

Cite this: *Dalton Trans.*, 2023, **52**, 14259

Cooperation towards nobility: equipping first-row transition metals with an aluminium sword

Sergio Fernández,  Selwin Fernando  and Oriol Planas *

The exploration for noble metals substitutes in catalysis has become a highly active area of research, driven by the pursuit of sustainable chemical processes. Although the utilization of base metals holds great potential as an alternative, their successful implementation in predictable catalytic processes necessitates the development of appropriate ligands. Such ligands must be capable of controlling their intricate redox chemistry and promote two-electron events, thus mimicking well-established organo-metallic processes in noble metal catalysis. While numerous approaches for infusing nobility to base metals have been explored, metal–ligand cooperation has garnered significant attention in recent years. Within this context, aluminium-based ligands offer interesting features to fine-tune the activity of metal centres, but their application in base metal catalysis remains largely unexplored. This perspective seeks to highlight the most recent breakthroughs in the reactivity of heterobimetallic aluminium–base-metal complexes, while also showcasing their potential to develop novel and predictable catalytic transformations. By turning the spotlight on such heterobimetallic species, we aim to inspire chemists to explore aluminium–base-metal species and expand the range of their applications as catalysts.

Received 21st August 2023,
Accepted 25th August 2023

DOI: 10.1039/d3dt02722h

rsc.li/dalton

1. Introduction

Noble metals (NM) are privileged elements that enable straightforward modification of organic molecules, thus establishing themselves as indispensable workhorses in the chemical industry. Their dominance stems from their ability to coordinate substrates and undergo well-understood two-electron organometallic steps, providing controllable and predictable chemical transformations.^{1–3} Yet, their limited availability, fluctuating costs and huge carbon footprint related to their production (*e.g.*, the production of 1 kg of pure Rh metal generates 32t of CO₂) represent major drawbacks to overcome if chemists aim at developing truly sustainable industrial synthetic processes.⁴ While catalysts based on abundant, inexpensive, and sustainable base metals (BM) have recently emerged as promising alternatives,^{5–9} their potential to replace noble metals in catalysis is hampered by their intrinsic differences in orbital overlap, bonding, and electronic structure. Contrary to 4d and 5d metals, reactivity on 3d metal centres often proceeds through one-electron redox events, resulting in difficulties in fully understanding, controlling and maintaining the function of the catalyst in chemical transformations.^{10–12} Fe, Cr, Ni and Co represent attractive

alternatives to pension off noble metals in key industrial processes,^{13–17} yet these metals demand adequate ligands to tame one of their several catalytically active redox couples.¹⁸ Indeed, fine-tuning their reactivity will certainly enable novel and challenging reactions that are unachievable with current technologies. Several strategies have been employed to control the chemistry of base metals in catalytic setups, including the use of redox active ligands,^{19–21} and metal–ligand cooperation.^{22,23} It is noteworthy that these approaches with Earth-abundant metals have been routinely employed by Nature in enzymes with metallic active centres to perform transformations that have no parallel in synthesis.²⁴

A particularly interesting approach that allows control of redox events in base metal centres is the use of chemical metal–ligand cooperation, in which *actor ligands* participate in bond-forming and breaking events. Combining base metals with ligands endowed with low-lying LUMOs, spanning from olefins to Lewis acids,²⁵ enables anchoring points for substrates during catalytic cycles, which facilitates their activation in a cooperative, predictable and fundamentally unique manner. In this vein, carbenes are a family of ligands endowed with ideal properties to engage base metals in cooperative two-electron events: they are strong σ -donors and can accept charge through their low-lying empty p-orbital.^{26,27} Nonetheless, their use in cooperative catalysis is prevented by the stability of the newly formed carbon–element bond after activation.^{28–30} Pairing base metals with heavier neighbours from group 14 is an approach that has been gaining increased

Queen Mary University of London, School of Physical and Chemical Sciences,
Department of Chemistry, Mile End Road, London E1 4NS, UK.
E-mail: o.planasfabrega@qmul.ac.uk



interest recently,^{31–34} as weaker bonding after activation with such elements promotes small molecule activation in an efficient and synergistic manner.^{35,36} A powerful alternative that has been recently exploited to overcome the reluctance of carbene–metal systems to undergo cooperative catalysis is the use of elements from group 13.^{37–39} Such elements possess unoccupied p-orbitals, in both high-valent and low-valent states, and coordination of ligands to such orbitals appears to be more labile, which facilitates bond scission and, subsequently, catalysis. Particularly attractive is the use of low-valent group 13 ligands, which are strong σ -donors.⁴⁰ Indeed, the use of boron-based anionic ligands (boryl-type ligands),⁴¹ which can be embedded in pincer scaffolds,⁴² has enabled cooperative catalytic processes with base metals,^{43–47} including Ni, Co and Cu.

The potential of such low valent group 13 species, combined with recent advances in low-valent aluminium (Al) chemistry,^{48–53} promoted a shift in interest towards the use of Al in cooperative catalytic transformations.⁵⁴ Al, the most abundant metal in the Earth's crust, is less electron-negative compared with B (1.47 vs. 2.01),⁵⁵ which endows Al(I) centres with a better σ -donor ability, resulting in the formation of stable, coordinatively unsaturated electron-rich metal centres. Furthermore, Al species possess highly acidic p-orbitals and a larger coordination number, key features that enable diverse substrate-coordination modes. Importantly, Al centres can be readily installed in a variety of scaffolds, which allows facile modification of their electronic and steric environment. All these attractive features have been recently employed to develop catalysts that pair Al with a noble metal, including Pd,⁵⁶ Ir,⁵⁷ and Rh,^{58–62} (species 1–4, Scheme 1) thus enabling the catalytic functionalization of C–X bonds and small molecules.

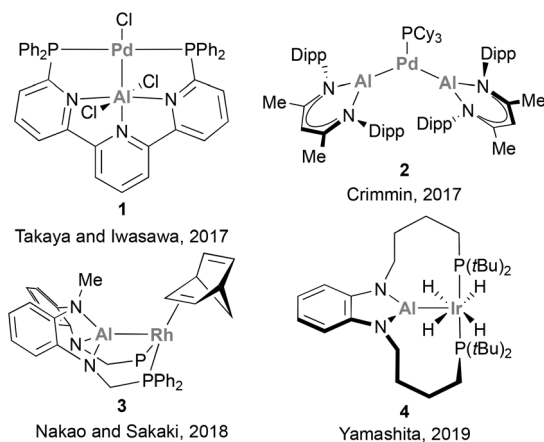
Despite these pioneering studies, heterobimetallic complexes in which Al is paired and directly bonded to a 3d metal remain underexplored, and their use in catalysis remains an uncharted territory. This is surprising considering the existing

synthetic methodologies to functionalise organic molecules that leverage from the synergistic interaction of base metals salts with aluminium-based additives.^{63–65} Thus, in this perspective we will highlight the current state-of-the-art regarding the synthesis and reactivity of main group metalloligands coordinated to base metals,^{66–68} focusing on aluminium–base-metal species that contain an aluminium–base-metal bond (Scheme 2). By doing so, we believe this perspective will encourage chemists to gain further insight into the reactivity of compounds that pair aluminium with abundant 3d metals. The heterobimetallic complexes described in the following section present an exceptional opportunity to tame and exploit unique cooperative two-electron processes, endowing *nobility* to base metals and take a step towards a more sustainable future.

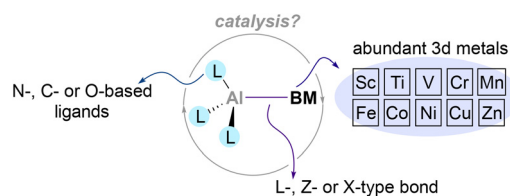
2. General synthesis of aluminium–base-metal complexes

The synthesis of heterobimetallic Al–BM complexes can be achieved from a wide range of precursors *via* two main transformations: salt elimination and ligand substitution. The former method is usually limited to the use of metal carbonyl salts (*e.g.*, Collman's reagent)⁶⁹ together with Al(III) halide precursors (Scheme 3, route 1a). However, the recent development of stable Al(I) anions has boosted the applicability of this approach using aluminyl salts together with metal carbonyls, which broadened the scope of BM precursors that can be employed to achieve heterobimetallic Al-supported species (Scheme 3, route 1b). The second methodology, ligand substitution, is particularly effective when the synthesis of Al–BM complexes with bridging H atoms is targeted, and starts from Al(III) hydride precursors (Scheme 3, route 2a). Neutral Al(I) species such as Cp*Al and Nacnac Al(I) complexes are also good L-type ligands capable of undergoing ligand substitution reactions (Scheme 3, route 2b). It is worth mentioning that substitution usually occurs with metal-carbonyl and metal-olefin species under thermal or photochemical conditions.

However, several aluminium–base-metal species have been also obtained following less explored approaches that, despite being rare and specific, are intriguing from a mechanistic point of view. These transformations would include a carboalumination reaction of a Cr–Cr quintuple bond,⁷⁰ a Ga(I)-to-Al(I) formal exchange,⁷¹ insertion reactions of Al(I) species into TM–N bonds,⁷² and direct coordination of AlX₃ species to TM



Scheme 1 Catalytically active aluminium–noble-metal species for the functionalisation of organic molecules.



Scheme 2 Heterobimetallic complexes pairing aluminium with base metals and their potential benefits.





Scheme 3 Typical procedures for the construction of Al–BM bonds.

complexes.⁷³ As this perspective is focused on the reactivity of heterobimetallic complexes bearing an aluminium–base-metal bond, the synthesis of these compounds will be briefly addressed in the following section, focusing on complexes whose reactivity has been explored.

Noteworthy, the reported synthetic approaches to forge intermetallic bonds between Al and first-row transition metals (TM) have been also employed with heavier group neighbours. While 4d and 5d metal complexes supported by aluminium-based ligands can be generated through salt elimination with anionic Al(i) species or simple coordination, they also present exclusive routes that are not applicable to 3d metals. These unique pathways to achieve aluminium–TM bonds include oxidative addition to Al–Cl or Al–H bonds, alkane elimination, addition of trialkylaluminium species to vinylidene complexes, and simple coordination followed by reduction.⁷⁴ It is worth noting that several formal oxidative addition examples have been reported for base metals and Al–Cl species; however, they are labelled as salt elimination (Scheme 3, route 1a, bottom) and limited to the use of supernucleophilic, low oxidation state metallic centres,^{69,75,76} in contrast to their heavier analogues.

3. Reactivity of aluminium–base-metal complexes

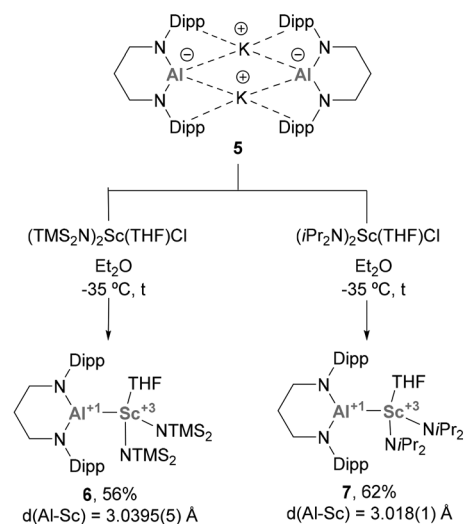
Heterobimetallic aluminium–base-metal complexes exhibit a wide range of reactivity patterns in small molecule activation and catalysis. This section will discuss such reactivity, as well as key structural features of some aluminium–base-metal species. This will be done following the first-row transition metals series and classifying reactivity patterns into specific

transformations, including small molecule activation (*e.g.* N₂, CO₂, CO, azides, *etc.*), C–H activation and oxidation reactions. Examples are presented in chronological order, except in cases where subsequent studies complemented earlier ones. It is important to emphasise that oxidation states will be explicitly depicted for those species that have undergone extensive investigation, and authors have unambiguously assigned them in their respective studies.

