Dalton Transactions



PERSPECTIVE

View Article Online
View Journal | View Issue



Cite this: *Dalton Trans.*, 2019, **48**, 12365

Low-coordinate first-row transition metal complexes in catalysis and small molecule activation

Laurence J. Taylor 📵 and Deborah L. Kays 📵 *

Enforcing unusually low coordination numbers on transition metals with sterically demanding ligands has long been an area of interest for chemists. Historically, the synthesis of these challenging molecules has helped to elucidate fundamental principles of bonding and reactivity. More recently, there has been a move towards exploiting these highly reactive complexes to achieve a range of transformations using cheap, earth-abundant metals. In this Perspective, we will highlight selected examples of transition metal complexes with low coordination numbers that have been used in catalysis and the activation of small molecules featuring strong bonds (N₂, CO₂, and CO).

Received 6th June 2019, Accepted 9th July 2019 DOI: 10.1039/c9dt02402f

rsc.li/dalton

Introduction

The stabilisation of transition metal complexes featuring unusually low coordination numbers remains a challenging and active area of research for synthetic inorganic chemists. These species are typically stabilised by sterically demanding ligands, which shield the metal from oligomerisation or further coordinates.

School of Chemistry, University of Nottingham, University Park, Nottingham, NG7 2RD, UK. E-mail: Deborah.Kays@nottingham.ac.uk

nation by Lewis basic ligands. To this end, a vast array of bulky ligands have been prepared and utilised for this purpose; including amides, alkyls, silyls, aryls, aryls, and carbenes. The development of such ligands has been instrumental in advancing the range of complexes featuring low-coordinate and highly reactive transition metal centres, and has facilitated the isolation of new classes of compounds, such as species featuring metal-metal multiple bonds.

As the chemistry of low-coordinate complexes has developed, there has been an increasing interest in the exploitation of their reactivity. With the 3d metals, which are cheap and



Laurence J. Taylor

Laurence graduated from the University of Cambridge with an MSci in Natural Sciences in 2013. He was then awarded the St Andrews 600th Anniversary Scholarship for his PhD studies at the University of St Andrews under Dr Petr Kilian. After completing his PhD studying airand moisture-sensitive pnictogen compounds, he took up a postdoctoral position at University of Nottingham under Professor Deborah Kays in late

2017. His current research focuses on the use of highly reactive complexes of the first-row transition and s-block metals as catalysts for organic transformations.



Deborah L. Kays

Deborah Kays received her MChem and PhD degrees in Chemistry from Cardiff University, Wales. After postdoctoral work, also at Cardiff University, she took up a Junior Research Fellowship at the University of Oxford. She has been at the University Nottingham since 2007, Lecturer in Inorganic Chemistry, she was promoted to Associate Professor in 2014 and to Professor in 2019. Her research

interests involve the investigation of the stabilisation of highly unsaturated complexes and their reactivity under stoichiometric and catalytic regimes. For her contributions to low-coordinate transition metal chemistry, Deborah was awarded the 2018 RSC Chemistry of the Transition Metals Award.

earth-abundant, ⁹⁻¹¹ enforcing low-coordinate geometries can drastically alter the reactivity and properties of these elements.⁷ For example, two-coordinate 3d metal complexes have been shown to act as single molecule magnets, ¹²⁻¹⁴ and have been employed in the synthesis of Zintl ions.¹⁵ In recent years, exciting examples of such species catalysing unusual reactions or promoting the activation of challenging substrates have started to appear in the literature. Our research group has become increasingly interested in this field, with some of our recent work looking at the use of *m*-terphenyl complexes in the

stoichiometric activation of small molecules and the catalysis

The term "low-coordinate" can be a rather tricky one to define when applied to the transition metals. One can argue, for example, that coordination numbers of four could be considered low for high oxidation state iron or vanadium compounds. One could also discuss how to define formal coordination number, and therefore depending on your definition "low-coordinate complexes" could cover a very broad range of species indeed. Given that Perspective articles are meant to be brief, we have restricted ourselves to considering complexes with a formal coordination number of three or less where, for example, ligands such as alkyls, aryls, amides, carbenes, phosphines, alkenes, $\it etc.$ are considered to occupy one coordination site, and $\eta^6\text{-}C_6H_6$ to occupy three. This is not meant to provide a rigorous definition of "low-coordinate" but has instead been chosen to keep this discussion focused.

Even with this restriction, there are still too many examples for this article to be an exhaustive account of the field. As such, this Perspective will cover selected examples of catalysis and small molecule activation by low-coordinate complexes as defined above.

Catalysis

Perspective

of chemical reactions.

Given the increasing importance of sustainability in catalysis, there has been a shift away from the use of platinum group metals (such as platinum, palladium, and rhodium) towards cheaper and more abundant first-row transition metals.9-11 In particular, the use of low coordination numbers for 3d transition metal complexes is affording catalysts for a wide range of reactions. A commonly employed ligand system used to support low-coordinate first-row transition metal catalysts is the β-diketiminate; bidentate monoanionic ligands that bind through nitrogen. This ligand has been complexed to numerous metals, and such species have been shown to catalyse a wide variety of transformations including polymerisation, 16,17 catalytic hydrodefluorination, 18,19 Z-selective alkene isomerisation, 20 nitrene transfer reactions, 21,22 the dehydrocoupling of phosphines,²³ phosphine-boranes and amine-boranes,²⁴ and the hydrophosphination²⁵ and hydroboration²⁶ of alkenes and alkynes. One example of such a catalyst is shown in Scheme 1, with the three-coordinate iron(π) catalyst (1), and the proposed mechanism by which it catalyses the dehydrocoupling of dimethylamine-borane.²⁴ The development of β-diketiminate 3d

Scheme 1 Example of a three-coordinate iron catalyst with a β -diketiminate ligand (1) and the proposed mechanism by which it catalyses the dehydrocoupling of dimethylamine-borane (Dipp = 2,6- $iPr_2C_6H_3$).²⁴

complexes as catalysts was covered in a recent Perspective article by Ruth Webster, and we recommend consulting this article for a more in-depth look at such species. Thus, this section will focus on low coordinate 3d metal catalysts that do not feature β -diketiminate ligands. The discussion will be subdivided by metal, to aid clarity.

Nickel

For the most part the metal complexes covered in this review are highly reactive species which have been forced into disfavoured low-coordinate geometries by sterically demanding ligands. However, nickel is something of an exception to this. Complexes of Ni(0) are found to readily adopt two- or three-coordinate geometries when combined with neutral donor ligands such as phosphines and N-heterocyclic carbenes (NHCs), and such species have proven competent catalysts for a broad range of organic transformations. In recent years, there have been examples of C-H activation with the alkenylation, ²⁸ alkylation, ²⁹⁻³¹ and stannylation of aromatic systems, C-O bond cleavage, ³³⁻³⁵ and catalytic cycloaddition chemistry. The catalysts are typically generated *in situ* from bis(cyclooctadiene)nickel(0) and the required ligand, with some

$$\begin{array}{c} 5 \text{ mol\% Ni(COD)}_2 \\ \hline 10 \text{ mol\% SIPr·HCI} \\ \hline H_2, \text{ NaO/Bu}, \\ 120 \, ^{\circ}\text{C}, \text{ m-xylene} \end{array} \begin{array}{c} \text{OH} \\ + \\ \end{array} \begin{array}{c} \text{H} \\ \text{SIPr·HCI} \\ \hline \\ \text{SIPr·HCI} \end{array}$$

Scheme 2 Hydrogenolysis of arvl ethers^{33,34} and alkenylation of arene rings²⁸ promoted by low coordinate Ni(0) catalysts

examples from the groups of Hartwig and Nakao shown in Scheme 2.^{28,33,34} There are too many examples to cover in detail here, and we recommend that readers interested in the field consult some of the excellent in-depth reviews on this topic. 37,38

Low-coordinate catalysts based on Ni(I) and Ni(II) are somewhat less common, although certainly not unheard of. For example, several low-coordinate Ni(1) complexes have been investigated as catalysts for Kumada cross-coupling reactions. 12,39-42 The Ni(0) species 2a and 2b were reacted with chlorobenzene or bromobenzene to generate the Ni(1) NHC complexes 3a-c (with elimination of biphenyl), 40,42 all of which catalyse the Kumada reaction (Scheme 3) between aryl magnesium bromides and aryl chlorides or bromides. Compounds 3b-c also catalysed the coupling of aryl boronic acids to aryl bromides (Suzuki coupling).42 These complexes are interesting as case studies for the reactivity of low coordinate metals, but more convenient and less sensitive nickel catalysts exist for such transformations. 43

Matsubara and co-workers found that treatment of 3a with triphenylphosphine afforded the mixed NHC/phosphine complex 4, which catalysed the Buchwald-Hartwig amination

