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Tuning steric and electronic effects in transition-metal β**diketiminate complexes**

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Abstract

β-Diketiminates are widely used supporting ligands for building a range of metal complexes with different oxidation states, structures, and reactivities. This Perspective summarizes the steric and electronic influences of ligand substituents on these complexes, with an eye toward informing the design of new complexes with optimized properties. The backbone and *N*-aryl substituents can give significant steric effects on structure, reactivity and selectivity of reactions. The electron density on the metal can be tuned by installation of electron withdrawing or donating groups on the β-diketiminate ligand as well. Examples are shown from throughout the transition metal series to demonstrate different types of effects attributable to systematic variation of β-diketiminate ligands.

Graphical Abstract

We summarize steric and electronic influences on structure, spectroscopy, and reactivity in transition metal β-diketiminate complexes.

Keywords

β-diketiminate; steric; electronic

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1. Introduction

The properties and reactions of metal complexes are highly dependent on the choice of supporting ligand, and this choice is one of the keys to successful coordination chemistry. Since its introduction in 1968,¹⁻³ the β -diketiminate (often called "nacnac" because of its addition of two nitrogen atoms to the common acac ligand) has gained great popularity as a supporting ligand. Unlike acetylacetonate (acac), the β-diketiminate ligand scaffold offers steric protection at the metal center through the choice of N-substituents; this makes βdiketiminates less labile and more suitable as spectator ligands. β-Diketiminate ligands are typically synthesized from condensation of a β-diketone and an amine, and chemists have only scratched the surface of the thousands of potential combinations.⁴

N-aryl β-diketiminate ligands have been most widely used, and they support a variety of metals in many oxidation states. Complexes of *N*-aryl β-diketiminates have shown great reactivity and selectivity for a variety of methodologies,^{4, 5} including polymerization and functionalization of alkenes and cross-coupling reactions. In addition, late transition metal βdiketiminate complexes have been used to build low coordinate metal centers, mimicking the active sites of metalloproteins.⁶⁻¹⁴ A vast number of ligand variations and different coordination modes have been reported, and some examples are shown in Figure 1.1. In this Perspective, the focus will be solely on complexes of the type shown in Figure 1.1 with *d* block transition metals in a η^2 binding mode. We summarize trends from systematic variations in these complexes with examples, though we make no claim that our coverage is complete. This Perspective is intended to serve as a guide to chemists who are interested in tuning the properties of β-diketiminate complexes to achieve their specific goals. We also refer the interested reader to another Perspective by Budzelaar which gives more depth on *N*-aryl β-diketiminate complexes of Ru, Os, Rh, Ir, Pd, and Pt.¹⁵

2. Nomenclature

In this Perspective, the ligand abbreviation $R^1L^{R2,R3}$ is used to specify the substituents on a β -diketiminate ligand. R1 refers to the substituent on the central backbone carbon (α-C), R2 refers to the substituents on the nitrogen-bearing carbon atoms (β-C), and R3 refers to the substituents on the *N*-aryl group. For the R3 aryl substituents *meta*- and *para*- substitutions of *N*-aryl are specified as *m*- and *p*-, respectively, while the common *ortho*-substituents are given without the *o*- abbreviation for convenience. Some other abbreviations can be found in Chart 1.1.

3. Steric effects on β**-diketiminates**

The steric demands of β-diketiminate ligands can be tuned by substitution of functional groups on the backbone (β-C) or the *N*-aryl substituents. Typical backbone (β-C) substituents are *tert*-butyl, phenyl, trifluoromethyl and methyl; unsubstituted (βdialdiminate) ligands are also known. Two approaches can be used to tune the sterics of the *N*-aryl groups: first, to change the size of *ortho*-substituents on the *N*-aryl; or second, to relocate the substituents from *ortho*- position to the *meta*- or *para*- position.

The modification of β-diketiminate steric hindrance can bring changes in the structure and reactivity. The structural differences include changes on the coordination number, bond angles and bond lengths, geometry and conformation of metal complexes. We highlight three types of reactivity differences: different structures of β-diketiminate complexes, different outcomes of stoichiometric reactions of β-diketiminate complexes, and different activity in catalytic reactions.

3.1. Steric effects on structural properties

Generally, using smaller substituents on the β-C and *N*-aryl, or relocation of the *N*-aryl substituents farther from the metal center, reduces the overall steric coverage of the metal coordination sphere. As a result, dimeric/polymeric metal complexes are more often formed with less sterically hindered β-diketiminate ligands. For example, comparisons with more hindered monomeric analogues were reported for $[LScCl_2]_n (L^{tBu,iPr}, ^{16}n=1; L^{Me,iPr}, ^{17}$ n=2), $[LSc(CH_3)_2]_n (L^{tBu,iPr}, ^{16}n=1; L^{Me,iPr}, ^{17}n=2)$, $[LFeCl]_n (L^{tBu,iPr}, ^{18}n=1;$ $L^{\text{Me,iPr}},^{19 \text{ Me}}L^{\text{Me,Me}},^{20}$ n=2), [LFeF]_n (L^{tBu,iPr}, n=1; L^{Me,iPr}, n=2),²¹ [LCoCl]_n (L^{tBu,iPr},²² $n=1; L^{Me,iPr},^{23} n=2$), $[LNiCl]_n (L^{tBu,iPr},^{22} n=1; L^{Me,iPr},^{24} L^{Me,Me},^{25} n=2)$, $[LNi(CO)]_n$, $(L^{tBu,iPr}, ^{26}L^{Me,iPr}, ^{27}n=1; L^{Me,Me}, ^{28}n=2), [L^{R,iPr}CuCl]_n (L^{Me,iPr}, ^{29}ClL^{Me,iPr}, ^{29}$ n=1; $\frac{\text{Ph}}{\text{L}}$ H,iPr, 30 $\frac{\text{Me}}{\text{L}}$ Me,Cl, 31 n=2), and $\frac{\text{L}}{\text{L}}$ (Leq(μ -OAc)]_n (L^{Me,iPr}, ³² n=1; L^{Me,H32} Cl_LMe,H₁33 n=2). The angle between the two β-diketiminate ligand planes in dimeric metal complexes is often influenced by the different substituents on the ligand (Table 3.1.1). However, there is no clear correlation between the substituent size and the angle, indicating that this angle is dependent on the bonding at the metal as well as steric interactions between the ligands on the two sides.

One trend that emerges is that higher coordination numbers can be achieved with smaller βdiketiminate supporting ligands. For example, more solvent molecules (THF, arene, etc.) and neutral ligands (CO, PPh₃, etc.) can be coordinated to a metal center with less sterically hindered β-diketiminate in LScCl₂(THF)_n (L^{tBu,iPr},¹⁶ n=0; L^{Me,iPr},⁴⁷ n = 1) LSc(CH₃)₂(THF)_n (L^{tBu,iPr},¹⁶ n=0; L^{Me,iPr},¹⁶ n = 1), LSc(Cl)(NHAr)(THF)_n L^{tBu,iPr},⁴⁸ n=0; L^{Me,iPr},⁴⁹ n = 1), [LSc(CH₃)(arene)_n]⁺ (L^{tBu,iPr},⁵⁰ n=0; L^{Me,iPr},⁵⁰ n = 1), $LTiCl₂(THF)_n(L^{tBu,iPr},⁵¹ L^{tBu,Me3},⁵² L^{Me,Tbt/Me3},⁵³ n=0; L^{Me,iPr},⁵⁴ n=1; L^{Me,H},⁵⁵ n=2),$ $\rm LVCl_2(THF)_n(L^{Me,iPr}, ^{52,~56}L^{Me,Et}, ^{34}L^{Me,Me3}, ^{34}L^{Ph,iPr}, ^{34}n=0; L^{Me,~H}, ^{55}n=2),$ [LCr(µ-Cl) $(Solvent)_{n}$]₂ (L^{tBu,iPr},³⁵ n=0; L^{Me,iPr},³⁶ L^{Me,Me},³⁷ n = 1; Solvent = THF, benzene), LFe(NHdipp)(THF)_n (L^{tBu,iPr},¹⁹ n=0; L^{Me,iPr},²⁸ n = 1), and LCu(PPh_{3)n} (^{Ph}L^{H,iPr},⁵⁷) L^{Me,Me},⁵⁸L^{Me,iPr},⁵⁹L^{Me,Me3},⁶⁰ n=1; ^{Ph}L^{H,Me},⁵⁷ L^{CF3,*m*-CF3_,61} n=2). Steric conflict between *N*-aryl substituents and metal can also push the metal center out of the β-diketiminate ligand plane in some metal complexes, especially for early transition metals (Table 3.1.2). However, exceptions can be found in $L^{R, Mes}$ TiCl₂,⁵² $L^{Me, RC}$ r(η ⁵-Cp),^{62, 63} L^{R,iPr}FeNNFeL,^{6, 41} [L^{Me,R}Ni(µ-Cl)]₂,^{24, 25} L^{Me,R}Cu(OAc),^{64, 65} [LCu(µ-OH)]₂,⁴⁴⁻⁴⁶ [LCu(μ -S)]₂,^{66, 67} and L^{R,iPr} Cu(CO).⁶⁸