3.1 Scandium

Heterobimetallic complexes bearing an Al–Sc bond are still rare and have received limited attention until a recent report from Yamashita and co-workers.⁷⁷ In this work, the authors synthesised a novel Al(i) anion (**5**, Scheme 4A), which was used to furnish two unprecedented complexes featuring Al(i)–Sc(III)

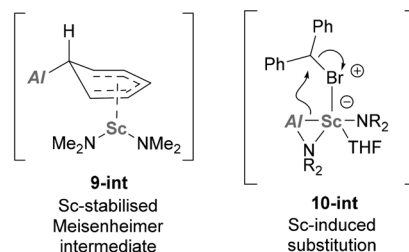
A. Synthesis of Al(i)–Sc(III) complexes



B. Reactivity of Al(i)–Sc(III) complexes



proposed activation pathways by computational studies



Scheme 4 Synthesis and reactivity of Al(i)–Sc(III) species.



bonds *via* salt elimination. Indeed, reaction of **5** with selected bis-amido Sc(III) chloride species furnished complexes **6** and **7**, which were isolated in good yields (Scheme 7A). Single Crystal X-Ray Diffraction (SC-XRD) analysis of such species showed an Al–Sc bond length of 3.0395(5) and 3.018(1) Å for **6** and **7**, respectively, thus lying between the sum of Al and Sc covalent (2.74 Å) and van der Waals (3.95 Å) radii.⁷⁸ This feature, which was previously observed in Al–Y complexes,⁷⁴ strongly suggests a partial ionic character of the intermetallic bond. Additionally, the Al–N distances in **6** and **7** are essentially identical compared with aluminium anion **5** (Al–N = 1.849(3) and 1.854(4) Å), which supports the +1-oxidation state on the Al centre.

Regarding the reactivity of such heterobimetallic complexes, **6** was found to be stable in the solid state. However, when dissolved in benzene, it slowly decomposed to complex **8** (Scheme 4B, top) *via* C–H bond activation of the methyl group of the btsa ligand. This transformation was proposed to proceed through oxidative addition at the Al(I) centre. Heterobimetallic Al–Sc systems were also tested towards S_N2-type reactivity. Complex **7**, which is stable in solution, was reacted with alkyl electrophiles (*e.g.* bromodiphenylmethane, Ph₂CHBr), resulting in the formation of species **9**, which exhibits a 1,4-dialuminiumcyclohexadiene motif. In addition, homocoupling product **10** was obtained in 74% yield (Scheme 4B, bottom). Interestingly, formation of species **9** does not occur in the absence of alkyl halide or when using toluene as solvent. Moreover, homocoupling product **10** was observed when reactions were run both in benzene and toluene, suggesting that they are not involved in its formation. Computational studies suggest complex **9** is formed *via* exchange of the coordinated THF in **7** for benzene, followed by a nucleophilic attack of the Al(I) lone pair involved in Al–Sc bonding. This would yield an Sc-stabilised Meisenheimer complex (**9-int**) as the key intermediate of this transformation. As for **10**, computational studies indicate a synergistic activation of the C–Br bond through a 2e[−] process. First, the halide coordinates to the Sc centre, which triggers a nucleophilic attack of the Al(I) atom that cleaves the intermetallic bond to furnish a new Al–C bond (**10-int**). This new organometallic bond can act as nucleophile and attack a Ph₂CHBr molecule, leading to the formation of **10**.

As observed, the chemistry of Al–Sc heterobimetallic complexes is still in its early stages of exploration and requires further development towards the development of novel small molecule activation pathways. Nevertheless, the aforementioned findings reported by Yamashita suggest that combining Al with Sc holds promise for the development of efficient systems in cooperative C–H and C–X bond activation and catalysis.

3.2 Titanium

Since the discovery of the Ziegler–Natta process,^{79,80} Ti complexes have been broadly applied in polymer and synthetic organic chemistry.⁸¹ Due to the significance of such polymerization processes and the practicality of Tebbe's reagent,⁸² the

combination of Ti systems with aluminium species has been an intense area of research.^{83–110} Although Ti and Al have been demonstrated to effectively work in synergistic processes, heterobimetallic systems featuring a Ti–Al bond remain largely underexplored,¹¹¹ and their application to catalytic transformations is a feature yet to be discovered.

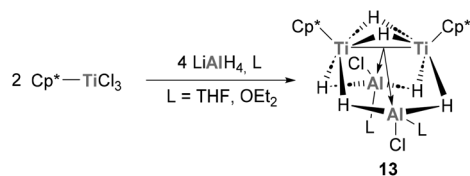
The chemistry of systems featuring a Ti–Al bond has also received attention from Yamashita and co-workers.¹¹² In this study, an anionic Al(I) complex (**11**) was reacted with Ti(OiPr)₄, leading to the formation of a system containing a Ti and an Al atom with no direct Ti–Al bond (Scheme 5). It is worth mentioning that, as a possible mechanism for the formation of **12**, two different mechanistic routes were proposed. Pathway 1 involves the formation of an open-shell Al(II) centre, with subsequent release of Ti(OiPr)₃, while pathway 2 entails the intermediacy of a heterobimetallic species containing a Ti–Al bond. In both cases, the reaction results in a concomitant redox step that leads to the formation of species **12**. Unfortunately, the reactivity of such species was not studied in further detail.

Jover and Yélamos have recently reported the synthesis of low-oxidation state Ti species with an intriguing Ti–Al bonding situation.¹¹³ Reaction of Cp*TiCl₃ with LiAlH₄ in the presence of donor solvents such as THF or diethyl ether results in the formation of [(TiCp*(μ-H))₂{(μ-H)₂AlCl(L)}₂] (**13**, Scheme 6). Interestingly, this family of mixed Ti–Al heterobimetallic clusters presents average titanium–aluminium distances of 2.644 (10) Å, which are smaller than the sum of covalent radii (2.81 Å) and shorter than those reported for heterometallic complexes with Ti(III)–H–Al units. Indeed, this short distance is explained by two strong interactions between the Ti–Ti bond and the empty s orbitals of Al, leading to unprecedented Ti–Ti → Al interactions in which the Lewis base donor is the pair of electrons involved in the Ti–Ti bond. The reactivity of such bonds is yet to be investigated, but holds the promise of unique transformations exploiting the rich redox ability of heterobimetallic Ti–Al species.



Scheme 5 Reaction between a low valent Al(I) and Ti(OiPr)₄.





Scheme 6 Synthesis of heterobimetallic Ti–Al clusters featuring Ti–Ti → Al interactions.

3.3 Vanadium

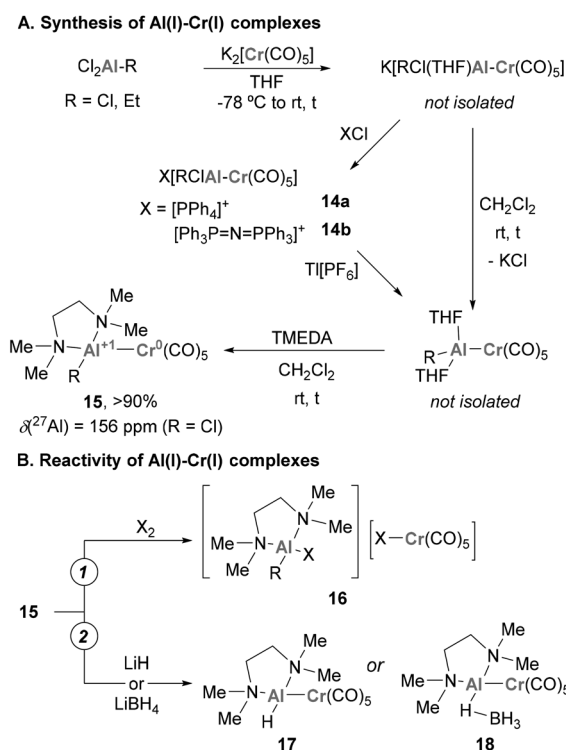
Similarly to its 3d neighbour from group 4, V salts have found broad application in polymer chemistry when combined with Al(III) organometallic additives.^{114–118} However, there is a lack of examples of heterobimetallic complexes featuring a V–Al bond, and the study of the reactivity at the interface of such species is yet to be meticulously explored.

3.4 Chromium

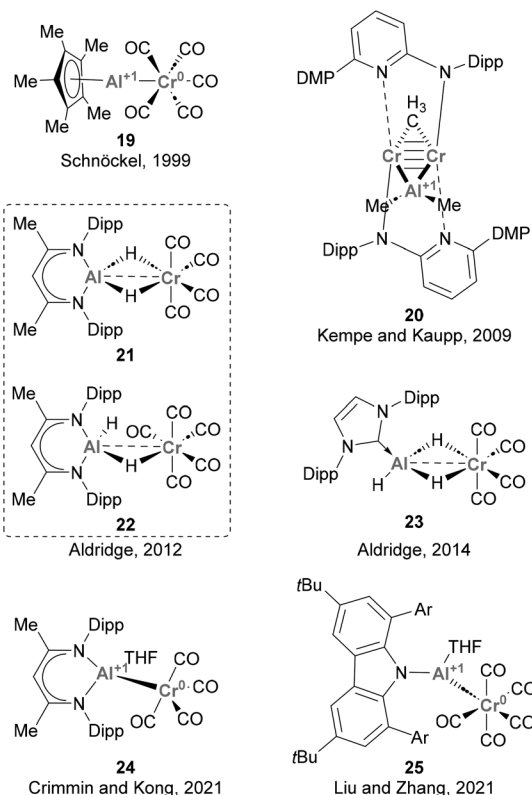
Pioneering examples of heterobimetallic complexes containing Cr–Al bonds were reported in 1996 by Fischer and co-workers.^{119–121} As shown in Scheme 7A, reacting common Al(III) halide precursors with $K_2[Cr(CO)_5]$ in THF, the authors synthesised a collection of anionic species $K[RCl(THF)Al-Cr(CO)_5]$ through salt elimination. These compounds can be isolated as solvent-free $X[RClAl-Cr(CO)_5]$ species (**14a–b**) through cation exchange ($X = PPh_4^+$ or $Ph_3P=N^+=PPh_3$). When DCM is added to these species together with a bidentate amine (TMEDA), a

second elimination of KCl takes place, yielding $[(TMEDA)Al-Cr(CO)_5]$ species **15**. Unfortunately, the authors were not able to analyse the structure of such species by SC-XRD, but their main features were assigned by comparison with its heavier Ga(i) analogue ($R = Me$). Spectroscopically, a significant shift from 210 to 156 ppm was observed by ^{27}Al -NMR when **15** was compared with Al(III) species, which indicates the presence of an Al(I) centre in **15**. The reactivity of heterobimetallic Cr–Al complex **15** was scrutinised towards the oxidative addition of halogens X_2 , resulting in the oxidative cleavage of the Cr–Al bond (**16**, Scheme 7B, pathway 1). Additionally, ligand exchange with hydride sources such as LiH and $LiBH_4$ produced hydride and borohydride analogues of complex **17** (Scheme 7B, pathway 2).

Since then, various synthetic approaches for accessing heterobimetallic species featuring Cr–Al bonds have been reported (Scheme 8). Studies from Schnöckel (**19**),¹²² Aldridge (**21–23**),^{123,124} Crimmin and Kong (**24**),¹²⁵ and Liu and Zhang (**25**)¹²⁶ achieved such heterobimetallic species by ligand substitution using low oxidation state Cr carbonyl species. In contrast, Kempe and Kaupp synthesised species **20** *via* carboalumination.⁷⁰ These works provided key insights into the synthesis and electronic structure of species featuring Cr–Al bonds, as well as the nature of such intermetallic bond. These studies have also provided information about the formal oxidation states of Cr and Al, which was unambiguously assigned to be Cr(0) and Al(I) in species **20**, **19**, **24** and **25**. Noteworthy,



Scheme 7 First examples of Al(I)–Cr(0) complexes synthesised by salt metathesis.



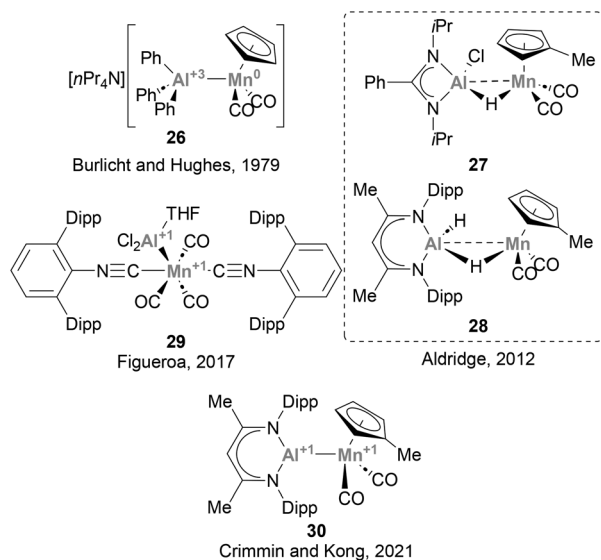
Scheme 8 Cr–Al complexes structurally characterised.



and as far as we know, the ability of species depicted in Scheme 8 towards small molecule activation and catalysis has not been explored so far. Hence, the chemical characteristics and reactivity of the Cr–Al bond are yet to be thoroughly investigated. This aspect is particularly intriguing considering the remarkable capacity of low oxidation state Cr species to activate a diverse range of chemical bonds,^{127,128} which ensures a rich redox reactivity for species featuring Cr–Al bonds and offers the opportunity for future investigations in this field.

