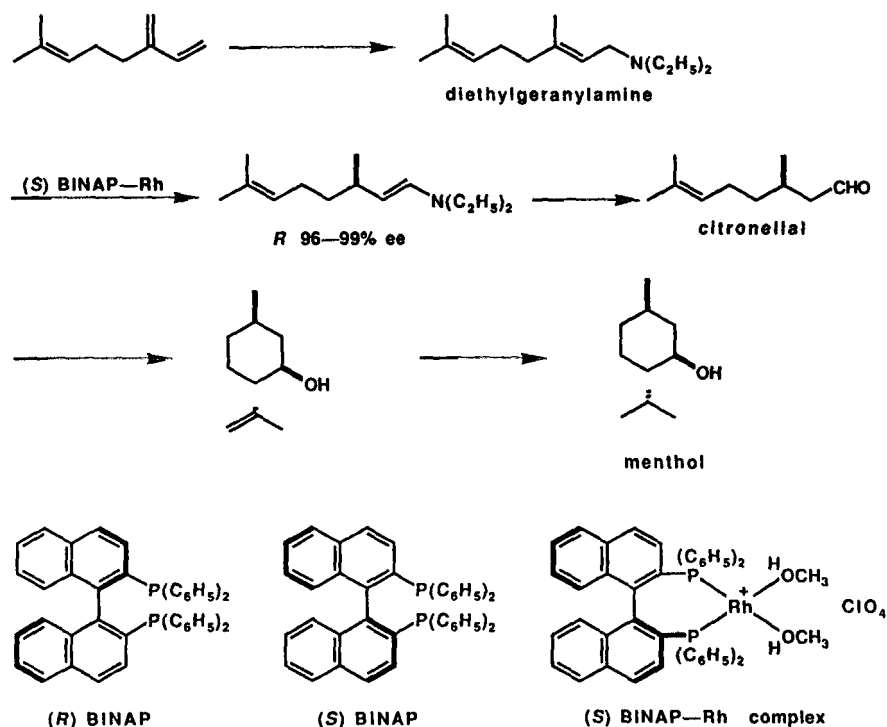
⁵ (a) A Miyashita, A Yasuda, H Takaya, K Toriumi, T Ito, T Souchi and R Noyori *J Am Chem Soc* 1980 **102** 7932 (b) A Miyashita, H Takaya, T Souchi and R Noyori *Tetrahedron* 1984 **40** 1245 (c) K Toriumi, T Ito, H Takaya, T Souchi and R Noyori *Acta Crystallogr Sect B* 1982 **38** 807 (d) S Inoue, M Osada, K Koyano, H Takaya and R Noyori *Chem Lett* 1985 1007

⁶ (a) K Tani, T Yamagata, S Otsuka, S Akutagawa, H Kumobayashi, T Taketomi, H Takaya, A Miyashita and R Noyori *J Chem Soc Chem Commun* 1982 600 (b) K Tani, T Yamagata, S Akutagawa, H Kumobayashi, T Taketomi, H Takaya, A Miyashita, R Noyori and S Otsuka *J Am Chem Soc* 1984 **106** 5208

⁷ (a) H Takaya, K Mashima, K Koyano, M Yagi, H Kumobayashi, T Taketomi, S Akutagawa and R Noyori *J Org Chem* 1986 **51** 629 (b) H Takaya, S Akutagawa and R Noyori *Org Synth* 1988 **67** 20

⁸ H Takaya, K Tani, S Otsuka, S Inoue, T Sato and R Noyori to be published

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Scheme 2

state model in the Rh-(S) BINAP catalysed reaction is illustrated by structure (1) (naphthalene rings are omitted for clarity)

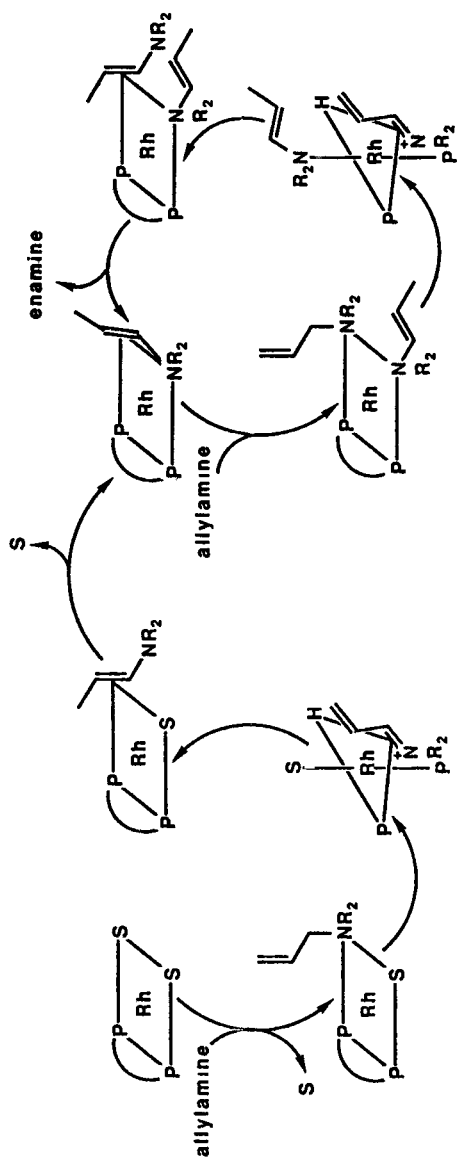
In principle any donor groups including olefinic bond heteroatom bases carbanions heteroanions etc are able to activate their adjacent C-H bonds through coordination to appropriate unsaturated transition metal centres. The resulting metal hydride complexes depending on the situations undergo unique chemical transformations. When racemic 4-hydroxy-2-cyclopentenone was exposed to 0.5 mol% of a cationic Rh-(R) BINAP complex⁹ in THF at 0 °C double bond isomerization occurred with 5:1 enantiomer discrimination to afford unreacted (R) hydroxy enone in 91% ee in 27% yield and 1,3-cyclopentanedione in 61% yield (Scheme 5)⁹

3 Ruthenium-catalysed Asymmetric Hydrogenation

Homogeneous asymmetric hydrogenation discovered in 1968¹⁰ has been one of the most exciting subjects in organic chemistry in the last two decades and a

⁹ M Kitamura, K Manabe, R Noyori and H Takaya *Tetrahedron Lett* 1987 28 4719

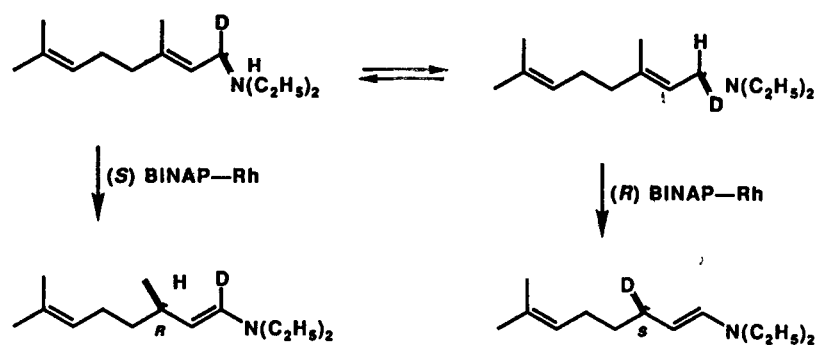
¹⁰ (a) L Horner, H Siegel and H Buthe *Angew Chem Int Ed Engl* 1968 7 942 (b) W S Knowles and M Sabacky *J Chem Soc Chem Commun* 1968 1445