When the backbone (β -C) substituent size increases (H < Me < CF₃ < tBu, Ph), the steric conflict between backbone (β-C) substituents and *N*-aryl groups escalates, pushing the *N*aryl rings closer to the metal and forcing them into a more rigid configuration. As a consequence of this "buttressing effect," the metal center often moves deeper into the β-

diketiminate binding pocket. This brings three changes to the structure: it typically increases the N-M-N bite angle, increases the C(aryl)-N-C(β) bond angle, and shortens the N-M bond length (see Table 3.1.3). Bulky substituents on the *N*-aryl may also affect the bonding to other ligands (see Table 3.1.4). Exceptions to this trend, however, are seen with $LTiCl₂$, 52 LZrCl₃,^{70, 87} [LCr(μ -Cl)]₂, and K₂[LFeNNFeL],^{6, 41} due to cation coordination or conformational changes at the metal center. The distances from the metal to the nondiketiminate co-ligand can also be affected by the backbone substituents (see ESI for details).

The choice of *N*-aryl substituent has a smaller influence on the bite angle, $C(\text{aryl})-N-C(\beta)$ bond angle and N-M bond length in most cases. However, changing *N*-aryl substituents can build up steric bulk above and below the N-M-N plane, which can significantly influence the distance from the metal to the other ligands. In general, more hindered *N*-aryl substituents lead to a longer M-L bonds (Table 3.1.4).

Other modifications of β-diketiminate ligands, including installation of functional groups on the backbone α-C, or on the *para*-position of the *N*-aryl substituents, have little influence on the core structural parameters of β-diketiminate metal complexes.

The geometry and conformation of metal complexes can also be changed with modification of the supporting β-diketiminate ligand. The zirconium center in $L^{Me,R}Zr(CH_2Ph)_3$ (R = *i*Pr, *p*-Me)⁹⁴ adopts a square pyramidal geometry with a crystallographic mirror plane passing through it. However, the relative orientation of the ligand planes shows differences (Figure 3.1.1). Without *ortho*-substitution on *N*-aryl, the β-diketiminate ligand plane in $L^{Me,pMeZr}(CH_2Ph)_3$ forms an angle of 67.7(3)° with the least squares plane defined by C(Bn)-C(Bn)-N-N. In contrast, the angle between the ligand planes in $L^{Me,Me}Zr(CH_2Ph)_3$ is only 7.0(3)°. Presumably, this difference is due to steric conflict between the benzyl and *N*aryl substituents. *N*-Aryloxy-β-diketiminate zirconium complexes also showed a different orientation depending on steric bulk (Scheme $3.1.1$).⁹⁵ Bridged aryloxides were observed with one *meta-t*Bu on the *N*-aryl, but the presence of a second *meta*-*t*Bu group gave steric conflict that resulted in the isolation of a $[LZrC1_2]_2$ dimer instead. In the same system, the L2Zr complexes also showed conformational differences where the bulkier ligand adopted a trigonal prismatic geometry (Figure 3.1.2).

The solution structure of the metal complex can be affected by different steric bulk as well. For example, two sets of peaks were observed in 1 H NMR and 125 Te NMR spectra of L^{tBu,iPr}Sc(TeCH₂TMS)₂,⁹⁶ suggesting *exo* and *endo* tellurolates that are static on the NMR time scale. In contrast, the two tellurolate groups are equivalent for L^{Me,iPr}Sc(TeCH₂TMS)₂,⁹⁶ indicating rapid *endo/exo* flipping. Thus, larger groups create more difficulty for Sc(TeR)₂ to flip through the channel restricted by the *N*-aryl groups. In another example, 1H NMR peaks of a molybdenum imido alkylidene supported by LMe,*m*-Me was broadened compared with that of its $L^{Me,Me}$ analogue, suggesting the relatively free rotation of *N*-aryl in the less sterically hindered *meta*-substituted ligand.

3.2. Steric effects on reactivity and product formation

Here, we highlight other cases where different choices of steric bulk of the supporting βdiketiminate ligand give structurally different products under the same reaction conditions. In general, bulkier groups restrict the available conformations. For example, treatment of L^{tBu,iPr}ScCl₂ or [L^{Me,iPr}ScCl(µ-Cl)]₂ with LiNHtBu in hexanes generated different products (Scheme 3.2.1).^{48, 49} The authors proposed that the less sterically hindered $L^{Me,iPr}$ allows the formation of a dimeric transition state that is necessary for ligand exchange and disproportionation.

Extrusion of Te(CH₂TMS)₂from L^{R,iPr}Sc(TeCH₂TMS)₂ (R = *t*Bu, Me) under photolysis formed different products depending on R (Scheme 3.2.2).⁹⁶ Crossover between $(LSc(TeCH₂SiMe₃)$ ₂ and $LSc(TeCH₂CH₂)$ ₂)₂ showed that the product came from a bimolecular process. It is likely that the tellurolate-telluride (LSc(TeCH₂TMS))₂(μ-Te) is an intermediate on the way to the bridging telluride complex. However, the greater steric bulk of $L^{tBu,iPr}$ stabilized the tellurolate-telluride species, preventing the loss of a second molecule of $Te(CH_2TMS)_2$.

Reduction of $L^{Me,R}VCl_2$ (R = Me, Et, anthracenyl) with 2 equivalents of KC₈ in THF gave dimeric vanadium(I) complexes, while reaction of $L^{Ph,iPr}VCl_2$ gave extrusion of the imido fragment from diketiminate under the same conditions (Scheme $3.2.3$).³⁴ This was not only from having an available arene for binding, because reduction of $L^{Me,iPr}VCl_2$ in toluene gave an inverted sandwich complex. Rather, the authors surmised that the steric conflict between *N*-aryl and backbone phenyl group twisted the *N*-aryl group, destabilizing the LV intermediate and bringing about the reductive C-N bond cleavage of the ligand.

In another example, oxidation of a chromium(II) complex gave a highly reactive chromium oxo complex. However, the attempt to generate a chromium oxo complex gave different products depending on the steric bulk of different β-diketiminate ligands (Scheme 3.2.4).³⁸ Reaction of LMe,MeCrCp or LMe,*m*-TIPPCrCp with pyridine *N*-oxide gave a μ-oxo dimer, while the bulkier $L^{Me,Et}$ Cr-Cp generated a product from hydrogen atom transfer. The sterically more hindered *ortho*-ethyl substituents may prevent the μ-oxo dimer from forming, and rather the highly reactive terminal α (L^{Me,Et}(Cp)Cr=O) can abstract a hydrogen atom from its own ligand, ultimately generating a new C-C bond.

