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Unlocking the potential of metal ligand cooperation for enantioselective transformations

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Metal-ligand cooperation, in which both the metal and the ligand of a transition metal complex actively participate in chemical transformations leading to enhanced reactivity or selectivity in chemical reactions, has emerged as a powerful and versatile concept in catalysis. This Viewpoint discusses the development trajectory of transition metal-based complexes as catalysts in (de)hydrogenative processes, in particular those cases where metal-ligand cooperation has been invoked to rationalise the observed high reactivities and excellent selectivities. The historical context, mechanistic aspects and current applications are discussed with the suggestion to explore the potential of the MLC mode of action of such catalysts in enantioselective transformations beyond (de)hydrogenative processes.

Introduction

The selection of a suitable ligand is a critical factor in the design of an effective homogeneous catalyst, as ligands can affect the reactivity, selectivity and the stability of a catalyst. Usually they remain chemically inert while all the key reaction steps take place at the metal centre. 1,2 Over the last few decades, however, an increasing number of systems have been developed in which the ligands in metal complexes have been proposed to play a more active co-operative role, leading to improved reactivities and selectivities. 1,2 These developments

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led to the introduction of the term 'metal-ligand cooperativity' (MLC), which can manifest itself in a variety of ways. In its most common use in catalysis, MLC refers to reactions catalysed by transition metal complexes that involve active participation of both ligand and metal in at least one mechanistically relevant bond-forming or bond-breaking step of the catalytic reaction.³ MLC catalysis with metal complexes bearing functional ligands, a design element not unknown to nature,4 allowed for significant developments in the field of catalytic hydrogenation including asymmetric versions.5

The earliest example of bond activation through MLC can be traced back to the work reported by Fujiwara and Moritani in 1967, describing a method to access stilbenes via aromatic C-H bond cleavage, that years later was shown to operate through a concerted metalation-deprotonation pathway.⁷⁻¹⁰ In



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Viewpoint Chem Soc Rev

the following years, Shvo and co-workers reported a ruthenium cyclopentadienone complex as a hydrogenation catalyst, capable of heterolytic cleavage of hydrogen-hydrogen bonds through MLC. 11 This catalyst class has been proven effective for (de)hydrogenations of C-O/C=O and C-N/C=N bonds and was mechanistically a forerunner of many follow-up methodologies involving (transfer-) hydrogenation (including asymmetric versions) that proposed the concept of MLC as a mechanistic rational. 12-14

Despite the success of this MLC concept in catalytic hydrogenations and the demonstration of its potential beyond dihydrogen activation (e.g. Si-H, B-H, and C-H), 15,16 its application in asymmetric catalysis remains largely limited to (de)hydrogenative transformations. Moreover, the reported mechanistic aspects of hydrogenative transformations have been revisited several times and the proposed 'classical' MLC pathway has been challenged following new computational and experimental data. 17-21 How catalytic mechanisms extend to systems beyond hydrogenation processes requires further investigation and it presents a great opportunity not only to develop new synthetic methodologies but also to test mechanistic hypotheses.

In order to assess the potential of already developed MLCtype hydrogenation catalysts and their structural analogues for applications in asymmetric methodologies beyond hydrogenative synthesis, here we review the development of relevant (de)hydrogenation catalysts and mechanistic proposals as well as recently developed non-hydrogenative transformations.

Before proceeding, it is worth noting that different literature reports use the terms 'MLC catalysis' and 'metal-ligand bifunctional catalysis' to describe a similar concept with subtle differences in the role of the ligand, with MLC catalysis being narrowed down to cases involving participation of both the metal and the ligand in the bond cleavage/formation events through their chemical modification, meaning the ligand becomes chemically non-innocent. 14,18,22-24 In this work, we will use these terms synonymously, implying that both terms cover cases with active ligand participation, including ligand deprotonation/protonation as well as ligand-driven substrate



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polarization or outer-sphere stabilization of the product intermediate via hydrogen bonding, thus involving both chemically innocent and non-innocent ligands. When discussing different mechanistic aspects, we will use these definitions interchangeably, trying to adhere to the terms described in the original reports.

Finally, the viewpoint does not aim to be comprehensive but rather focuses on the examples relevant for its conclusions.