P-P = BINAP
S = donor solvent

Scheme 3

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number of impressive chemistries have been presented¹¹ In addition the catalysis is of practical significance (*S*) DOPA a drug for the treatment of Parkinson's disease has been prepared at Monsanto Co USA¹² and VEB Isis Chemie DDR by using hydrogenation of a (*Z*) (α acetamido)cinnamic ester with soluble Rh complex catalysts possessing a chiral phosphine or phosphinite ligand The same method was used for commercial production of (*S*) phenylalanine a component of the non nutritive sweetener Aspartame (Amic S p A Italy)¹³ Thus a variety of natural and unnatural amino acids are now available in >90% e.e. by enantioselective hydrogenation but unfortunately the scope of the Rh catalysed reaction is not very wide For example [Rh(binap)(CH₃OH)₂]ClO₄ caused hydrogenation of dehydroamino acid derivatives (Scheme 6) with nearly perfect enantioselectivities^{5a,b} whereas optical yields of the reactions of geraniol or nerol with varying conditions did not exceed 70%.^{5d} In view of the general importance of hydrogenation in organic synthesis we have been intrigued by the possibility of developing a catalyst system capable of adopting a wide range of olefinic substrates In this context recent invention of Ru-BINAP dicarboxylate complexes¹⁴ extended the utility of asymmetric hydrogenation to a great extent (Figure 1)

The Ru dicarboxylate complexes undergo ligand exchange reaction with $\alpha\beta$ or $\beta\gamma$ unsaturated carboxylic acids resulting in highly enantioselective (80–100%) hydrogenation¹⁵ Thus with many substrates the highest enantioselectivities

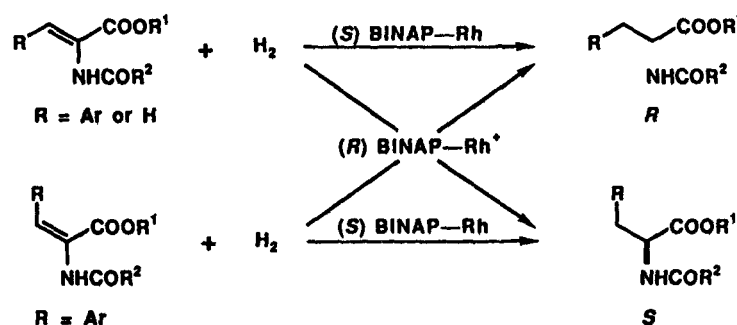
¹¹ Pertinent reviews (a) J Halpern in *Asymmetric Synthesis* Vol 5 ed J D Morrison Academic Press New York 1985 Chapter 2 (b) K E Koenig *ibid* Chapter 3

¹² W S Knowles *J Chem Educ* 1986 63 222

¹³ H B Kagan *Bull Chem Soc Fr* 1988 846

¹⁴ T Ohta, H Takaya, and R Noyori *Inorg Chem* 1988 27 566

¹⁵ T Ohta, H Takaya, M Kitamura, K Nagai, and R Noyori *J Org Chem* 1987 52, 3174



Scheme 6

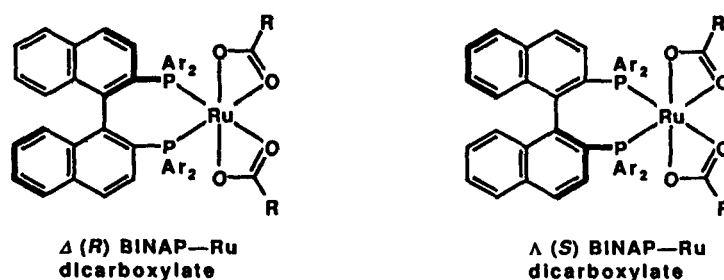


Figure 1

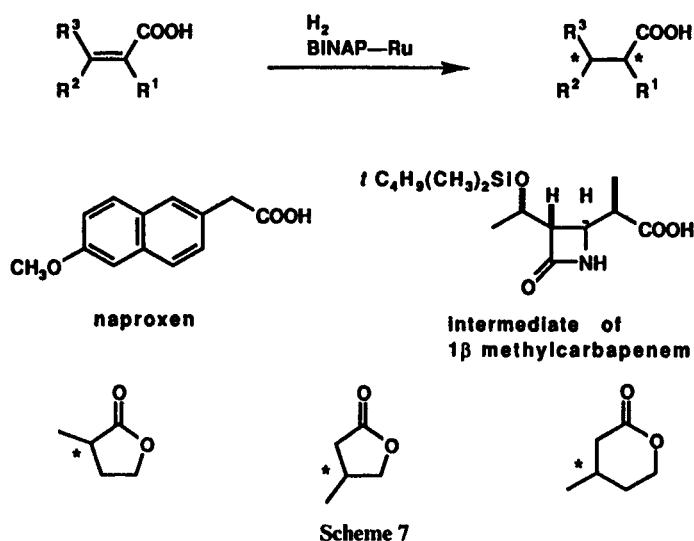
have been recorded Methyl esters are inert to the hydrogenation Alcohols are the solvents of choice The sense and extent of the asymmetric induction are highly dependent on the substitution pattern of the substrates and reaction conditions particularly the hydrogen pressure Anti inflammatory (*S*) naproxen was prepared in 97% ee under a high pressure condition This method is also applicable to synthesis of a 1 β methylcarbapenem precursor and some optically active methylated γ and δ lactones (Scheme 7)

Olefins containing certain neutral donor functionalities are also hydrogenated in a satisfactory manner¹⁶ The Ru-BINAP catalysed hydrogenation of *N* acyl (*Z*) 1 benzylidene 1,2,3,4 tetrahydroisoquinolines in a mixture of ethanol and dichloromethane leads consistently to (*1R*) or (*1S*) benzyltetrahydroisoquinolines in nearly quantitative yield and in 95–100% ee¹⁷ With Rh complexes such as [Rh(binap)(cod)]ClO₄ or [Rh(binap)(CH₃OH)₂]ClO₄ the hydrogenation proceeded in lower optical yield (*ca* 75%) and with opposite enantioselection The asymmetric hydrogenation followed by removal or modification of the *N* acyl groups gave tetrahydropapaverine laudanosine norreticuline (biogenetic precursor of morphine) tretoquinol (bronchodilating agent) *etc*

¹⁶ For a review on stereoselective olefin hydrogenation directed by functional groups see J M Brown *Angew Chem Int Ed Engl* 1987 26 190

¹⁷ R Noyori M Ohta Y Hsiao M Kitamura T Ohta and H Takaya *J Am Chem Soc* 1986 108 7117

Centenary Lecture



which became homochiral by single recrystallization. The reaction of the simple 1-methylene substrate affords, after deacylation, *salsolidine* in 96% ee. This procedure is applicable to the synthesis of natural morphine, various benzo-morphans, analogues such as *metazocine* and *pentazocine*, morphinans including *dextromethorphan* (antitussive agent) *etc.* (Figure 2).¹⁸ This discovery has thus realized a general asymmetric synthesis of isoquinoline alkaloids.¹⁹

The Ru-catalysed hydrogenation of prochiral allylic alcohols exhibits unprecedented efficacy. Thus *geraniol* and *nerol* are hydrogenated in methanol containing a Ru-BINAP dicarboxylate complex to give (*S*) or (*R*) *citronellol* in 96–99% ee.²⁰ Initial hydrogen pressure higher than 30 atm gave satisfactory results. Either natural or unnatural forms can be made flexibly by changing the chirality of the catalyst or geometry of the olefinic substrates. The enantiomeric purity of the synthetic *citronellol* exceeds the highest value of the natural product (92%). The substrate/catalyst mole ratio is extremely high and in certain cases the efficiency of the chiral multiplication, defined as [major enantiomer – minor enantiomer] (in mole)/chiral source (in mole), approaches 48 500! Notably, in this hydrogenation, only allylic C(2)–C(3) double bonds are saturated and nonallylic C(6)–C(7) double bonds remain intact. *Homogeraniol* was hydrogenated in 92% optical yield with the same enantioselection, but the bis-homologue was inert to the standard reaction conditions. This hydrogenation is usable for the stereoselective synthesis of side chains of *vitamin E* and *K₁* (Scheme 8).