Ar Ni⁰
$$X = CI, Br$$
 Ar $Ni^1 = X$ Ar $Ni^1 = X$ Ar $Ni^1 = X$ Ar $Ni = Ni$ Ar $Ni = Ni$ Ar $Ni = Ni$ Ar Ni Ar Ni

Scheme 3 Synthesis of Ni NHC complexes for Kumada and Suzuki cross-couplings, and Buchwald-Hartwig amination (Dipp = 2,6 $iPr_2C_6H_3$, Mes = 2,4,6-Me₃C₆H₂).⁴⁰⁻⁴²

of aryl halides (chlorides, bromides and iodides) by diphenylamine under mild conditions (40-70 °C), affording yields ranging between 41-98% dependent on substrate. 41 The catalyst showed good tolerance of ketone, alkene, and nitro functional groups; providing the first example of a nickel catalyst capable of coupling diphenylamine to aryl halides to afford triphenylamine derivatives (Scheme 3).41

Following on from this work, the Matsubara group very recently published an improved Ni(1) amination catalyst (5a) which coupled a range of primary and secondary arylamines to 4-bromobenzophenone in yields ranging from 58-96%. 44 The initial pre-catalyst 5a is four-coordinate, but the active species is believed to be the two-coordinate Ni(1) amide species 5b (Scheme 4). This species was also synthesised, isolated, and found to be catalytically active.44 An exchange reaction with 2,2'-biquinoline revealed that the 2,2-bipyridine ligand on 5a is labile and readily exchanged, allowing for the generation of the active two-coordinate species. 44 EPR spectroscopy of a stoichiometric mixture of 5b and 4-bromoanisole provided evidence of both Ni(1) and Ni(111) species in solution, providing direct support for the involvement of a Ni(III) intermediate in the catalytic cycle. Such Ni(1)/Ni(111) redox cycles have previously been proposed for a number of nickel catalysts, 45-47 but this study provides some of the best direct evidence for the existence of a Ni(III) intermediate. 44 The catalytic cycle proposed by Matsubara et al. is presented in Scheme 4.

Nickel(1) catalysts bearing NHCs of different ring sizes (Scheme 5) were investigated for their efficacy in Kumada cross-coupling reactions, with EPR spectroscopy revealing that the magnetic properties of the complexes were strongly affected by ring size. The NHCs with the smallest ring size gave the best catalysts, although no correlation was observed with the magnetic parameters. The combination of a six-membered ring and mesityl flanking groups was found to give the best catalytic performance, with biaryl yields of 51-83% obtained at room temperature. 12,39

A two-coordinate Kumada cross-coupling catalyst is seen in the Ni(II) bis(silylamide) complex 6a reported by Lipschutz and Tilley (Scheme 6).48-50 This catalyst was more effective for electron-poor aryl halides and, notably, promoted couplings to pyridine-based heterocycles. Stoichiometric reaction of 6a with MeMgBr resulted in the rapid formation of an alkylated Ni(II) species, 6b.48 This compound was found to react over

Perspective Dalton Transactions

Scheme 4 Proposed mechanism for Buchwald-Hartwig amination by pre-catalyst 5a proceeding via two-coordinate Ni(i) amide 5b. 44

$$Ar = Mes Apis o Tol$$

Scheme 5 Synthesis of Ni(i) NHC complexes with varying NHC ring size for Kumada cross-coupling (Anis = 2-MeOC_6H_4 ; o-Tol = 2-MeC_6H_4). ^{12,39}

Scheme 6 Proposed reaction mechanism for Kumada cross-coupling catalysed by low-coordinate Ni(ii) species $6a.^{48}$

45 minutes with 1-iodonaphthalene to afford the coupled product (1-methylnaphthalene) in low yield (13%) along with 1,1'-binaphthalene and Ni(III) methyl complex 6c. Reacting 1-iodonaphthalene with 2 equivalents of 6b, by contrast, gave

clean and rapid conversion to 1-methylnaphthalene in 98% vield. 48 It is believed that the formation of 1,1'-binaphthalene in the stoichiometric reaction is due to the formation of naphthyl radicals, and the second equivalent of 6b is required to trap these. The radical nature of this step is supported by a radical clock experiment coupling (iodomethyl)cyclopropane to PhMgBr, which resulted in 4-phenyl-1-butene, the expected product of a radical rearrangement. 48 Finally, an anionic Ni(1) species (6e) can be obtained by chemical reduction of 6a, and the reaction of 6c and 6e results in a redox equilibrium affording 6b and 6a. Based on the results of these various stoichiometric transformations, Lipschutz and Tilley proposed the catalytic cycle shown in Scheme 6.48 Species 6a is also an effective catalyst for the hydrosilylation of 1-octene with diphenylsilane, affording (*n*-octyl)diphenylsilane as the sole product after 2 h at room temperature (5 mol% catalyst loading).⁵⁰

Treatment of the Ni(II) chloride pincer complexes 7a-b with lithium triethylborohydride affords the four-coordinate Ni(II) hydrides 8a-b.51 These complexes can reversibly lose hydrogen, and under ambient pressure exist predominantly as the three coordinate Ni(1) species 9a-b (Scheme 7a, Fig. 1). The dehydrogenation reaction is second order with respect to 8b, and likely proceeds via a biomolecular elimination reaction.⁵¹ These compounds catalyse hydrodehalogenation reactions, such as the dehalogenation of geminal dihalogenides,⁵¹ and defluorination of geminal difluorocyclopropanes to fluoroalkenes. 52 Such reactions are regarded as a promising route to partially halogenated compounds from readily available perhalogenated species.⁵³ Both reactions are stereoselective, with moderate ee (enantiomeric excess) values for the monohalides of 20-74%, ⁵¹ and high Z-selectivity in the formation of alkenes from cyclopropanes.⁵² Reactions with (2,2,6,6-tetramethylpiperidin-1-yl)oxyl (TEMPO) resulted in catalyst inhibition, suggesting that the mechanism is radical in nature. 51,52 This was further supported by reactions with radical clock reagents. 51 The proposed catalytic cycle for the dehalogenation of geminal dihalogenides is shown in Scheme 7b. 51

b)
$$[Ni^{l}] = 9a \text{ or } 9b$$

$$Ar \longrightarrow R$$

$$R \longrightarrow R$$

$$[Ni^{l}] \longrightarrow R$$

Scheme 7 (a) Synthesis of the three-coordinate Ni(i) species **9a** and **9b**. (b) Proposed mechanism for catalytic, stereoselective hydrodehalogenation of geminal dihalogenides by **9a** and **9b**. LiBHEt₃ is used in excess as a reductant and hydride source. ^{51,52}

The reaction between proligand **10**, nBuLi, and trans-[Ni(PPh₃)₂(Ph)Cl] resulted in the formation of Ni(i) complex **11**, with concomitant elimination of biphenyl (Scheme 8a).⁵⁴ This spontaneous reduction of the nickel centre presumably occurs to reduce the steric demands arising from the bulky bidentate ligand. This three-coordinate species was found to be a remarkably active catalyst for the polymerisation of norbornene in the presence of a methylaluminoxane (MAO) co-catalyst (Scheme 8b), affording high molecular weight polynorbornene (M_w ca. 10^6 g mol⁻¹) with catalytic activities of up to 2.82×10^7 g_{PNP} mol⁻¹_{Ni} h^{-1} .⁵⁴

Cobalt

Several two- and three-coordinate cobalt catalysts bearing N-heterocyclic carbene (NHC) ligands have been published by Deng and co-workers. The three-coordinate Co(I) species 12 was shown to catalyse the hydrosilylation of terminal alkynes, showing broad functional group tolerance and moderate to good selectivity for formation of the *E*-alkene (Scheme 9a). Subsequently, the Co(II) amide species 13, bearing an asymmetric NHC ligand, catalysed the hydrosilyl-

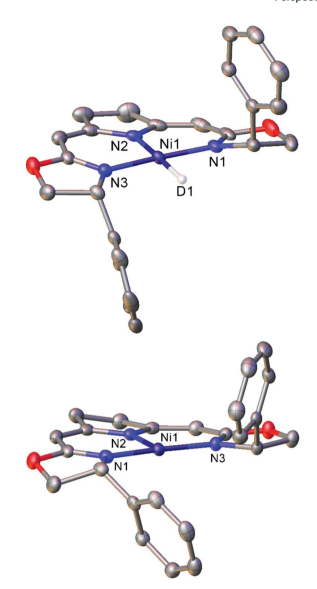


Fig. 1 Molecular structures of the deuterium analogue of **8b** (top) and three-coordinate Ni(i) complex **9b** (bottom). Hydrogen atoms and second molecule in asymmetric unit (**8b**) omitted for clarity. Anisotropic displacement ellipsoids are set at 50% probability.⁵¹

