1 The Primary Organometallic in Copper-Catalyzed Reactions

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1.1 Scope and Introduction

In this chapter, the term primary organometallic will mean both the terminal organometallic (RM) selected for a desired asymmetric transformation and those Cu-species that result once the RM is combined with a suitable copper precursor. A significant advantage in copper-promoted chemistry is the ability to access a very wide library of $M[CuXR_nL]$ species ($M$, main group metal; $X$, halide or pseudohalide; $R$, organofunction; $L$, neutral ligand) by simple variation of the admixed reaction components. Normally, the derived cuprate mixture is under rapid equilibrium such that if one species demonstrates a significant kinetic advantage, highly selective reactions can be realized. The corollary to this position is that deconvoluting the identity of such a single active species from the inevitable “soups” that result from practical preparative procedures can prove highly challenging. In this review, we concentrate on asymmetric catalytic systems developed in the last 10 years, but where necessary, look at evidence from simpler supporting achiral/racemic cuprates. Our aim is to try and present a general overview of bimetallic (chiral) cuprate structure and reactivity. However, given this extremely wide remit, the coverage herein is necessarily a selective subset from the personal perspective of the author. There are a number of past books of general use (either totally or in part) that provide good primers for this area [1]. Additionally, because of its relevance the reader is advised to also consult Chapter 12, which deals with mechanism.
1.2

Terminal Organometallics Sources Available

One partial representation of the totality of asymmetric processes promoted by copper and main group organometallic mixtures is given in Figure 1.1; where the height of the bar indicates published activity \( (n = \text{number of papers, etc.}) \) and the percentage in the black roundel is the fraction published in the last 5 years (2007–2012).

The seven metals identified (Li, Mg, Zn, B, Al, Si, and Sn) form the basis of this overview. It should be noted that (i) the dominance of magnesium is due to numerous simple addition reactions where the resultant stereochemistry is controlled only by a chiral substrate; (ii) asymmetric reactions of the organometallics of the lower periods are still largely unreported; (iii) while all areas have developed, there has been especial interest in some metalloids in recent years (e.g., organoboron reactions); and (iv) the use of silicon organometallics is over reported in Figure 1.1 by the extensive use of silanes as reducing agents. The general properties of the organometallics used in asymmetric copper-promoted reactions are given in Table 1.1, compared with a generalized \( \text{LCuR} \) fragment. A common feature is their

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1) The data arises from a Scifinder search (24 April 2012) of the terms: “copper, asymmetric, and the various organoelement terms (e.g., organolithium, etc.).” For Mg and Si, the more common terms Grignard and silane were used. Manual screening of the derived dataset indicated the applicability of the references to this chapter. A similar ratio of use was attained from substructure searching of asymmetric reactions of RM with generalized substrates.
### Table 1.1 Properties of main group organometallics used in asymmetric Cu-promoted processes in order of element electronegativity

<table>
<thead>
<tr>
<th>Organometallic type(s)</th>
<th>M–Me bond energy (kcal mol(^{-1}))</th>
<th>M–C bond length (Å)</th>
<th>M–O bond energy (kcal mol(^{-1}))</th>
<th>Electronegativity(^b)</th>
<th>Oxophilicity(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiR</td>
<td>64 [267]</td>
<td>2.31</td>
<td>112 [470]</td>
<td>0.98</td>
<td>1.7</td>
</tr>
<tr>
<td>RMgX, MgR(_2)</td>
<td>60 [253]</td>
<td>2.15</td>
<td>113 [471]</td>
<td>1.31</td>
<td>2.3</td>
</tr>
<tr>
<td>AlR(_3), AlR(<em>n)Y(</em>{1−n})</td>
<td>68 [283]</td>
<td>1.97</td>
<td>101 [418]</td>
<td>1.61</td>
<td>1.6</td>
</tr>
<tr>
<td>ZnR(_2), RZnX</td>
<td>68 [285]</td>
<td>1.93</td>
<td>91 [381]</td>
<td>1.65</td>
<td>2.4</td>
</tr>
<tr>
<td>LCuR(^d)</td>
<td>57 [238]</td>
<td>1.98</td>
<td>49 [204]</td>
<td>1.90</td>
<td>0.7</td>
</tr>
<tr>
<td>SiR(_4), R(^1)SiR(_2)(_1)</td>
<td>77 [320]</td>
<td>1.85</td>
<td>100 [419]</td>
<td>1.90</td>
<td>2.0</td>
</tr>
<tr>
<td>SnR(_4), R(^1)SnR(_2)(_3)</td>
<td>63 [262]</td>
<td>2.16</td>
<td>49 [203]</td>
<td>1.96</td>
<td>1.0</td>
</tr>
<tr>
<td>BR(_3), RBY(_2)</td>
<td>89 [374]</td>
<td>1.58</td>
<td>124 [519]</td>
<td>2.04</td>
<td>1.5</td>
</tr>
</tbody>
</table>

\(^a\)Most data in Table 1 from Ref. [2], 1 kcal mol\(^{-1}\) is 4.19 kJ mol\(^{-1}\); X = halide, Y = OR.

\(^b\)Pauling’s scale, data from Ref. [3].

\(^c\)E(M – O)/E(M – S) is often taken to correlate to a metal’s oxophilic HSAB character, see Ref. [2].

\(^d\)Generalized data from additional citations in Ref. [2].

Tendency to form strong bonds with oxygen, providing strong thermodynamic driving forces for additions to carbonyl-containing substrates. This tendency can be correlated to their relatively low electronegativities and high oxophilicities (“hardness” defined here as E(M – O)/E(M – S)). The published Sn–O bond energy, derived from density functional theory (DFT) calculation, is probably somewhat underestimated in this respect. The reactivity of main group organometallics in Table 1.1 is reinforced by their weak M–C bonds. In fact, M–Me values are often upper limits – the bond energies of the higher homologs are frequently lower by 5–10 kcal mol\(^{-1}\) meaning that in mixed R\(^1\)MMe\(_n\) the methyls can be used as a potential nontransferable groups. Similarly, significant increases in the reactivity of organoelement compounds across the series M(alkyl), M(aryl), and M(allyl) are observed. At least in the allyl case, this is correlated to the M–C bond strength, which is typically >10 kcal mol\(^{-1}\) lower than M–Me.