¹⁸ M. Kitamura, Y. Hsiao, R. Noyori and H. Takaya, *Tetrahedron Lett.* 1987, **28**, 4829.

¹⁹ For synthesis *via* stoichiometric enantioselective alkylation, see A. I. Meyers, *Aldrichimica Acta* 1985, **18**, 59.

²⁰ H. Takaya, T. Ohta, N. Sayo, H. Kumobayashi, S. Akutagawa, S. Inoue, I. Kasahara and R. Noyori, *J. Am. Chem. Soc.* 1987, **109**, 1596–4129.

Noyori

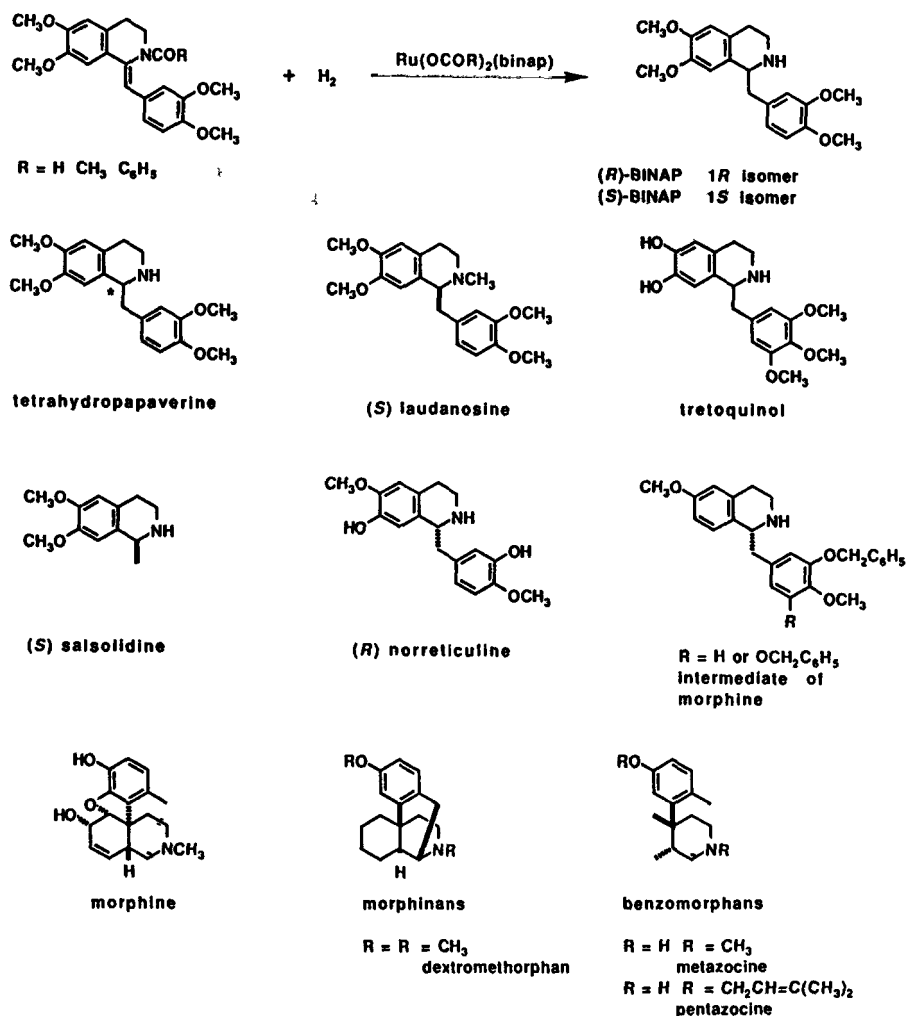
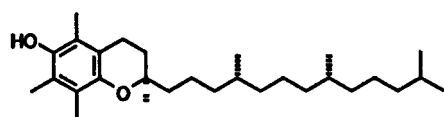
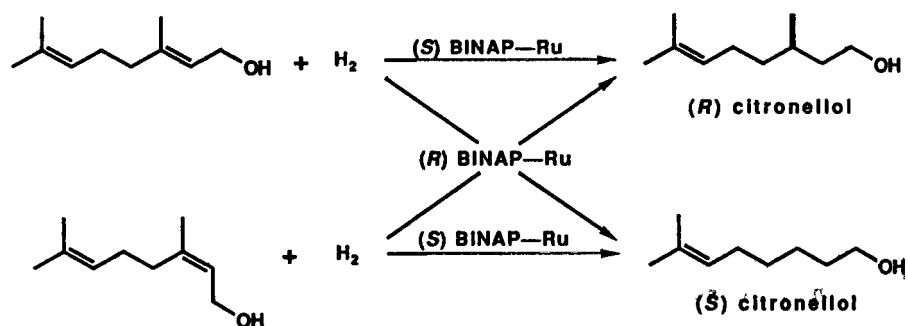


Figure 2

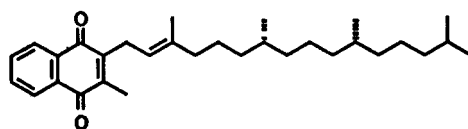
Chiral allylic secondary alcohols can be resolved efficiently by homogeneous hydrogenation catalysed by the Ru-BINAP diacetate complexes.²¹ The combined effects of intramolecular and intermolecular asymmetric induction give up to 76:1 differentiation between the enantiomeric cyclic unsaturated alcohols. For instance when racemic 3-methyl-2-cyclohexenol (Figure 3) was hydrogenated with the Ru-(R)-BINAP complex in methanol at 46% conversion, R,R-configured *trans*-3-methylcyclohexenol was obtained in 95% ee. At 54% conversion the slow reacting S

²¹ M. Kitamura, I. Kasahara, K. Manabe, R. Noyori, and H. Takaya, *J. Org. Chem.* 1988, 53, 708.

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vitamin E



vitamin K₁

Scheme 8



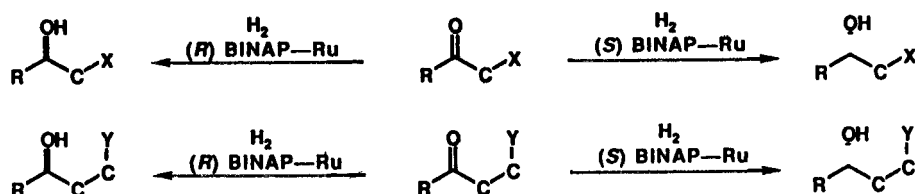
Figure 3

enantiomer was recovered in >99% e.e. A significant application includes a practical resolution of 4-hydroxy-2-cyclopentenone (Figure 3) an important building block for the three-component coupling prostaglandin synthesis.²²