Upon addition of O_2 , copper(I) complexes supported by different β-diketiminate ligands form different products (Scheme 3.2.5). More sterically hindered $L^{tBu,iPr}Cu(NCCH_3)$ and $L^{Me,iPr}Cu(NCCH_3)$ formed a copper(II) peroxo $LCu(O_2)$ while less bulky $R^2L^{H,R}Cu$ (R = iPr, Me, Et; R' = H, Ph) complexes gave a bis(μ -oxo)dicopper(III) complex.^{8, 30} These reactivity differences between the two systems were attributed to the steric effect of the backbone (β-C) substituents, which rigidify the *N*-aryl substituents and prevent the dimer from forming.

The dinitrogen ligand in $L^{R,iPr}FeNNFeL^{R,iPr}$ ($R = tBu$, Me) can be replaced by other neutral ligands like carbon monoxide or isocyanide. 41 When exposing with excess CO, $L^{Me,iPr}$ FeNNFeL converted to square pyramidal $L^{Me,iPr}$ Fe(CO)₃, while the $L^{tBu,iPr}$ analogue gave a mixture of $L^{tBu,iPr}Fe(CO)$ ₃ and $L^{tBu,iPr}Fe(CO)$ ₂. Since the two *N*-dipp substituents

are closer in L^{tBu,iPr}, binding the third axial CO may bring steric tension between *i*Pr and CO, which explains the formation of square planar $L^{tBu,iPr}Fe(CO)_{2}$. Similarly, N₂ exchange in $L^{R,iPr}$ FeNNFe $L^{R,iPr}$ is much more rapid with R=Me than R=^tBu, implying that transient species with axial N₂ are also accessible but only with the smaller $R = Me⁴¹$ In a more deep-seated difference in reactivity, attempts to make analogous MeLMe,MeFeNNFe^{MeLMe,Me} complexes gave N₂ cleavage to a tetra-iron bis(nitride) complex, with complete cleavage of the N-N bond (Scheme $3.2.6$).²⁰ The authors proposed that the smaller supporting ligand allows access to an intermediate in which three LFe units can interact simultaneously with the same molecule of N_2 .

3.3. Steric effect on activity of metal complexes

Varying the steric bulk of the β-diketiminate ligand has a significant effect on activity of metal complexes in both stoichiometric and catalytic reactions. In most cases, a more sterically hindered β-diketiminate ligand builds up steric tension in transition states or intermediates, which raises the activation barrier and slows the reaction rates. However, the added steric bulk has advantages because it can enable the isolation of transient intermediates.

The single-electron oxidative addition of organic halides to chromium(II) complexes (Scheme 3.3.1) illustrated the steric effect of *ortho*-substituents on the *N*-aryl group.62, 72, 97 The less hindered asymmetric $L^{Me,iPr/p \to Y}Cr(Cp)$ gave a rate constant of 0.5-1.0 M⁻¹s⁻¹ (depending on the electronic properties of Y; see section 4.2 below), 97 whereas $L^{Me,iPr}Cr(Cp)$ and its $L^{Me,Me}$, $L^{Me,Me}$, and $L^{Me,Et}$ analogues gave rate constants that were more than an order of magnitude smaller, ranging from 0.02-0.03 $M^{-1}s^{-1}$.⁷² Thus, removing the *ortho*-alkyl groups from one of the *N*-aryl groups greatly enhanced the reactivity of chromium(II) by increasing the accessibility of methyl iodide.

Catalytic 1-hexene isomerization and dimerization was reported with $[L^{Me,R}NiBr]_2$ (R = *i*Pr, Me), where the less sterically hindered $[L^{Me,Me}NiBr]_2$ gave higher conversions under the same conditions.⁹⁸ The authors proposed that a β-diketiminate nickel hydride complex was the active catalyst, which would proceed through insertion, β-hydride elimination and chain walking to generate internal alkenes. This makes sense if β -hydride elimination is the ratelimiting step, because larger β-diketiminate substituents would prevent the increase in coordination number. In a demonstration of this idea in a stoichiometric reaction, L^{tBu,iPr}Fe*t*Bu isomerized to $L^{tBu,iPr}Fe-CH_2iBu$ only at elevated temperatures, while $L^{Me,iPr}Fe-*tBu*$ isomerized at room temperature to $L^{Me,iPr}$ Fe-CH₂*i*Bu (Scheme 3.3.2).⁷⁶

The mechanism of alkyne insertion was also studied in detail with isolated β-diketiminate iron hydride complexes. The rate of alkyne insertion was first order in [FeH] and zero order in [alkyne], with $k_{obs} = 1.7(2) \times 10^{-3}$ s⁻¹ for [L^{Me,iPr}FeH]₂⁹⁹ and 5.0(5) × 10⁻⁴ s⁻¹ for $[L^{tBu,iPr}FeH]_2$ ⁴⁰ again the less hindered complex had higher reactivity. In a related B-C bond cleavage reaction, two mechanisms were proposed: the less hindered iron complex undergoes single iron-hydride opening followed by insertion, while the more hindered $L^{tBu,iPr}$ system can completely dissociate to a reactive monomer.⁷⁴

β-Diketiminate iron imido complexes are prone to hydrogen atom transfer (HAT) from the *ortho* isopropyl substituents of the supporting ligand. To solve the problem, $L^{Me, Ph3}Fe=NR$ was prepared.¹⁰⁰ The second-order rate constants for hydrogen atom transfer to LFe=NAd from 1,4-cyclohexadiene in C₆D₆ were 2.0(2) × 10⁻² M⁻¹ s⁻¹ for L^{Me,Ph3}Fe=NAd, 1.4(2) × 10−4 M−1 s−1 for LMe,iPrFe=NAd and ~0 for LtBu,iPrFe=NAd (Scheme 3.3.3). Clearly the most bulky L^{tBu,iPr}Fe=NAd gave the slowest HAT reactivity. However, the relative sizes of LMe,iPr and LMe,Ph3 were not obvious. The authors measured the size using the *G* parameter, which estimates the fraction of the metal overshadowed by the ligand.¹⁰¹ The results indicated very similar *G* parameter for $L^{Me,iPr}$ Fe=NAd (G = 63.8%) over $L^{Me,Ph3}$ Fe=NAd (G = 62.2%), but different shapes (Figure 3.3.1). The different orientation of *N*-aryl with respect to the ligand backbone shows more opening above the imido nitrogen, which results in a larger binding pocket for hydrocarbon substrates (Figure 3.3.2).

Increasing the steric bulk of the β-diketiminate can also prevent formation of certain metal complexes due to steric blocking. In an example, β-diketiminate zirconium tribenzyl complex $(L^{Me,p-Me}Zr(CH_2Ph)_3)$ can be synthesized through alkane elimination between tetra-alkyl zirconium (IV) and β-diketimines. For its bulkier analogue $L^{Me, iPr}Zr(CH_2Ph)_3$, sterically hindered *i*Pr groups prevent $Zr(CH_2Ph)_4$ from accessing the β -diketiminate binding pocket. Therefore, it was necessary to develop a different synthetic method for $L^{Me,iPr}Zr(CH_2Ph)$ ₃ involving salt metathesis of LLi and $ZrCl_4$ followed by alkylation (Scheme 3.3.4).⁹⁴ In another example, $L^{Me,iPr}$ FeNNFe $L^{Me,iPr}$ releases the labile dinitrogen ligand immediately in aromatic solvents forming $L^{Me,iPr}Fe(\eta^6-C_6H_6)$. However, the more sterically hindered L^{tBu,iPr}FeNNFeL^{tBu,iPr} retains its structure in C_6H_6 up to 100 °C, without coordination of benzene.⁴¹