### 1.3 Coordination Motifs in Asymmetric Copper Chemistry

Copper-promoted asymmetric reactions frequently attain high enantioselectivity through reduction in substrate conformational mobility via two-point binding, as in the general copper(I) cuprates \(1a,b\) or by \(\eta^2\)-binding at chiral Cu\(^{II}\) complexes, generalized by 2 (Scheme 1.1). As copper(II) does not form organometallic species, and readily undergoes reduction to Cu\(^{I}\) in the presence of RM, we concentrate here mainly on the former (overviews of activation by 2 can be found through the work
of Rovis and Evans [4]). One other common scenario is use of a chiral Lewis acid fragment linked to a simple heterocuprate via a bridging ligand 3.

Clearly, in attaining enantioselective transition states for asymmetric reactions, the nature of the bridging ligand (typically a halide or pseudohalide) is at least as important as the identification of an effective chiral ligand (L*) in attaining effective docking of substrates in 1–3.

1.3.1 Classical Cuprate Structure and Accepted Modes of Reaction

1.3.1.1 Conjugate Addition

Owing to their initial discovery, an enormous degree of activity has focused on the structures and reactivity of the Gillman-type homocuprates (LiCuR₂) and their heterocuprate analogs (LiRCuX, where X is the halide of pseudohalide [5]). In general, while these systems have provided underlying understanding of the basics of copper(I)/copper(III) organometallic chemistry they have not directly provided reagents that give highly selective catalytic asymmetric methodology. It is instructive to ask, “Why is this the case?” — a question that modern computational DFT understanding of the reaction course can cast light on. In classic conjugate addition, dimeric [LiCuMe₂]₂ reacts with cyclohexenone via transition state 4 [6], in which the enone-bound copper is formally at the +3 oxidation state (Scheme 1.2). One issue is that coordination of the d⁸ Cu³⁺ center with an additional neutral chiral ligand (e.g., a phosphine) has to compete with excess strong σ-ligands in the solution (e.g., Me⁻), and also from intramolecular donation from the enolate π-bond (which renders the Cu-center coordinatively saturated). Another issue is associated with the lability of any (R–Cu–R)Lιₙ bridge. Although Li–O contacts in organocuprates are essentially covalent, an ionic formulation for 4 has been used here to emphasize the propensity of such units to exchange and associate. Such behavior is nicely demonstrated by the diffusion NMR studies of Gschwind [7], which measure the “size” of cuprates in solution allowing estimations of their identities. These studies show how easily cuprate-based bridges are readily displaced by tetrahydrofuran (THF) (leading to a catastrophic enone inactivation by loss of the Oenone···Li Lewis acid contact) or, alternatively, promotion of multiple
species through aggregation of 4 in less polar solvents. This ease of displacement of homocuprate bridging groups by “lithium-liking” pseudohalides (e.g., alkoxide impurities in RLi or derived from halides in CuX precursors) sparked, even in the earliest days, ideas of avoiding such problems through heterocuprates LiRCuX* use (where X* provides a rigid ordered bridge promoting strong transfer of stereochemistry). The proposal of Dieter and Tokles 5 [8] (Scheme 1.2) is one such case. In fact, related species have been characterized in solution by NMR, of which 6 is a nice example [9].

Chiral heterocuprates of types 5 and 6, and other species using alkoxide-based units, or related units controlling chirality through motif 3 (e.g., the sparteine-based reagents of Dieter [10]) have all provided rich structural chemistry [11] and effective asymmetric stoichiometric reagents for target synthesis [8, 10]. However, such chiral heterocuprate approaches have not transferred well to catalytic applications. Even the most effective system of van Koten [12] (Scheme 1.3) provides only a modest 76% ee_{max} in the addition of MeMgI to benzylidene acetone. This is not due to a failure in chiral recognition by the cuprate derived from 7 but due to unavoidable transfer of the chiral thiolate donor to the terminal main group RM source, which is facilitated by the excess of MeMgI present in the catalytic system. This provides inactive magnesium chiral thiolates and highly debilitating racemic catalysis through small amounts of MgBr[CuMe₂]. Davies et al. [13] has demonstrated an explicit case of the failure of a related “nontransferable” amido group in 8 as this undergoes rapid exchange on the NMR timescale leading, ultimately, to a mixture of homo and heterocuprates at 0 °C (Scheme 1.3).
Scheme 1.3  Exchange of hetero groups in chiral heterocuprates and effects on selectivity.

Scheme 1.4  DFT modeling of $S_N2'$ allylation regiochemistry (key bond lengths in angstrom next to $\text{C}_3$–Cu).
1.3 Coordination Motifs in Asymmetric Copper Chemistry

1.3.1.2 S_N2’ Allylation Reactions
Despite early successes using chiral-leaving groups, before 1999 [14], Cu\(L^\ast\)-mediated asymmetric displacements of \(X (X = \text{halide, OAc, OP(O)(OEt)}_2\text{, etc.})\) from \((E)-\text{RCH}=\text{CHCH}_2\text{X}\) proved highly challenging. While it was clear that the selective transition state(s) were normally associated with anti \(\gamma\)-attack of the nucleophile on allyl electrophile, the structures were rather too reactive to be identified. Using DFT approaches, Nakamura has put forward the most useful picture of the reaction coordinate (Scheme 1.4) [15]. Oxidative addition of allyl acetate to Gilman’s reagent defines the enantioface of the electrophile coordinated and through transition state 9 and delivers the symmetrical \(\pi\)-allyl complex 10. For clarity, the \(\gamma\)-carbon is emphasized (●). The symmetrical nature of 10-\(\pi\) and its ability to undergo classic \(\pi-\sigma\) interconversion indicates that, in substituted allylic systems, control of regiochemical issues is likely to be at least as great a challenge as inducing high levels of asymmetric selectivity. Support for these ideas comes from the interaction of theory and experiment. Replacing the Cu\(\text{Me}_2\)− fragment by MeCuCN− in the DFT modeled oxidative addition reveals two factors: (i) a slower oxidative addition, but (ii) high polarization of the Cu \(d_{xz}\)-based highest occupied molecular orbital (HOMO) providing greater electron density trans to the CN group. This leads to more developed C\(\gamma\)–Cu bond in transition state 11, which is retained in the resulting Me(CN)Cu\(\text{III}\) (allyl) intermediate (the analog of 10). Faster reductive elimination of this species is also predicted, minimizing \(\pi-\sigma\) interconversions. The predictions nicely account for the change of regioselectivity observed in the reactions of substrate 12 (Scheme 1.4).