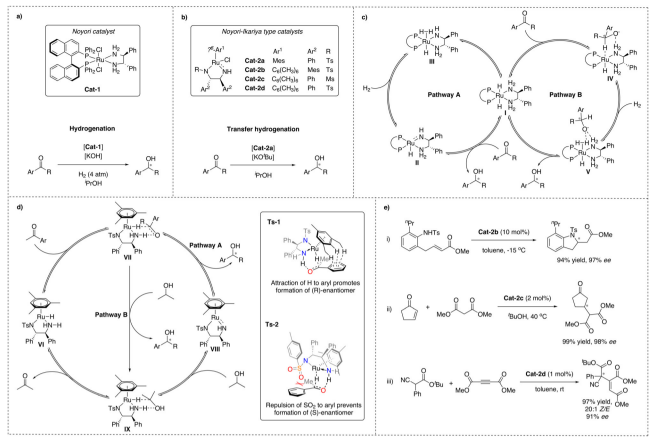
MLC in asymmetric (transfer) hydrogenation

In 1995, Noyori and co-workers described a ruthenium-based complex, Cat-1, bound to both a chiral diphosphine and a diamine ligand, which turned out to be a highly efficient catalyst for the enantioselective hydrogenation of arylketones with excellent stereoselectivity and favouring hydrogenation of carbonyl groups over olefin groups with remarkable chemoselectivity (Scheme 1a). 25,26 Later, Noyori and Ikariya developed another ruthenium catalyst that contains a similar diamine functionality together with an η^6 -coordinated arene, Cat-2, which shows exceptional performance in asymmetric transfer hydrogenation (ATH) reactions (Scheme 1b).27 Catalysts based on these original designs have found numerous applications in the pharmaceutical industry due to their high reactivity, chemo- and stereoselectivity.28 Because of this remarkable contribution to the field of asymmetric catalysis, Noyori was one of the Nobel prize laureates in 2001. 28-30

For both catalyst types, the effect of the NH-functionality was found to be crucial for the catalytic performance in hydrogenations and transfer hydrogenations, as tertiary amines were found to be inactive, pointing at an MLC mode of action and chemical non-innocence of the ligand. Extensive effort was made to understand the mechanistic details responsible for the high (up to 99%) C=O/C=C chemo- and enantioselectivities, exhibited by Noyori and Ikariya catalysts and the role of the NH-moiety of the ligand. This has led to the identification of two main scenarios (Scheme 1c and d).27,31,32

In the first scenario (pathway A), based on early studies by Noyori and Ikariya, the NH motif of the diamine ligand undergoes protonation and deprotonation during the course of the reaction. It was proposed that in the case of hydrogenation, the H₂ cleavage would occur via cooperative action of the metal and a deprotonated amide ligand formed in the presence of a base, leading to amine-hydride complex I.33 In the case of transfer hydrogenation, the Noyori-Ikariya catalyst (Scheme 1b) would transform into the active hydride species VI in the presence of a sacrificial hydrogen donor, e.g. isopropanol, often introduced as a solvent, and a catalytic amount of an alkoxide base. Subsequent concerted hydride and proton transfer to the substrate was proposed to occur next. Finally, the active form of the catalysts is regenerated from the resulting Ru-amido complex II and VIII upon reaction with H₂ for the former and isopropanol for the latter. 34,35 This scenario implies that hydrogenation and transfer hydrogenation take place through the outer-sphere of the metal catalysts via concerted transfer of both the metal RuH and the protic NH from the ligand to the carbonyl group via a six-membered transition state. This is opposite to the at the

Chem Soc Rev Viewpoint



Scheme 1 (a) Asymmetric hydrogenation using Noyori catalyst; (b) asymmetric transfer hydrogenation using Noyori–lkariya type catalysts; (c) simplified mechanisms for asymmetric hydrogenation using Noyori–lkariya type catalysts with stereodetermining catalyst–substrate interactions **TS-1** and **TS-2**; (e) selected examples of CH and NH bond activation by Noyori–lkariya type catalysts.

time more commonly accepted inner-sphere mechanism of transition metal-based hydrogenation catalysts. $^{36}\,$