However, more sterically hindered metal complexes are favored in some cases because a sterically crowded environment can facilitate intramolecular reactions or increase the concentration of key unsaturated species. An example comes in reactions where metalation of ligand C-H bonds involves intramolecular C-H insertion. Upon heating in aromatic solvent, the four-coordinate dialkyl complexes $L^{R,iPr}ScR'_{2}(R = tBu, Me; R' = alkyl)$ (Scheme 3.3.5) underwent C-H metalation and eliminated alkane. The half-life of $L^{Me,iPr}$ ScR₂ in metalation was significantly longer than its $L^{tBu,iPr}$ analogue, suggesting lower reactivity with the less sterically hindered metal complex.¹⁶

 $L^{R,R'}$ NiBr, $L^{R,R'}$ NiPh(PPh₃) and $L^{Me,R'}$ Ni(alkyl) (R = CF₃, Me; R' = *i*Pr, Me) were reported to be active catalysts for ethylene, 102 , 103 styrene, 104 norbornene 105 , 106 polymerization and their copolymerization.^{107, 108} The polymer yield was significantly higher with more hindered ligand systems. Presumably, alkyl insertion into coordinated alkene is greatly facilitated by the more sterically hindered coordination environment.¹⁰⁵

Reductive elimination is another process facilitated by a crowded coordination environment. With a β-diketiminate-supported Pd(II) methyl phosphine complex, catalytic Castro-Stephens coupling, 109 Stille coupling 110 and Hiyama coupling 111 were more rapid with a more sterically hindered β-diketiminate ligand ($L^{Me,Me}$ vs. $L^{Me,H}$) which gave faster reductive elimination.

In addition, homolysis is influenced by ligand size. Since chromium(III) alkyl mediated radical polymerization often involves homolysis of the Cr-C bond to gain chain growth, more sterically hindered β-diketiminate ligand increases the Cr-C bond distances (see Table 3.1.4), giving a lower BDE, and increasing the rate of homolysis and thus rate of polymerization.112, 113

Catalytic carbodiimide formation from isocyanide and organic azide with a diketiminateiron(I) catalyst gave significantly higher yields with a more sterically bulky catalyst (L^{tBu,iPr} $>L^{Me, Ph3} > L^{Me, iPr}$). The proposed mechanism involves loss of one molecule of coordinated isocyanide before turning over the catalytic cycle. Not surprisingly, more hindered complexes favor a lower coordination number, which facilitates the loss of isocyanide, production of an active site, and turnover of the catalytic reaction.¹¹⁴

LCrCp catalyzed oxygen atom transfer reaction³⁸ (eq 3.3.1) and LCu(2-methylpyridine)catalyzed alkene azirdination¹¹⁵ (Scheme 3.3.6) are also more rapid with more hindered complexes because the smaller catalysts have more rapid rates for corresponding side reactions. Upon formation of catalytically active $[LCr=O]$ intermediate, $L^{Me,Me}Cr$ -Cp generates $L^{Me,Me}Cr(Cp)(\mu-O)Cr(Cp)L^{Me,Me}$ which is inactive towards catalytic oxygen atom transfer from O_2 to PPh₃. In contrast, more hindered L^{Me,Et}Cr(Cp)=O is less reactive towards formation of the μ-oxo complex and more catalytically active. Under catalytic aziridination conditions, smaller $L^{Me,Me}Cu(2-methylpyridine)$ underwent a side reaction generating TsNH₂, which lowered the reactivity and yield of aziridination compared with $L^{Me,Me/IPr}Cu(2-methylpyridine).$

Ethylene polymerization with L_2Ticl_2 complexes supported by different ligands have been studied. $L^{Me,iPr}2TiCl_2$ amd $L^{CF3,iPr}2TiCl_2$ showed significantly higher activity than their corresponding LMe,Me,LMe,H and LCF3,Me analogues. In this case, it is possible that bulky *N*aryl substituents can prohibit β-hydride elimination and thus maintain chain growth.¹¹⁶ In contrast, LTiMe₂ showed a different steric effect, where the less hindered $L^{Me,Me3}$ TiMe₂ was an order of magnitude more reactive than its more hindered $L^{tBu,Me3}$ TiMe₂ and $L^{Me,iPr}$ Me₂ analogues.⁵²

The steric effect for C-P cross-coupling catalyzed by LCrCp complex is another interesting example, because the influence is different depending on the relative rate of oxidative addition and Cr-C homolysis.117 For more reactive alkyl bromide substrates, more hindered $L^{Me,Me}$ CrCp or $L^{Me,Me}$ Cr(Cp)Br gave higher yields than less hindered asymmetric LMe,iPr/*p*-MeCrCp and LMe,iPr/*p*-MeCr(Cp)Br. Because these substrates undergo rapid single electron oxidative addition, the rate determining step is homolysis of the Cr-C bond. As previously mentioned, the Cr-C BDE is lower with more hindered ligands, so these ligands speed the catalytic rate. On the other hand, for less active substrates like Cy-Cl, oxidative addition is rate limiting, and the rate is faster with a less sterically hindered coordination environment.

3.4. Steric effects on selectivity of metal complexes

Changing steric bulk can also influence the selectivity of reactions of β-diketiminate complexes. This is due to the conformational differences in the energy of the intermediate/

As mentioned in section 3.3.3, changing the steric bulk can affect the reactivity of alkene polymerization and isomerization catalyzed by [LNiBr]2. Less bulky supporting ligands lead to more rapid β-hydride elimination, giving polyethylene with more branching. In alkene isomerization, the steric hindrance of the ligand can have important influences on the selectivity between *cis* and *trans* alkene products. More sterically hindered [L^{Me,iPr}NiBr]₂ gave more *cis* product (44%) compared with $[L^{Me,Me}NiBr]_2$ (28%).⁹⁸ It is believed that the crowded coordination environment restricted the rotation of C-C bond in Ni-alkyl complex, hindering the formation of *trans*-transition states. A bulkier L^{tBu,iPr}Co-alkyl complex isomerized alkenes with much higher *cis* selectivity, often greater than 6:1 *cis/trans*, but the LMe,iPrCo analogue gave poor selectivity. In this cobalt(II) system, the preference of the $L^{tBu,iPr}$ complex for isomerization of terminal alkenes to only the 2 position was also attributed to the bulk of the ligand above and below the N_2 Co plane.⁸⁸

4. Electronic effects on β**-diketiminate complexes**

To tune the electronic properties of β -diketiminate ligands, various groups have been installed on the backbone (α-C and β-C) or on the *N*-aryl substituents. These modify the electron density at the metal center, which can affect the redox potential, IR frequency of other ligands, UV-Vis absorption maxima, and NMR chemical shifts. In addition, these electronic changes can also affect the reactivity through perturbation of the energy of transition states or intermediates. It should be borne in mind that many of the substituents used to change the electronic effects can also influence sterics as well, particularly on the backbone (β-C) and *ortho* positions of *N*-aryl groups.

4.1. Electronic effects on electron density and core structure of the metal center

Changes in electron density on the metal center can be monitored by various methods. Often, electron-withdrawing groups lead to more positive redox potentials, lower field chemical shifts in NMR spectra, and less backbonding into coordinated ligands, consistent with less electron density at the metal ion.