3.5 Manganese

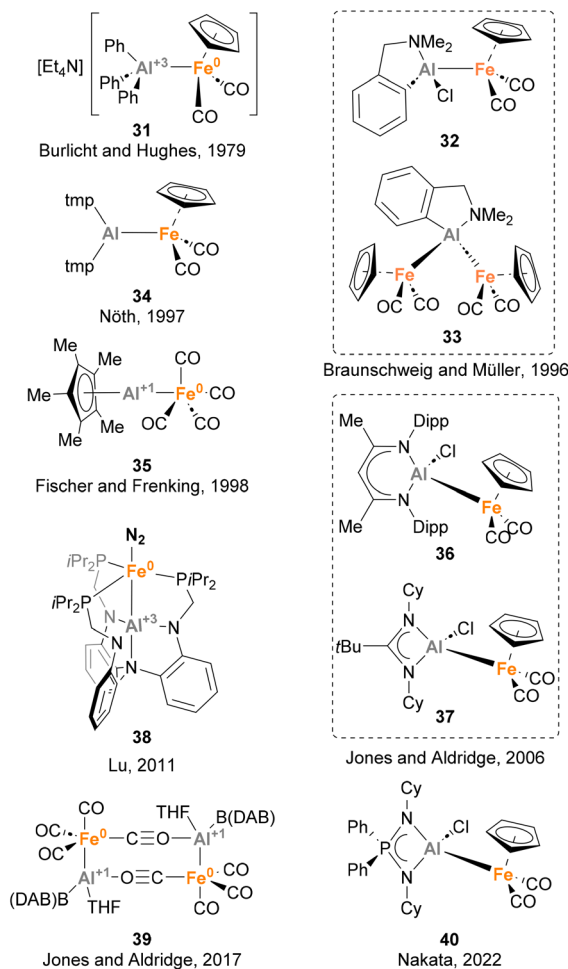
The lightest element of group 7, Mn, has been largely underexplored in heterobimetallic complexes bearing a direct Mn–Al bond, and only a handful of examples have been described so far.^{73,123,125,129,130} Burlitch, Hughes and co-workers pioneered the field, reporting the first complex featuring a direct Mn–Al bond (**26**, Scheme 9) in 1979.⁷³ In this species, AlPh₃ acts as a Z-type ligand, accepting charge from the anionic [CpMn(CO)₂][−] fragment. Complexes **27** and **29** were synthesised following salt elimination routes with Mn(I) and Mn(−I) precursors, respectively, while **28** and **30** were accessed *via* decarbonylative ligand substitution. While the intermetallic bond in **27** and **28** was not discussed in detail, species **29** and **30** were unambiguously assigned as low oxidation state Al(I)–Mn(I) heterobimetallic systems by means of computational and experimental studies. Unfortunately, and similarly to its group 6 neighbour, the reactivity of Mn–Al species towards the activation of small molecules and strong bonds has not received attention from the chemical community. Given the abundance of Mn, its potential to catalyse a wide range of organic reactions, including C–H activation and oxidation,^{131–133} as well as its ability to engage in visible-light-induced transformations,¹³⁴ the use of heterobimetallic Mn–Al species in small molecule activation and catalysis represents a promising opportunity towards more sustainable organic synthesis.



Scheme 9 Mn–Al species structurally characterised.

3.6 Iron

Iron is the 3d metal that has been more studied in heterobimetallic complexes that feature an Fe–Al bond, thus forming metal complexes containing the most abundant metals in the Earth's crust. Nowadays, a plethora of heterobimetallic Fe–Al complexes have been described (Scheme 10).^{73,129,135–151} These complexes attracted the interest of the inorganic community due to their particular bonding situation, as well as their reactivity in cooperative small molecule activation. To the best of our knowledge, the first heterobimetallic complex bearing an Fe–Al (**31**) bond was reported in 1979 by Burlitch, Hughes and co-workers.⁷³ Similarly to Mn species **26**, this Fe–Al species features a Z-type coordination of AlPh₃, which acts as acceptor. The most exploited route to synthesise such complexes is the salt metathesis approach, which was used by Braunschweig and Müller (**32** and **33**),¹³⁶ Nöth (**34**),¹³⁸ Jones and Aldridge (**36**, **37** and **39**),^{140,145} and Nakata (**40**),¹⁴⁸ while ligand substitution has been reported by Fischer and Frenking¹³⁷ and Lu¹⁴² for the synthesis of **35** and **38**, respectively. While the reactivity of complexes depicted in Scheme 10 was not investigated, the potential of Fe–Al bonds in small molecule activation and cata-



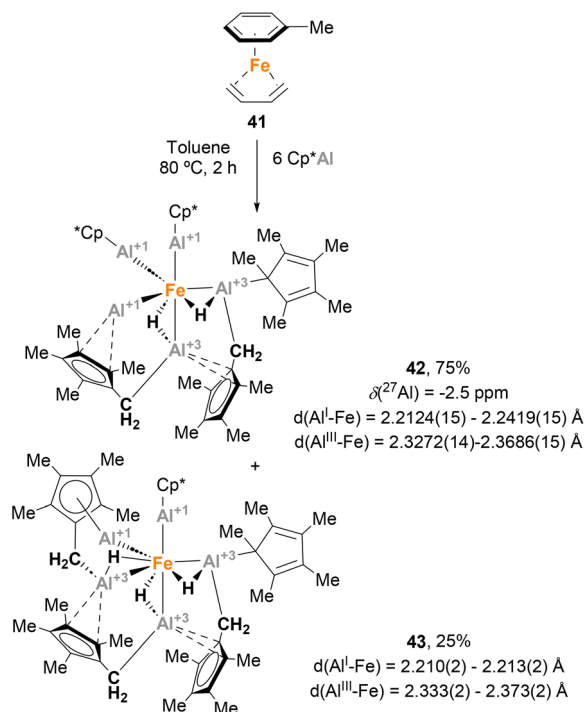
Scheme 10 Structurally characterised Fe–Al complexes.



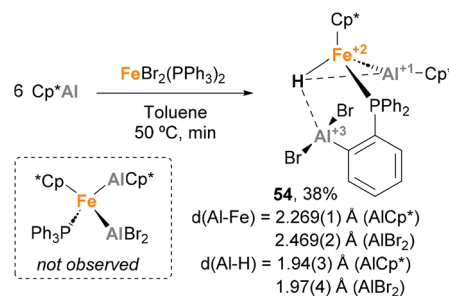
lysis has been studied in detail using other heterobimetallic complexes, including serendipitous transformations. These reactions, which are detailed in the section below, include C–H bond cleavage and small molecule activation processes, as well as catalytic reduction of N₂ to NH₃.

3.6.1 C–H bond activation. Trying to test the limits of Fe(0) complexes in terms of number of Cp*Al coordinated to the metal, Fischer and co-workers focused their efforts on the synthesis of [Fe(AlCp*)₅] complexes, which were predicted to be stable by DFT calculations.^{37,152} With this goal in mind, [Fe(η⁶-toluene)(η⁴-1,3-butadiene)] **41** was reacted with a large excess of Cp*Al (Scheme 11). However, instead of [Fe(AlCp*)₅], the reaction resulted in a 3 : 1 mixture of two isomers, **42** and **43**, respectively.¹³⁹ SC-XRD analysis of **42** showed Fe–Al distances in agreement with Fe(0)–Al(I) oxidation states for the AlCp* moieties (2.2124(15)–2.2419(15) Å)¹²⁰ and Fe(0)–Al(III) oxidation states for the [HAL(CH₂C₅Me₄)] fragments (2.3272(14)–2.3686(15) Å). SC-XRD analysis of **43** exhibited similar structural parameters. The formation of these structures can be explained through ligand substitution to generate [Fe(AlCp*)_n] species, followed by a C–H bond activation step involving methyl groups of the Cp* ligands. As far as we know, this study involving the serendipitous Csp³–H activation of Cp* ligands represents the first example of the ability of heterobimetallic Fe–Al species to perform C–H bond activation.

Fischer and co-workers observed a similar transformation while studying the insertion of Cp*Al into Fe–Br bonds.¹⁴¹ While reacting ECp* (E = Ga, In) with [FeBr₂(PPh₃)₂] (X = Cl, Br) resulted in a Cp* transfer process, when [FeBr₂(PPh₃)₂] was



Scheme 11 First example of C–H bond activation mediated by an Fe–Al complex.



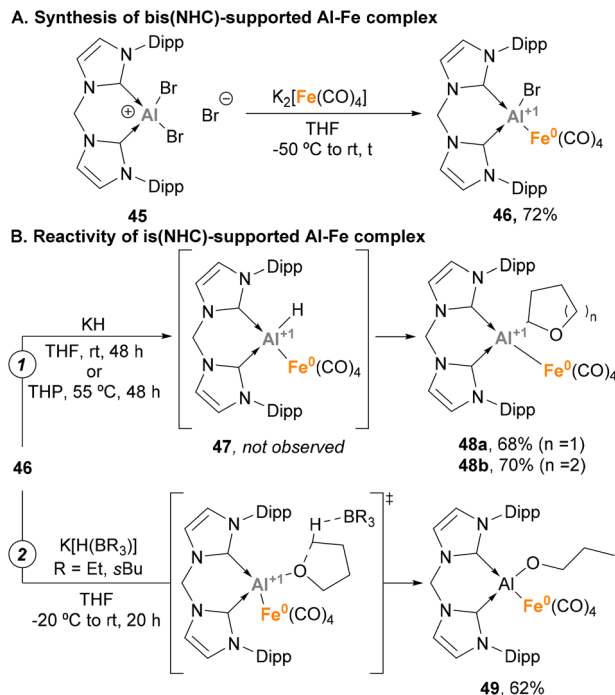
Scheme 12 C–H bond insertion in the synthesis of a Fe–Al complex reported by Fischer.

treated with Cp*Al, complex **44** was obtained in 38% yield (Scheme 12). In this case, SC-XRD analysis also shows different Fe–Al bond lengths of 2.269(1) Å (Fe–AlCp*) and 2.469(2) Å (Fe–AlBr₂–Ph), in agreement with the previously reported values for Fe(0)–Al(I) and Fe(0)–Al(III) bonds, respectively.^{73,120,139} In this example, formation of **44** can be rationalised by a Csp²–H bond activation of the phenyl residue from the phosphine ligand after formation of putative [FeCp*(AlCp*)(AlBr₂)(PPh₃)] species, which were not observed. This unique reactivity of Fe–Al bonds towards C–H activation, which was not observed when using Ga/In Cp* analogues, can be attributed to the higher acidity of Al, which leads to more reactive ambiphilic systems.

Taking advantage of the steric protection that bis-NHC ligands provide, Driess and co-workers focused on the synthesis of low oxidation state Al(I)–H species embedded in a Fe–Al unit.¹⁴³ To this end, a salt elimination-reduction strategy was employed starting from CNC^{DiPP}AlBr₃ (**45**) and Collman's reagent analogue K₂[Fe(CO)₄], resulting in a heterobimetallic Fe(0)–Al(I) complex **46** (Scheme 13A). With the goal of achieving Al–H species, different hydride sources were employed (Scheme 13B). Surprisingly, when KH was added to **46** in tetrahydrofuran (THF) or tetrahydropyran (THP), Al(I)–H species **47** were not detected (Scheme 13B, pathway 1). However, complexes **48a–b** were formed in high yields. This transformation is a good example of α -metalation of O-heterocycles with concomitant H₂ extrusion, as evidenced by spectroscopic and computational studies. Usually, α -metalation processes with organolithium or organopotassium reagents lead to the cleavage and polymerisation of the heterocycle,^{153,154} which makes this reaction particularly interesting. On the contrary, when K[H(BR₃)] species were employed as hydride sources a ring-opening reaction of THF was observed, yielding complex **55** (Scheme 13B, pathway 2). This transformation, which represents a pioneering example of C–O cleavage with heterobimetallic Fe–Al species, is proposed to start by THF coordination to the Al(I) LUMO, followed by hydride-mediated ring-opening of THF.