Later studies of these metal-ligand bifunctional catalytic systems revealed a second scenario, constituting an alternative, more complex pathway (pathway **B**) for both Noyori hydrogenation and Noyori–Ikariya transfer hydrogenation catalysts. According to this revised mechanism, the ketone substrate is in both cases predominantly reduced *via* consecutive outersphere hydride transfer from the catalyst's metal center to the unbound substrate, followed by proton transfer from an η^2 -H₂ ligand and/or protic solvent. This also suggests that the NH functionality of the ligands is involved in the catalytic reaction *via* stabilization of the determining transition states through N-H···O hydrogen-bonding interactions, rather than *via* the originally proposed reversible proton transfer, and therefore, the ligand in this case would be chemically innocent. The second se

Both scenarios have been studied experimentally and *in silico*, leading to the conclusion that a stepwise process is energetically more favorable in solution. ^{14,17,18,31,33,34,37,38} Furthermore, it was suggested that the source of protonation of the product intermediate in the revised mechanism of Noyori–Ikariya transfer hydrogenation is dependent on the

reaction conditions. Solvents such as water or formic acid mixtures favor proton transfer from the protic solvent (pathway **B**), while use of isopropanol as solvent or increased acidity of the NH group of the ligand favor pathway **A**, although not in a concerted, but stepwise manner. It cannot be ruled out that both protonation pathways contribute to the overall reaction profile. ^{17,32,37}

Finally, to explain the experimentally observed acceleration of the reaction rate in the presence of high loadings of an inorganic base, for Noyori's catalyst an additional variation of pathway **B** has been proposed to contribute as well, in which the NH hydrogen atom is replaced by a metal cation, able to stabilise the alkoxylate *via* electrostatic forces, thereby replacing the hydrogen bond interaction.^{17,38,39}

In both scenarios, hydride transfer from the ruthenium center to the ketone is considered to be the stereodetermining step. 33,37,40 For the Noyori–Ikariya catalyst enantiodiscrimination has been mainly attributed to attractive π -CH interactions between the aryl moiety of the arylketone and the η^6 -arene moiety of the ruthenium complex (Scheme 1d, **TS-1**) as well as to repulsive interactions between the π -electrons of the arylketone and the lone pair on the SO₂ moiety of the diamine ligand (Scheme 1d, **TS-2**). 40,41

Chem Soc Rev Viewpoint

C-H acidic heterolytic bond activation for C-C and C-N couplings

Importantly, in addition to hydrogenative transformations, Novori-Ikariya type catalysts have also been reported to be effective for enantioselective C-C and C-N bond formations via activation of CH and NH acidic substrates (Scheme 1e). Specific examples are the intramolecular aza-Michael reaction (i),42 conjugate addition of 1,3dicarbonyl substrates to Michael acceptors (ii)43,44 and addition of α-cyanoacetates to acetylenic esters (iii)⁴⁵ and dimethylazodicarboxvlates.46 A common mechanistic characteristic of these examples are substrate activation by deprotonation via the amido ligand. Interestingly, the reaction with acetylenic esters shows a high Z-selectivity, which has been rationalised by a stereospecific protonation of the coordinated reaction intermediate via the NH functionality of the catalyst.47 This indicates that the NH moiety might have two functions: facilitating stereoselective coordination of the substrate to the metal and serving as a donor for stereospecific protonation. Nevertheless, the scope of the reported non-hydrogenative asymmetric transformations catalysed by Noyori-Ikariya type catalysts is mainly limited to these few examples.

Pincer complexes

Noyori's discovery has also influenced the development of another class of catalyst structures that contain a tridentate, predominantly meridionally coordinated ligand and also an acidic functionality capable of engaging in MLC upon base activation.24,48 This class of catalysts, referred to as 'pincer' complexes, a terminology introduced by van Koten et al., exhibits unique properties. 49 The general structure of pincer complexes consists of a central donor atom (e.g. NH, N_{Pvridine}, C) connected to two flanking donor atoms (P, N, S,O, C) (Scheme 2a). 50 This blueprint allows for facile tuning of ligand properties. Where the flanking donor atom can be varied to stabilise different metal ions, tune electron density at the metal center and introduce steric hindrance, the nature of the central donor ligand is pivotal for the reactivity of the corresponding complex. 48 These remarkable characteristics of pincer complexes have resulted in the development of numerous methodologies, including (de)hydrogenations and dehydrogenative couplings relevant for industry and sustainable synthesis efforts. 51-54