Copper and nickel complexes supported by β-diketiminate ligands bearing different electronic properties have been studied with cyclic voltammetry (Table 4.1.1). Judging from the redox potentials in Table 4.1.1, $NO₂$ and $CF₃$ have the strongest electronic effect, followed by CN and 3,5-bis(trifluroromethyl)phenyl substituents. In addition, greater electronic effects result from substitutions on α-C and β-C, and less with *N*-aryl substituents. This is reasonable because the aryl ring is roughly perpendicular to the MN_2C_3 plane, and thus there is little conjugation of the π -systems. In contrast, backbone substituents are in the plane of the ligand backbone, and thus can have a greater impact on the electron density of

the metal center. The exception is the relatively small electronic effect from 3,5 bis(trifluoromethyl)phenyl substituents on the backbone (α-C), which is presumably again from lack of conjugation between the perpendicular π-systems. However, the electronic influence of *N*-aryl substituents is not negligible. For example, alkyl substituents on the *N*aryl behaved as electron-donating groups when ^{Ph}L^{H,iPr}-supported copper complexes had a more negative redox potential than $P^h L^{H, Me}$ and $P^h L^{H,Et}$ (Table 4.1.1).³⁰

Another consequence of the changing redox potentials is the relative stability of certain oxidation levels. In $\rm L_2Cu$ complexes, irreversible reductions were observed with $\rm {^{Me}L^{H,H}}$ and ${}^H L^{H,H}$ while reversible redox couples were observed in ${}^{CN} L^{H,H}$ and ${}^{NO2} L^{H,H}$, suggesting that the reduced Cu(I) state of the bis(β-diketiminate) complex is unstable in the complexes with more electron rich ligands. In contrast, with $LCu(NCCH_3)$ complexes, the $Cu(II)$ state was less stable with a more electron withdrawing group.⁴⁶ Ruthenium(II) complexes of $L^{CF3,m-CF3}Ru(Cl)(Ar)$ (Ar = arene ligand) were studied to determine the electronic effects of the supporting ligand on the metal and the other coordinating ligands in comparison to analogous complexes with the $L^{Me,m-Me}$ supporting ligand.⁸⁵ Interestingly, there was no clear trend between the Ru^{II}/Ru^{III} redox potentials from the cyclic voltammograms through the series L^{Me,Me}, L^{Me,m-Me}, L^{CF3,m-Me}, and L^{CF3,m-CF3}, indicating that other factors also play a role.⁸⁶

Electronic modification can also have an impact on the positions of the maxima in electronic absorption (UV-Vis) spectra. β-Diketiminate complexes typically have a $\pi \rightarrow \pi^*$ transition in the 300-400 nm region, which shifts to shorter wavelength with more electron-withdrawing substituents in $LCu(NCCH₃)$.³⁰ This suggests that electron-withdrawing groups lower the energy of the π orbital more than they do the π^* orbital. The positions of *d-d* transitions was also studied in L_2 Cu complexes, where the d-d absorption bands shift toward shorter wavelength with electron withdrawing backbone substituents (α-C) and shift to longer wavelength with more electron donating substituents on the *N*-aryl group.46 It is proposed that the ligand field was enhanced with electron donating substituents and thus affected the UV-Vis absorptions.

IR and Raman peaks on coordinated diatomic ligands is another traditional method for quantifying the relative electron density of a metal center. The ν(CO) in LCu(CO) complexes and $v(OO)$ in $LCu(O₂)$ each shift to higher frequency when electron withdrawing CF_3 groups were installed on the backbone β -C.⁶⁸ This is attributable to a less electron rich metal center that has weaker back-donation into ligand antibonding orbitals. The influence of *m*-CF₃ groups on the *N*-aryl substituents was less, again indicating a smaller influence from *N*-aryl substitution.

Due to the shielding or deshielding effect of substituents, the chemical shift in NMR spectra also indicates the electron density on metal center. For example, the chemical shift of the backbone (α -C) proton shifted downfield when CF₃ was substituted for CH₃ on backbone and for *meta*- positions on the *N*-aryl.⁸⁵ This is correlated to the deshielding effect with more electron withdrawing groups attached directly to the π system.

Though the introduction of electron withdrawing groups hardly affects the metal ligand core structure, it can affect the coordination number as well as bonding properties in some cases. For example, when $NO₂$ was installed on backbone (α -C) of LCu-OAc, one molecule of methanol coordinated to the metal center, but no coordinated methanol was observed with $\rm {CN_LH,iPr}$ and $\rm {Ph_LH,iPr}$. This is consistent with the stronger Lewis acidity of metal center when its supporting ligand has an electron withdrawing $NO₂$ substituent.⁹⁰ Ru-Cl bond lengths and Ru-arene distances in $LRu(Cl)(\eta^6$ -arene) are shorter with $L^{CF3,m-CF3}$ compared with L^{Me,*m*-Me}, suggesting an increase in Lewis acidity of the metal with more electronwithdrawing substitutents.⁸⁵

4.2. Electronic effects on reactivity of metal complex

Changes of electron density on the metal center can have a significant effect on reactivity of metal complexes. For example, the oxidative addition of methyl iodide to mixed-aryl LCrCp complexes (Scheme 3.3.1) is affected by electronic substituents on *para-N*-aryl (OMe, Me, H, CF_3).⁹⁷ There was a correlation between the *para*-substituent and the rate constant, with the rate constant decreasing two-fold from most electron-donating (*para*-OMe, k_{obs} =(9.80±0.3) × 10⁻¹ M⁻¹ s⁻¹) to most electron-withdrawing (*para*-CF₃, k_{obs} =(4.96±0.3) \times 10⁻¹ M⁻¹ s⁻¹) substituent. Even though the solid structures indicate that the *N*-aryl planes are aligned roughly perpendicular to the metal-ligand plane, the authors noted that the lack of *ortho*-substituents may allow the *N*-aryl to rotate closer to the diketiminate plane in solution, enabling some conjugation. In this way, the more electron-donating substituents can stabilize the chromium(III) product, which could lower the barrier if Hammond's postulate holds.

In another example, catalytic oxidation of alkanes to alcohols and ketones was reported with LCu(OAc) as a catalyst.⁹⁰ When LCu(OAc) was supported by a more electron-withdrawing β-diketiminate ligand, the catalytic reactivity was higher. The results were rationalized through a mechanistic model where the reactions proceed through a metal-based oxidant, based on the observed kinetic isotope effect and regioselectivity.120 Thus, more electron withdrawing groups would give more unstable and energetic high-valent copper intermediates that are more reactive toward the alkane.

Atom transfer radical addition (ATRA) and atom transfer radical cyclization (ATRC) are particularly interesting for organic synthesis. Using β-diketiminate ruthenium complexes (LRu(Cp*)Cl and LRu(Cp*)), lower conversions were observed with LMe,Me, LMe,*m*-Me, and LMe,*m*-CF3, while the addition of electron-withdrawing substituents in LCF3,*m*-Me and L^{CF3,*m*-CF3} gave higher reactivity.⁸⁶ No simple correlation between catalytic reactivity and redox potential of the ruthenium complexes was observed, but the addition of the CF_3 groups also rendered the complexes air-stable in solution and solid state. Likewise, in the copper(I) complexes mentioned above, $L^{Me,iPr}Cu(NCMe)$ and $L^{CF3/Me,iPr}Cu(NCMe)$ react with O_2 , but L^{CF3,iPr}Cu(NCMe) does not react with O_2 . This agrees with the more positive redox potential with an electron-withdrawing group.⁶⁸

The previously mentioned nickel catalyzed polymerization of styrene and norbornene (see section 3.3) showed a strong influence of the β-diketiminate ligand electronic properties. The substitution of backbone methyl with trifluoromethyl significantly improved the

catalytic reactivity.104, 105, 121 This can be explained if the more electrophilic nickel center has a lower activation energy for alkene insertion during rate-limiting chain growth.

5. Conclusions

The examples in this Perspective support the idea that β-diketiminate ligands have great tunability in terms of both steric and electronic effects, and they point future chemists in the directions that could benefit their own chemistry. The β-C and *N*-aryl ortho substituents are most important for steric effects, whereas the α-C and β-C positions are most influential for electronic effects. *N*-aryl groups can have a small electronic influence, but this has been best documented when there are no *ortho*-substituents and the *N*-aryl group can rotate closer to planarity with the ligand backbone. In contrast, the steric effects are more varied, because they can change the structure and transition states in different ways depending on the specific coordination number, reaction, and co-ligands. However, the ability of relatively small changes to cause structural, spectroscopic, and reactivity differences suggests that further tuning will uncover multitudes of new chemistry. We note particularly that chiral substituents have only been used in β-diketiminate ligands with *N*-benzyl substituents,¹²²⁻¹²⁵ and incorporation of chiral anilines should be a fruitful area for preparation of *C*1 and *C*² symmetric complexes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

6. Acknowledgments

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Biography

Chi Chen received his Bachelor of Science degree at Peking University in 2009 and did additional research at the University of Texas - Arlington before starting graduate research at the University of Rochester in 2011. In a joint project with Daniel Weix and Patrick Holland, he is developing and studying new β-diketiminate supported cobalt catalysts for alkene transformations such as isomerization and hydrosilylation. In 2013, he moved to Yale University where he is completing his PhD research.