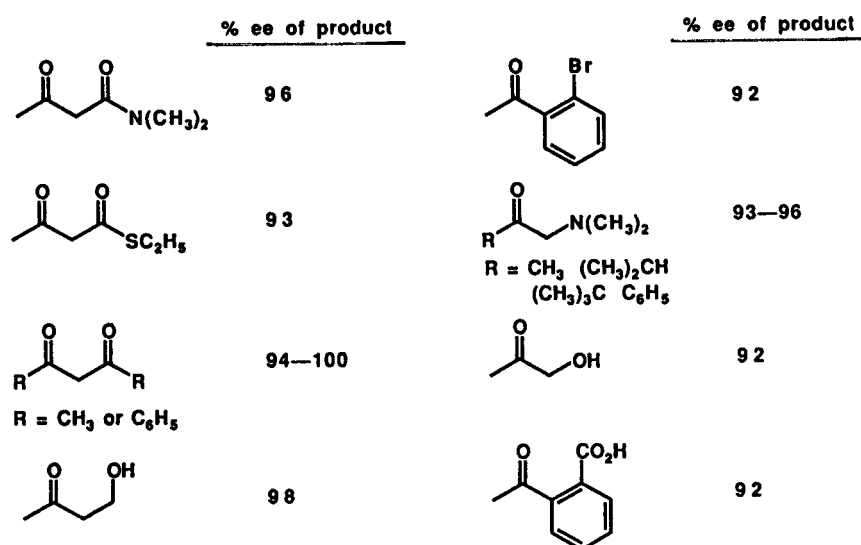
Homogeneous asymmetric hydrogenation of ketones has remained far less fruitful than the catalysis of olefinic substrates. Now, however, a variety of functionalized ketones can be hydrogenated with synthetically useful enantioselectivities and in a predictable manner with the aid of $RuX_2(\text{binap})$ [empirical formula $X = Cl, Br, I$] prepared by mixing $Ru(\text{OCOCH}_3)_2(\text{binap})$ and HX in a

²² (a) R. Noyori and M. Suzuki, *Angew Chem Int Ed Engl* 1984, 23, 847; (b) M. Suzuki, A. Yanagisawa and R. Noyori, *J Am Chem Soc* 1988, 110, 4718; (c) Y. Motita, M. Suzuki and R. Noyori, *J Org Chem* 1989, 54, 1785.

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X Y = heteroatom
C = sp^2 or nonstereogenic sp^3 carbon



Scheme 9

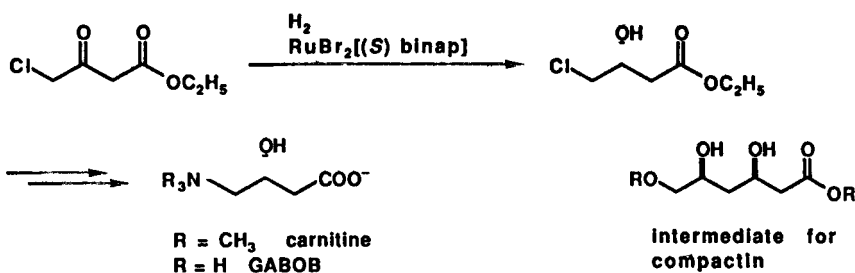
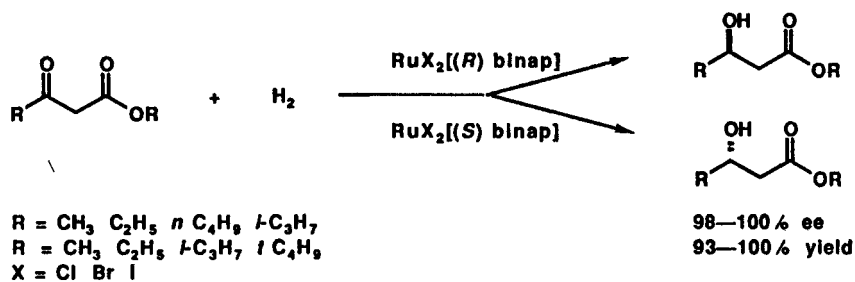
1:1 mole ratio]^{23, 24} or $Ru_2Cl_4(binap)_2(C_2H_5)_3N$ ²⁵ The general sense of the asymmetric induction indicates that the key factor in the enantioface differentiation is the simultaneous coordination of the carbonyl oxygen and heteroatom X or Y to the Ru atom forming a five- and six-membered chelate ring respectively. Some nitrogen- and oxygen-containing directive groups include dialkylamino, hydroxyl, silyloxy, keto, alkoxycarbonyl, alkylthiocarbonyl, dialkylaminocarbonyl, carboxyl, etc.²² To our surprise, halogen atoms were revealed to facilitate the carbonyl hydrogenation and to direct the stereochemical outcome. Thus, *o*-bromoacetophenone gave the corresponding alcohol in 92% ee and 97%

²³ R. Noyori, T. Ohkuma, M. Kitamura, H. Takaya, N. Sayo, H. Kumobayashi, and S. Akutagawa, *J. Am. Chem. Soc.* 1987, **109**, 5856.

²⁴ M. Kitamura, T. Ohkuma, S. Inoue, N. Sayo, H. Kumobayashi, S. Akutagawa, T. Ohta, H. Takaya, and R. Noyori, *J. Am. Chem. Soc.* 1988, **110**, 629.

²⁵ T. Ikariya, Y. Ishii, H. Kawano, T. Arai, M. Saburi, S. Yoshikawa, and S. Akutagawa, *J. Chem. Soc. Chem. Commun.* 1985, 922.

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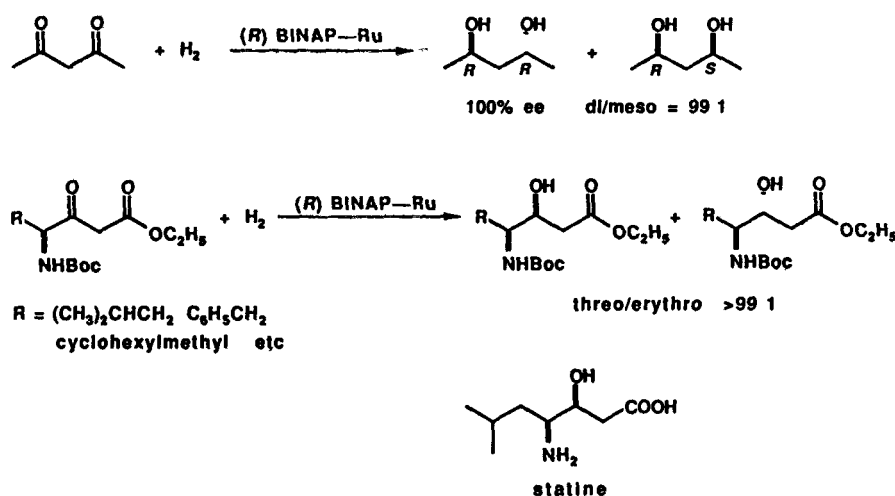
Scheme 10

yield although unsubstituted acetophenone and the *m* or *p* bromo derivative failed to be hydrogenated in a satisfactory manner (Scheme 9)

This method is particularly useful for enantioselective access to β hydroxy carboxylic esters which serve as important intermediates for natural product synthesis. A wide variety of prochiral β keto esters having flexible structures are hydrogenated consistently in nearly quantitative yields and with extremely high (up to 100%) enantioselectivities.²³ Esters of methyl, primary, secondary and tertiary alcohols as well as α alkylated and $\alpha\alpha$ dialkylated substrates were equally employable. Thus synthetic organic chemists no longer need envy bakers yeast in this context (Scheme 10). This procedure allowed the first efficient chemical synthesis of GABOB and (*R*) carnitine, a carrier of long chain fatty acids through the mitochondrial membrane.²⁶ Hydrogenation of ethyl 4-chloro-3-oxobutanoate aided by the (*S*) BINAP catalyst under the conditions effecting the reaction of 3-oxobutanoate in 99.4% optical yield (ethanol, room temperature, 100 atm, 10–40 h) afforded the desired (*R*) hydroxy-chloro ester in only <70% ee. The inefficient enantiofacial differentiation is perhaps due to the competitive directing effect of the ester group and halogen atom present in the same molecule. However, a surprising chiral efficiency was obtained by the high temperature short period reaction (100 °C, <5 min) affording the *R* enantiomer in 97% ee in 97% chemical yield. The same technique has been used for the synthesis of a component of compactin, an HMG CoA reductase inhibitor (Scheme 10).