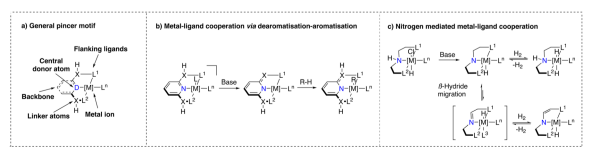
Bond activation with pincer complexes through MLC

Pincer complexes (Scheme 2a) that were suggested to engage in MLC can be divided into two major classes: (i) ones that bear ligands consisting of a pyridine core with either a CH254 or an NH⁵⁵ linker, which we will refer to as 'Milstein pyridyl' type pincer complexes (Scheme 2b) and (ii) ones that bear a central secondary amine donor atom, 24,50 which we will refer to as 'NHonly' type pincer complexes (Scheme 2c). In the presence of a base, the former class undergoes a dearomatisation process initiated by deprotonation of the linker, whereas the latter creates a basic amido moiety. Upon deprotonation, the corresponding complexes can activate reactants through MLC.

In 2005, Milstein and co-workers were the first to report on pincer complexes that can undergo reversible dearomatisation.⁵⁶ These complexes were found to be effective for activating various bonds and it is proposed that the linker of the dearomatised ligand plays a crucial role as nucleophilic acceptor to activate an electrophilic substrate followed by re-aromatisation.^{3,54} This MLC action mode has attracted much interest from both experimentalists and theoreticians who have extensively studied the mechanism involving the aromatisation/dearomatisation process during the catalytic cycle of the corresponding reactions. However, several recent studies have also pointed out that the originally proposed mechanism of hydrogenation/dehydrogenation reactions with Milstein type catalyst may not necessarily follow the 'classical' MLC pathway observed under stoichiometric conditions due to likely differences in speciation/conditions when comparing stoichiometric and catalytic reactions. 17,57-61

In case of the NH-only type pincer complexes (Scheme 2c), their ability to engage in MLC bond activation was first proposed by Fryzuk in the 1980s for the hydrogenation of olefins. 62-64 The proposed reactivity pattern of these complexes is strikingly similar to that of the Novori-Ikariya catalysts.

Deprotonation of the secondary amine results in a basic amido moiety, capable of stabilizing a coordinatively unsaturated metal species and activating H-H bonds. Depending on the nature of the metal and ligands, a β -hydride migration step, followed by subsequent dihydrogen elimination (or deprotonation) can lead to an enamine modification of the pincer ligand, which influences the geometry and hence the ligand field distortion (Scheme 2c).65 This structure shows similarities to the dearomatised Milstein-pyridyl complexes described earlier. NH-only type pincer complexes have been shown to exhibit high catalytic activity and stability in numerous hydrogenbased transformations.66-69 However, further heterolytic HX bond activations are scarce.⁷⁰



Scheme 2 (a) General pincer complex structure; (b) base induced dearomatisation-aromatisation MLC activation mode of Milstein pyridyl type pincer complexes; (c) base induced MLC activation modes of NH-only type pincer complexes

Chem Soc Rev Viewpoint

MLC pincer complexes in enantioselective reactions

As the field progressed, asymmetric transformations using chiral MLC pincer complexes were explored as well. For the Milstein pyridyl type complexes, chirality was introduced primarily at the linker moiety and the flanking donor ligand (Scheme 3a). In 2015, Castillón and Díaz reported a ruthenium-based complex bearing P-stereogenic flanking donors for the hydrogenation of arylketones.⁷¹ Good results with enantioselectivities up to 95% were obtained only at low temperature (-40 °C). It is suggested that conformational equilibria of the catalyst may account for this temperature effect.⁷² In addition, Mezzetti and co-workers reported in 2018 an earth abundant variant using iron which was subjected to enantioselective hydrogenation of acetophenone.⁷³ In contrast to the work of Castillón and Díaz, rapid conformational interconversion could not be prevented, resulting in mediocre enantiomeric excesses for this transformation.