1.3.2 Motifs in Copper-Main Group Bimetallics and Substrate Binding
The ground-state structures of active copper reagents, especially those “loaded” with reaction substrates, are normally too labile to be isolated. While major progress has clearly been made through computational (DFT) approaches above, scanning crystallographic databases2) reveals a significant number of model compounds that provide insight into substrate binding. For example, a handful of CuOTf complexes indicate similar binding modes for alkenes, dienes, and alkynes (all Cu–C 2.05–2.22 Å with C=C \(\sim\)1.4 Å or C≡C \(\sim\)1.2 Å), phosphines (Cu–P 2.19–2.28 Å), and triflate (Cu–O 2.04–2.43 Å). Particularly, structures OFANAK [16] and HIZCIC [17] provide tantalizing hints that typical bridging ligands such as triflate and acetate will produce highly ordered bimetallic structures when presented with suitable RM (Scheme 1.5). For example, in addition to clearly showing the poorer binding of the trisubstituted alkene, stripping out the core of OFANAK provides a key CuOTf core well predisposed to bind enones and RM. Similarly, HIZCIC

2) Search of the Cambridge Crystallographic database conducted (25 April 2012). The complexes analyzed were CAFQUV, CEJGEE, COMMAS, FUTRIV, GEKZOL, HIZCEY, HIZCIC, JUPXUN, MIMCAN, MOHLIE, OFANAK, REXJOU, TACYAX, and VIFSEJ.
reveals the potentially tight and ordered structure that a bridging acetate can engender. For clarity, the $C_{40}H_{58}N_{2}O_{4}$ linker in HIZCIC is not shown. No structural data for carboxylate bridging between Cu(I) and boron, aluminium, or silicon were found.

An additional range of X-ray structures are available providing insights into M-halide–Cu(I) motifs in nonisolable catalytic intermediates (M = Li, Mg, Zn, Si). These model compounds, together with selected inter Cu–X–M atomic distances are given in Scheme 1.6. In the case of Mg and Zn, only copper(II) models could be identified. By taking published average binding modes for the docking of carbonyl oxygen species with various main group metals [2] and a value of 2.15 Å for Cu–C alkene binding, then rough estimates of the optimal distance ($d_2$) the bridging ligand which should separate the Cu···M pair by for an s-trans-enone 13 (such as cyclohexenone), are attained (table in Scheme 1.6).

As can be seen, for the table within Scheme 1.6, single halogen atom bridges typically place the key M···Cu bimetallic pair a little closer than the idealized binding mode. It can be surmised that the success of larger bridging groups (OAc, OTf, thiophene carboxylate, etc.) in asymmetric catalysis is due to their ability to increase the Cu/M separation into an optimal range.

In the sections that follow, we focus on information that is available in real-world copper-based asymmetric reagents focusing on evidence that points to the structure of the primary cuprate (or other) species involved. In looking for common features, the reactivity has been grouped by metal rather than by transformation. Again,
1.4 Asymmetric Organolithium–Copper Reagents

Only very recently (2011) have organolithium reagents finally yielded to ligand-promoted asymmetric catalysis. Use of TaniaPhos $\mathbf{L_A}$ with CuBr·SMe$_2$ allows some utterly remarkable additions of alkyl organolithiums to (E)-cinnamyl bromides 14 (Scheme 1.7) with near perfect enantioselectivity [19]. This catalyst is able to promote the γ-selective allylation reaction with many desirable features: (i) ArBr exchange with nBuLi is avoided (run 3), (ii) alkyl substrates with poor steric profiles are tolerated (run 4), and (iii) even normal electrophiles for RLi (Boc groups and esters) are tolerated (run 6). The success of this chemistry is due to the

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Scheme 1.6 X-ray crystal structures of M–X–Cu model complexes (interatomic distances in angstrom) and their relationship to an idealized Cu···M bound s-trans-enone ($d_2$ calculated from J mol models).

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a number of reviews, especially those concentrating on catalytic chemistry are pertinent [18].

1.4 Asymmetric Organolithium–Copper Reagents

Scheme 1.6 X-ray crystal structures of M–X–Cu model complexes (interatomic distances in angstrom) and their relationship to an idealized Cu···M bound s-trans-enone ($d_2$ calculated from J mol models).

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Scheme 1.6 X-ray crystal structures of M–X–Cu model complexes (interatomic distances in angstrom) and their relationship to an idealized Cu···M bound s-trans-enone ($d_2$ calculated from J mol models).
reaction conditions – RLi is added slowly into a very nonpolar solvent mix at low temperature. Under these conditions, only compound 16 is formed and this is stable to excess RLi in solution. The use of nonpolar solvent is critical. Replacement of CH$_2$Cl$_2$ by Et$_2$O results in the very fragile bimetallic 17 that readily expels the chiral ligand forming achiral LiCuMe$_2$, a process that can be followed by $^{31}$P NMR spectroscopy. This decomposition pathway means that only low enantioselectivities are realized (28%) in Et$_2$O. In CH$_2$Cl$_2$–hexane, the ee values realized for nBuLi addition to 14 are largely independent of the copper source used (CuBr·SMe$_2$, CuCl, CuI, Cu(TC); TC, 2-thiophene carboxylate), suggesting that this is a rare example of direct reaction of a ligated organocopper reagent (17) without a ligand bridge (halide or alkyl) to Li-activated 14 (cf. Scheme 1.4). This idea is supported by the observation that similar excellent behavior is achieved for hindered iPr and sBu additions to 14 (X = Cl) using a simple phosphoramidite ligand. Clearly, mechanistic and calculative studies are needed to address this point (Scheme 1.7).