Scheme 8 (a) Synthesis of three-coordinate Ni(ı) complex $\bf 11$ with elimination of biphenyl from a Ni(ıı) source. (b) Polymerisation of norbornene by $\bf 11$ in the presence of MAO co-catalyst. 54

Scheme 9 Cobalt NHC complexes in hydrosilylation reactions. (a) Complex 12, featuring an NHC and phosphine ligand, selectively gives E-alkenes from the reaction of terminal alkynes with triphenylsilane. (b) Co(||) amide NHC complex 13 gives anti-Markovnikov products in the reaction of triethoxysilane with terminal alkenes. (c) Different Co(||) NHC complexes afford different selectivities in the reaction of diphenylsilane with terminal alkenes. Ad = 1-Adamantyl. $^{55-58}$

ation of terminal alkenes affording predominantly anti-Markovnikov products (Scheme 9b).⁵⁷ This catalyst was most effective for reactions with triethoxysilane; reactions with other silanes, including triphenylsilane and triethylsilane, gave poor conversions and low yields.⁵⁷ Following this investigation, a selection of two-, three-, and four-coordinate Co(i) NHC complexes were able to achieve different selectivity (Markovnikov, anti-Markovnikov, or hydrogenation) for the reaction of diphenylsilane with terminal alkenes, dependent on the choice of catalyst (Scheme 9c).⁵⁸ While cobalt hydrosilylation catalysts are widely known, those displaying Markovnikov or hydrogenation selectivity are relatively rare, ^{58,60,61} so these cobalt NHCs provide a valuable addition to the field.

More recently, the three-coordinate Co(0) complexes 14 and 15 were shown to catalyse the dehydrocoupling of primary arylphosphines to the corresponding diphosphanes (Scheme 10),

Dipp SiMe₃ Cy N Co⁰ SiMe₃ Cy N SiMe₃ SiMe₃ 14 15 15
$$2 \text{ ArPH}_2 = \frac{14 \text{ or } 15 \text{ (5 mol\%)}}{80 \text{ °C, } C_6 D_6} \text{ ArP(H)—(H)PAr } + \text{H}_2$$

Scheme 10 Dehydrocoupling of primary aryl phosphines by Co(0) NHC complexes 14 and 15. 59

an unusual example of a cobalt catalyst for such a transformation.⁵⁹ The catalysts afforded the coupled products in moderate to good yields (47–73%), but were ineffective with the secondary phosphine Ph₂PH and primary alkyl phosphine *t*BuPH₂ (7% and 8% yields, respectively).⁵⁹

Manganese and iron

In recent work by our research group, a series of Mn(II) and Fe(II) m-terphenyl complexes were found to catalyse the cyclotrimerisation of primary aliphatic isocyanates (Scheme 11).⁶² The reactions proceeded cleanly under mild conditions to afford isocyanurates, which find use as polymer additives to improve their physical properties, 63,64 in addition to applications in microporous materials,65,66 selective ion bonding,67 and drug delivery. 68-71 Two different flanking aryl groups were employed in the m-terphenyl ligands; [2,6-Mes₂C₆H₃], and the less sterically encumbering $[2,6-Tmp_2C_6H_3]^-$ (Tmp = 2,4,5-Me₃C₆H₂), giving catalytically active manganese and iron complexes (16-19, Scheme 11a). The Mes-substituted complexes are twocoordinate, while the less bulky Tmp-substituted ligands stabilise three-coordinate complexes, with metal-coordination by an additional THF ligand. While 17 and 19 showed no significant difference in reactivity, two-coordinate 16 was found to give a faster reaction rate than 18. Catalyst poisoning and radical trap experiments showed no evidence for the formation of catalytic nanoparticles or the involvement of radical processes, and, as a result, the mechanism is postulated to proceed via a homogeneous pathway. Kinetic experiments in the reaction of 16 with ethyl isocyanate showed a first order

Scheme 11 Cyclotrimerisation of isocyanates with low coordinate m-terphenyl metal complexes. (a) Metal complexes used as precatalysts in the cyclotrimerisation reaction. (b) General reaction scheme for cyclotrimerisation of primary aliphatic isocyanates. (c) Proposed Lewis acid mechanism for reaction.⁶²

rate dependence in both 16 and substrate. Based on these observations, a mechanism was proposed involving Lewis acid catalysis (Scheme 9c).

The manganese complexes 17, 19 (Scheme 11a) and the xylyl-substituted analogue catalyse the dehydrocoupling of dimethylamine-borane (Scheme 12), 72 which is of interest in the chemical storage and release of hydrogen. 73-76 While the reaction was slow at room temperature, increasing the reaction temperature to 60 °C provided relatively clean conversion to the cyclic dimer [Me2NBH2]2 in high yields and reasonable timeframes. The linear species Me₂NH-BH₂-NMe₂-BH₃ was also observed as a side product and intermediate in the reaction. While reactions with the two-coordinate catalyst 17 (and its xylyl analogue) showed no appreciable change upon addition of elemental mercury, reactions with the three-coordinate 19 showed a significant drop in activity. Furthermore, reaction of 19 with Me2NH·BH3 results in the rapid formation of a dark red suspension, while 17 remains a clear yellow solution. This evidence suggests that the reaction with 19 is heterogeneous in nature, involving the formation of catalytic nanoparticles, while reactions with 17 proceed via a homogeneous

Scheme 12 Dehydrocoupling of Me₂NH·BH₃ catalysed by low coordinate Mn(II) terphenyl complexes.⁷²

route. This was confirmed by isolation of the manganese nanoparticles from reactions between 19 and Me₂NH·BH₃, followed by characterisation by transmission electron microscopy (TEM), scanning transmission electron microscopy (STEM) and energy dispersive X-ray spectroscopy (EDX).⁷² This reaction serves as an example of how small changes in the steric properties of the flanking aryl groups of m-terphenyl ligands can result in significant differences in reactivity.

The iron(π) *m*-terphenyl complexes **16** and **18** (Scheme 11a) are effective catalysts for the hydrophosphination of isocyanates.⁷⁷ This reaction affords a mixture of phosphinocarboxamide and phosphinodicarboxamide products, corresponding to both mono- and diinsertion of the isocyanate into the P-H bond (Scheme 13a). Such diinsertion reactions are rare⁷⁸ and the double insertion of an isocyanate into a P-H bond has led to a family of phosphinodicarboxamide compounds. By changing the reaction conditions, this reaction can be made selective to afford either the phosphinocarboxamide (reaction in THF solvent) or phosphinodicarboxamide (through the reaction in benzene or toluene solutions in the presence of Et₃N·HCl) (Scheme 13b).

Reaction monitoring through in situ ¹H and ³¹P NMR spectroscopy reveals that the three-coordinate complex 18 shows significantly higher activity than 16, which may be due to the presence of a labile THF ligand. Interestingly, reactions with 16 show an induction period of ca. 2 h, while no such induction period was observed in reactions catalysed by 18. Monitoring of stoichiometric reactions by IR spectroscopy suggest that an iron amidate complex may be the active catalytic species in this reaction (Scheme 13c). Poisoning experiments have suggested that the reaction is homogeneous and does not involve radical processes; and a catalytic cycle where the transition metal acts as a Lewis acid was proposed to account for these observations (Scheme 13c). 77

a)
$$R_{N=C=O} + Ph_2PH \xrightarrow{5 \text{ mol}\% 16/18} Ph_2P \xrightarrow{N}_{R} + Ph_2P \xrightarrow{N}_{R}_{R} + Ph_2P-PPh_2$$

Scheme 13 Iron(II) catalysed hydrophosphination of isocyanates. (a) Initial hydrophosphination conditions affording a mixture of products. R = Ph, p-Tol ($p-Tol = 4-MeC_6H_4$), 3,5-(OMe)₂ C_6H_3 , 4-BrC₆H₄, nHex, Cy, iPr, tBu. (b) Optimised conditions for selective formation of monoinsertion (20) or diinsertion (21) product. (c) Proposed catalytic cycle for formation of observed products.