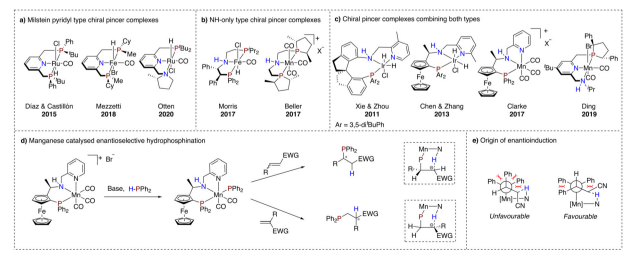
The conformational problems of these types of complexes were further described by Otten and co-workers, showcasing two challenges for stereoretention of dearomatised asymmetric ruthenium hydride PNN pincer complexes with a chiral pyrrolidine flanking donor (Scheme 3a). 74 The issues are associated with inversion of stereochemistry on the chiral amine and epimerisation of a metal-hydride bond facilitated by the dearomatised linker moiety.

On the other hand, NH-only type pincer complexes have successfully been employed in asymmetric (transfer) hydrogenation of ketones and imines. Remarkably, earth abundant metal complexes have not fallen short within this development trajectory (Scheme 3b). In fact, chirality introduced in an iron complex via substitution on the ethyl linker, 75,76 as well as in a manganese complex through chiral phospholane flanking donor substituents, has led to chiral alcohol and amine products with enantiomeric excesses exceeding 80%.⁷⁷

The two types of reviewed pincer motifs display almost complementary advantages. The Milstein pyridyl type complexes have been suggested to engage in cooperative activation of various bonds, but are less developed for asymmetric transformations. On the other hand, NH-only type pincer complexes are known to catalyse asymmetric (transfer) hydrogenations, but are underexplored in terms of further HX bond activation. Thus, combining these two different ligand motifs in one pincer complex potentially offers more opportunities in asymmetric catalysis (Scheme 3c).

In this context, a very effective iridium complex for the hydrogenation of arylketones with both a dearomatisable pyridine ring and a central NH moiety was reported by Xie and Zhou in 2011. 78 MLC was suggested to operate via the NH moiety, while the pyridine ring was proposed to simply add stability to the catalyst. Following up on this work, in 2013, Chen and Zhang reported an enantioselective hydrogenation reaction with a similar iridium complex, containing planar chirality as a chiral element in addition to a carbon stereocenter. 79 In 2017, Clarke and co-workers reported the first example of a manganese catalysed enantioselective hydrogenation using a similar ligand structure. 80 Interestingly, a facial coordination was found for this complex. Another example of a manganese-based catalyst that merges structural features of both types of MLC-capable pincer complexes was reported by Ding and co-workers.81 Mechanistic studies in this case suggest an outer sphere type mechanism, mediated by the NH functionality of the flanking donor.

Surprisingly, to date only the manganese complex reported by Clarke has been demonstrated to catalyse different asymmetric transformations. In 2021, the group of Harutyunyan reported the first enantioselective hydrophosphination of unsaturated conjugated nitriles using this catalyst. 82 It is noteworthy that both internal and terminal unsaturated nitriles yielded enantiomerically pure products, where the former results from enantioselective phosphination and the latter from stereospecific protonation, which was attributed to an MLC mode of action of the catalyst through the NH functionality of the ligand (Scheme 3d).



Scheme 3 (a) Chiral Milstein pyridyl type pincer complexes; (b) chiral NH-only type pincer complexes; (c) chiral pincer complexes with merged structural features of both MLC active pincer types; (d) enantioselective hydrophosphination of α , β -unsaturated compounds catalysed by Clarke's manganese catalyst; (e) proposed origin of enantioinduction for the manganese catalysed enantioselective hydrophosphination