### Scheme 1.7
Asymmetric $S_N$2′ allylation with organolithium species.

The same simple phosphoramidite $L_B$ has very recently allowed highly enantioselective desymmetrization reactions of exobicyclic alkenes (Scheme 1.8) [20]. However, in this case, no NMR studies were carried out on the primary organometallic, but clearly at least 1 equiv of Et$_2$O is tolerated by the catalyst structure.
1.5 Asymmetric Grignard–Copper Reagents

While traditional magnesium cuprates “MgX[CuR₂]·MgX₂” are easily prepared by stoichiometric reaction of CuX with 2RMgX; extension to chiral heterocuprate versions of such chemistry have distinct limitations (see Section 1.1.1). The dramatic growth in this area in the last 10 years has come from utilization of chiral diphosphine-based copper(I) catalysts in nonpolar solvents, typically tBuOMe or CH₂Cl₂ [21], a concept whose genesis can be traced to the seminal use of *monophosphines* by Tomioka [22]. The primary organometallics involved in conjugate addition reactions have become unmasked through NMR studies. For example, the isolated Josiphos complex [CuBrL₄]₂ readily catalyzes the addition of a wide range of Grignard reagents to various α,β-unsaturated carbonyl compounds with good to excellent enantioselectivity (Scheme 1.9). Addition of excess MeMgBr to [CuBrL₄]₂ in CH₂Cl₂ led to formation of a major new species that was assigned structure 18 with a Cu–Me signal present at about δ₁H −0.3, compared to the free
Table 1.2  Primary organometallics in asymmetric 1,4 RMgX additions at about \(-60^\circ\text{C}\).

<table>
<thead>
<tr>
<th>Reaction conditions</th>
<th>Major species present</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuBr·SMe(_2) + L(_C) + MeMgBr(^a)</td>
<td>Mixture containing no 18</td>
</tr>
<tr>
<td>[CuBr(_{2})Cl(_2)] + excess MeMgBr (CH(_2)Cl(_2), E(_2)O, or toluene)</td>
<td>18</td>
</tr>
<tr>
<td>[CuBr(_{2})Cl(_2)] + excess MeMgCl (CH(_2)Cl(_2))</td>
<td>19</td>
</tr>
<tr>
<td>[CuBr(_{2})Cl(_2)] + excess MeMgI (CH(_2)Cl(_2))</td>
<td>Exchanging mixtures</td>
</tr>
<tr>
<td>[CuBr(_{2})Cl(_2)] + excess MeMgBr (THF)</td>
<td>19</td>
</tr>
<tr>
<td>[CuBr(_{2})Cl(_2)] + &gt;2 equiv MeLi (CH(_2)Cl(_2))</td>
<td>(L_C + LiCuMe_2)</td>
</tr>
</tbody>
</table>

\(^a\)Reaction solvent not defined.

MeMgBr at \(\delta_H 1.64\) [23]. The Cu:Me ratio in 18 was determined by integration of the methyl signal against the aryl region of the ligand. Traces of a second species were identified by \(^{31}\text{P}\) NMR but its nature could not be ascertained; however, 18 was entirely converted to this entity when it was exposed to air at \(-60^\circ\text{C}\). Conversely, when [CuBr\(_{2}\)Cl\(_2\)] was reacted with MeLi in CH\(_2\)Cl\(_2\), only 19 was formed. The presence of MgBr\(_2\) in 18 was confirmed by its precipitation as its dioxane coordination polymer. In stoichiometric reactions with (E)-BuCH=CHCOMe, each of the three species behave differently: 18 gives mostly 1,4-addition in 92% ee; 19 only traces of 1,4 addition in 62% ee; and the unknown species (\(\delta_p +13.6, -19.1\)) 1,4-addition in 90% ee. In general, each of the species 18 and 19 is rather fragile and small changes in the reaction conditions dramatically change their populations, as can be seen in Table 1.2.

The rate of conjugate addition of RMgBr to Michael acceptors does not completely correlate to the concentration of 18 (specifically, its higher ethyl homolog) and this points to the involvement of a second species (as in the uncharacterized \(\delta_p +13.6/-19.1\) entity of Scheme 1.9). The presence of two cuprates in these Grignard-based systems is supported by the chemistry of Loh [24] (Scheme 1.10).

Scheme 1.10  Primary organometallics in conjugate additions promoted by \((+)-2,2'\text{-bis(di-p-tolylphosphino)}-1,1'\text{-binaphthyl (Tol-BINAP) } L_D\).
Unfortunately, further structural data was not extractable from the $^{31}$P NMR experiments – the signals in Loh’s experiments are rather broad (such behavior is expected [25]) and no other nuclei were studied. On the basis of the (+)-ESI mass spectrum of the NMR sample, showing a cluster of isotopic peaks at around $m/z$ 1445, the two species were assigned to ion-pairing isomers of the formula $[\text{Cu}(\text{L}_D)_2][\text{MgMeBrI}]$. However, this may be an artifact of the electrospray MS technique as such a latter complex is predicted to show $m/z$ 1665 ($C_{97}H_{83}BrCuIMgP_4$) and the applicability of ESI-MS to sampling labile Mg/Cu organometallic systems has not been widely demonstrated. In contrast to the species of Feringa (18 and 19), it is clear that the catalytically competent primary organometallic must contain at least two L$_D$ as the system shows a nonlinear product ee dependence when the ee of L$_D$ is varied.