20

The two-coordinate Fe(II) amide 22 catalyses the hydrosilylation of ketones, affording the corresponding silyl ethers in good to quantitative yields (Scheme 14b).⁷⁹ The catalyst was

a)

i. $Et_3N \cdot HCI$ ii. $[^{Dipp}NHC]$ iii. KC_8 Dipp

Me₃Si

Dipp

22

23

b)

Ph₂SiH₂

R

Ph₂SiH₂

22 (0.01–2.7 mol%)

23 °C, C₆D₆

R

R

R

R

R

R

R

Scheme 14 (a) One-pot synthesis of Fe(i) heteroleptic complex 23 from Fe(ii) diamide 22. [$^{\mathrm{Dipp}}$ NHC] = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene. (b) Hydrosilylation of ketones catalysed by 22. (c) Cyclotrimerisation of alkynes catalysed by 23. $^{79.81}$

effective for the reaction of diphenylsilane with a range of ketones, proceeding cleanly at room temperature with low catalyst loadings (0.01–2.7 mol%). However, the reaction failed with tertiary silanes or silanes with bulky substituents, presumably due to steric effects. ⁷⁹ Species 22 represents an early example of an iron catalyst for this industrially relevant transformation, although it has since been supplanted by more convenient systems. ⁸⁰

Complex 22 was later used as the precursor for the (one-pot) synthesis of the first heteroleptic two-coordinate Fe(I) complex 23 (Scheme 14a), which was shown to catalyse the cyclotrimerisation of alkynes to arenes (Scheme 14c). This catalyst was effective at loadings of 2–5 mol% at room temperature, but had limited scope, showing poor reactivity towards substrates with bulky or electron-withdrawing substituents. Nonetheless, the reactivity of this complex is comparable to the handful of iron-based catalysts known for this reaction. Relations

Small molecule activation

The activation of small molecules such as N₂, CO₂, and CO poses significant but exciting challenges. Utilising these molecules generally involves overcoming large energy barriers

owing to their high bond strengths and, in some cases, low polarity.⁸⁴ However, the activation of such molecules is of significant industrial importance in reactions such as the Haber process and Fischer-Tropsch catalysis. 85 The development of homogeneous systems for functionalising these relatively inert species is thus an area of considerable research interest. In

$$R_3P$$
 $Ni-N=N-Ni$
 PR_3

24a: R = Cy; Jolly and Jonas, 1968
24b: R = i Pr; Johnson, 2013

$$(N_3N)Mo-N=N$$
 Fe
 $N_3NMo-N=N$
 Fe
 $N_3NMo-N=N$
 N_3NMo-N
 N

Fig. 2 Notable early examples of 3d metal complexes which bind dinitrogen.

Scheme 15 Synthesis and subsequent reactivity of three-coordinate Fe and Co dinitrogen bridged complexes. 91-97

this section, we will outline some examples of low-coordinate first-row transition metal complexes that are able to bind and activate these small molecules.

Nitrogen activation

An early example of a low-coordinate 3d metal binding and activating dinitrogen is the nickel phosphine complex 24a, which was first synthesised in 1968,86 and crystallographically characterised in 1971 (Fig. 2). 87,88 In this structure, the N₂ unit is protected by a cage of cyclohexyl rings,87 and a related compound featuring PiPr₃ ligands (24b) was reported in 2013.⁸⁹ Another notable early example of N2 binding is seen in the mixed Fe/Mo complex 25, which was the first complex to feature a three-coordinate iron centre coordinated entirely by N₂-derived ligands. 90

Scheme 16 Synthesis of chromium complex 34, featuring side-on N2 coordination.98

Dipp
$$Fe^0$$
 $\pm N_2$ Dipp Ee^0 $\pm N_2$ N—Dipp $\pm N_2$ N—Dipp N—

Scheme 17 Reversible binding of N2 by Fe(CAAC) species 35 and subsequent reduction by KC₈. 102

Some elegant examples of N₂ fixation and activation with low-coordinate 3d metals are seen in the series of iron and cobalt β-diketiminate complexes investigated by the Holland group. Reduction of the metal(II) chloride species (26a-c) using KC₈ under a nitrogen atmosphere affords the dinuclear N₂-bridged complexes 27a-c (Scheme 15). 91-95 Of these N₂bridged species, complex 27b was shown to be an effective precatalyst for the synthesis of asymmetric carbodiimides from organoazides and isocyanates. 21 The reduction of a THF solution of 26c with Reike magnesium (Mg*) under a N2 atmosphere afforded the highly unusual complex 28c, which features a magnesium bridging between two N₂ ligands. 96 This complex was sufficiently stable for isolation and characterisation by single crystal X-ray diffraction. The corresponding Mg* reductions with 26a and 26b afforded the metastable Fe complexes 28a and 28b, which could only be characterised in

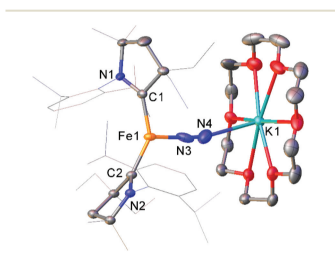


Fig. 3 Molecular structure of 36. Hydrogen atoms omitted and flanking groups of CAAC ligands are depicted as wireframe for clarity. Anisotropic displacement ellipsoids are set at 50% probability. 102

THF solution by infrared ($\nu_{NN} = 1808 \text{ cm}^{-1} \text{ for } 28a; 1818 \text{ cm}^{-1}$ for 28b) and Mössbauer spectroscopy.

Complexes 27a–c can also be reduced by potassium to afford the metal(0) species 29a–c, which features two K^+ ions coordinated to the aromatic Dipp rings. $^{91-95}$ The side-on coordination of K^+ results in elongated N=N bonds [29a 1.233(6) Å; 29b 1.215(6) Å; 29c 1.220(2) Å], $^{91-95}$ compared to those observed in 27a–c [27a 1.189(4) Å; 27b 1.186(7) Å; 27c 1.139(2) Å]. $^{91-95}$ Treatment of the reduced iron species 29b with 18-crown-6 affords a new four-coordinate complex 30, in which potassium is coordinated by an N_2 ligand. This species is active towards silylation, reacting with Me₃SiI to afford the three-coordinate Fe(III) hydrazido species 31 along with Fe(II) iodide 32. The should be noted that reacting 29b with Me₃SiI directly affords 31 in low yields (ca. 5%) with long reaction times.

Complexes 27a-c can be compared with the closely related chromium complex 34, prepared by Theopold and coworkers. Despite having a very similar ligand framework, in 34 the two nickel atoms bind dinitrogen side-on rather than end on as in 27a-c. Complex 34 was prepared by magnesium reduction of the $Cr(\pi)$ iodide 33 in THF solution under a nitrogen atmosphere (Scheme 16). This side-on coordination results in greater lengthening of the N-N bond (1.249(5) Å in 34)⁹⁸ compared with the end-on coordination of 27a-c.

While the binding and partial reduction of dinitrogen in this manner presents exciting possibilities, the full reduction of nitrogen to ammonia by a homogeneous catalyst remains a challenging prospect. One example of a low-coordinate complex which is capable of reducing N_2 in this manner is the Fe cyclic alkyl amino carbene (CAAC) complex 35, recently published by Peters, Ung and co-workers. This two-coordinate complex was capable of binding N_2 reversibly at low temperatures (ca. -78 °C) which resulted in significant changes in the UV/Vis spectra of a solution in pentane. The reaction between this complex and KC₈ at -95 °C in the presence of

Scheme 18 CO₂ fixation and activation by dinitrogen-bridged nickel complexes 37 and 38.^{109–111}

18-crown-6 facilitated the isolation of the Fe(-I) complex 36 (Scheme 17), the structure of which was confirmed by X-ray crystallography (Fig. 3). Attempting this reaction at temperatures above -78 °C resulted in decomposition to a complex mixture of products. The treatment of 35 with an excess of both KC_8 and $HBAr^F_4 \cdot OEt_2$ ([BAr F_4] $^-$ = tetrakis(3,5-bis(trifluoromethyl)phenyl)borate) at −95 °C in diethyl ether under a dinitrogen atmosphere allowed for the catalytic generation of ammonia, albeit with modest turnover numbers (3.3 ± 1.1

Fig. 4 Molecular structure of 40. Hydrogen atoms and co-crystallised hexane omitted and carbon atoms of β -diketiminate ligands are depicted as wireframe for clarity. Anisotropic displacement ellipsoids are set at 50% probability. 112

CO2 Dipp Dipp Dipp aaid Dipp Dipp 43 tBu Dipp Dipp *t*Ru Dipp Dipp

Scheme 19 Reactivity of "slipped" Co(i) β-diketiminate complex 42 with CO_2 and $N_2O.^{114,115}$

equivalents of NH₃ per Fe). Reactions at temperatures above -95 °C were ineffective, which is attributed to the poor binding of N₂ to 35 at elevated temperatures. 102

Scheme 20 (a) Formation of isocyanate from reaction of CO2 with iron-silylamide complex 46. (b) Transposition reaction of Ni complex 47 with CO_2 to generate Ni(ı) isocyanate. 124,125