Sarina Bellows received her Bachelor of Science degree from Syracuse University in 2008, and pursued PhD research at the University of Rochester with Patrick Holland. In her research, she synthesized iron complexes of new β-diketiminate ligands, and also performed computations to explain the mechanisms of their reactions. Since receiving her PhD in 2014, she has been a postdoctoral fellow at Rochester with Thomas Cundari and William Jones through the Center for Enabling New Technologies through Catalysis.

Patrick Holland completed an AB at Princeton University, and a PhD at UC Berkeley with Richard Andersen and Robert Bergman. In postdoctoral work at Minnesota with William Tolman, he learned to love β-diketiminates through the synthesis of copper complexes. In his independent career, he has explored the use of β-diketiminate complexes of iron, cobalt and nickel, as applied to N_2 reduction, C-H oxidation, redox-active ligands, new bonding environments, and novel reactivity. He was on the faculty at the University of Rochester from 2000-2013, and is now a Professor of Chemistry at Yale University.

References

- 1. McGeachin SG. Can. J. Chem. 1968; 46:1903–1912.
- 2. Bonnett R, Bradley DC, Fisher KJ. Chem. Commun. 1968:886–887.
- 3. Parks JE, Holm RH. Inorg. Chem. 1968; 7:1408–1416.
- 4. Bourget-Merle L, Lappert MF, Severn JR. Chem. Rev. 2002; 102:3031–3066. [PubMed: 12222981]
- 5. Tsai Y. Coord. Chem. Rev. 2012; 256:722–758.
- 6. Smith JM, Lachicotte RJ, Pittard KA, Cundari TR, Lukat-Rodgers G, Rodgers KR, Holland PL. J. Am. Chem. Soc. 2001; 123:9222–9223. [PubMed: 11552855]
- 7. Holland PL, Tolman WB. J. Am. Chem. Soc. 1999; 121:7270–7271.
- 8. Spencer DJE, Aboelella NW, Reynolds AM, Holland PL, Tolman WB. J. Am. Chem. Soc. 2002; 124:2108–2109. [PubMed: 11878952]
- 9. Aboelella NW, Lewis EA, Reynolds AM, Brennessel WW, Cramer CJ, Tolman WB. J. Am. Chem. Soc. 2002; 124:10660–10661. [PubMed: 12207513]
- 10. Aboelella NW, Gherman BF, Hill LMR, York JT, Holm N, Young VG, Cramer CJ, Tolman WB. J. Am. Chem. Soc. 2006; 128:3445–3458. [PubMed: 16522125]

- 11. Vela J, Stoian S, Flaschenriem CJ, Münck E, Holland PL. J. Am. Chem. Soc. 2004; 126:4522– 4523. [PubMed: 15070362]
- 12. Tonzetich ZJ, Do LH, Lippard SJ. J. Am. Chem. Soc. 2009; 131:7964–7965. [PubMed: 19459625]
- 13. Randall DW, George SD, Holland PL, Hedman B, Hodgson KO, Tolman WB, Solomon EI. J. Am. Chem. Soc. 2000; 122:11632–11648.
- 14. Brown EC, York JT, Antholine WE, Ruiz E, Alvarez S, Tolman WB. J. Am. Chem. Soc. 2005; 127:13752–13753. [PubMed: 16201771]
- 15. Zhu D, Budzelaar PHM. Dalton Trans. 2013; 42:11343–11354. [PubMed: 23787915]
- 16. Hayes PG, Piers WE, Lee LWM, Knight LK, Parvez M, Elsegood MRJ, Clegg W. Organometallics. 2001; 20:2533–2544.
- 17. Hayes PG, Piers WE, Parvez M. J. Am. Chem. Soc. 2003; 125:5622–5623. [PubMed: 12733887]
- 18. Smith JM, Lachicotte RJ, Holland PL. Chem. Commun. 2001:1542–1543. DOI: 10.1039/ B103635C.
- 19. Eckert NA, Smith JM, Lachicotte RJ, Holland PL. Inorg. Chem. 2004; 43:3306–3321. [PubMed: 15132641]
- 20. Rodriguez MM, Bill E, Brennessel WW, Holland PL. Science. 2011; 334:780–783. [PubMed: 22076372]
- 21. Vela J, Smith JM, Yu Y, Ketterer NA, Flaschenriem CJ, Lachicotte RJ, Holland PL. J. Am. Chem. Soc. 2005; 127:7857–7870. [PubMed: 15913376]
- 22. Holland PL, Cundari TR, Perez LL, Eckert NA, Lachicotte RJ. J. Am. Chem. Soc. 2002; 124:14416–14424. [PubMed: 12452717]
- 23. Wei Gao YM, Li Guang-hua, Liu Xiao-ming, Su Qing, Yao Wei. Chem. Res. Chin. Univ. 2005; 21:240.
- 24. Eckert NA, Bones EM, Lachicotte RJ, Holland PL. Inorg. Chem. 2003; 42:1720–1725. [PubMed: 12611544]
- 25. Wiencko HL, Kogut E, Warren TH. Inorg. Chim. Acta. 2003; 345:199–208.
- 26. Horn B, Pfirrmann S, Limberg C, Herwig C, Braun B, Mebs S, Metzinger R. Z. Anorg. Allg. Chem. 2011; 637:1169–1174.
- 27. Eckert NA, Dinescu A, Cundari TR, Holland PL. Inorg. Chem. 2005; 44:7702–7704. [PubMed: 16241116]
- 28. Wiese S, Aguila MJB, Kogut E, Warren TH. Organometallics. 2013; 32:2300–2308.
- 29. Jazdzewski BA, Holland PL, Pink M, Young VG, Spencer DJE, Tolman WB. Inorg. Chem. 2001; 40:6097–6107. [PubMed: 11703106]
- 30. Spencer DJE, Reynolds AM, Holland PL, Jazdzewski BA, Duboc-Toia C, Le Pape L, Yokota S, Tachi Y, Itoh S, Tolman WB. Inorg. Chem. 2002; 41:6307–6321. [PubMed: 12444774]
- 31. Wiese S, Badiei YM, Gephart RT, Mossin S, Varonka MS, Melzer MM, Meyer K, Cundari TR, Warren TH. Angew. Chem. Int. Ed. 2010; 49:8850–8855.
- 32. Hadzovic A, Song D. Inorg. Chem. 2008; 47:12010–12017. [PubMed: 19006292]
- 33. Hadzovic A, Song D. Organometallics. 2008; 27:1290–1298.
- 34. Chang K-C, Lu C-F, Wang P-Y, Lu D-Y, Chen H-Z, Kuo T-S, Tsai Y-C. Dalton Trans. 2011; 40:2324–2331. [PubMed: 21152635]
- 35. Fan H, Adhikari D, Saleh AA, Clark RL, Zuno-Cruz FJ, Sanchez Cabrera G, Huffman JC, Pink M, Mindiola DJ, Baik M-H. Journal of the American Chemical Society. 2008; 130:17351–17361. [PubMed: 19035634]
- 36. Gibson, Vernon C.; Newton, C.; Redshaw, C.; Solan, Gregory A.; White, Andrew J. P.; Williams, David J. Eur. J. Inorg. Chem. 2001; 2001:1895–1903.
- 37. Charbonneau F, Oguadinma PO, Schaper F. Inorg. Chim. Acta. 2010; 363:1779–1784.
- 38. MacLeod KC, Patrick BO, Smith KM. Inorg. Chem. 2012; 51:688–700. [PubMed: 22175660]
- 39. Sadique AR, Gregory EA, Brennessel WW, Holland PL. J. Am. Chem. Soc. 2007; 129:8112–8121. [PubMed: 17564444]
- 40. Smith JM, Lachicotte RJ, Holland PL. J. Am. Chem. Soc. 2003; 125:15752–15753. [PubMed: 14677959]