What the fate of the chiral-ligated cuprates in Scheme 1.9 and Scheme 1.10 is, after addition of suitable Michael acceptors, is unclear – no new compounds are detected in solution, which just begins generating enolate product. By analogy with Scheme 1.1 and Scheme 1.2, the presence of $\pi$-bound enone substrates and a transient Cu$^{III}$ intermediate has been postulated. From a practical point of view, it is worth noting that equivalent additions of sp$^2$ (vinyl/aryl) Grignard reagents to enones have proved much more challenging for asymmetric catalysis suggesting a greater lability. Systems using moderately polar 2-methyl-THF and ligand L$_E$ have proved the most effective (Scheme 1.11) but little is known about their intimate structure.

![Scheme 1.11](image)

Similarly, while effective methods for catalytic asymmetric S$_N$2’ allylation of RMgX reagents have recently become available [21], no attempts to reveal their exact primary structure through NMR has been successful thus far – again they appear highly labile. One clever approach to improving enantioselectivities is to build into the substrate a potential catalyst-directing group – an approach popularized by Breit and Schmidt [26]. However, using allylic dichloride 20 allows the product to retain a much more useful, and low mass, electrophilic site (Scheme 1.12) [27]. Coordination of one chloro group, either directly to Cu or via a magnesium bridge, orders the transition state derived from 21, simultaneously delivering high enantioselectivity and Z selectivity in 22. Falciola and Alexakis [28] have similarly proposed that a suitable (filled) ligand orbital stabilizes the Cu$^{III}$ transition state 24 in selective transformation to 25 using the (S,S,S) diastereomer of L$_E$ and other closely related ligands. The proposed interactions of Scheme 1.12 are closely akin to those discovered computationally in the cuprate 11 (Scheme 1.4).
The Primary Organometallic in Copper-Catalyzed Reactions

Scheme 1.12 Proposed direct (n → d) or MgX-bridged contact control in asymmetric S_N2′ allylation reactions.

1.6 Asymmetric Organozinc–Copper Reagents

Although publications on asymmetric copper-catalyzed additions of organozinc reagents have become legion over the last 20 years, relatively few of these explicitly set out to probe the structure of the active copper reagent. Of the primary zinc organometallics, diorganozinc (ZnR_2) reagents are by far the most useful; the highly sluggish reactivity of organozinc halides (RZnX) limits their asymmetric applications in copper chemistry despite the early success of "Knochel-type" reagents (RZnCuCNX) [29]. While RZnX can be turned into more reactive ZnR_2 via Lewis acid-promoted Schlenk processes (Equation 1 and Equation 2, Scheme 1.13) [30] the presence of these additional promoters can cause problems – as in the formation of (±)-26 during attempted asymmetric additions of BnZnBr. Nevertheless, such approaches can be moderately successful, as in BuZnCl minimization/activation in S_N2′ chemistry (Scheme 1.13) [31].

Information on the nature of the organocopper species formed after transmetalation with ZnR_2 is not helped by the paucity of kinetic and NMR solution studies of such systems. In a seminal study, Noyori used ReactIR to obtain high quality kinetic data for ZnEt_2 1,4-addition to enones promoted by very low levels of a 1 : 1
1.6 Asymmetric Organozinc–Copper Reagents

\[
\begin{align*}
RZnBr + MR & \rightleftharpoons ZnR_2 + MBr \\
2RZnBr + LA & \rightleftharpoons ZnR_2 + LA \cdot ZnBr_2
\end{align*}
\]

(1) (2)

\[
MR = \text{AlMe}_3, \text{ZnMe}_2 \\
LA = \text{Lewis acid promoter}
\]

\includegraphics[width=\textwidth]{scheme13}

Scheme 1.13 Promoted zinc Schlenk equilibria processes as routes to diorganozincs.

mix of PhSO₂NHBN and CuX (X = OTf, CN) [32]. This kinetic order dependence in this system is \([\text{Cu}+\text{L}]^1[\text{ZnEt}_2]^1[\text{enone}]^1\) and the reaction was modeled by a rate law involving rapid preassociation of the components followed by rate-limiting conversion of an “activated complex” 27. Unfortunately, supporting NMR data for 27 could not be attained. The unknown CuR₃Zn motif in 27 was assumed to be required to form a cuprate of sufficient reactivity for 1,4-insertion. Such analyses exclude any bridging ligand roles for triflate/cyanide – which could avoid the need for the unprecedented CuR₃Zn “higher order” cuprate in 27. For example, structure 28 could equally be in accord with the kinetic data. In an asymmetric version of this chemistry (via modification of the sulfonamide ligand), Piarulli [33] determined the rate dependence of the components to be \([\text{Cu}+\text{L}]^1[\text{ZnEt}_2]^0[\text{enone}]^1\), implying rapid transmetallation in this system. For more typical chiral PY₃ donors (phosphoramidites, phosphites), the exact composition of the organometallic resulting from mixtures of enone, ZnR₂, CuX, and used in asymmetric 1,4-additions is also contentious. Accurate kinetic data for these systems is notoriously hard to attain and the only published study fits almost equally well first- and second-order analyses of \([\text{Cu}_{\text{cat}}]\) and \([\text{ZnEt}_2]\) [5e, 34]. Direct NMR data of the “loaded” π-enone complexes is hard to attain owing to their high reactivity and the fast ligand exchange of Cu²⁺ d¹⁰ centers. In the absence of direct evidence, proposals 29–31 have arisen (Scheme 1.14). Intermediate 29 has its origins in the original suggestion of Noyori that (fast) kinetic association of a second molecule of ZnR₂ is necessary to provide a cuprate of sufficient reactivity. Complex 31 has some supporting evidence based on diffusion-ordered NMR spectroscopy (DOSY) studies of the precatalytic mixtures of CuBr and LG [35]. As the molecular volume of the complex is ligand dominated, the measured diffusion coefficients correlate to the number of ligands present. At 1 : 1 CuBr:L₆ ratios, (CuBrL₆)₃ of unknown structure is present. At greater relative ligand concentrations (typically the Cu:L is 1 : > 1.5 in many asymmetric reactions), the tetrahedral–trigonal dimer Cu₂Br₂(L₆)₃ is formed (cf. 31, Scheme 1.14) which
Scheme 1.14 Mechanistic proposals for active cuprate structure in conjugate addition of ZnR₂; L* is a generalized PY₃ donor, of which L_C is just one specific example.

is in equilibrium with other species via ligand association processes. The cleanest speciation was observed in CDCl₃, in more typical conjugate addition solvents (toluene, THF), a more complex mixture of exchanging complexes was formed. However, these DOSY studies have not yet been extended to real catalytic mixtures containing organometallics.