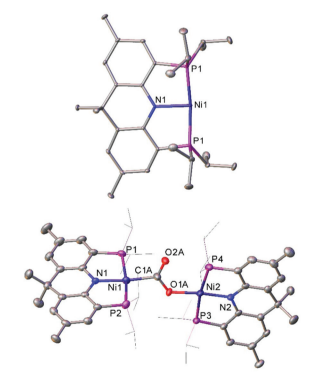


Fig. 5 Molecular structure of T-shaped Ni(1) complex 48 (top), and the species obtained after treating with CO2 (49, bottom). Hydrogen atoms, co-crystallised naphthalene (48) and disorder in carboxylate group (49) omitted, and iPr groups (49) are depicted as wireframe for clarity. Anisotropic displacement ellipsoids are set at 50% probability. 127

Scheme 21 (a) Reaction of chromium dinitrogen complex 34 with CO to give bridged isocarbonyl complex 50.98 (b) Oxidation of CO to carbonate by reaction with O2 or N2O with 51.112

Carbon dioxide activation

One common example of carbon dioxide activation by lowcoordinate 3d metals is the co-polymerisation of CO2 and epoxides by Zn β-diketiminate complexes, with a significant number of reports on this subject. 103-108 The reaction is postulated to be an example of Lewis acid catalysis, and recent work has shown that introducing electron withdrawing CF3 groups into the ligand framework can drastically increase turnover frequency for these catalysts. 106–108

A Ni complex (37), closely related to the Fe and Co complexes 27a-c shown in Scheme 15, was synthesised by Limberg and co-workers by reduction of the corresponding nickel(II) bromide with potassium triethylborohydride under a nitrogen atmosphere. 109 This complex can be further reduced to the Ni(0) species 38 by reaction with KC8. Both complexes were able to activate carbon dioxide, undergoing reductive coupling and cleavage to generate Ni(I)CO (39), Ni(II)CO₃ (40), and $Ni(II)C_2O_4Ni(II)$ (41) species (Scheme 18). Complex 40 forms a macrocyclic structure in the solid state, consisting of six nickel and six potassium cations, which was characterised by X-ray diffraction (Fig. 4). All of the Ni(II) centres in this structure are square planar and possess low spin configurations. Complex 40 was also synthesised by reacting a Ni(0)CO complex with either N2O or O2, a highly unusual example of CO oxidation at nickel. 112 The iron dinitrogen complex 27a (Scheme 15) reacted with CO2 in a similar manner, affording the first four-coordinate iron dicarbonyl complex, and a carbonate-bridged diiron complex. 113

Similar reactivity towards CO2 was demonstrated by the Co(1) β-diketiminate complex 42, which features a highly unusual slipped $\kappa N, \eta^6$ -arene coordination mode. The reaction between this compound and CO2 affords the monocarbonyl complex 43 and dicobalt carbonate complex 44 (Scheme 19). The mechanism has been probed by DFT calculations, and is considered to proceed via the oxo-bridged dimer 45, which was synthesised independently by reaction of 43 with N₂O. The reaction between 45 and CO₂ afforded 44 as the sole product (Scheme 19). 115

Reactions between low-coordinate metal amides and carbon dioxide can afford isocyanates, carbodiimides, or (often) a mixture of the two. While the reaction has been performed with metals from the s-, 116,117 p-, 118-120 and f-block, 121,122 zinc- and iron-based systems have shown some of the greatest selectivities, 123,124 with a recent iron silvlamide (46) affording the corresponding isocyanate with >95% selectivity at CO₂ pressures as low as 0.01 atm (Scheme 20a). 124 A related reaction is seen with the PNP-pincer Ni(1) complex 47,¹²⁵ which undergoes a transposition of the ligand N atom on reaction with CO2 to afford a POP-pincer and Ni(1) isocyanate (Scheme 18b).126

Ar

$$Ar$$
 Ar
 Ar

Scheme 22 Reactions of Co m-terphenyl complexes with CO to afford sterically encumbered ketones. Naph = 1-Naphthyl, $1-C_{10}H_7$. ¹³¹

b) Naph
$$Fe^{U}$$
 Naph SCO Naph

Scheme 23 Reactions of Fe(ii) m-terphenyl complexes with CO. (a) Mes and Xyl substituted compounds (16 and 55) react to give squaraines (56a-b), iron carboxylates (57a-b), and Fe(CO)₅. (b) Naphthyl substituted 58 reacts with CO to afford iron-carbene 59. (c) Proposed mechanism for the reaction between 16 or 55 with CO. 132

The T-shaped Ni(I) complex **48**, which features a rigid acridane-based ligand, was recently investigated by Yoo and Lee. This species was obtained by reduction of the corresponding Ni(II) chloride by sodium naphthalenide, and proved capable of activating a diverse range of small molecules under mild conditions. Notably, **48** reduced carbon dioxide under ambient conditions, affording the carboxylate-bridged species **49** (Fig. 5). The complex can also reduce ethene to an ethane bridge, and causes homolytic bond cleavage of a range of molecules; including dihydrogen, methanol, phenol, diphenyl disulfide, methyl iodide, hydrazine, and acetonitrile. ¹²⁷

Reactivity towards carbon monoxide

The activation of carbon monoxide by low-coordinate 3d species is less explored than that of CO_2 or N_2 , with the majority of low-coordinate complexes simply binding CO to give the corresponding metal carbonyls. Examples of more interesting reactivity include the chromium dinitrogen complex 34, which reacts with CO to afford the bridged

complex $\mathbf{50}$, ⁹⁸ a relatively rare example of an isocarbonyl complex. ¹²⁸ Compound $\mathbf{50}$ features activated CO molecules bridging through both the carbon and oxygen atoms with a diamagnetic mixed valent Cr(II)/Cr(0) core (Scheme 21a). ⁹⁸ The three-coordinate Ni(0) carbonyl complex $\mathbf{51}$ (Scheme 21b) is capable of oxidising CO in the presence of N₂O or O₂ to give the macrocyclic carbonate complex $\mathbf{40}$, ¹¹² previously seen in Scheme 18 and Fig. 4 as the result of CO₂ activation. ¹¹⁰

Of the low-coordinate 3d complexes investigated for CO activation, *m*-terphenyl complexes have arguably shown the most promise. There are examples of such complexes undergoing CO insertion reactions to afford carbonyl complexes, such as metal acyl species. ^{129,130} The cobalt(II) complexes **52a** and **52b** reacted with CO to afford sterically encumbered ketones. ¹³¹ These reactions proceeded cleanly at room temperature affording either a benzophenone (53) or a keto-fluorenone (54) depending on the flanking aryl group (Scheme 22). ¹³¹

The Fe(II) complexes 16 and 55 reduce CO with complete scission of the C \equiv O bond, affording the highly unusual squar-

Perspective **Dalton Transactions**

aines 56a-b, with concomitant formation of the Fe(II) carboxylate complexes 57a-b and Fe(CO)₅ (Scheme 23a, Fig. 6). ¹³² This reaction proceeds cleanly at room temperature and 1 bar pressure, and is both the first example of reductive cleavage of CO by a low-coordinate iron complex and C4 ring formation from CO with complete C=O bond cleavage. 135 The squaraines 56a-b feature broken conjugation due to the steric bulk of the arvl substituents, which forces them out-of-plane with the C₄O₂ ring. This results in less delocalisation into the aromatic rings, giving an unusually high C=O IR stretching frequency (56a = 1673 cm⁻¹) and carbonyl chemical shift (56a $\delta_{\rm c}$ = 269.7 ppm) in comparison to other squaraines. ^{136–139} Reactions with ¹³CO proved that all four carbon atoms in the central squaraine ring are derived from CO, and monitoring by

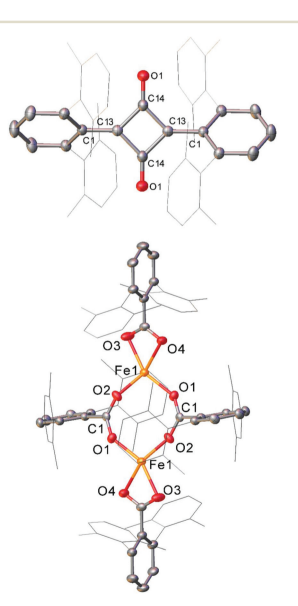


Fig. 6 Molecular structure of sterically encumbered squaraine 56b (top), and iron carboxylate 57b (bottom). Hydrogen atoms omitted and flanking xylyl groups depicted as wireframe for clarity. Anisotropic displacement ellipsoids are set at 50% probability. 132

IR spectroscopy has suggested that ketene or ketenyl (C=C=O) intermediates may be formed during the reaction. The analogous reaction using the related *m*-terphenyl complex 58, which features flanking 1-naphthyl substituents, has facilitated the isolation of Fe(II) carbene 59 (Scheme 23b), which is postulated to be an intermediate in the formation of squaraines from 16 and 55. It is proposed that the reaction halts at 59 due to the increased steric demands of the flanking naphthyl groups, which prevent further reaction. Indeed, naphthyl-substituted m-terphenyl complexes are known to display conformational isomerism due to restricted rotation, 140 and the results of DFT calculations support the notion that these flanking groups halt the reaction at species 59. Based upon these observations, a mechanism was proposed that accounts for the formation of these products, and fits all the available mechanistic data (Scheme 23c). 132

Conclusions and outlook

Although the use of low-coordinate, first-row metal complexes in catalysis and small molecule activation is a relatively young field, this strategy shows considerable promise for achieving reactivity that would be challenging by other means. Already, a vast array of diverse and exciting transformations has been discovered, and we feel confident that this research area will continue to grow and develop in the future. One thing we noted while composing this perspective is the relatively narrow range of 3d metals that dominate this area. Cobalt, nickel and iron are by far the most thoroughly investigated elements, with manganese, chromium and copper141 appearing less frequently, for example. It is possible that the less-explored elements could yield as-yet unseen reactivity, and we hope to see more work in this area in the future.