Over the past year, Crimmin and co-workers made important contributions to the development of heterobimetallic Fe–Al species and their application in C–H activation. In 2022, they successfully synthesised and studied the reactivity of





Scheme 13 C–H metalation and C–O bond cleavage mediated by Fe–Al species.

novel low-spin κ^2 -hydride Fe(II)–Al(I) complexes.¹⁵⁰ These species are readily accessed in excellent yields *via* a two-step procedure (Scheme 14A) that involves a first ligand exchange with FeBr₂ with NacnacAlH₂ and phosphine ligands (**50a–b**) followed by reduction with Mg turnings (**51a–b**). SC-XRD analysis unveiled Fe–Al bond distances close to the sum of their covalent radii for **50a–b** (2.453(1)–2.459(1) Å *vs.* a covalent radii of 2.43 Å),⁷⁸ prompting the authors to define these systems as σ -alane iron(II) complexes. When reduced with Mg, Fe–Al bond significantly shortened in species **51a–b** (2.2176(13)–2.194(1) Å), which indicates a reduction of the Al centre to a +1 oxidation state. Computational studies of species **51a–b** show these entities reside in an intermediate state between an aluminylene–Fe(II) complex and a highly polarized system comprising a cationic Al unit and an anionic Fe dihydride fragment. Additionally, SC-XRD analysis revealed that the Fe centre possesses a distorted octahedral geometry, in which the aluminyl ligand bends away from the axial position by 45°. This results in an unusual ground state destabilisation that strongly impacts their chemical behaviour.

In contrast to traditional, fully saturated d⁶ low spin Fe complexes, which tend to be stable and unreactive, it was found that complexes **51a–b** were capable of performing C–H bond activation. Indeed, when **51a** is heated in toluene solution, cyclometalated product **52** is formed *via* Csp³–H activation (Scheme 14B, pathway 1). More remarkably, complexes **51a–b** were able to selectively activate the C₂–H bonds of pyridines with a variety of *para*-substituents, leading to κ^3 -trihydride Fe–Al species **53** (Scheme 14B, pathway 2).

Computational studies indicate that the key step for such C–H bond activation pathways involves a cooperative reductive deprotonation reaction in which the Fe–Al bond behaves as a frustrated Lewis-pair (FLP), where the HOMO (Fe–Al σ -bonding orbital) acts as donor and the LUMO (p-orbital on Al) as acceptor.

In a subsequent study, Crimmin and co-workers further tested the reactivity of species **51** towards *meta*- and *ortho*-substituted pyridines.¹⁵⁵ This investigation unveiled that *meta* substituents hamper C₂–H bond activation, leading to C₆-aluminated heterocycles. Interestingly, these complexes are not stable upon heating, which leads to a nucleophilic attack of the organometallic Al–C_{pyridine} bond to the Nacnac ligand and the formation of a tricoordinated Al fragment (**54**). On the other hand, when 2-alkyl pyridines were used (Scheme 14B, pathway 3), Csp³–H activation occurs under mild conditions, leading to species **55** after nucleophilic attack on the ligand. The thermal stability of complex **55** was also investigated. Interestingly, these studies revealed a series of rearrangements (when R = H) and deprotonations (when R = Me) involving the Nacnac ligand, with concomitant extrusion of the initial 2-alkyl pyridine and formation of the corresponding κ^3 -hydride Fe–Al complex **56**. This mechanism was supported by computational studies, showing a feasible energy activation barrier of 25.3 kcal mol^{–1} for the process depicted in pathway 3. More recently, Gorgas and Crimmin reported a double Csp³–H activation of CH₃CN using **57** (Scheme 14B, pathway 4).¹⁵⁶ SC-XRD analysis exhibited a μ^2 - κ_C - κ_N -[CHCN]^{2–} bis-anion as a bridging unit through the Al atoms (**57**). Computational studies indicate that the monodeprotonated acetonitrile complex acts as intermediate, while the distorted octahedral geometry of the Fe species contributes towards making possible this unique reactivity.

3.6.2 Small molecule activation. Given the ability of tripodal, tetradentate Fe–B species to catalyse N₂-to-NH₃ reduction,¹⁵⁷ Peters and Fajardo decided to investigate the impact of heavier Al congeners in N₂ activation reactions.¹⁴⁷ Thus, starting from FeBr₂ and ligand **58** scaffold in the presence of Na(Hg) amalgam, complex **59** was achieved in 67% yield (Scheme 15A). Remarkably, the Fe–Br remained intact in **59**, in contrast to metal–aluminium complexes in which halogen abstraction by the Al centre takes place.¹⁵⁸ In **59**, SC-XRD analysis unveiled an Fe–Al bond length of 2.6619(1) Å, slightly longer than the sum of covalent radii. The reduction of **59** using Na(Hg) under N₂ atmosphere led to the formation of complex **60**, which shows an Fe–Al bond length 2.5393(5) Å, suggesting that **59** induces a closer proximity between the two metal atoms compared with AltraPhos (depicted in Scheme 10, complex **38**).¹⁴² The Fe–Al bonding in **59** and **60** was described as a dative Fe → Al(III) interaction, where aluminium acts as a Z-type ligand that stabilises the σ -(Fe–Br/N₂) orbital of Fe d_{2z} parentage. For this reason, further reduction of **60** with Na(Hg) leads to a well-defined Fe(–I)–Al(III) species **61**, which shows an even shorter Fe–Al bond length (2.485(1) Å). This clearly suggests a stronger stabilising donor–acceptor interaction with the Al centre, as aforementioned. After characterisation of



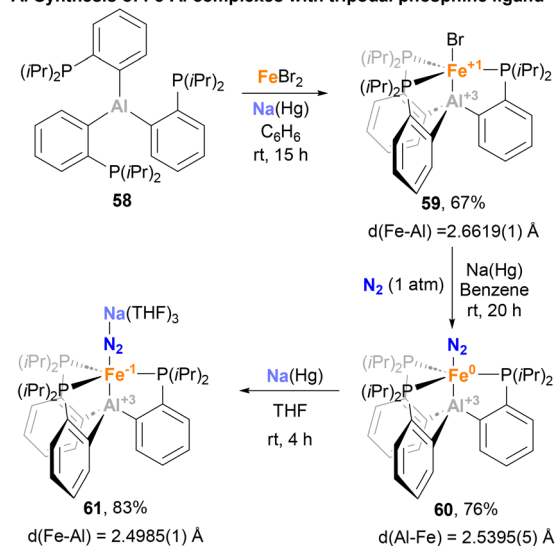
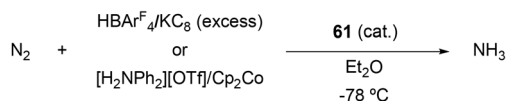
A. Synthesis of NacNacAl-Fe complexes



B. Reactivity of NacNacAl-Fe complexes

Scheme 14 C–H bond activation mediated by a low-spin d^6 Al–Fe complex.

A. Synthesis of Fe–Al complexes with tripodal phosphine ligand

B. N_2 -to- NH_3 reaction catalysed by Fe–Al species 61Scheme 15 Al–Fe complexes employed in catalytic N_2 -to- NH_3 reduction.

complex **61**, its ability to catalytically activate N_2 was studied (Scheme 15B). Cyclic voltammetry and FT-IR experiments were conducted to identify the extent of N_2 activation in species **61** and its corresponding analogues containing group 13 elements B and Ga. The results revealed a comparable degree of N_2 activation among these species, suggesting that the presence of the group 13 element bonded to the Fe centre has minimal impact. However, the efficiency of **61** towards N_2 reduction was found to be lower compared with the B analogue with two different sets of proton source and reductant combinations. The authors attributed this lower activity to a higher preference of species **61** for competing Hydrogen Evolution Reaction (HER), particularly when the reductant employed was milder (Cp_2Co vs. KC_8), preventing the regeneration of the active species. It is noteworthy to mention that, to the best of our knowledge, this process represents the sole reported reaction wherein an Fe–Al system functions as a catalyst.

Similar reactivity towards N_2 activation was described by Lu and co-workers using AltraPhos-supported Fe–Al species **38** (Scheme 16).¹⁵⁹ After one-electron reduction in THF, species **62** is readily prepared. As expected, spectroscopic and SC-XRD studies indicate that species **62** presents a weaker N–N bond compared with **38** due to stronger π -back donation. In addition, sizeable changes also occur in the Fe–Al bond, which shortens 0.23 Å upon reduction. These features suggest that reduction occurs at the Fe centre, which adopts a formal oxidation state value of -1 . Species **62** allowed functionalisation





Scheme 16 Al–Fe complexes employed in catalytic N_2 -to- NH_3 reduction.

of the bound dinitrogen by reacting it with 1,2-bis(chlorodimethylsilyl)ethane in the presence of KC_8 . The reaction occurs smoothly, and generates diamagnetic Fe(II) species **63** in a transformation that involves double substitution of the distal nitrogen atom as well as four-electron reduction of the N–N bond. Contrary to the previous example, catalytic functionalisation was not explored, but **63** represents an exceptional opportunity to explore complete N–N bond scission with release of amines.

Recently, Mankad and co-workers described the synthesis of a novel Fe–Al heterobimetallic complex, and extensively investigated its reactivity.¹⁴⁹ By reacting a Nacnac Al(III) iodide complex with $[\text{K}[\text{FeCp}(\text{CO})_2]]$, complex **64** could be obtained through a salt elimination route (Scheme 17A). SC-XRD analysis exhibited a tetracoordinated Al centre with distorted tetrahedral geometry, and the Fe–Al bond length was comparable to the Fe–Al species reported by Aldridge (Scheme 10, species **36** and **37**).¹²⁹ Computational insight into the atomic charge and bonding nature of species **64** unambiguously determined a +3 oxidation state for Al and a zero oxidation state for Fe. Regarding its reactivity, **64** generates adduct **65** upon reaction with CO_2 under mild reaction conditions (Scheme 17B, pathway 1), representing the first example of Al(III)–base-metal-mediated CO_2 activation. Interestingly, irradiation of complex **65** with UV light triggered a formal 1,3-methyl shift, leading to the formation of species **66**. Species **64** is also able to ring-open epoxides, such as cyclohexene oxide, yielding product **67** (Scheme 17B, pathway 2). Such reactivity studies were complemented with a computational analysis of the reaction profile, aiming at the elucidation of the cooperative pathways governing CO_2 and C–O activation. Surprisingly, computational studies revealed a low energetic barrier for the homolytic dissociation of the Fe–Al bond in complex **64**, which at the same time is similar to the experimental barrier obtained. Based on this, a mechanism in which the Fe–Al bond dissociates to generate a highly reactive metalloradical Fe(I)–Al(II) pair was proposed. Further evidence of the intermediacy of a metalloradical Fe(I)–Al(II) pair was gathered when **64** was mixed with benzophenone (Scheme 17B, pathway 3), leading to the formation of the NMR-silent species **68**. It is worth noting that the origin of the low energetic barrier for homolytic Fe–Al cleavage

is attributed to the significant steric hindrance of the Nacnac^{DIPP} ligand. In contrast, the less bulky $[\text{Nacnac}^{\text{Ph}}\text{Al}(\text{Me})\text{FeCp}(\text{CO})_2]$ displays an absence of reactivity towards CO_2 activation, further underscoring the influence of the Nacnac^{DIPP} ligand's steric hindrance.

This first study prompted Mankad and co-workers to further explore the reactivity of complex **64** under thermal and photochemical conditions.¹⁵¹ The more intricate cooperative ring-opening reaction of THF was subjected to testing (Scheme 17B, pathway 4), yielding species **69** where Fe and Al moieties were found to be connected by a linear four-carbon chain. This transformation represents an uncommon radical cleavage of C–O bonds of THF, as typical mechanisms with other Al or Fe complexes involve cationic intermediates.^{160,161} When an episulfide was reacted with **64**, rather than forming ring-opening product, the reaction resulted in a sulfur atom transfer into the Fe–Al bond (**70**, Scheme 17B, pathway 5). Species with weak and labile bonds, such as TMS-N_3 , were also activated, yielding **71** with concomitant dimerization of the $[\text{FeCp}(\text{CO})_2]$ metalloradical fragment (Scheme 17B, pathway 6). Finally, dimerization reaction and CO insertion into the Fe–Al bond was also observed when **64** was irradiated with UV light, forming species **72**, which can be formulated as having a Fe=Fe double bond (Scheme 17B, pathway 7).