Double stereodifferentiation is a powerful mechanism to enhance a degree of

²⁶ M. Kitamura, T. Ohkuma, H. Takaya, and R. Noyori, *Tetrahedron Lett.* 1988, 29, 1555.



Scheme 11

stereoselection²⁷ When prochiral symmetrical β diketones were subjected to the Ru catalysed hydrogenation mixtures of *dl* and *meso* 1,3 diols were formed (Scheme 11). The *dl* isomers were dominant and their ees were uniformly high. For instance, the reaction of 2,4-pentanedione catalysed by the (*R*)-BINAP catalyst proceeded by way of (*R,R*)-4-hydroxy-2-pentanone in 98.5% ee, but the ultimate product was a 99:1 mixture of (*R,R*)-2,4-pentanediol in nearly 100% ee and (*R,S*)-2,4-pentanediol. The minor (*S*)-hydroxy ketone intermediate was washed away by intramolecular 1,3-chirality transfer, giving the *meso* diol, and the calculated *R/R,S,S* ratio in the *dl* type diol was ca. 900:1.²⁴ Diastereoselective hydrogenation of *N*-protected γ -amino β -keto esters catalysed by the (*R*)-BINAP catalyst provides an efficient entry to statine, a key component of the aspartic proteinase inhibitor pepstatin.²⁸ The efficiency of the catalyst to substrate chirality transfer (catalyst control >33:1) and the intramolecular 1,2-asymmetric induction (substrate control 3:1) cooperate to form the natural *threo* series in >99:1 diastereoselectivity. A number of statine analogues are obtainable by this method using double asymmetric induction (Scheme 11).

Thus the present Ru catalysed hydrogenation exhibits wider scope than reactions with any other chiral transition metal complexes so far designed. A range of optically active compounds of either chirality sense are now accessible, providing a versatile tool in stereoselective organic synthesis. This homogeneous hydrogenation procedure is superior to the heterogeneous version and compares well with the biochemical transformations whose yields and enantioselectivities are often variable. The hydrogenation method is clean, operationally simple, economical.

²⁷ S. Masamune, W. Choy, J. S. Petersen, and L. R. Sita, *Angew. Chem. Int. Ed. Engl.* 1985, 24, 1.

²⁸ T. Nishi, M. Kitamura, T. Ohkuma, and R. Noyori, *Tetrahedron Lett.* 1988, 29, 6327.

Centenary Lecture

and hence is capable of conducting a large scale reaction using high (up to 50%) substrate concentration^{29 30}

Now one may raise questions (1) What is the major difference between the Ru chemistry and well studied Rh catalysed hydrogenation? (2) Why does BINAP ligand work so effectively? The mechanism of the Ru-BINAP catalysed reaction remains to be elucidated. However d^6 Ru^{II} chemistry differs from d^8 Rh^I chemistry distinctly. First Ru^{II} complexes utilize higher coordination numbers up to six in an octahedral structure than Rh^I complexes which normally have a square planar geometry. Second reaction of a Ru^{II} complex with hydrogen generates Ru monohydride species³¹. In contrast to the Rh promoted reaction occurring by way of the metal dihydride intermediate¹¹. Such characteristics would reflect on the marked difference in reactivity-selectivity profiles in the hydrogenation. In the BINAP chemistry the degeneracy caused by C_2 chirality of the diphosphine minimizes the number of the diastereomeric reactive intermediates and transition states. Flexible atropisomeric skeletal backbone of BINAP can produce a conformationally unambiguous metal chelate ring without concomitant increase of strain energy². In addition phenyl rings attached to the phosphorus atoms can suitably modulate stabilities of the intermediary complexes and transition states. Molecular structure of Ru-(*S*) BINAP dipivalate complex determined by single crystal X ray analysis is given in Figure 4¹⁴. The whole structure approximates C_2 chirality. The dissymmetry of (*S*) BINAP fixes the delta conformation of the seven membered chelate ring containing the diphosphine and Ru. This cyclic structure is highly skewed and this geometry in turn determines the chiral disposition of the phenyl rings on the phosphorus atoms. Two phenyl substituents are oriented in axial directions and the others in equatorial directions. These equatorial phenyl rings exert profound steric influence on the equatorial coordination sites of Ru. Consequently the bidentate ligation of the pivalate moieties to Ru occurs stereoselectively leading to exclusive formation of the Λ diastereomer. This diastereomeric differentiation of the two sets of quadrant space sectors is made in such a way as to avoid nonbonded interactions between the sterically demanding equatorial phenyl substituents and the carboxylate ligands. This is merely a ground state structure of a catalyst precursor but whatever the detailed reaction mechanism is such an argument should also be applicable to the transition state or intermediates. Actual chemical transformations take place at the oxygen coordinated sites and we believe that this is the steric origin of the high level of enantioselection. Stability of the transition structure (1) in the Rh-BINAP chemistry is also understandable in such a way⁸.

²⁹ R. Noyori *Chimia* 1988 42 215

³⁰ For related work see (a) ref 25 (b) H. Kawano, Y. Ishii, T. Ikariya, M. Saburi, S. Yoshikawa, Y. Uchida and H. Kumobayashi *Tetrahedron Lett* 1987 28 1905 (c) T. Tsukahara, H. Kawano, Y. Ishii, T. Takahashi, M. Saburi, Y. Uchida and S. Akutagawa *Chem Lett* 1988 2055 (d) H. Kawano, Y. Ishii, M. Saburi and Y. Uchida *J Chem Soc Chem Commun* 1988 87