It is highly likely that the preparation of new bulky ligands with different electronic properties will be key to the development of future low-coordinate catalysts and reagents. Of particular interest are the relatively new class of cyclic (alkyl) (amino)carbenes (CAACs), which are more σ-donating and π -accepting than N-heterocyclic carbenes; ¹⁴² and the recently developed $[2,6-(2,4,6-tBu_3C_6H_2)_2C_6H_3]^-$, an incredibly sterically encumbering *m*-terphenyl ligand, 143 which was recently exploited in the synthesis of several Sn-Sn bonded compounds.144

Finally, we note the increasing interest in mechanistic investigations of these catalysts, with more researchers undertaking kinetic measurements of these systems rather than simply viewing the reaction as a "black box". Hopefully this will lead to a greater understanding of the factors that underpin the reactivity of low-coordinate metal species and allow for the future rational design of improved catalytic systems.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We gratefully acknowledge the Engineering and Physical Sciences Research Council [grant number EP/R004064/1] and the University of Nottingham for their support.

References

- 1 D. L. Kays, Chem. Soc. Rev., 2016, 45, 1004-1018.
- 2 P. P. Power, J. Organomet. Chem., 2004, 689, 3904-3919.
- 3 D. L. Kays, Dalton Trans., 2011, 40, 769-778.
- 4 D. L. Kays, in *Organometallic Chemistry*, 2010, vol. 36, pp. 56–76.
- 5 L. Falivene, A. Poater and L. Cavallo, in *N-Heterocyclic Carbenes: Effective Tools for Organometallic Synthesis*, ed. S. P. Nolan, Wiley-VCH, 1st edn., 2014, pp. 25–38.
- 6 A. Schmidt, S. Wiechmann and C. F. Otto, in *Advances in Heterocyclic Chemistry*, ed. E. F. V. Scriven and C. A. Ramsden, Elsevier Ltd, 2016, vol. 119, pp. 143–172.
- 7 P. P. Power, Chem. Rev., 2012, 112, 3482-3507.
- 8 T. Nguyen, A. D. Sutton, M. Brynda, J. C. Fettinger, G. J. Long and P. P. Power, *Science*, 2005, 310, 844–847.
- 9 P. Chirik and R. Morris, Acc. Chem. Res., 2015, 48, 2495– 2495.
- 10 L. L. Schafer, P. Mountford and W. E. Piers, *Dalton Trans.*, 2015, 44, 12027–12028.
- 11 S. Enthaler, K. Junge and M. Beller, *Angew. Chem., Int. Ed.*, 2008, 47, 3317–3321.
- 12 M. J. Page, W. Y. Lu, R. C. Poulten, E. Carter, A. G. Algarra, B. M. Kariuki, S. A. MacGregor, M. F. Mahon, K. J. Cavell, D. M. Murphy and M. K. Whittlesey, *Chem. – Eur. J.*, 2013, 19, 2158–2167.
- 13 J. M. Zadrozny, D. J. Xiao, M. Atanasov, G. J. Long, F. Grandjean, F. Neese and J. R. Long, *Nat. Chem.*, 2013, 5, 577–581.
- 14 Y. S. Meng, Z. Mo, B.-W. Wang, Y.-Q. Zhang, L. Deng and S. Gao, *Chem. Sci.*, 2015, 6, 7156–7162.
- 15 B. Zhou, M. S. Denning, D. L. Kays and J. M. Goicoechea, J. Am. Chem. Soc., 2009, 131, 2802–2803.
- 16 V. C. Gibson, E. L. Marshall, D. Navarro-Llobet, A. J. P. White and D. J. Williams, J. Chem. Soc., Dalton Trans., 2002, 4321–4322.
- 17 M.-S. Zhou, S.-P. Huang, L.-H. Weng, W.-H. Sun and D.-S. Liu, *J. Organomet. Chem.*, 2003, **665**, 237–245.
- 18 J. Vela, J. M. Smith, Y. Yu, N. A. Ketterer, C. J. Flaschenriem, R. J. Lachicotte and P. L. Holland, J. Am. Chem. Soc., 2005, 127, 7857–7870.
- 19 N. M. Hein, F. S. Pick and M. D. Fryzuk, *Inorg. Chem.*, 2017, **56**, 14513–14523.
- 20 C. Chen, T. R. Dugan, W. W. Brennessel, D. J. Weix and P. L. Holland, *J. Am. Chem. Soc.*, 2014, **136**, 945–955.
- 21 R. E. Cowley, M. R. Golder, N. A. Eckert, M. H. Al-Afyouni and P. L. Holland, *Organometallics*, 2013, 32, 5289–5298.
- 22 S. Wiese, M. J. B. Aguila, E. Kogut and T. H. Warren, *Organometallics*, 2013, 32, 2300–2308.

- 23 A. K. King, A. Buchard, M. F. Mahon and R. L. Webster, Chem. - Eur. J., 2015, 21, 15960-15963.
- 24 N. T. Coles, M. F. Mahon and R. L. Webster, *Organometallics*, 2017, **36**, 2262–2268.
- 25 M. Espinal-Viguri, A. K. King, J. P. Lowe, M. F. Mahon and R. L. Webster, ACS Catal., 2016, 6, 7892–7897.
- 26 M. Espinal-Viguri, C. R. Woof and R. L. Webster, *Chem. Eur. J.*, 2016, 22, 11605–11608.
- 27 R. L. Webster, Dalton Trans., 2017, 46, 4483-4498.
- 28 Y. Nakao, N. Kashihara, K. S. Kanyiva and T. Hiyama, J. Am. Chem. Soc., 2008, 16170–16171.
- 29 J. S. Bair, Y. Schramm, A. G. Sergeev, E. Clot, O. Eisenstein and J. F. Hartwig, *J. Am. Chem. Soc.*, 2014, **136**, 13098– 13101.
- 30 S. Okumura, S. Tang, T. Saito, K. Semba, S. Sakaki and Y. Nakao, J. Am. Chem. Soc., 2016, 138, 14699–14704.
- 31 Y. Schramm, M. Takeuchi, K. Semba, Y. Nakao and J. F. Hartwig, J. Am. Chem. Soc., 2015, 137, 12215–12218.
- 32 S. A. Johnson, Dalton Trans., 2015, 44, 10905-10913.
- 33 N. I. Saper and J. F. Hartwig, *J. Am. Chem. Soc.*, 2017, **139**, 17667–17676.
- 34 A. G. Sergeev and J. F. Hartwig, *Science*, 2011, 332, 439-443.
- 35 J. Cornella, E. Gómez-Bengoa and R. Martin, *J. Am. Chem. Soc.*, 2013, 135, 1997–2009.
- 36 A. Thakur and J. Louie, Acc. Chem. Res., 2015, 48, 2354-2365.
- 37 B. M. Rosen, K. W. Quasdorf, D. A. Wilson, N. Zhang, A.-M. Resmerita, N. K. Garg and V. Percec, *Chem. Rev.*, 2011, 111, 1346–1416.
- 38 N. A. Harry, S. Saranya, S. M. Ujwaldev and G. Anilkumar, *Catal. Sci. Technol.*, 2019, **9**, 1726–1743.
- 39 E. Carter and D. M. Murphy, *Top. Catal.*, 2015, **58**, 759–768.
- 40 S. Miyazaki, Y. Koga, T. Matsumoto and K. Matsubara, *Chem. Commun.*, 2010, **46**, 1932–1934.
- 41 S. Nagao, T. Matsumoto, Y. Koga and K. Matsubara, *Chem. Lett.*, 2011, **40**, 1036–1038.
- 42 K. Zhang, M. Conda-Sheridan, S. R. Cooke and J. Louie, *Organometallics*, 2011, **30**, 2546–2552.
- 43 S. Z. Tasker, E. A. Standley and T. F. Jamison, *Nature*, 2014, **509**, 299–309.
- 44 T. Inatomi, Y. Fukahori, Y. Yamada, R. Ishikawa, S. Kanegawa, Y. Koga and K. Matsubara, *Catal. Sci. Technol.*, 2019, **9**, 1784–1793.
- 45 L. Iffland, A. Petuker, M. van Gastel and U.-P. Apfel, *Inorganics*, 2017, 5, 78.
- 46 I. Kalvet, Q. Guo, G. J. Tizzard and F. Schoenebeck, *ACS Catal.*, 2017, 7, 2126–2132.
- 47 X. Lin and D. L. Phillips, *J. Org. Chem.*, 2008, 73, 3680–3688.
- 48 M. I. Lipschutz and T. D. Tilley, *Angew. Chem., Int. Ed.*, 2014, 53, 7290–7294.
- 49 A. N. Vedernikov, ChemCatChem, 2014, 6, 2490-2492.
- 50 M. I. Lipschutz and T. D. Tilley, *Chem. Commun.*, 2012, **48**, 7146–7148.