3.6.3 Miscellaneous reactivity. In 1994, Fischer and Priermeier reported a collection of heterobimetallic complexes featuring an Fe–Al bond, aiming at the synthesis of Al-containing thin films for microelectronics.¹³⁵ Using a salt metathesis approach, distinctly substituted haloaluminanes were reacted with $[\text{K}[\text{CpFe}(\text{CO})_2]]$, obtaining species **73-R** and **74** in moderate to excellent yields (Scheme 18A). The structure of such heterobimetallic complexes was unambiguously assigned by SC-XRD analysis (**73-iBu**), showing an unsupported Fe–Al unsupported bond. The Fe–Al distance (2.456(1) Å) in **73-iBu** was similar to the distance reported for the adduct $[\text{Ph}_3\text{Al-FeCp}(\text{CO})_2]^-$ **31**,⁷³ which led the authors to suggest a Z-type donor–acceptor intermetallic bond. Nonetheless, in this case no formal oxidation states were unambiguously assigned to the metal centres. Albeit these species were not tested towards small molecule activation, complex **73-Br** can undergo ligand exchange at the Al centre with alkyllithium reagents or LiBH_4 , resulting in the formation of the corresponding alkyl or borohydride derivative (Scheme 18B). Additionally, species **74** can also undergo Al-centred ligand exchange transformations to form alumacycles **73-R** when mixed with $\text{Li}-(\text{CH}_2)_3\text{-NMe}_2$. As mentioned above, these heterobimetallic Fe–Al species were employed to obtain thin films based on Al and Fe, and their reactivity towards small molecule activation and cooperative cleavage of strong bonds was not explored.

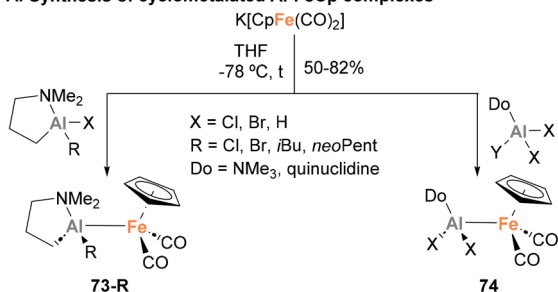
All the reported heterobimetallic Fe–Al species described so far, with exemption of Nöth's¹³⁸ and Crimmin's¹⁵⁰ examples, include a four-coordinated, fully saturated Al centre. An example of unsaturated Fe–Al species was recently introduced by Tokitoh and co-workers.^{144,146} To do so, Tbb– $\text{AlBr}_2\text{-OEt}_2$ **75** was employed as precursor (Tbb = 2,6-bis(bis(trimethylsilyl)methyl)-4-(*tert*-butyl)phenyl), which was treated with the





Scheme 17 Stoichiometric small molecule activation by a heterobimetallic Fe–Al complex via metal-radical pairs.

A. Synthesis of cyclometalated Al-FeCp complexes



Scheme 18 Salt-metathesis-based synthesis and derivatisation of Al–Fe systems.

anionic carbonyl salt $\text{K}[\text{FeCp}(\text{CO})_2]$ to obtain complex **76** in 48% yield after salt elimination (Scheme 19A). Analysis of this complex after reaction completion by ^{27}Al NMR revealed no signal, suggesting a tricoordinated structure of the Al centre that was unambiguously confirmed by SC-XRD analysis.¹⁶² Although the nature of the Fe–Al bond was studied by means of computational techniques, no discussion about the formal oxidation states of each metal was provided. Regarding its reactivity, when **76** was mixed with EBr_3 (E = Al, Ga) in toluene, a formal exchange of the aluminyl moiety occurred, generating $\text{Br}_2\text{Al-Fe}$ complex **77** (Scheme 19B, pathway 1) and $\text{Br}_2\text{Ga-Fe}$ complex **80** as dimeric species (Scheme 19B, pathway 3), in both cases with subsequent formation of Al(III) species **78**. When the reaction with AlBr_3 was performed in a coordinating solvent such as diethyl ether, monomeric complex **79** was generated (Scheme 19B, pathway 2). Contrarily, reaction of **76** with GaBr_3 in diethyl ether led to dimeric complex **80**, which can be explained invoking the higher Lewis acidity of the Al centre, making a better acceptor for weakly coordinating ligands. It is worth mentioning that **77** and **79** were analysed in depth, including Al^{27} NMR and characterisation by SC-XRD. Fe–Al bonds (**77**: 2.304(1) Å, **79**: 2.3487(8) Å) were revealed to be similar to parent Fe–Al species **76** (2.319(1) Å), and both com-





Scheme 19 Synthesis of bulky Al-Fe system and its reactivity with Lewis acids.

plexes were completely NMR silent. Computational studies of the intermetallic bond in **76** were performed using Natural Bond Order (NBO) analysis, and although formal oxidation states were not explicitly assigned, they strongly suggest the existence of an Fe(II)-Al(I) bond, where aluminium functions as an X-type donor ligand.

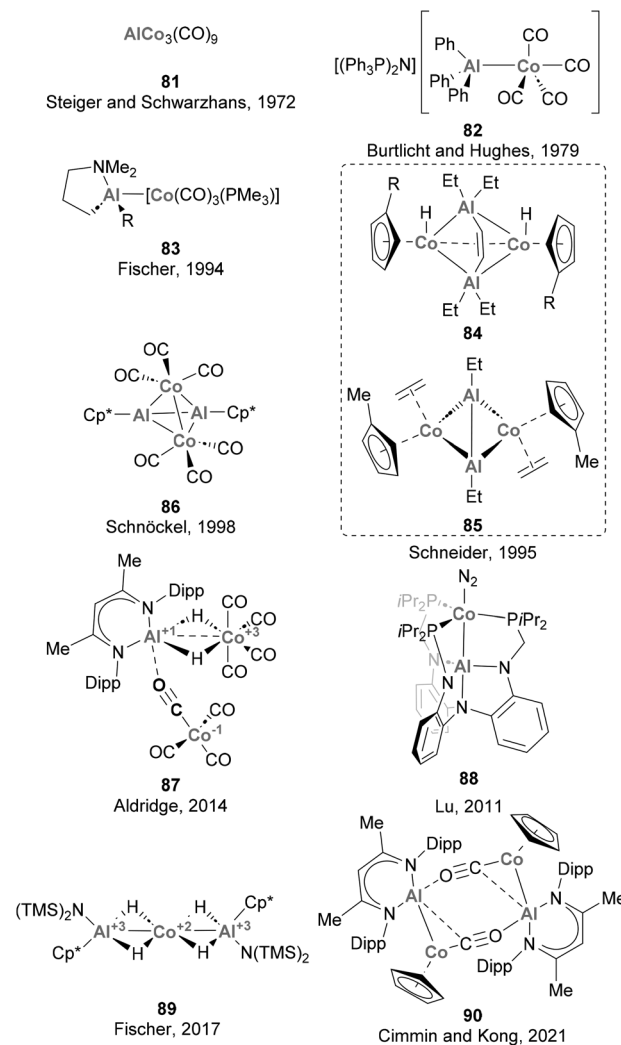
Overall, heterobimetallic species containing an Fe-Al bond have undergone extensive study since their discovery in the late 1970s. Initially considered structural curiosities, their reactivity did not attract significant attention until recent years. Notably, Driess, Crimmin, Mankad, Peters, and others have demonstrated the potential of these species in small molecule activation, including the cleavage of strong C-H bonds, leading to their application in crucial processes such as catalytic N₂-fixation reactions. Nonetheless, taking into account the importance of Fe in organic synthesis,¹⁶³ chemists have just scratched the surface of the actual potential of heterobimetallic Fe-Al species in small molecule activation. Considering the abundance and environmentally benign properties of both Fe and Al, the combination of these elements in heterobimetallic species capable of performing cooperative catalytic transformations represents a promising avenue that is likely to garner increasing attention in the future.

3.7 Cobalt

Steiger and Schwarzzhans pioneered the field of heterobimetallic aluminium-base-metal complexes in 1972 with the first example of a Co complex containing an Al-based ligand (**81**,

Scheme 20).¹⁶⁴ While this represented a milestone in the field, the full characterisation of a heterobimetallic species featuring a Co-Al had to wait more than 20 years, when Schneider and co-workers reported a SC-XRD analysis of species **84** and **85**.¹⁶⁵ Furthermore, it is worth noting that the majority of the Co-Al systems reported subsequent to these pioneering examples,^{72,73,124,125,135,142,164-166} depicted in Scheme 20, have predominantly remained as structural curiosities without significant exploration in the realms of small molecule activation and catalysis. Among these complexes, only **81** and **83** were synthesised through salt elimination, prevailing the use of direct coordination of low-valent Al species and ligand substitution reactions.

Recently, heterobimetallic species featuring a Co-Al bond have received attention from the organometallic community, and their reactivity has been explored in C-H bond and small molecule activation. However, it is worth noting that the catalytic application of such species remains very limited, as Co-Al



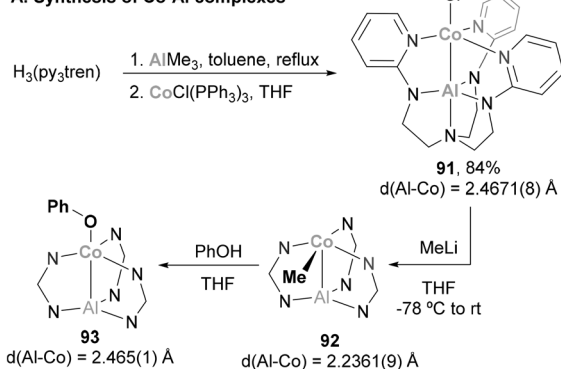
Scheme 20 Structurally characterised Co-Al complexes.



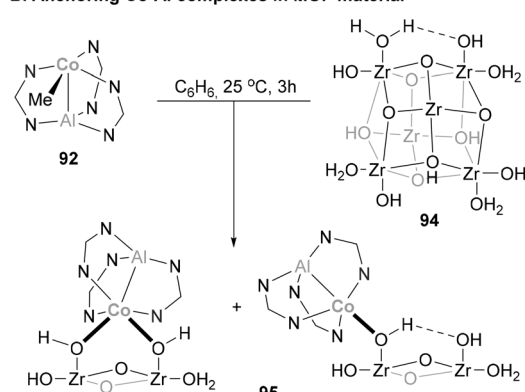
complexes find application as catalysts in CO₂ hydrogenation and oxidation of alcohols (see below).

3.7.1 Oxidation of alcohols. Taking advantage of a double-decker ligand, Lu and co-workers synthesised a heterobimetallic Al(III)–Co(I) complex for MOF functionalisation.¹⁶⁷ Starting from H₃(py₃tren), the Al–Co complex **91** was readily obtained in a 2-step procedure (Scheme 21A). The structure of **91** was unambiguously confirmed by SC-XRD experiments, featuring a Co–Al bond length of 2.4671(8) Å, 5% shorter than the sum of Al and Co covalent radii (2.60 Å),⁷⁸ indicating a donor–acceptor interaction. Transmetalation reaction with MeLi results in the formation of organometallic Co–Me species **92**. This complex was anticipated to be a good precursor for the functionalisation of metal organic frameworks (MOF), since its reaction with –OH residues on the MOF surface (**94**) was anticipated to be straightforward. In this process, organometallic species **92** was reacted with phenol, producing phenoxide complex **93** under mild conditions. Encouraged by these results, **92** was mixed with MOF **94**, obtaining Co–Al single sites on a MOF (Scheme 21B, **95**, simplified structures).