³¹ D. Evans, J. Osborn, J. A. Jardin and G. Wilkinson *Nature* 1965 208 1203

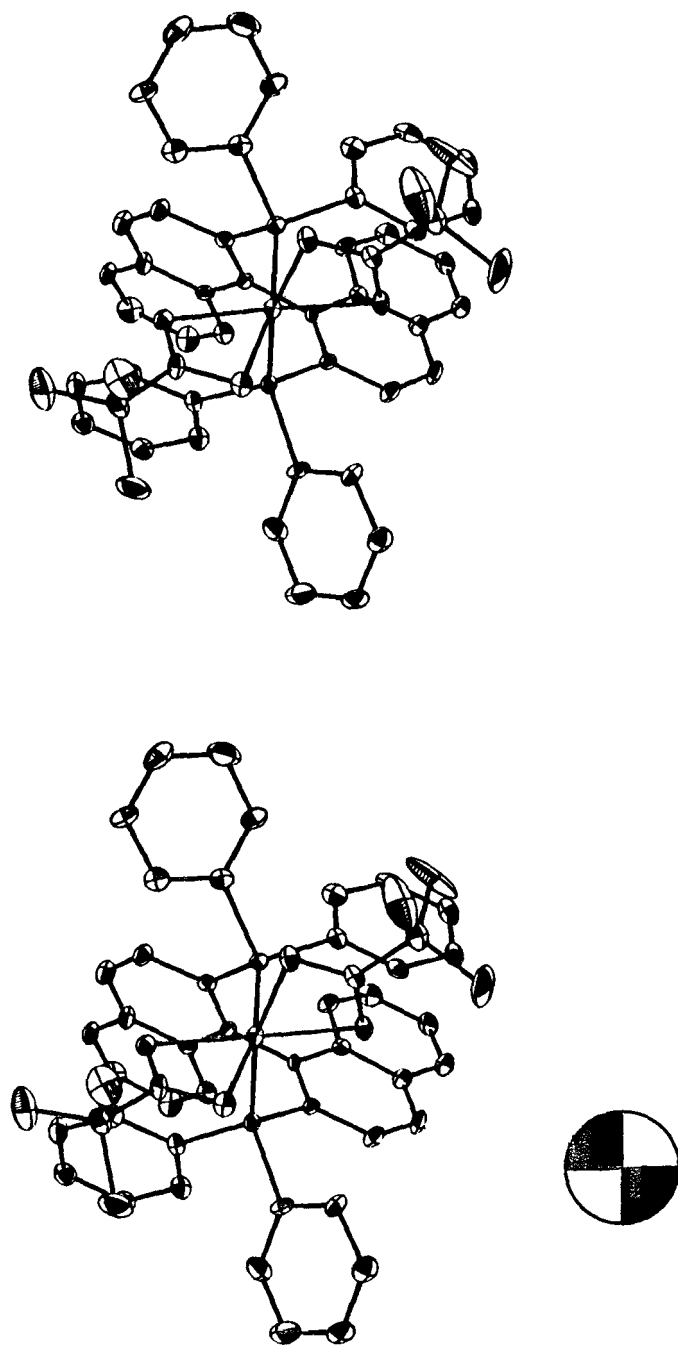


Figure 4 ORTEP drawings of Λ Ru[(S) binap](OCO t C₆H₉)₂

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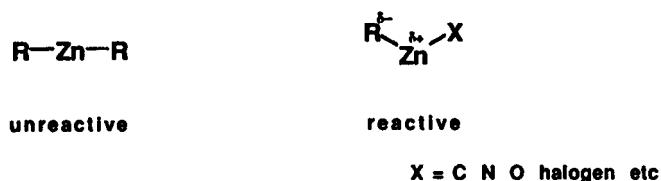


Figure 5

4 Asymmetric Alkylation of Carbonyl Compounds

Enantioselective alkylation of aldehydes by organometallic reagents is a fundamental problem in organic synthesis. Although there have been reports of several successful examples of this type of reaction^{32,33} a high degree of enantioselection is achievable by using a stoichiometric or even excess amount of chiral auxiliary. Certain ligands may accelerate the nucleophilic alkylation but the difference in rates of the catalysed and uncatalysed reactions is not large enough to lead to a practical asymmetric catalysis.^{32b} In this context dialkylzincs, the oldest organometallic compounds, generate a variety of new unprecedented chemistries opening a novel domain of asymmetric catalysis. Monomeric dialkylzincs having a linear geometry are inert to carbonyl compounds but the structural modification by appropriate ligands or auxiliaries forming a coordinatively unsaturated bent structure increases the acceptor character of the Zn atom and donor property of the alkyl group thereby increasing the reactivity toward carbonyl substrates (Figure 5). Here some chirally well designed auxiliary should also direct the stereochemical outcome in an absolute sense as well. Thus in the presence of a catalytic amount of (–) 3 *exo* (dimethylamino)isoborneol (DAIB) reaction of dialkylzincs and benzaldehyde in nonpolar solvents is accelerated markedly to give after hydrolysis the corresponding *S* alcohols in high (up to 99%) enantiomeric purity (Scheme 12).^{34,35} Various *p* substituted benzaldehydes and certain α, β unsaturated and aliphatic aldehydes can also be alkylated with a high level of enantioselectivity. Dimethyl, diethyl, and di-*n*-butylzinc are employable as alkylating agents.

The catalytic cycle is illustrated in the scheme where the DAIB structure is simplified.³⁶ Reaction of (–) DAIB and dialkylzinc in a 1:1 molar ratio

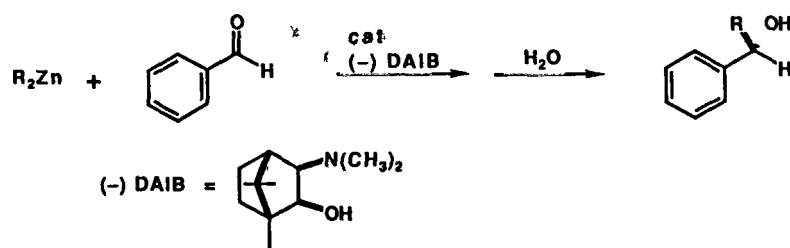
³² (a) G. Soladie in *Asymmetric Synthesis*, Vol. 2A, ed. J. D. Morrison, Academic Press, New York, 1983, Chapter 6; (b) J. P. Mazaleyrat and D. J. Cram, *J. Am. Chem. Soc.* 1981, **103**, 4585; (c) T. Mukaiyama, K. Soai, T. Sato, H. Shimizu, and K. Suzuki, *J. Am. Chem. Soc.* 1979, **101**, 1455; (d) D. Seebach, A. K. Beck, S. Roggo, and A. Wonnacott, *Chem. Ber.* 1985, **118**, 3673; (e) M. T. Reetz, T. Kukenhohner, and P. Weing, *Tetrahedron Lett.* 1986, **27**, 5711.

³³ R. Noyori, S. Suga, K. Kawai, S. Okada, and M. Kitamura, *Pure Appl. Chem.* 1988, **60**, 1597.

³⁴ M. Kitamura, S. Suga, K. Kawai, and R. Noyori, *J. Am. Chem. Soc.* 1986, **108**, 6071.

³⁵ Related works: (a) N. Oguni and T. Omi, *Tetrahedron Lett.* 1984, **25**, 2823; (b) A. A. Smaardijk and H. Wynberg, *J. Org. Chem.* 1987, **52**, 135; (c) K. Soai, A. Ookawa, K. Ogawa, and T. Kaba, *J. Chem. Soc. Chem. Commun.* 1987, 467; (d) S. Itsuno and J. M. J. Frechet, *J. Org. Chem.* 1987, **52**, 4140; (e) P. A. Chaloner and S. A. R. Perera, *Tetrahedron Lett.* 1987, **28**, 3013; (f) E. J. Corey and F. J. Hannon, *Tetrahedron Lett.* 1987, **28**, 5233, 5237; (g) K. Soai, A. Ookawa, T. Kaba, and K. Ogawa, *J. Am. Chem. Soc.* 1987, **109**, 7111; (h) W. Oppolzer and R. N. Radinov, *Tetrahedron Lett.* 1988, **29**, 5645. See also D. A. Evans, *Science* 1988, **240**, 420.

³⁶ M. Kitamura, S. Okada, S. Suga, and R. Noyori, *J. Am. Chem. Soc.* in press.