Perspective

51 C. Rettenmeier, H. Wadepohl and L. H. Gade, *Chem. – Eur. J.*, 2014, **20**, 9657–9665.

- 52 J. Wenz, C. A. Rettenmeier, H. Wadepohl and L. H. Gade, *Chem. Commun.*, 2016, **52**, 202–205.
- 53 M. F. Kuehnel, D. Lentz and T. Braun, *Angew. Chem., Int. Ed.*, 2013, **52**, 3328–3348.
- 54 H.-Y. Wang, X. Meng and G. Jin, *Dalton Trans.*, 2006, 2579–2585.
- 55 J. Sun and L. Deng, ACS Catal., 2016, 6, 290-300.
- 56 Z. Mo, J. Xiao, Y. Gao and L. Deng, J. Am. Chem. Soc., 2014, 136, 17414–17417.
- 57 Y. Liu and L. Deng, J. Am. Chem. Soc., 2017, 139, 1798-1801.
- 58 Y. Gao, L. Wang and L. Deng, ACS Catal., 2018, 8, 9637–9646.
- 59 D. Wang, Q. Chen, X. Leng and L. Deng, *Inorg. Chem.*, 2018, 57, 15600–15609.
- 60 B. Raya, S. Jing, V. Balasanthiran and T. V. RajanBabu, *ACS Catal.*, 2017, 7, 2275–2283.
- 61 B. Raya, S. Biswas and T. V. Rajanbabu, ACS Catal., 2016, 6, 6318–6323.
- 62 H. R. Sharpe, A. M. Geer, H. E. L. Williams, T. J. Blundell, W. Lewis, A. J. Blake and D. L. Kays, *Chem. Commun.*, 2017, 53, 937–940.
- 63 J. M. Kenny, L. Torre and L. M. Chiacchiarelli, J. Appl. Polym. Sci., 2015, 132, 42750.
- 64 G. Wegener, M. Brandt, L. Duda, J. Hofmann, B. Klesczewski, D. Koch, R. Kumpf, H. Orzesek, H.-G. Pirkl, C. Six, C. Steinlein and M. Weisbeck, *Appl. Catal.*, A, 2001, 221, 303–335.
- 65 Y. Zhang, S. N. Riduan and J. Y. Ying, *Chem. Eur. J.*, 2009, 15, 1077–1081.
- 66 E. Preis, N. Schindler, S. Adrian and U. Scherf, ACS Macro Lett., 2015, 4, 1268–1272.
- 67 M. Mascal, I. Yakovlev, E. B. Nikitin and J. C. Fettinger, Angew. Chem., Int. Ed., 2007, 46, 8782–8784.
- 68 P. Gibbons, D. Love, T. Craig and C. Budke, *Vet. Parasitol.*, 2016, 218, 1–4.
- 69 A. Bosco, L. Rinaldi, G. Cappelli, A. Saratsis, L. Nisoli and G. Cringoli, *Vet. Parasitol.*, 2015, **212**, 408–410.
- 70 A. P. Murray and M. J. Miller, J. Org. Chem., 2003, 68, 191– 194.
- 71 M. Ghosh and M. J. Miller, *J. Org. Chem.*, 1994, **59**, 1020–1026.
- 72 H. R. Sharpe, A. M. Geer, T. J. Blundell, F. R. Hastings, M. W. Fay, G. A. Rance, W. Lewis, A. J. Blake and D. L. Kays, *Catal. Sci. Technol.*, 2018, 8, 229– 235.
- 73 S. Bhunya, T. Malakar, G. Ganguly and A. Paul, *ACS Catal.*, 2016, **6**, 7907–7934.
- 74 E. M. Leitao, T. Jurca and I. Manners, *Nat. Chem.*, 2013, 5, 817–829.
- 75 A. Staubitz, A. P. M. Robertson and I. Manners, *Chem. Rev.*, 2010, **110**, 4079–4124.
- 76 A. Staubitz, A. P. M. Robertson, M. E. Sloan and I. Manners, *Chem. Rev.*, 2010, **110**, 4023–4078.

- 77 H. R. Sharpe, A. M. Geer, W. Lewis, A. J. Blake and D. L. Kays, *Angew. Chem., Int. Ed.*, 2017, 56, 4845–4848.
- 78 Y. Sun, Z. Zhang, X. Wang, X. Li, L. Weng and X. Zhou, *Dalton Trans.*, 2010, **39**, 221–226.
- 79 J. Yang and T. D. Tilley, Angew. Chem., Int. Ed., 2010, 49, 10186–10188.
- 80 B. A. F. Le Bailly and S. P. Thomas, *RSC Adv.*, 2011, 1, 1435–1445.
- 81 M. I. Lipschutz, T. Chantarojsiri, Y. Dong and T. D. Tilley, J. Am. Chem. Soc., 2015, 137, 6366–6372.
- 82 A. Casitas, H. Krause, R. Goddard and A. Fürstner, *Angew. Chem.*, *Int. Ed.*, 2015, **54**, 1521–1526.
- 83 B. A. Frazier, V. A. Williams, P. T. Wolczanski, S. C. Bart, K. Meyer, T. R. Cundari and E. B. Lobkovsky, *Inorg. Chem.*, 2013, 52, 3295–3312.
- 84 W. B. Tolman, Activation of Small Molecules: Organometallic and Bioinorganic Perspectives, Wiley, 1st edn, 2006.
- 85 J. Hagen, *Industrial Catalysis: A Practical Approach*, Wiley, 3rd edn, 2015.
- 86 P. W. Jolly and K. Jonas, Angew. Chem., 1968, 80, 705.
- 87 P. W. Jolly, K. Jonas, C. Krüger and Y.-H. Tsay, J. Organomet. Chem., 1971, 33, 109–122.
- 88 This initial complex was followed by some other interesting nickel N₂ complexes by the same authors; see: (a) K. Jonas, Angew. Chem., Int. Ed. Engl., 1973, 12, 997–998; (b) C. Kruger and Y.-H. Tsay, Angew. Chem., Int. Ed. Engl., 1973, 12, 998–999; (c) K. Jonas, D. J. Brauer, C. Kruger, P. J. Roberts and Y.-H. Tsay, J. Am. Chem. Soc., 1976, 98, 74–81.
- 89 R. Beck, M. Shoshani, J. Krasinkiewicz, J. A. Hatnean and S. A. Johnson, *Dalton Trans.*, 2013, 42, 1461–1475.
- 90 M. B. O'Donoghue, W. M. Davis, R. R. Schrock and W. M. Reiff, *Inorg. Chem.*, 1999, 38, 243–252.
- 91 J. M. Smith, R. J. Lachicotte, K. A. Pittard, T. R. Cundari, G. Lukat-Rodgers, K. R. Rodgers and P. L. Holland, *J. Am. Chem. Soc.*, 2001, **123**, 9222–9223.
- 92 J. M. Smith, A. R. Sadique, T. R. Cundari, K. R. Rodgers, G. Lukat-Rodgers, R. J. Lachicotte, C. J. Flaschenriem, J. Vela and P. L. Holland, J. Am. Chem. Soc., 2006, 128, 756–769.
- S. F. McWilliams and P. L. Holland, Acc. Chem. Res., 2015, 48, 2059–2065.
- 94 P. L. Holland, Acc. Chem. Res., 2008, 41, 905-914.
- 95 K. Ding, A. W. Pierpont, W. W. Brennessel, G. Lukat-Rodgers, K. R. Rodgers, T. R. Cundari, E. Bill and P. L. Holland, *J. Am. Chem. Soc.*, 2009, **131**, 9471–9472.
- 96 T. R. Dugan, K. C. MacLeod, W. W. Brennessel and P. L. Holland, *Eur. J. Inorg. Chem.*, 2013, 3891–3897.
- 97 S. F. McWilliams, E. Bill, G. Lukat-Rodgers, K. R. Rodgers, B. Q. Mercado and P. L. Holland, *J. Am. Chem. Soc.*, 2018, 140, 8586–8598.
- 98 W. H. Monillas, G. P. A. Yap, L. A. Macadams and K. H. Theopold, *J. Am. Chem. Soc.*, 2007, **129**, 8090–8091.
- 99 K. C. MacLeod and P. L. Holland, *Nat. Chem.*, 2013, 5, 559–565.