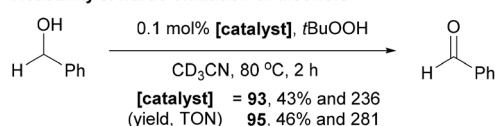
A. Synthesis of Co–Al complexes



B. Anchoring Co–Al complexes in MOF material



C. Reactivity towards oxidation of alcohols

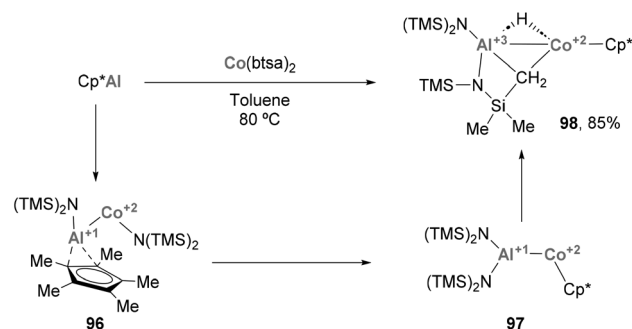


Scheme 21 Catalytic oxidation of benzyl alcohol performed by Co–Al systems anchored in MOF material (simplified structure drawn for clarity).

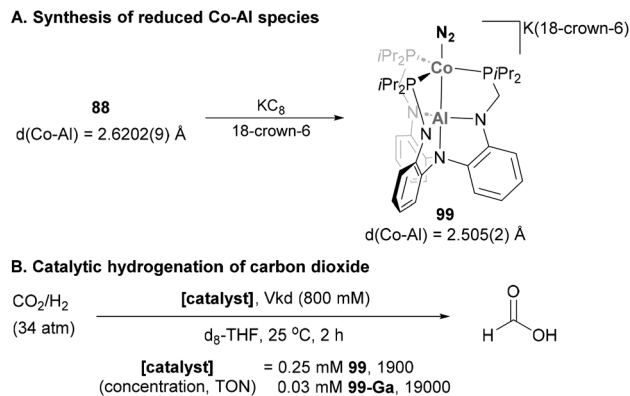
Species **93** and **95** were tested in alcohol oxidation reactions, and albeit they present similar activity towards the formation of benzaldehyde, species **94** decomposes to an ill-defined structure and is not amenable for catalyst recycling.

3.7.2 C–H activation. Heterobimetallic species featuring a Co–Al bond have also been involved in the activation of relatively inert C–H bonds. Fischer and co-workers recently reported how low valent ECp* (E = Al, Ga) species trigger Csp³–H activation of –Me residues when mixed with Co(bt₃a)₂ salts (Scheme 22), resulting in complex **98**.⁷² SC-XRD analysis of this heterobimetallic complex presents a Co–Al distance of 2.402(2) Å, slightly longer than the previously described examples^{124,165,166} but shorter than reported AlCo alloys,¹⁶⁸ while the Co–Cp* distance (1.709 Å) is in agreement with a Co(II) oxidation state (1.714 Å for Cp*₂Co vs. 1.623 Å for [Cp*₂Co][PF₆]).^{169,170} Formation of species **98** can be rationalised by a sequence of events, which is illustrated in Scheme 22. The reaction starts with the insertion of Cp*Al into a Co–N bond (**96**), followed by Cp* migration from Al to Co leading to **97** and a subsequent C–H activation of the one of the methyl groups of the bt₃a ligands, resulting in the formation of **98** in 85%. This example illustrates the still underexplored ability of systems featuring a Co–Al bonds to cleave and functionalise C–H bonds.⁶⁴ Taking into account the ability of Co to perform such transformations,^{171–174} together with precedents with heavier group 9 elements,^{58,60} intensive investigation of Co–Al systems will be crucial to develop novel methods for functionalizing C–H bonds.

3.7.3 Small molecule activation. Lu and co-workers exploited the use of AltraPhos in heterobimetallic Co–Al species containing a Z-type Co → Al interaction (Scheme 23A).¹⁴² Indeed, such interaction allowed the synthesis of stable Co–Al species **99** with a formal Co(–I) centre after one-electron reduction of complex **88** with KC₈.¹⁵⁹ Similarly to its Fe analogue (see Scheme 16), the bonded dinitrogen molecule in **99** also presents an enlarged N–N bond compared with its precursor as a result of stronger π-back donation. In addition, the Co–Al bonds in **99** is 0.12 Å shorter than in **88**, thus indicating a Co-centred reduction event that yields a reduced Co atom in a formal oxidation state value of –1. In this case, and in contrast to highly reduced Fe species



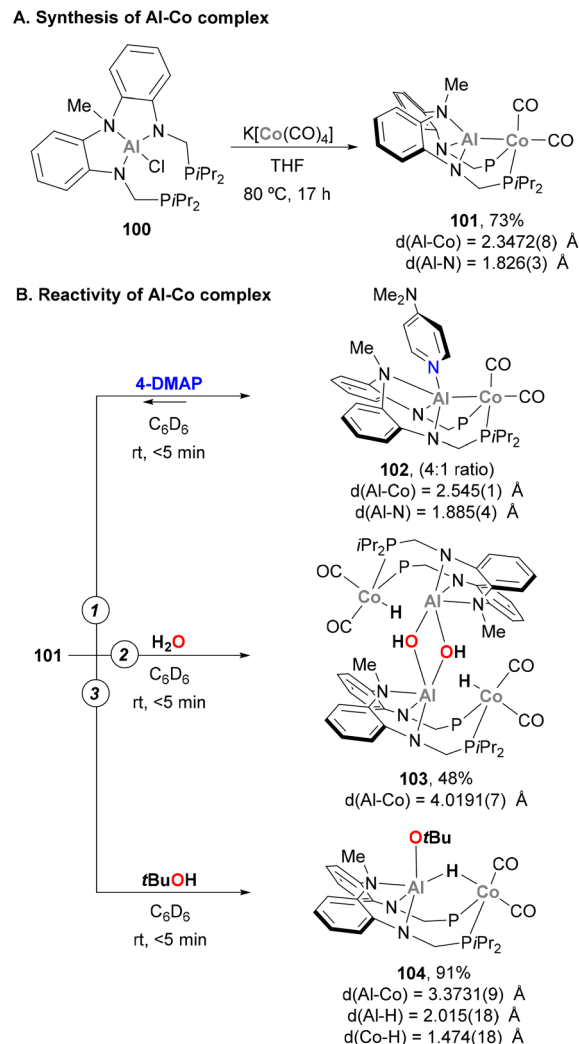
Scheme 22 Formation of Co–Al species via a C–H activation mechanism.



Scheme 23 Synthesis of reduced Co–Al species and their use in catalytic reduction of CO₂.

62, the bound dinitrogen molecule could not be functionalised with 1,2-bis(chlorodimethylsilyl)ethane as this reaction led to an intractable mixture of products. Nonetheless, the highly reduced Co–Al species **99** proved to be excellent at binding dihydrogen, a property that was exploited in catalysis to hydrogenate CO₂ (Scheme 23B).¹⁷⁵ The mechanism of CO₂ hydrogenation was only studied in depth for the Ga analogue **99-Ga**, resulting in an unprecedented Co(-I)/Co(I) catalytic cycle with a key oxidative addition of Co(-I) into the H–H bond. Hydrogenation with species **99** is proposed to proceed through an analogous mechanism, representing a unique catalytic hydrogenation example of a heterobimetallic Co–Al species.

The activation of E–H bonds using Co–Al species has also been explored by Nakao and co-workers.¹⁷⁶ Using a PAIP pincer and following a salt metathesis approach, Al(III) chloride precursor **100** was reacted with K[Co(CO)₄], yielding heterobimetallic complex **101** in 73% yield (Scheme 24A). SC-XRD analysis indicates that the Al and Co centres adopt tetrahedral and bipyramidal configurations, respectively. In addition, Co–Al (2.3472(8) Å) and Al–N_{amide} (1.826(3) Å) distances in **101** are similar to previously reported Co(I)–Al(I) complexes, suggesting a low oxidation state Co centre. Two main transformations were studied with heterobimetallic complex **101**. Firstly, the coordination ability of 4-dimethylaminopyridine (4-DMAP) was found to be reversible, illustrating the high Lewis acidity of the Al centre (**102**, Scheme 24B, pathway 1). SC-XRD analysis of **102** shows an expected elongation of Co–Al (2.545(1) Å) and Al–N_{amide} (1.885(4) Å) distances after coordination of DMAP, as well as a trigonal bipyramidal Al centre. Inspired by a recent report from Ozerov,¹⁷⁷ the ability of **101** for O–H bond activation was tested with H₂O and *t*BuOH. This resulted in the isolation of dimeric complex **103** (Scheme 24B, pathway 2) and complex **104** in 91% yield (Scheme 24B, pathway 3). It is important to note that in both cases, the integrity of the Co–Al bond was lost, as evidenced by the long Co–Al distances after O–H bond activation (4.0191(7) Å for **103** and 3.3731(9) Å for **104**), generating an Al alkoxide and a Co hydride. In complex **104**, Al–H and Co–H bond lengths suggest that the μ -H ligand is a Co hydride, with partial coordination to the Al centre. It is



Scheme 24 O–H bond activation by a PAIP Co–Al complex.

important to note that treatment of complex **101** with PhNH₂ or *n*-BuNH₂, in contrast to Ozerov's work, did not afford N–H bond activation products. This fact indicates that the O–H activation by complex **101** does not proceed *via* a cooperative mechanism and can be described as a protonation of the low oxidation state Co centre.

Contrary to heavier heterobimetallic M–Al (M = Rh and Ir) group 9 analogues,^{57–59,61,178} cooperative small molecule activation with heterobimetallic Co–Al species needs further investigation if these species are to be applied to catalytic transformations. This is an exceptional opportunity to unlock the full potential of the field and pave the way for the development of predictable and efficient transformations using inexpensive and benign elements of the d and p blocks.

3.8 Nickel

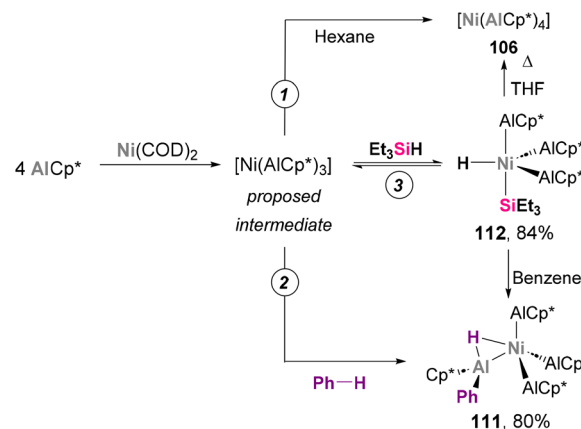
A long story of success is shared by Ni and Al when combined together, starting with their role in polymerisation reactions and the so-called “Nickel effect”,¹⁷⁹ which almost one century



ago set the basis for the discovery of the Ziegler catalysts in Mülheim an der Ruhr.^{79,180} Despite this early discovery and the potential intermediacy of Ni–Al species in hydroalumination reactions and polymerisation,^{181,182} well-defined heterobimetallic species with Ni–Al bonds remained elusive until 1995, when Schnöckel and co-workers described species **105** (Scheme 25).¹⁸³ While Ni has also been extensively used in combination with Al additives in catalytic processes, the synthesis and reactivity of heterobimetallic species with Ni–Al bonds has not received the same attention. Pioneering examples of heterobimetallic species with Ni–Al bonds whose reactivity has not been studied are depicted in Scheme 25. As shown, Ni centres coordinated to AlCp* fragments have dominated this realm from 1995 to 2014 (**105–109**),^{71,183–187} with major contributions by Fischer and co-workers.

The reactivity of Ni–Al bonds has been studied in a more systematic manner over the past five years, and relevant works are exemplified below. Among the transformations investigated, stoichiometric small molecule activation has been the most extensively reported. Notably, a recent report highlights a unique example in which a heterobimetallic complex featuring two Ni–Al bonds is used as catalyst for the dearomatisation of N-heterocycles.