- 41. Smith JM, Sadique AR, Cundari TR, Rodgers KR, Lukat-Rodgers G, Lachicotte RJ, Flaschenriem CJ, Vela J, Holland PL. J. Am. Chem. Soc. 2006; 128:756–769. [PubMed: 16417365]
- 42. Yao S, Xiong Y, Milsmann C, Bill E, Pfirrmann S, Limberg C, Driess M. Chem. Eur. J. 2010; 16:436–439. [PubMed: 19937871]
- 43. Spencer DJE, Reynolds AM, Holland PL, Jazdzewski BA, Duboc-Toia C, Le Pape L, Yokota S, Tachi Y, Itoh S, Tolman WB. Inorganic Chemistry. 2002; 41:6307–6321. [PubMed: 12444774]
- 44. Hong S, Hill LMR, Gupta AK, Naab BD, Gilroy JB, Hicks RG, Cramer CJ, Tolman WB. Inorg. Chem. 2009; 48:4514–4523. [PubMed: 19425614]
- 45. Dai X, Warren TH. Chem. Commun. 2001:1998–1999. DOI: 10.1039/B105244F.
- 46. Shimokawa C, Yokota S, Tachi Y, Nishiwaki N, Ariga M, Itoh S. Inorg. Chem. 2003; 42:8395– 8405. [PubMed: 14658893]
- 47. Lee LWM, Piers WE, Elsegood MRJ, Clegg W, Parvez M. Organometallics. 1999; 18:2947–2949.
- 48. Knight LK, Piers WE, Fleurat-Lessard P, Parvez M, McDonald R. Organometallics. 2004; 23:2087–2094.
- 49. Basuli F, Tomaszewski J, Huffman JC, Mindiola DJ. Organometallics. 2003; 22:4705–4714.
- 50. Hayes PG, Piers WE, Parvez M. Chem. Eur. J. 2007; 13:2632–2640. [PubMed: 17171731]
- 51. Basuli F, Bailey BC, Watson LA, Tomaszewski J, Huffman JC, Mindiola DJ. Organometallics. 2005; 24:1886–1906.
- 52. Budzelaar PHM, van Oort AB, Orpen AG. Eur. J. Inorg. Chem. 1998; 1998:1485–1494.
- 53. Hamaki H, Takeda N, Tokitoh N. Organometallics. 2006; 25:2457–2464.
- 54. Basuli F, Bailey BC, Tomaszewski J, Huffman JC, Mindiola DJ. J. Am. Chem. Soc. 2003; 125:6052–6053. [PubMed: 12785824]
- 55. Kim W-K, Fevola MJ, Liable-Sands LM, Rheingold AL, Theopold KH. Organometallics. 1998; 17:4541–4543.
- 56. Tsai Y-C, Wang P-Y, Lin K-M, Chen S-A, Chen J-M. Chem. Commun. 2008:205–207. DOI: 10.1039/B711816C.
- 57. Li X, Ding J, Jin W, Cheng Y. Inorg. Chim. Acta. 2009; 362:233–237.
- 58. York JT, Young VG, Tolman WB. Inorg. Chem. 2006; 45:4191–4198. [PubMed: 16676981]
- 59. Reynolds AM, Lewis EA, Aboelella NW, Tolman WB. Chem. Commun. 2005:2014–2016. DOI: 10.1039/B418939F.
- 60. Badiei YM, Warren TH. J. Organomet. Chem. 2005; 690:5989–6000.
- 61. Carrera N, Savjani N, Simpson J, Hughes DL, Bochmann M. Dalton Trans. 2011; 40:1016–1019. [PubMed: 21152532]
- 62. Doherty JC, Ballem KHD, Patrick BO, Smith KM. Organometallics. 2004; 23:1487–1489.
- 63. Champouret Y, MacLeod KC, Baisch U, Patrick BO, Smith KM, Poli R. Organometallics. 2010; 29:167–176.
- 64. Inosako M, Kunishita A, Shimokawa C, Teraoka J, Kubo M, Ogura T, Sugimoto H, Itoh S. Dalton Trans. 2008:6250–6256. DOI: 10.1039/b808678h. [PubMed: 18985258]
- 65. Yokota S, Tachi Y, Itoh S. Inorg. Chem. 2002; 41:1342–1344. [PubMed: 11896697]
- 66. Brown EC, Aboelella NW, Reynolds AM, Aullón G, Alvarez S, Tolman WB. Inorg. Chem. 2004; 43:3335–3337. [PubMed: 15154793]
- 67. Brown EC, Bar-Nahum I, York JT, Aboelella NW, Tolman WB. Inorg. Chem. 2007; 46:486–496. [PubMed: 17279827]
- 68. Hill LMR, Gherman BF, Aboelella NW, Cramer CJ, Tolman WB. Dalton Trans. 2006:4944–4953. DOI: 10.1039/b609939d. [PubMed: 17047744]
- 69. Kenward AL, Ross JA, Piers WE, Parvez M. Organometallics. 2009; 28:3625–3628.
- 70. Basuli F, Kilgore UJ, Brown D, Huffman JC, Mindiola DJ. Organometallics. 2004; 23:6166–6175.
- 71. Kakaliou L, Scanlon WJ, Qian BX, Baek SW, Smith MR, Motry DH. Inorganic Chemistry. 1999; 38:5964–5977. [PubMed: 11671302]
- 72. MacLeod KC, Conway JL, Tang L, Smith JJ, Corcoran LD, Ballem KHD, Patrick BO, Smith KM. Organometallics. 2009; 28:6798–6806.