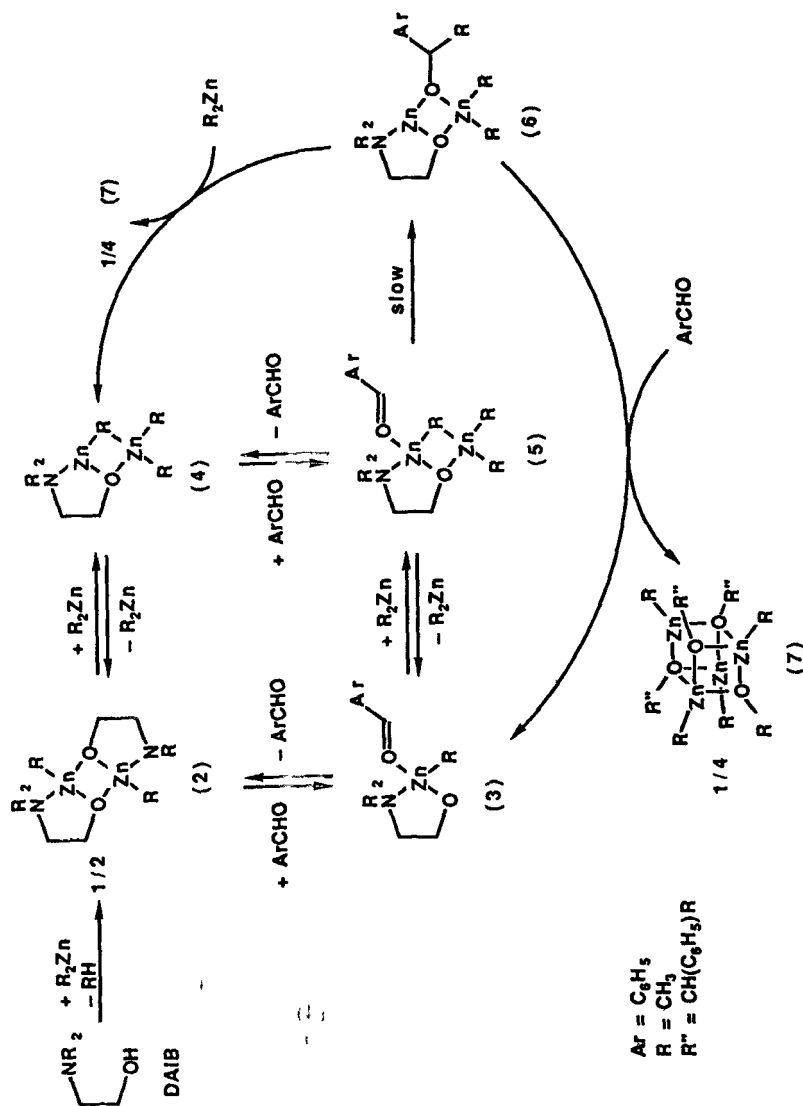
Scheme 12

produces a single Zn chelate complex (2) which does not alkylate benzaldehyde but acts as catalyst precursor. Significantly the alkylation proceeds *via* a dinuclear Zn species (5) containing DAIB auxiliary aldehyde ligand and three alkyl groups. The resulting bridged alkoxide (6) upon exposure to benzaldehyde or dialkylzinc undergoes instantaneous decomposition to the stable cubic tetramer (7) regenerating (3) and (4) respectively (Scheme 13). Under the catalysis conditions complexes (2)–(5) are equilibrating on a soft energy surface consistent with the fact that when two different dialkylzincs are used a statistical distribution of the possible products is obtained regardless of the order or way of mixing the two alkylzincs. Here only relative reactivity of alkyl groups is important. Kinetic measurements and temperature effects on the enantioselectivity indicate that the alkyl transfer process (5) \rightarrow (6) is the turnover limiting and stereo determining step.

The DAIB aided enantioselective alkylation exhibits enormous nonlinearity in terms of optical purity of the chiral source and alkylation products.^{33, 36–38} Typically when benzaldehyde and diethylzinc are reacted in the presence of 8 mol % of (–) DAIB in 15% ee in toluene (*S*) 1-phenylpropyl alcohol is produced with 95% ee, a value close to 98% obtained using enantiomerically pure DAIB. The nonlinear effect is clear in Figure 6 which shows the ees of (*S*) products as a function of the ee of (–) DAIB. Under certain conditions turnover efficiency of the chiral catalyst system is >600 times greater than that of the coexisting achiral catalyst system. This unusual phenomenon is a result of a marked difference in chemical properties of the diastereomeric dinuclear complexes formed from dialkylzincs and DAIB. Reaction of equimolar amounts of dimethylzinc and enantiomerically pure (–) DAIB affords a dinuclear chelate complex with C_2 chirality which dissociates readily to catalytically active monomeric species. By contrast dimethylzinc and racemic DAIB generate a more stable but much less reactive dinuclear complex possessing *meso* C structure rather than a racemic mixture of the chiral complexes. Molecular structures of these complexes determined by single crystal X-ray analyses are illustrated in Figure 7.³⁶

³⁷ N. Oguni, Y. Matsuda, and T. Kaneko, *J. Am. Chem. Soc.* 1988, 110, 1877.

³⁸ (a) C. Puchot, O. Samuel, E. Dunach, S. Zhao, C. Agami, and H. B. Kagan, *J. Am. Chem. Soc.* 1986, 108, 2353; (b) C. Agami, J. Levisalles, and C. Puchot, *J. Chem. Soc., Chem. Commun.* 1985, 441.



Scheme 13

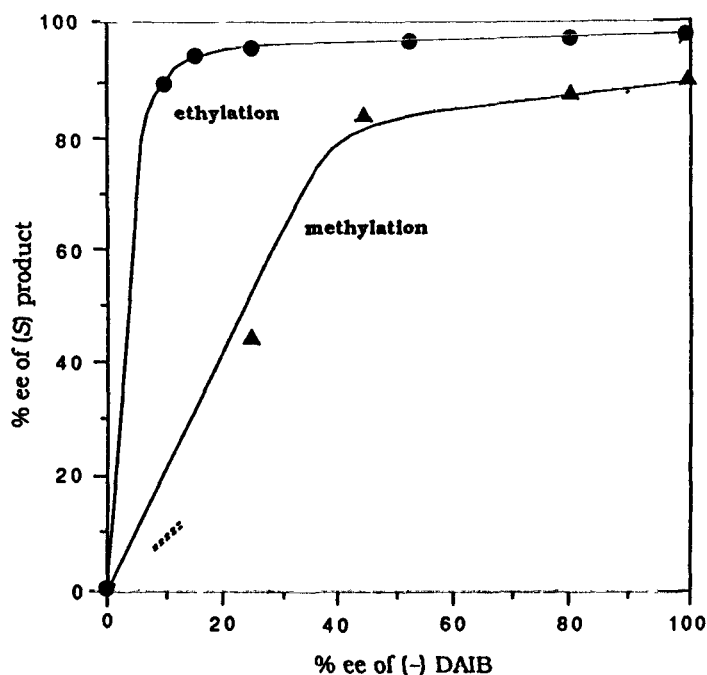


Figure 6 Correlation between the *ee* of the alkylation product and the *ee* of the chiral auxiliary ● Reaction using 0.42 M $(C_2H_5)_2Zn$, 0.42 M C_6H_5CHO and 34 mM (-) DAIB in toluene at 0 °C ▲ 0.47 M $(CH_3)_2Zn$, 0.49 M C_6H_5CHO , 47 mM (-) DAIB in toluene d_8 at 32 °C

Alkyl transfer from the mixed ligand complex (5) is conceived to occur *via* a folded bicyclic transition state (8) featuring a tricoordinate structure of the migrating R group. The kinetic bias leading to the *S* configured alkoxide derives primarily from a nonbonded repulsion between the carbonyl substrate (Ar and H) and a terminal R group attached to Zn_B atom.

Organometallic chemistry of homo- or hetero multinuclear compounds is increasing the synthetic importance and the nonclassical dinuclear mechanism which has been theoretically advanced³⁹ can provide reasonable explanations for various stereoselective reactions. In order to create a single reactive species we designed binaphthol modified Li/Mg binary organometallic reagents having empirical formula of (9) and found that they undergo stoichiometric enantioselective alkylation with aldehydes (Scheme 14) to give the corresponding secondary alcohols in high *ee*s.³³ For example, 1-phenylpropyl alcohol was produced in up to 92% *ee*. The possible transition state resulting in the *S/S* auxiliary/alcohol asymmetric induction is illustrated by the structure (10) (*S* = solvent, naphthalene rings are omitted in (10a)). Chiral reducing agent BINAL.

³⁹ A planar bicyclic transition state has been proposed for reaction of methyllithium dimer and formaldehyde. E. Kaufmann, P. von R. Schleyer, K. N. Houk, and Y. D. Wu, *J. Am. Chem. Soc.* 1985, **107**, 5560.