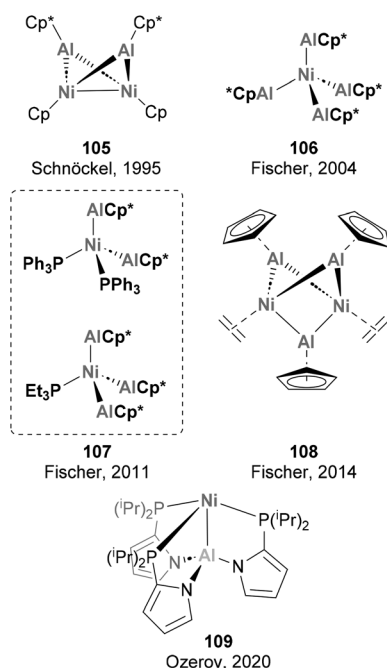
3.8.1 C–H activation. The cleavage of C–H bonds has been an extensive area of research in heterobimetallic aluminium-base-metal complexes. In 2004, Fischer and co-workers discovered several E–H (E = C, Si) activation transformations aiming to synthesise species **106**, which represented innovative reactivity examples of Ni–Al complexes.¹⁸⁴ As previously reported, treatment of Ni(COD)₂ with 4 equivalents of AlCp* in hexane results in the formation of [Ni(AlCp*)₄] **106** (Scheme 26,



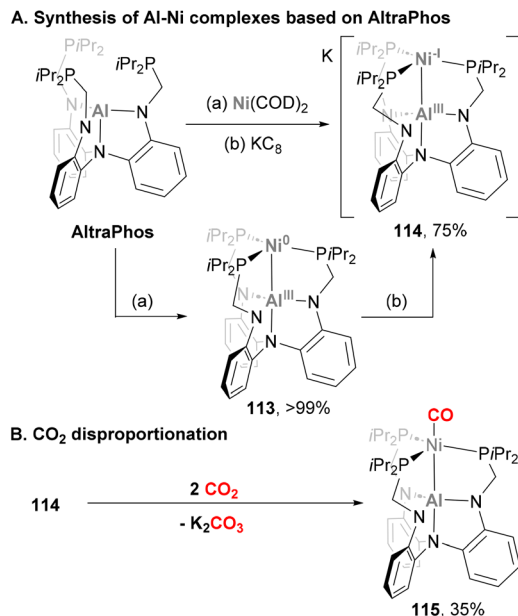
Scheme 26 Reactivity of Ni–Al complexes towards E–H (E = C, Si) bonds.

pathway 1). In an attempt to improve the yield of this transformation, the authors employed benzene as solvent, observing no traces of **106** but quantitative formation of **111** (Scheme 26, pathway 2). This product results from C–H bond cleavage of the benzene, which illustrates the ability of Ni–Al bonds to perform this transformation. SC-XRD of **111** exhibits a Ni–Al bond distance of 2.2105(11) Å (AlCp* fragment) and 2.2912(11) Å (for HAlPhCp* fragment), which suggest a +3 and +1 oxidation state for the Al centres in the H-bridged Al atom and the AlCp* ligands, respectively. Alternative control experiments indicate a 16-electron complex with chemical formula [Ni(AlCp*)₃] as intermediate, as tetracoordinated Ni species **106** was found to be chemically inert. Interestingly, when 3 equivalents of Et₃SiH were added to the mixture of Ni(COD)₂ and AlCp*, complex **112** was obtained in high yield after a reversible Ni-centred Si–H bond insertion (Scheme 26, pathway 3). SC-XRD analysis of **112** reveals a distorted trigonal bipyramidal geometry, with Ni–Al distances between 2.180(7) and 2.208(10) Å. In a later work by the same authors, this reversibility was employed for the selective synthesis of [Ni(PEt₃)(AlCp*)₃] **107**, which cannot be accessed through conventional one-pot procedures.¹⁸⁶

3.8.2 Small molecule activation. Complexes bearing a Ni–Al bond have also been applied to small molecule activation, albeit most of their reactivity so far involves ligand exchange studies. An example of such processes was reported by Lu and co-workers using AltraPhos as ligand (Scheme 27),¹⁴² similarly to Fe and Co complexes **38** and **88**, respectively. Reaction of Ni(COD)₂ with AltraPhos yields a diamagnetic, formally Ni(0)–Al(III) complex **113**.¹⁸⁸ Interestingly, ²⁷Al NMR studies together with a Ni–Al distance of 2.450(1) Å suggests direct Ni–Al bonding, presumably *via* a Ni → Al dative bond. A complex recently reported by Ozerov and co-workers (**109**, Scheme 25) displays similar features compared with **113**, although in this work a tris(*N*-pyrrolyl)aluminium ligand was used.¹⁸⁷ Contrary to its Ga and In analogues, **113** was shown to be inactive in catalytic olefin hydrogenation. In a subsequent study from Lu and co-workers, reduction of Ni–alane species **113** with KC₈



Scheme 25 Structurally characterised Ni–Al complexes.



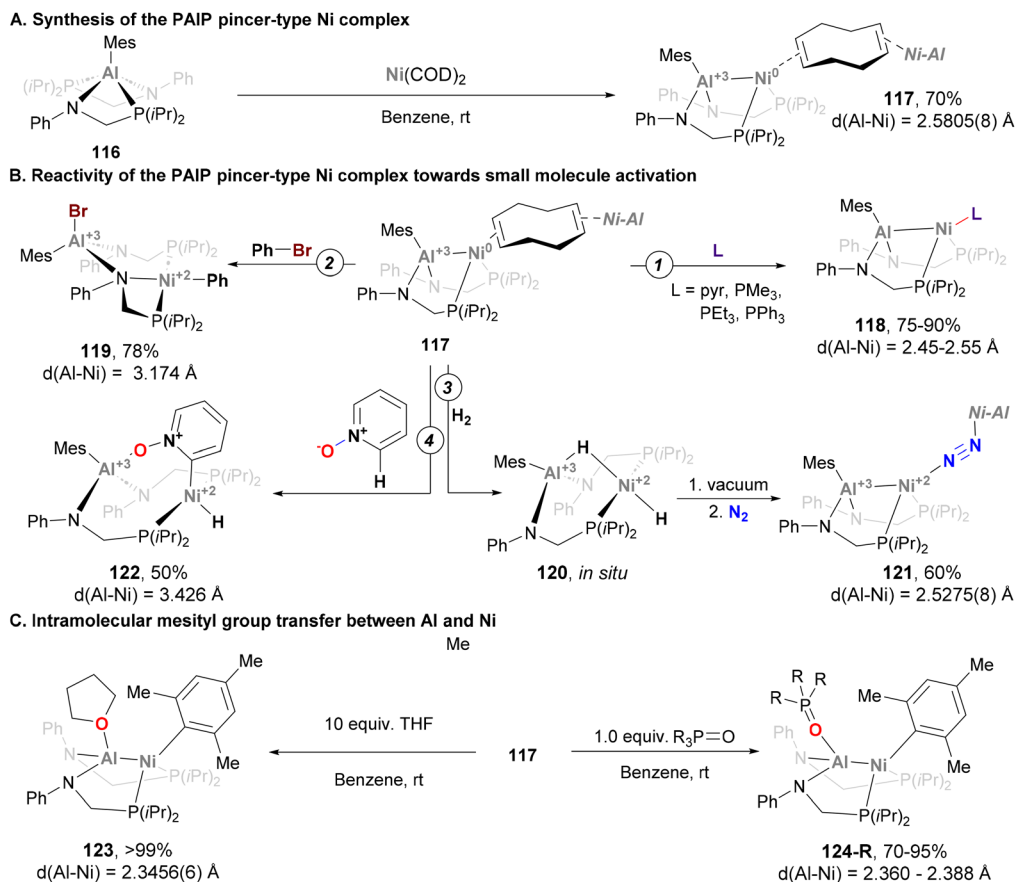
Scheme 27 Synthesis and reactivity of Ni-alane species based on AltraPhos.

resulted in a dark red solution containing radical anion **114**,¹⁸⁹ which was crystallised in the presence of cryptand crypt-222. SC-XRD studies revealed that the Ni–Al bond distance in **114** contracts to 2.389(1) Å as a result of a Ni-centred reduction and a subsequent stronger Ni → Al donation. Furthermore, combined experimental and computational studies of such species support its formulation as formally Ni(–I)–Al(III), with an unexpected stabilisation arising from a one-electron polar σ –(Ni–Al) and a two-electron $(3d_{z^2}) \rightarrow \text{Al}$ dative bond. The reactivity of this open-shell Ni–Al species towards CO₂ activation was then tested.¹⁹⁰ Stirring **114** under a CO₂ atmosphere yielded K₂CO₃ and a 1 : 1 mixture of **113** and CO adduct **115** via a disproportionation reaction. Formation of such species was rationalised via a nucleophilic attack of metalloradical **114** to CO₂, followed by a reduction mediated by another molecule of species **114** to generate a formally dianionic nickel diolate. The latter species is proposed to insert a second CO₂ molecule to generate a Ni di-CO₂ intermediate, which decays to CO₃^{2–} and heterobimetallic complex **115**. This species was independently synthesized by treatment of **114** with ethyl formate, confirming its structural features. SC-XRD analysis resulted in a Ni–Al bond distance of 2.626(1) Å, which suggests the Ni–Al bond in **115** is weaker compared with **114**, probably due to the strong π -acceptor ability of CO as ligand.

In the example depicted in Scheme 27, the Al atom is buried, and its role is limited to indirectly adjusting the reactivity of the Ni centre. In a recent example, Lu and co-workers designed a Ni–Al system featuring a ligand platform that enables substrates to interact with both Ni and Al centres (Scheme 28), and promote cooperative activation of substrates.¹⁹¹ Treatment of the corresponding Al-based metalloligand **116** with Ni(COD)₂ yields a dimer heterobimetallic pincer

species (**117**) bridged by cyclooctadiene (Scheme 28A). Complex **117** has been fully characterised, and its SC-XRD analysis revealed an intriguing Ni–Al distance of 2.5805(8) Å, which is longer than the sum of covalent radii. This bond length, together with the observed pyramidalization of the Al centre suggest a weak Z-type Ni → Al interaction.^{78,192} After characterisation of heterobimetallic species **117**, its reactivity towards ligand substitution with pyridine and phosphines was tested, resulting in the formation of species **118** (Scheme 28B, pathway 1). This species was also fully characterised, revealing that ligand substitution occurs exclusively at the Ni centre, showing virtually identical Ni–Al bond lengths compared with parent complex **117**. More interestingly, cooperative activation of small molecules was investigated taking advantage of the open coordination environment of the bimetallic Ni–Al unit. As the authors point out in their study, it is important to note that the Ni → Al interaction breaks after substrate activation as a consequence of Ni(0) being oxidised to Ni(II). As an example of C–X activation, Ph–Br was reacted with **117**, furnishing species **119** (Scheme 28B, pathway 2). This example, which represents the first Ph–Br oxidative addition into an aluminium–base–metal bond, shows ligation of the halide to the Al centre, while the phenyl group coordinates the Ni atom. Addition of H₂ to **117** generates species **120**, which features a bridged Al–H–Ni and a terminal hydride Ni–H moieties, similarly to previously reported Ni → B species.¹⁹³ Interestingly, complex **120** releases H₂ upon exposure to vacuum or Ar and converts to **121** when stored under a N₂ atmosphere (Scheme 28B, pathway 3). This reactivity shows that the Ni → Al bond interaction not only facilitates cooperative activation of substrates, but also promotes reductive elimination. The open coordination environment of the bimetallic Ni–Al unit also allows the Al centre to direct substrates for selective C–H activation at the Ni site. Indeed, this reactivity was demonstrated by the reaction of **117** with pyridine *N*-oxide, which resulted in the *ortho* C–H oxidative addition product **122** in 50% yield (Scheme 28B, pathway 3). Spectroscopic evidence and SC-XRD analysis is consistent with a Ni centre coordinated by a terminal hydride ligand, as well as a C₂-cyclometalated pyridine *N*-oxide moiety. Importantly, the Al centre engages in a strong O → Al interaction (1.814(4) Å) with the pyridine *N*-oxide ligand, which indicates a coordination prior to the C–H oxidative addition event. This transformation is highly relevant, as it represents a unique example of selective *ortho* C–H activation of pyridine *N*-oxides with well-defined heterobimetallic complexes pairing a base metal with Al. An intriguing reaction occurred when species **117** was reacted with weak donors such as THF or phosphine oxides (**123** and **124**, respectively; Scheme 28C). SC-XRD revealed a Ni–Al complex where the mesityl group shifted from Al to Ni, with a significant contraction of the Ni–Al distance from 2.5805(8) Å to 2.3456(6) Å. These features suggest a ligand-responsive conversion of a Ni-alane to a Ni–aluminyl complex, which is reversible upon addition of more COD ligand. Computational analysis shows that the Ni–Al in **123** and **124** lies in between a Ni(0) → Al(III) interaction and a polarised covalent Ni(I)–Al(II) bond. Overall,





Scheme 28 Synthesis and reactivity of PAIP Ni–Al complex.