- 73. Huang Y-B, Jin G-X. Dalton Transactions. 2009:767–769. DOI: 10.1039/B820798B. [PubMed: 19156267]
- 74. Yu Y, Brennessel WW, Holland PL. Organometallics. 2007; 26:3217–3226. [PubMed: 18725998]
- 75. Cowley RE, Elhaïk J, Eckert NA, Brennessel WW, Bill E, Holland PL. J. Am. Chem. Soc. 2008; 130:6074–6075. [PubMed: 18419120]
- 76. Vela J, Vaddadi S, Cundari TR, Smith JM, Gregory EA, Lachicotte RJ, Flaschenriem CJ, Holland PL. Organometallics. 2004; 23:5226–5239.
- 77. Vela J, Smith JM, Lachicotte RJ, Holland PL. Chem. Commun. 2002:2886–2887. DOI: 10.1039/ B209389H.
- 78. Stoian SA, Yu Y, Smith JM, Holland PL, Bominaar EL, Munck E. Inorg. Chem. 2005; 44:4915– 4922. [PubMed: 15998018]
- 79. Yu Y, Smith JM, Flaschenriem CJ, Holland PL. Inorg. Chem. 2006; 45:5742–5751. [PubMed: 16841977]
- 80. Panda A, Stender M, Wright RJ, Olmstead MM, Klavins P, Power PP. Inorg. Chem. 2002; 41:3909–3916. [PubMed: 12132915]
- 81. Oguadinma PO, Schaper F. Inorg. Chim. Acta. 2009; 362:570–574.
- 82. Hill LMR, Gherman BF, Aboelella NW, Cramer CJ, Tolman WB. Dalton Transactions. 2006:4944–4953. DOI: 10.1039/B609939D. [PubMed: 17047744]
- 83. Phillips AD, Zava O, Scopelitti R, Nazarov AA, Dyson PJ. Organometallics. 2010; 29:417–427.
- 84. Phillips AD, Laurenczy G, Scopelliti R, Dyson PJ. Organometallics. 2007; 26:1120–1122.
- 85. Schreiber DF, Ortin Y, Müller-Bunz H, Phillips AD. Organometallics. 2011; 30:5381–5395.
- 86. Phillips AD, Thommes K, Scopelliti R, Gandolfi C, Albrecht M, Severin K, Schreiber DF, Dyson PJ. Organometallics. 2011; 30:6119–6132.
- 87. Kakaliou L, Scanlon, Qian B, Baek SW, Smith MR, Motry DH. Inorg. Chem. 1999; 38:5964– 5977. [PubMed: 11671302]
- 88. Chen C, Dugan TR, Brennessel WW, Weix DJ, Holland PL. J. Am. Chem. Soc. 2014; 136:945– 955. [PubMed: 24386941]
- 89. Young J, Yap GA, Theopold K. J. Chem. Crystallogr. 2009; 39:846–848.
- 90. Shimokawa C, Teraoka J, Tachi Y, Itoh S. J. Inorg. Biochem. 2006; 100:1118–1127. [PubMed: 16584781]
- 91. Bernoud E, Oulié P, Guillot R, Mellah M, Hannedouche J. Angew. Chem. Int. Ed. 2014; 53:4930– 4934.
- 92. Huang H, Hughes RP, Rheingold AL. Polyhedron. 2008; 27:734–738.
- 93. Annibale VT, Tan R, Janetzko J, Lund LM, Song D. Inorg. Chim. Acta. 2012; 380:308–321.
- 94. Qian B, Scanlon, Smith MR, Motry DH. Organometallics. 1999; 18:1693–1698.
- 95. Dulong F, Thuéry P, Ephritikhine M, Cantat T. Organometallics. 2013; 32:1328–1340.
- 96. Knight LK, Piers WE, McDonald R. Chem. Eur. J. 2000; 6:4322–4326. [PubMed: 11140961]
- 97. Zhou W, Tang L, Patrick BO, Smith KM. Organometallics. 2011; 30:603–610.
- 98. Zhang J, Gao H, Ke Z, Bao F, Zhu F, Wu Q. J. Mol. Catal. A: Chem. 2005; 231:27–34.
- 99. Yu Y, Sadique AR, Smith JM, Dugan TR, Cowley RE, Brennessel WW, Flaschenriem CJ, Bill E, Cundari TR, Holland PL. J. Am. Chem. Soc. 2008; 130:6624–6638. [PubMed: 18444648]
- 100. Cowley RE, Holland PL. Inorg. Chem. 2012; 51:8352–8361. [PubMed: 22800175]
- 101. Guzei IA, Wendt M. Dalton Trans. 2006:3991–3999. DOI: 10.1039/B605102B. [PubMed: 17028708]
- 102. Zhang J, Ke Z, Bao F, Long J, Gao H, Zhu F, Wu Q. J. Mol. Catal. A: Chem. 2006; 249:31–39.
- 103. Li Y, Wang L, Gao H, Zhu F, Wu Q. J. Appl. Organomet. Chem. 2006; 20:436–442.
- 104. Li Y, Gao M, Wu Q. J. Appl. Organomet. Chem. 2008; 22:659–663.
- 105. Li Y, Gao M, Wu Q. J. Appl. Organomet. Chem. 2007; 21:965–969.
- 106. Li Y, Jiang L, Wang L, Gao H, Zhu F, Wu Q. J. Appl. Organomet. Chem. 2006; 20:181–186.
- 107. Li Y, Wu Q, Shan M, Gao M. J. Appl. Organomet. Chem. 2012; 26:225–229.
- 108. Li Y, Gao M, Gao H, Wu Q. Eur. Polym J. 2011; 47:1964–1969.

- 109. Lee D-H, Kwon Y-J, Jin M-J. Adv. Synth. Catal. 2011; 353:3090–3094.
- 110. Lee D-H, Qian Y, Park J-H, Lee J-S, Shim S-E, Jin M-J. Adv. Synth. Catal. 2013; 355:1729– 1735.
- 111. Lee D-H, Jung J-Y, Jin M-J. Chem. Commun. 2010; 46:9046–9048.
- 112. Champouret Y, Baisch U, Poli R, Tang L, Conway JL, Smith KM. Angew. Chem. Int. Ed. 2008; 47:6069–6072.
- 113. MacLeod KC, Conway JL, Patrick BO, Smith KM. J. Am. Chem. Soc. 2010; 132:17325–17334. [PubMed: 21070039]
- 114. Cowley RE, Golder MR, Eckert NA, Al-Afyouni MH, Holland PL. Organometallics. 2013; 32:5289–5298.
- 115. Amisial LD, Dai X, Kinney RA, Krishnaswamy A, Warren TH. Inorg. Chem. 2004; 43:6537– 6539. [PubMed: 15476347]
- 116. Li Y, Gao H, Wu Q. J. Polym. Sci., Part A: Polym. Chem. 2008; 46:93–101.
- 117. Zhou W, MacLeod KC, Patrick BO, Smith KM. Organometallics. 2012; 31:7324–7327.
- 118. Rajendran NM, Maheswari K, Reddy ND. Polyhedron. 2014; 81:329–334.
- 119. Takaichi J, Morimoto Y, Ohkubo K, Shimokawa C, Hojo T, Mori S, Asahara H, Sugimoto H, Fujieda N, Nishiwaki N, Fukuzumi S, Itoh S. Inorg. Chem. 2014; 53:6159–6169. [PubMed: 24884152]
- 120. Costas M, Chen K, Que L Jr. Coord. Chem. Rev. 2000; 200–202:517–544.
- 121. Gao H, Pei L, Li Y, Zhang J, Wu Q. J. Mol. Catal. A: Chem. 2008; 280:81–86.
- 122. Oguadinma PO, Schaper F. Organometallics. 2009; 28:4089–4097.
- 123. El-Zoghbi I, Latreche S, Schaper F. Organometallics. 2010; 29:1551–1559.
- 124. Binda PI, Abbina S, Du G. Synthesis. 2011; 2011:2609–2618.
- 125. Ellis WC, Jung Y, Mulzer M, Di Girolamo R, Lobkovsky EB, Coates GW. Chem. Sci. 2014; 5:4004–4011.

Figure 1.1. Substituent patterns in β-diketiminate ligands.

Figure 3.1.2.

Structural differences between bis(ligand) complexes on zirconium, with different ortho substituents.

Figure 3.3.1.

Differences in ligand coverage in L^{Me,iPr} vs. L^{Me,Ph3} in iron(III) imido complexes. The G parameter quantifies the ligand coverage, as describe in ref. 100. Thus, even though the overall coverage is similar between the two ligands, the shape of the coverage is different.

Figure 3.3.2.

Side view of the complexes in Fig. 4, showing the greater access to the Fev=N bond when using LMe,Ph3 .

Scheme 3.2.1.

Scheme 3.2.2.

Scheme 3.2.3.

Scheme 3.2.4.

Scheme 3.2.5.

Scheme 3.2.6.

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Scheme 3.3.1.

Scheme 3.3.2.

Scheme 3.3.3.

Scheme 3.3.4.

 R' = t Bu, Me; R" = H, Me, Ph, TMS, t Bu

Scheme 3.3.5.

Scheme 3.3.6.

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Scheme 4.2.1.

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Scheme 4.2.2.

Chart 1.1

Abbreviations used in this Perspective

Selected examples of steric effects on ligand plane orientation of bimetallic complexes complexes

Selected examples of steric effect on distance of metal to ligand plane

Steric effects of backbone (β-C) substituents on structural properties

Steric effects of *N*-aryl substituents on structural properties

Table 4.1.1

Dependence of Reduction Potential on Substituents

a Bu4NPF6 was used as electrolyte.

 b All values reported with Fc/Fc⁺ in CH₃CN.

 c_{AII} values reported with Fc/Fc⁺ in THF.