Viewpoint Chem Soc Rev

In addition to the hydrophosphination of nitriles, the catalyst system was also successfully applied for the enantioselective hydrophosphination of α,β-unsaturated esters, ketones, carboxamides, phosphine oxides and trifluoromethyl substituted substrates. 83-85 Based on initial experimental and theoretical studies, it was proposed that activation of the H-P bond occurs through MLC, initiated by treatment of the manganese catalyst with a base (Scheme 3d). The transfer of the resulting phosphide from the manganese atom to the alkene was put forward as the stereodetermining step of this transformation. Steric interactions between the alkene substrate and the phenyl substituents at the phosphorus atom, as well as by hydrogen bonding between the NH functionality of the catalyst and the α-carbon of the alkene, are the main processes that control the stereochemistry (Scheme 3e). The observed enantioselectivity for terminal alkenes was rationalised by the subsequent same face stereospecific intramolecular protonation step, taking place after the transfer of the diphenylphosphide from the manganese center to the β-carbon of the alkene.82 Crucial for the successful H-P bond activation and subsequent enantioselective hydrophosphination are the interplay between the metal and suggested non-innocence of NH and pyridine functionalities of the ligand.

Conclusions and outlook

Since its discovery, MLC has become a versatile tool for catalysis, especially in the field of (transfer) hydrogenation. Noyori's first reports described a bifunctional ruthenium catalyst with a diamine ligand bearing an NH functionality for the enantioselective (transfer) hydrogenation of aryl ketones. This work foreshadowed the development of MLC-active tridentate pincer complexes, which subsequently found widespread application in catalysis. The two most investigated MLC activation modes either operate via dearomatisation of a pyridyl moiety with subsequent rearomatisation upon bond activation or contain a central noninnocent NH functionality that can act as a hydrogen bond donor or proton relay. The former activation mode has been shown to activate various bonds, such as NH, CN and CH, but is underdeveloped for enantioselective transformations. In contrast, pincer scaffolds with a central NH functionality have been successfully used in various enantioselective (transfer) hydrogenation methodologies by introducing chirality to the linker and flanking donor framework, but their ability to activate other chemical bonds has not been studied in depth.

Both types of ligand motifs (a dearomatisable pyridyl flanking donor and a secondary NH functionality) have also been combined, leading to, among others, the highly efficient facially coordinated manganese pincer complex reported by Clarke. Apart from catalysing enantioselective (transfer) hydrogenations, this complex enabled the development of a general catalytic enantioselective hydrophosphination method for a wide range of activated terminal and internal alkenes. The interplay between the two modes of activation of the MLC present within the single pincer complex was suggested to be a crucial feature for the observed broad substrate scope in this transformation. The various reactivity patterns underline the potential of MLC in pincer complexes for versatile enantioselective bond activation.

Neither ligand nor coordinated metal exploration has been exhausted so far. New MLC chiral pincer complexes with earth abundant metal ions and ligand variations could further expand the type of bonds that can be activated with these complexes, particularly with respect to heterolytic HX bond activation.86 Opportunities lie, for instance, in exploration of the reactivity of unsaturation in the ligand backbone in conjunction with NH-only functionalities. Furthermore, the tunable charge density of an activated amido functionality could provide control over electrostatic interactions and substrate polarisation, potentially relevant for nucleophilic substitution reactions. Finally, the reversibility of the discussed MLC activation modes could result in proton relay, facilitating isomerisation reactions and the stabilisation of reactive species.⁸⁷

Nevertheless, the mechanistic picture of operating baseactivated MLC pincer complexes remains intricate. Mechanistic details on the role of the NH functionality in (transfer-) hydrogenations, for example, are still under investigation, and different mechanistic routes are possible depending on the reaction conditions. Especially for chiral pincer ligands, formation of several isomers of the resulting complexes, including facial and meridional coordination must be taken into account. 70,75 Despite this ambiguity, the initially proposed MLC concept has resulted in the development of a large number of methodologies, used in many useful applications. However, opportunities arising from this mechanistic framework go hand in hand with low predictability of reactivity profiles. Hence, more work is required on rational design and mechanistic understanding of chiral pincer complexes, especially for structures combining different MLC activation modes. This development direction may allow the promising potential of MLC pincer complexes to be fully exploited in various asymmetric transformations, including for instance heterolytic bond activation of phosphines, amines, alcohols and thiols, as well as activation of CH acids, nucleophilic substitution reactions and isomerisations.88

Conflicts of interest

There are no conflicts to declare.

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