Centenary Lecture

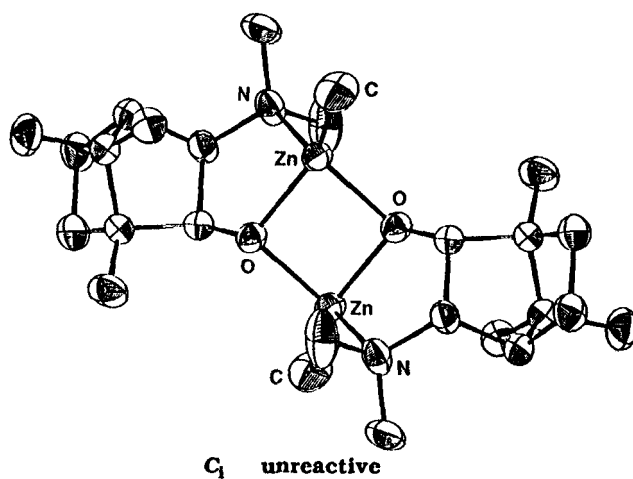
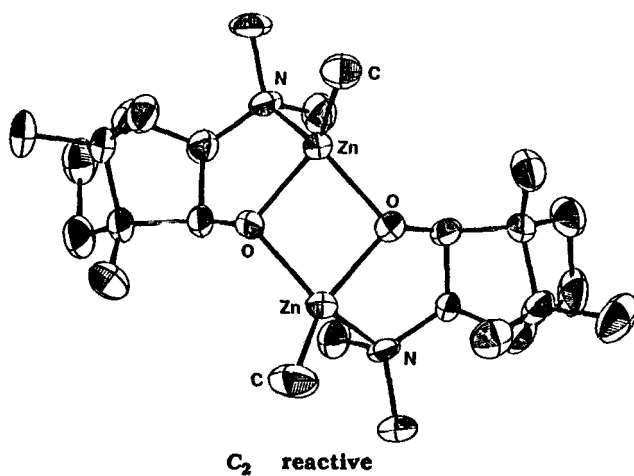
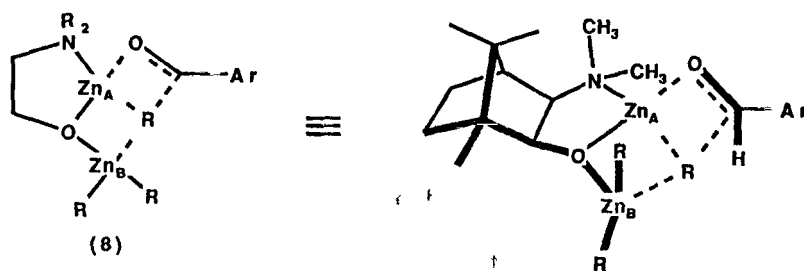
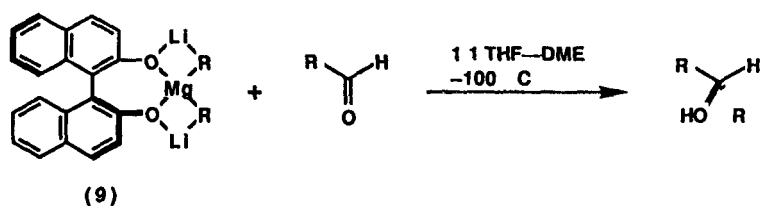
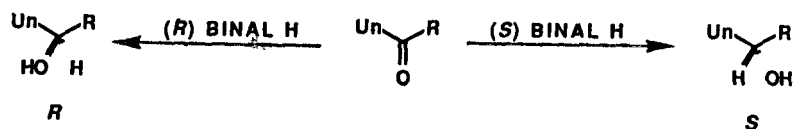


Figure 7 ORTEP drawings of complexes formed from equimolar amounts of dimethylzinc and (-) DAIB (upper) and dimethyl zinc and (±) DAIB (lower)

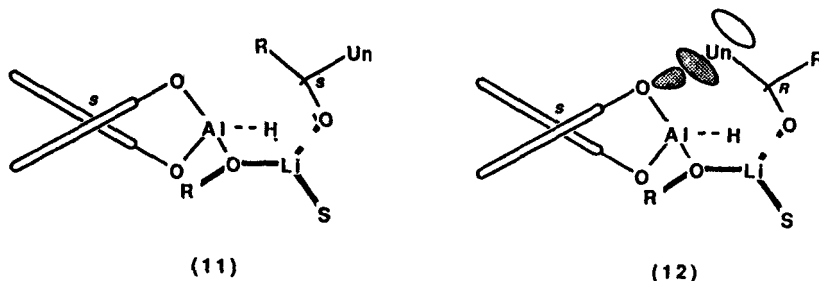
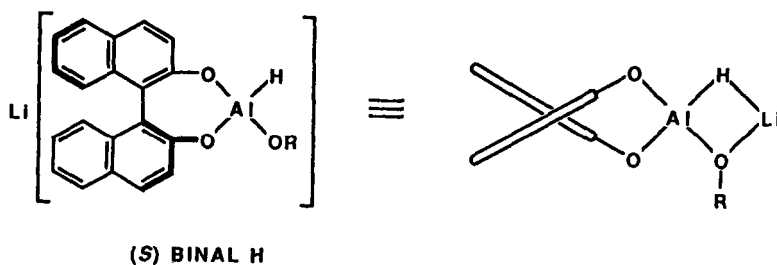




Scheme 14



Un = aryl alkenyl
alkynyl etc
R = alkyl H

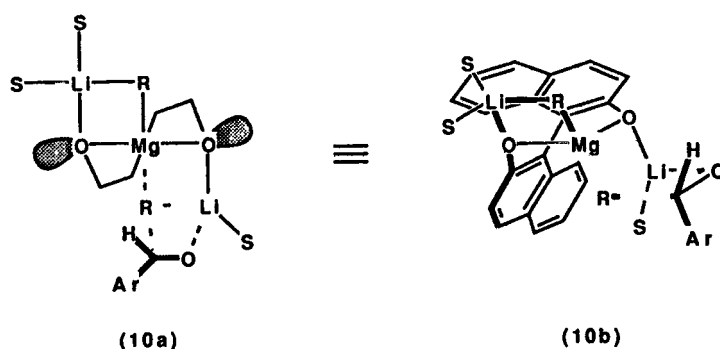


Scheme 15

H exhibits exceptionally high enantioface differentiating ability in the stoichiometric reduction of prochiral ketones having an aromatic olefinic or acetylenic substituent⁴⁰ With many carbonyl substrates *e.e.s* greater than 90% are obtainable where the enantioselection is governed primarily by electronic factors. Now a new model is presented to explain the general binaphthol/carbinol

⁴⁰ (a) R. Noyori, I. Tomino, Y. Tanimoto, and M. Nishizawa, *J. Am. Chem. Soc.* 1984, **106**, 6709. (b) R. Noyori, I. Tomino, M. Yamada, and M. Nishizawa, *J. Am. Chem. Soc.* 1984, **106**, 6717.

Centenary Lecture



configurational relationship (*S/S* or *R/R*) which is independent of the relative bulkiness of unsaturated and alkyl groups flanking the carbonyl moiety. In the (*S*) BINAL H reduction the *S* generating transition structure (11) is favoured over the diastereomeric *R* generating structure (12) because the latter is destabilized by the substantial n/π type electronic repulsion between a binaphthoxyl oxygen and the unsaturated moiety. The oxygen/*R* steric repulsion in (11) becomes significant by increasing the bulkiness of *R* but cannot overcome the overwhelming electronic influence (Scheme 15).

5 Epilogue

The development of homogeneous asymmetric catalysis using chiral metal complexes has provided the straightforward solutions to many challenging problems proving the validity of the chemical conception. I would conclude that asymmetric catalysis is a four dimensional chemistry which consists of two fundamental elements in Nature: chirality and circularity. High efficiency is obtainable by creation of ideal three dimensional structures (x, y, z) coupled with appropriate kinetics (t). This is a frontier of organic chemistry full of promise.¹

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