In the absence of direct experimental observation of enone π-complexes in this area, DFT computational studies have been carried out, which lend support to proposal 31. A computationally less expensive PBE1PBE hybrid functional approach allows rapid in silico screening of the accessibility of 29–31, among others, and this indicates that the latter is highly favored – much higher energy barriers to the formation of other species are encountered. In particular, computational attempts to attain the association of additional ZnMe₂ (i.e., motif 29) failed – only facile dissociation of the zinc resulted. The reaction coordinate of the addition of ZnMe₂ to acetal-protected 32 was studied in detail and a viable transition state for the insertion of Cu–Me into the π-complex 33 was detected, whose calculated energy barrier was close to the experimentally observed value (Scheme 1.15) [36]. Interestingly, a closely related geometry has been calculated recently for (admittedly Cu-free) addition of CH₂(ZnCl)₂ to acrolein [37]. In copper-catalyzed asymmetric 1,4-additions of ZnR₂, most interest has focused on the screening of libraries of chiral ligands, of which huge ranges now exist [38]. However, owing to the bimetallic nature of the active catalyst, the nature of the bridging ligand may be at least as important to attaining high enantioselectivities, even though it is itself normally achiral. Evidence for this can be seen in the large range of ee values attained from
1.6 Asymmetric Organozinc–Copper Reagents

Scheme 1.15 A calculated π–complex in the 1,4-addition of ZnMe₂ to diene 32.

the same chiral ligand as a function of CuX precursor in ZnEt₂ additions in typical experimental optimizations [39]. Such effects become increasingly apparent when the potential bridging ligands are “built into” the fabric of the chiral ligand. Two instructive recent examples are the extended carboxylate in \( \text{L}_H \) [40], which allows stereocompelled 1,6-additions (affording 34); and \( \text{L}_I \), which allows enantiofacial reversal depending on the copper precursor used [41] (Scheme 1.16). While the structures of the key transition states in these reactions are not known, they clearly point to effects related to the structures of Scheme 1.5.

<table>
<thead>
<tr>
<th></th>
<th>( R^1 )</th>
<th>( R^2 )</th>
<th>( R^3 )</th>
<th>Yield (%)</th>
<th>ee (34) (%)</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>Me</td>
<td>H</td>
<td>( n\text{Bu} )</td>
<td>79</td>
<td>97</td>
</tr>
<tr>
<td>2</td>
<td>Me</td>
<td>H</td>
<td>( i\text{Pr} )</td>
<td>71</td>
<td>97</td>
</tr>
<tr>
<td>3</td>
<td>( n\text{Bu} )</td>
<td>H</td>
<td>Et</td>
<td>63</td>
<td>95</td>
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<td>5</td>
<td>Me</td>
<td>Me</td>
<td>( n\text{Bu} )</td>
<td>75</td>
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<tr>
<td>6</td>
<td>Me</td>
<td>Me</td>
<td>( i\text{Pr} )</td>
<td>84</td>
<td>99</td>
</tr>
</tbody>
</table>

Scheme 1.16 Internal bridging ligand outcomes in ligands \( \text{L}_H \) and \( \text{L}_I \).
Direct experimental information on primary transmetallation events in $S_N2'$ allylic substitution is sparse – mechanistic speculation being the norm. Through a series of chiral ligand modifications, Hoveyda was able to argue that reactive intermediates 35a,b are responsible for the high enantioselectivity observed in $\gamma$-selective reaction of ZnEt$_2$ of allylic phosphates with his dipeptide-based ligands (Scheme 1.17). It is the terminal amide that plays the important organizational role – changing R from H to CH$_2$C(O)O$_t$Bu in the initial amino acid has only a minor effect. The requirement for pseudohalide-to-copper bridging is not so clear cut in $N$-heterocyclic carbene (NHC)-based catalysts, while this has been proposed for noncopper systems [42]. Direct stoichiometric reaction of 36a with enantiopure propargylic phosphates affords allenes with almost perfect chirality transfer to 37 [43]. Two points are of note: (i) racemization of the product allene 37 is minimal and (ii) in the published structures of isolated versions of such complexes, no indications of additional binding to the copper have been seen, the geometry being perfectly linear [44].

**Scheme 1.17** Stereochemical observations in $S_N2'$ allylations with peptidic and NHC-based ligands.

1.7 Asymmetric Organoboron–Copper Reagents

The widespread commercial availability of the pinacol-based diboron reagent (pin)B–B(pin) (pin = pinacolato = OMe$_2$CMe$_2$O) since about 2005, has fostered
a meteoric rise in interest in asymmetric borylation due, in part, to the known utility of B–C bonds for further manipulation with retention of stereochemistry (e.g., oxidation and cross-coupling applications) [45]. The high strength of the B–O bond (Table 1.1) predicts that additions to carbonyls, Michael acceptors, and allylic electrophiles with oxy-leaving groups should all be favored, if electronenegativity issues can be overcome in the transmetallation to copper (see Table 1.1). One successful strategy is the use of NHC ligands on copper, to obtain strongly stabilized Cu–B species, such as 38 (Scheme 1.18). These show remarkable stoichiometric transformations to stable (and, in many cases, crystallographically characterized) products, which, in many cases, can be rendered catalytic [46]. While this area is still in its infancy, asymmetric versions have been developed using chiral NHC ligands such as $L_j$ that allows access to the equivalent Cu–B motifs (Scheme 1.18).