100 Y. Roux, C. Duboc and M. Gennari, *ChemPhysChem*, 2017, **18**, 2606–2617.

Dalton Transactions

- 101 G. Ung, J. Rittle, M. Soleilhavoup, G. Bertrand and J. C. Peters, *Angew. Chem.*, *Int. Ed.*, 2014, 53, 8427–8431.
- 102 G. Ung and J. C. Peters, *Angew. Chem., Int. Ed.*, 2015, **54**, 532–535.
- 103 M. Cheng, E. B. Lobkovsky and G. W. Coates, *J. Am. Chem. Soc.*, 1998, **120**, 11018–11019.
- 104 D. R. Moore, M. Cheng, E. B. Lobkovsky and G. W. Coates, *J. Am. Chem. Soc.*, 2003, **125**, 11911–11924.
- 105 C. M. Byrne, S. D. Allen, E. B. Lobkovsky and G. W. Coates, J. Am. Chem. Soc., 2004, 126, 11404–11405.
- 106 S. Kissling, M. W. Lehenmeier, P. T. Altenbuchner, A. Kronast, M. Reiter, P. Deglmann, U. B. Seemann and B. Rieger, *Chem. Commun.*, 2015, 51, 4579–4582.
- 107 M. Reiter, A. Kronast, S. Kissling and B. Rieger, *ACS Macro Lett.*, 2016, 5, 419–423.
- 108 M. Reiter, S. Vagin, A. Kronast, C. Jandl and B. Rieger, Chem. Sci., 2017, 8, 1876–1882.
- 109 S. Pfirrmann, C. Limberg, C. Herwig, R. Stößer and B. Ziemer, *Angew. Chem., Int. Ed.*, 2009, 48, 3357–3361.
- 110 B. Horn, C. Limberg, C. Herwig and B. Braun, *Chem. Commun.*, 2013, 49, 10923–10925.
- 111 P. Zimmermann and C. Limberg, *J. Am. Chem. Soc.*, 2017, 139, 4233–4242.
- 112 B. Horn, C. Limberg, C. Herwig, M. Feist and S. Mebs, *Chem. Commun.*, 2012, **48**, 8243–8245.
- 113 A. R. Sadique, W. W. Brennessel and P. L. Holland, *Inorg. Chem.*, 2008, 47, 784–786.
- 114 T. R. Dugan, X. Sun, E. V. Rybak-Akimova, O. Olatunji-Ojo, T. R. Cundari and P. L. Holland, *J. Am. Chem. Soc.*, 2011, 133, 12418–12421.
- 115 L. Roy, M. H. Al-Afyouni, D. E. DeRosha, B. Mondal, I. M. DiMucci, K. M. Lancaster, J. Shearer, E. Bill, W. W. Brennessel, F. Neese, S. Ye and P. L. Holland, *Chem. Sci.*, 2019, 10, 918–929.
- 116 V. U. Wannagat, H. Kuckertz, C. Krüger and J. Pump, *Z. Anorg. Allg. Chem.*, 1964, **333**, 54–61.
- 117 D. A. Dickie, K. B. Gislason and R. A. Kemp, *Inorg. Chem.*, 2012, **51**, 1162–1169.
- 118 L. R. Sita, J. R. Babcock and R. Xi, *J. Am. Chem. Soc.*, 1996, 118, 10912–10913.
- 119 J. R. Babcock and L. R. Sita, *J. Am. Chem. Soc.*, 1998, **120**, 5585–5586.
- 120 C. A. Stewart, D. A. Dickie, B. Moasser and R. A. Kemp, *Polyhedron*, 2012, **32**, 14–23.
- 121 H. Yin, P. J. Carroll and E. J. Schelter, *Chem. Commun.*, 2016, **52**, 9813–9816.
- 122 C. Camp, L. Chatelain, C. E. Kefalidis, J. Pécaut, L. Maron and M. Mazzanti, *Chem. Commun.*, 2015, **51**, 15454–15457.
- 123 A. M. Felix, B. J. Boro, D. A. Dickie, Y. Tang, J. A. Saria, B. Moasser, C. A. Stewart, B. J. Frost and R. A. Kemp, *Main Group Chem.*, 2012, 11, 13–29.

- 124 D. L. J. Broere, B. Q. Mercado and P. L. Holland, *Angew. Chem., Int. Ed.*, 2018, 57, 6507–6511.
- 125 M. J. Ingleson, B. C. Fullmer, D. T. Buschhorn, H. Fan, M. Pink, J. C. Huffman and K. G. Caulton, *Inorg. Chem.*, 2008, 47, 407–409.
- 126 B. C. Fullmer, H. Fan, M. Pink and K. G. Caulton, *Inorg. Chem.*, 2008, 47, 1865–1867.
- 127 C. Yoo and Y. Lee, *Angew. Chem., Int. Ed.*, 2017, **56**, 9502–9506.
- 128 Isocarbonyl Complexes in Encyclopedia of Inorganic Chemistry, ed. R. B. King, R. H. Crabtree, C. M. Lukehart, D. A. Atwood and R. A. Scott, 2006.
- 129 H. Lei, B. D. Ellis, C. Ni, F. Grandjean, G. J. Long and P. P. Power, *Inorg. Chem.*, 2008, 47, 10205–10207.
- 130 C. Ni and P. P. Power, *Chem. Commun.*, 2009, 5543-5545.
- 131 B. M. Gridley, A. J. Blake, A. L. Davis, W. Lewis, G. J. Moxey and D. L. Kays, *Chem. Commun.*, 2012, 48, 8910–8912.
- 132 H. R. Sharpe, A. M. Geer, L. J. Taylor, B. M. Gridley, T. J. Blundell, A. J. Blake, E. S. Davies, W. Lewis, J. McMaster, D. Robinson and D. L. Kays, *Nat. Commun.*, 2018, 9, 3757.
- 133 Z. Mo and L. Deng, *Coord. Chem. Rev.*, 2017, **350**, 285–299.
- 134 D. Benito-Garagorri, I. Lagoja, L. F. Veiros and K. A. Kirchner, *Dalton Trans.*, 2011, **40**, 4778–4792.
- 135 It should be noted that the formation of squarates $(C_4O_4)^{2-}$ from CO is known, but this does not require the cleavage of C=O bonds. See: N. Tsoureas, O. T. Summerscales, F. G. N. Cloke and S. M. Roe, Organometallics, 2013, 32, 1353-1362.
- 136 G. Xia and H. Wang, *J. Photochem. Photobiol.*, *C*, 2017, 31, 84–113.
- 137 V. Maltese, S. Cospito, A. Beneduci, B. C. De Simone, N. Russo, G. Chidichimo and R. A. J. Janssen, *Chem. – Eur. J.*, 2016, 22, 10179–10186.
- 138 G. Chen, H. Sasabe, Y. Sasaki, H. Katagiri, X.-F. Wang, T. Sano, Z. Hong, Y. Yang and J. Kido, *Chem. Mater.*, 2014, 26, 1356–1364.
- 139 T. Maeda, S. Mineta, H. Fujiwara, H. Nakao, S. Yagi and H. Nakazumi, *J. Mater. Chem. A*, 2013, **1**, 1303–1309.
- 140 B. M. Gridley, G. J. Moxey, W. Lewis, A. J. Blake and D. L. Kays, *Chem. Eur. J.*, 2013, **19**, 11446–11453.
- 141 It should be noted that, while not mentioned in this review, there are some interesting examples of low-coordinate copper β -diketiminate complexes in catalysis. Please refer to Webster's recent Dalton Perspective (ref. 27).
- 142 M. Soleilhavoup and G. Bertrand, *Acc. Chem. Res.*, 2015, 48, 256–266.
- 143 K. V. Bukhryakov, R. R. Schrock, A. H. Hoveyda, P. Müller and J. Becker, *Org. Lett.*, 2017, 19, 2607–2609.
- 144 L. G. Perla, J. M. Kulenkampff, J. C. Fettinger and P. P. Power, *Organometallics*, 2018, 37, 4048–4054.