Scheme 1.18 Reactive entities in borylation reactions and associated catalytic processes.

The requirement for additional solvolytic alcohols or Lewis acids to attain catalytic turnover is a common feature of these processes – most notably in the work of Yun (use of ROH) and Shibasaki (use of in situ LiX) [45]. Similar approaches have been demonstrated [47]. DFT calculations provide insights on the need for these additives by indicating that the borylation of $\alpha,\beta$-unsaturated substrates proceeds via
borocupration, rather than direct 1,4-addition, affording $\text{39}$ as the kinetically favored product due to the high $\text{B–C}$ bond strength (about 85 kcal mol$^{-1}$) (Scheme 1.19) [48]. Conversion of the C-enolate $\text{39}$ is rapid if $\text{Y} = \text{aryl, H but slow if Y = OR}$. Rapid protonolysis of $\text{39}$ is given credence by the high deuterium incorporation observed in related experiments using MeOD [49] and by very recent studies demonstrating association of ROH with $\text{39}$ by ESI-MS [50]. The Lewis acid promoters are thought to coordinate the carbonyl function of $\text{39}$ activating it to transmetallation – but no theoretical study of this process has been made. The calculations of Scheme 1.19 were carried out with $\text{L} = \text{PMe}_3$ and these have not been repeated with the typical chiral ligands (e.g., $\text{L}_\text{C}$) used in asymmetric catalysis. While computationally more expensive, such an approach would be useful in identifying potential key stereo-defining substrate–catalyst interactions.

Applications of (pin)$\text{B–B(pin)}$ to asymmetric $\text{S}_\text{N}2^\prime$ allylation are also attractive as these directly provide allylboranes – classic stalwarts of asymmetric synthesis. Again, chiral diphosphines and NHC complexes have proved the most attractive promoters and both of these also provide clues into the nature of the primary organometallics. Recently, unprecedented stereoconvergent transformations of both acyclic and cyclic allylic ethers have been demonstrated (Scheme 1.20) [51]. The stereoconvergence of these reactions to single enantiomers with very high ee values is hard to explain by classic copper(III) $\pi$-allyl chemistry (cf. Scheme 1.4) that is expected to lead to observation of some $\alpha$-coupled products, which is not the case. In light of the observations of Scheme 1.19, one alternative explanation is cuproboration via $\text{40}$ and $\text{42}$ leading to a transient $\sigma$-copper alkyl that then expels (pin)BOR on attack of the external diboron species. Such a mechanism would lead the selectivity to arise only through complexation of one $\pi$-face of the alkene avoiding the need for a range of different diastereomeric transition states to act kinetically very similarly. Finally, it is interesting to speculate if the features of
1.8 Asymmetric Organoaluminium–Copper Reagents

Although alanes, AlR₃, have become recognized as some of the most effective terminal organometallics for asymmetric conjugate addition and S_N2' allylation strategies [53], information on the intimate structure of the primary organocopper species involved in these reactions is distinctly sparse. They are popularly assumed to be “mechanistically analogous” to additions of the equivalent diorganozinc reagents, yet there are significant differences – wildly differing enantiofacial selectivities; under near identical conditions to their zinc counterparts, the observation that the Cu⁰ source [Cu(MeCN)₄]BF₄ (which contains no strong “bridging ligand”) is often effective points to the “zinc-like” analogy for Cu/Al systems being a poor one. Early low level PM3 studies on the rather too labile sulfide L₄ has suggested structures 43 and 44 for the major π-enone precursor and conjugate addition pathway (Scheme 1.21) [54] but this ligand is unrepresentative of most of the “modern” generation.

Scheme 1.20  Stereoconvergent substrate-binding modes in S_N2’ borylation reactions.

Scheme 1.20 will extend into S_N2’ additions of ArB(OR)₂ to allylic substrates for which good, nonasymmetric, precedents already exist [52].

1.8 Asymmetric Organoaluminium–Copper Reagents
Some definitively characterized Cu–R species have been prepared via organoaluminium reagents (Scheme 1.22). Significantly, air-sensitive 45 has been characterized both in solution and crystallographically [45a, 55] and these are of clear relevance to organoalanine 1,4 and 2,2′ reactions catalyzed by chiral NHC complexes. The B3LYP/6-31G(d) calculated bond dissociation energy (BDE) of the Cu–Me in 45 is \( \sim 80 \text{ kcal mol}^{-1} \). The high oxophilicity of AlR₃ reagents can cause problems with ligand and stability. For example, phosphoramidites coordinate AlMe₃ in nonpolar toluene or CH₂Cl₂ leading to adducts of type 46 and rapid cleavage to 47 (Scheme 1.22) [56]. Related processes in NHC chemistry are known, as in the formation of 48, but this is slow (20 h) and less likely to compete in the presence of Cu(I) sources [42].
Finally, it is important to realize that, the role of bridging ligands in asymmetric cuprates, while recognized as important, is not very well understood. Thus, although simple examples of such motifs are well characterized – as in the crystal structures of XARDUR and XARFON (Scheme 1.23) extrapolation to predicted selective asymmetric intermediate, such as 49, is presently just mechanistic speculation and further computational and spectroscopic studies are sorely needed here.

Scheme 1.23  Bridging ligand roles; distances in angstrom.

1.9  Asymmetric Silane and Stannane Copper-Promoted Reagents

Organosilane reagents engage in a wide range of copper-catalyzed transformations of enones, allylic(pseudo)halides, and carbonyl compounds via primary organometallics of the types: LCuR, LCuH, and LCuSiR₃. While the majority of the asymmetric processes that have arisen use chiral phosphines to produce the desired induction, the intermediates in such reactions have proved too labile to allow study. Conversely, NHC species (although mostly achiral) allow insights into structure and reactivity. Scheme 1.24 shows crystallographically characterized examples derived from (NHC)CuX transmetallation with appropriate organosilanes [57]. Direct comparison of their Cu–Cα electron density via their ¹³C NMR spectra is unfortunately not possible as this was only provided for 52. The formation of the latter may be a good model for similar processes occurring in the catalytic enone reduction chemistry of Lipshutz and Krause [58] (Scheme 1.24). The presence of (n-octyl)SiF₃ is vital for effective catalysis, but how this aids transmetallation is presently unknown. Similarly, 50 directly enters into catalytic 1,2-additions to aldehydes and is therefore clearly related to imine 1,2 allylation, of which formation of 53 is a nice example [59].

Recent contributions have been made in understanding of asymmetric reductions of carbonyl compounds. These reactions are believed to involve initial formation of a chiral copper hydride that inserts into the C=O bond to deliver a product alkoxide
that then undergoes transmetallation to close the catalytic cycle (Scheme 1.25).
DFT modeling of this proposed pathway suggests that the rate determining step
for the process is transmetallation of copper alkoxide intermediate. The absence
of any nonlinear effect (NLE) in the product ee when scalemic mixtures of a chiral
ligand are used point strongly to mononuclear catalysis. However, kinetic analysis
of the reaction of acetophenone and Ph₂SiH₂, under CuCl/NaO\text{tBu}/1,1’-binaphthyl
(BINAP) catalysis yields the component dependency: [ketone]^{1}[silane]^{1}[CuL]^{0.5}. The
half dependence in catalyst concentration is believed to indicate that the resting
state of the catalyst is the copper dimer \textit{54} \textsuperscript{[60]}. Very recently an example (2011,
structure INIRIH) of such a species has been characterized where the bridging
hydrides were crystallographically located \textsuperscript{[61]}. The cycle of Scheme 1.25 is also in
accord with recent (2011) studies using \textit{C}_2-chiral NHC ligands \textsuperscript{[62]}. Two further
interesting facts are (i) the calculated energy barriers for the cleavage of the
initial mononuclear alkoxide are unexpectedly high and (ii) that test stoichiometric
reactions of LCuH species, in the absence of excess silane, fail. Kleeberg \textit{et al.} \textsuperscript{[63]}
has explicitly characterized \textit{55}, which results from (NHC)Cu-SiMe₂Ph addition to
TolCHO. This undergoes slow cleavage with (pin)B-SiMe₂Ph to give the equivalent
silyl product.
Until comparatively recently copper-catalyzed 1,4- and $S_N2'$ additions of Cu-SiR$_3$ have resisted attempts to render them efficient asymmetric processes. In essence the issue is that the generalized asymmetric copper(I) silyl must compete against ligand free silyl cuprates or other main group silyl anions for addition to the substrate (Scheme 1.26) [18k]. The latter species are easily formed in even slightly
“ionizing” reaction mixtures and their rates of addition are comparable (or even better!) than addition of most examples of 56. Successful solutions to this problem center around use of the reagents PhMe\(_2\)Si-SiMe\(_2\)Ph 57 and PhMe\(_2\)Si-B(pin) 58 to afford clean transmetallation to 56 with the formation of only covalent (non promoting) by-products. A simple model for 56 is available in crystallographically characterized SAKRUR, which was attained from the reaction of CuCl, PMe\(_3\), and Ph\(_3\)SiLi in early work [64]. Once the restrictions of Scheme 1.26 were realized, then ligand screening soon revealed a range of chiral phosphines and NHC ligands able to deliver good to excellent levels of enantioselectivity for conjugate 1,4-additions (for details see Ref. [18k]).

The situation is even more problematic in S\(_\text{N}2\)’ allylation chemistry. While transmetallation to copper from a variety of “R\(_3\)Si−” sources is facile, Oestreicher has shown that facile σ−π−σ interconversion occurs in allylcopper species derived from Zn(SiPhMe\(_2\))\(_2\), containing LiCl, leading to rapid erosion of and induced stereochemistry [65]. While use of reagent 58 allows effective “ligand free” systems for silylation of (E)-cinnamyl chloride type allylic electrophiles, the use of BINAP or Josiphos ligated equivalents gave only low conversions to racemic products [66].

Well characterized species involved in cuprostannation remain essentially unreported, apart from the (IPr)CuSnPh\(_3\) complex of Sadighi which shows an almost linear C\(_\text{NHC}−\)Cu−Sn motif with a Cu−Sn bond distance of 2.47 Å [67]. This species shows a strong tendency to react with EX electrophiles fashioning EPh and SnPh\(_2\). The relationship of such isolated complexes to tin-based enantioselective catalysis largely remains to be defined. For example, in the reaction of allylstannanes with 2-nitrosopyridine no transmetallation to copper(I) is proposed, the latter acting only as a Lewis acid [68].

1.10 Conclusions

Even within the short space of this overview, it is clearly seen that, while the intimate structures of the primary organometallics present in enantioselective cuprate reagents are becoming better defined, a very wide range of factors can perturb their constitutions. In particular, the simplest reaction parameter – the solvent, can have a profound effect in directing the chiral cuprate from simple ligated organocopper species to complex bridged structures containing a wide range of gegenion Lewis acids.\(^3\) Controlling such effects for effective asymmetric catalysis requires a mixture of “discovery” and “design” approaches [54].

\(^3\) The author is indebted to his coworkers past and present for input into this document and to Dr Ross Denton (University of Nottingham) for DFT calculations on CuXMe (X = O, S).
References


2. Most data from: (a) Woodward, S. (2002) *Tetrahedron*, 58, 1017–1050 For the generalised LCuR data see: structures ESOXAL and JAXSIK in the Cambridge Crystallographic database (Cu–Me distances); (b) Ziegler, T., Tschinke, V., and Becke, A. (1987) *J. Am. Chem. Soc.*, 109, 1351–1358 (Cu–Me bond energy). The bond enthalpies for CuYMe (Y = O gives 48.8 kcal mol$^{-1}$; Y = S gives 72.8 kcal mol$^{-1}$) were calculated by the same DFT methods used for the other metals.


21. For an overview of this area see: reference 18a.


References


45. For an overview see reference 18j and references therein.