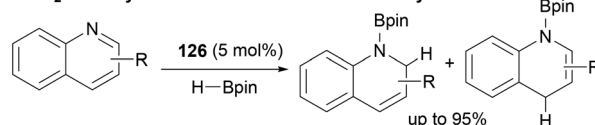
the rich reactivity in cooperative activation of substrates, together with its reversible redox processes, represent key features that make species **117** a good candidate to be applied in catalytic transformations.

Very recently, Shoshani and co-workers developed a well-defined heterobimetallic Ni–Al complex (**126**) with catalytic application,¹⁹⁴ which was synthesised through metalation of the Al(III)–H precursor **125** with Ni(COD)₂ (Scheme 29A). SC-XRD analysis unveiled a trimetallic NiAl₂(μ₂-H)₂, in which both Al atoms are bridged to the Ni centre by a hydride ligand. In addition, the Ni centre adopts a distorted tetrahedral geometry consistent with an 18 electron Ni(0) centre. In addition, Ni–Al distances were found to be short (2.3629(7) and 2.3658(6) Å), indicating a strong interaction between both atoms. Indeed, these distances are consistent with previously reported Ni–alane species,^{142,187} and are shorter than the sum of Ni and Al covalent radii.⁷⁸ Based on these structural features, the oxidation state of the metal centres was assigned as +3 for Al and 0 for Ni. Considering the importance of hydride species in hydrogenation and hydroboration of heterocyclic compounds, the ability of **126** towards the catalytic hydroboration of quinoline was tested (Scheme 29B). Using HBpin as hydride source and catalytic quantities of **126** (5 mol%), quinoline was hydrogenated quantitatively. Catalytic amounts of starting Al(III)–H **125** species or Ni(COD)₂ resulted in lower yields and slower

A. Synthesis of Al₂Ni complex



B. Al₂Ni-catalysed dearomatisation of N-heterocycles



Scheme 29 Synthesis and reactivity of Ni–alane species featuring two Ni–Al bonds.

reaction rates, which highlights the synergistic effect of the Ni–Al bond and its potential in catalysis. Nonetheless, the authors highlight that further studies are needed to identify the responsible catalytically active species and to determine the true nature of the Ni–Al bond. Albeit the field is still in its infancy, this pioneering study opens the door to a new horizon



of catalytic reactivity and the application of heterobimetallic Ni–Al complexes in organic synthesis and catalytic small molecule functionalisation.

3.8.3 Miscellaneous reactivity. Heterobimetallic Ni–Al species have also been used to produce metallic clusters. In 2014, Fischer and co-workers described the synthesis of novel all-hydrocarbon ligand-stabilised binuclear clusters of metal-core composition Ni₂Zn₇Al (Scheme 30).⁷¹ To do so, species **127**, which is synthesised through an unconventional pathway involving transmetallation, was reacted with 8.0 equiv. of ZnMe₂. After Al/Zn ligand exchange, a 55:45 mixture of organometallic **128** and **129** species was obtained and products were fully characterised by computational studies, spectroscopic techniques and SC-XRD, revealing short Ni–Zn distances. Furthermore, an open-shell configuration was assigned to species **129**, which makes it an exceptional example within mononuclear Zn-rich molecules that strictly follow closed-shell 18 valence electron counting. Furthermore, the discovery of such Ni₂Zn₇Al species from heterobimetallic Ni–Al complexes opens the door to new strategies for the synthesis of larger heterometallic aggregates.

In summary, the synthesis and reactivity of heterobimetallic species featuring a Ni–Al bond have opened new avenues in C–H functionalization, small molecule activation, production of new metallic clusters, and, importantly, catalytic processes for the functionalization of organic molecules. It is worth mentioning that, although not emphasized in this review, Ni–Al species have also shown promise as efficient catalysts for polymerization.¹⁹⁵ The pioneering studies reported in this section have laid the foundation for further exploration. In fact, we believe future research in this area will likely focus on developing catalytic systems and exploring the potential of heterobimetallic Ni–Al species in synthetic transformations.

3.9 Copper

Forging Cu–Al bonds has attracted increasing attention in recent years, particularly since the discovery of aluminyl anions.^{48,49,51} To date, more than a dozen of examples have been reported, although most of them only contributed to

expanding the collection of structures featuring a Cu–Al intermetallic bond, and their reactivity was not further investigated (Scheme 31). The synthesis of compounds featuring Cu–Al bonds can be tracked down to a study from 2013 by Bourissou, Bouhadir, Miqueu and co-workers,¹⁵⁸ who reported two complexes featuring Z-type Cu(I)–Al(III) interactions, **130** and **131**. After this groundbreaking example, a series of complexes featuring Cu–Al bonds was recently synthesised and fully characterised by Crimmin (**132** and **135**),^{125,196} Power (**133**),¹⁹⁷ Liu (**134**),¹²⁶ Lu (**135**)¹⁹⁸ and Muñoz-Hernández (**137**).¹⁹⁹ Synthesis of such heterobimetallic Cu–Al species was achieved through conventional methods, such as ligand substitution and direct complexation of neutral low-valent Al(I) species, while salt metathesis has been more extensively utilised when Cu was paired to Al starting from Al(I) anions, as reported by Hill and McMullin and Aldridge.^{200,201} While the species depicted in Scheme 31 were not tested towards small molecule activation, the reactivity of Cu–Al species with X-type Al(I) ligands has seen significant advances during the last couple of years (see below), demonstrating promising potential for catalysis.

3.9.1 Small molecule activation. In 2014, a report on the synthesis and reactivity of the heterobimetallic Cu₆Al₆ cluster was published by Fischer and co-workers (Scheme 32).²⁰² Species **138**, which is synthesised by treating [(Ph₃PCu)₆H₆] with Cp*Al in a 2:3 ratio, was revealed to contain a Cu₆ unit

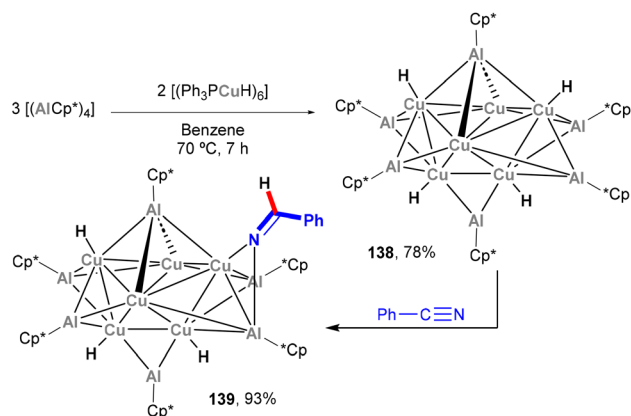


Scheme 30 Reactivity of Ni₂Al₃ cluster towards transmetallation with ZnMe₂.



Scheme 31 Structurally characterised Cu–Al complexes.





Scheme 32 Synthesis of Cu₆Al₆ clusters and their surface reactivity with PhCN.

inside an octahedral Al₆ moiety, with the resulting Cu₆Al₆ core being wrapped in a shell of Cp* and H ligands. Interestingly, species **138** offers the opportunity to investigate the surface reactivity of the (Cu₆Al₆)H₄ unit towards hydrogenation of unsaturated organic functional groups. Indeed, **138** was shown to react with benzonitrile, as depicted in Scheme 32, furnishing **139** in quantitative yield as the 1 : 1 insertion product. The reactivity of these clusters was not studied in further detail, but their hydrogenation ability exemplifies the potential of complexes bearing Cu–Al bonds in small molecule activation and functionalisation of chemical feedstock.

McMullin, Hill and co-workers have largely contributed to this area during the last three years. Inspired by Aldridge's breakthrough in the field of anionic Al(I) species,⁴⁸ the authors designed a novel seven-membered heterocyclic potassium diamidoaluminum salt (**140**, Scheme 33) with formula [K{Al(SiN^{Dipp})₂}]₂.²⁰³ Through a salt elimination route, anionic Al(I) species **140** was treated with Cu(I) chloride carbene complexes, [(NHC^{iPr})CuCl] and [(Me²CAAC)CuCl],²⁰⁰ which acted as precursors to synthesise heterobimetallic Cu–Al species **141** and **142** in 89% and 79% yield, respectively (Scheme 33A, pathways 1 and 2). In a later work by Hill, Mahon and McMullin, the same methodology was employed for the synthesis of Cu phosphine complex **143** in 78% yield.²⁰⁴ The three complexes were isolated and characterised through SC-XRD, unveiling Cu–Al (**141**: 2.3449(4) Å, **142**: 2.4028(7) Å, **143**: 2.3755(3) Å) and Al–N (**141**: 1.8464(10) Å, **142**: 1.8607(18) Å, **143**: 1.848(1) Å) distances in agreement with previously reported complexes (Scheme 31), suggesting a dative X-type Al(I) → Cu(I) bond. The nature of the Cu–Al intermetallic bond was further studied by testing the reaction of Cu–Al complexes with heteroallenes. Thus, when **141–143** were treated with *N,N'*-diisopropylcarbodiimide, complexes **144–146** were obtained in excellent yields (Scheme 33B, in blue). Noteworthy, as shown in Scheme 36B, the insertion mode of *N,N'*-diisopropylcarbodiimide into the Cu–Al bond is modulated by the ligand attached to the Cu centre, furnishing a Cu–C bond (**145** and **146**) or a Cu–N bond (**144**). Reaction of heterobimetallic species featuring a Cu–Al bond with CO₂

resulted in **147–149** in good yields (Scheme 33B, in red). Interestingly, CO₂ inserts into the Cu–Al bond, which cleaves after activation with concomitant oxidation of Al(I) to Al(III). These results highlight the ambiphilic character of the aluminyl–Cu species. Indeed, computational mechanistic insight into these transformations indicates a polarised Cu^{δ−}–Al^{δ+} bond, in which the Al centre acts as an electrophilic site and the Cu atom as a nucleophile. Decomposition of species **149** results in oxo-bridged species **150**, which reacts with another molecule of CO₂ to eventually furnish carbonate-bridged species **151**. Further computational studies by Lin, Sheong and co-workers²⁰⁵ concluded that the rate-determining insertion of CO₂ and carbodiimide is followed by several rearrangements, supporting McMullin and Hill's work. The insertion, however, is proposed to be initiated by the nucleophilic attack of the σ-(Cu–Al) bond on the heteroallene coupling partner, providing valuable insight for developing future Cu–Al-mediated reactivity.

Indeed, Hill and McMullin recently investigated the reactivity of Cu–Al species towards terminal alkynes.²⁰⁶ When complex **141** was treated with three equivalents of a terminal alkyne RCCH, bis(alkynyl)aluminates **155** or **156** were obtained with concomitant extrusion of the corresponding alkene (Scheme 33B, in green). An in-depth study allowed the authors to determine the sequence of intermediates involved in this transformation. Thus, after a first insertion of the Cu–Al bond into the alkyne C–H bond, cuprous (hydrido)(alkynyl)aluminates **152** and **153** are formed, in a transformation that could be classified as a Csp–H activation event. This step is followed by a hydroalumination reaction of a second molecule of alkyne, which furnishes (*E*-alkenyl)(alkynyl)aluminates. These intermediates could be isolated and characterised crystallographically only when alkynes bearing bulky aryl substituents were used (**154**). Finally, the last RCCH equivalent undergoes σ-bond metathesis with the alkenyl fragment of the (*E*-alkenyl)(alkynyl)aluminates *via* a Cu-acetylide intermediate, releasing the corresponding alkene and bis(alkynyl)aluminates **155** and **156** after Cu-to-Al alkyne migration. It is important to note that the experimental study was fully supported by DFT calculations, which provided further insight into the mechanism of this unprecedented reaction. The overall process represented a distinctive scenario of alkyne transfer semi-hydrogenation, in which the combination of the protic acetylenic C–H bond and the reducing ability of the σ-(Cu–Al) bond furnishes the necessary hydride source.

Aldridge and Goicoechea took advantage of their groundbreaking nucleophilic aluminium anion reported in 2018,⁴⁸ and together with Zhao and Frenking contributed to the development of heterobimetallic complexes pairing aluminium with Cu. Following McMullin and Hill's work on Cu–Al species,²⁰⁰ Aldridge reported the synthesis of xanthene-supported Al(I)-coinage metal bimetallics and their reactivity towards CO₂ insertion reactions (Scheme 34). The xanthene-supported Al(I) anion **157** was treated with equimolar amounts of copper–phosphine complex Ph₃PCuI, affording the bisaluminyl cuprate complex **158** in 50% yield after salt elimination





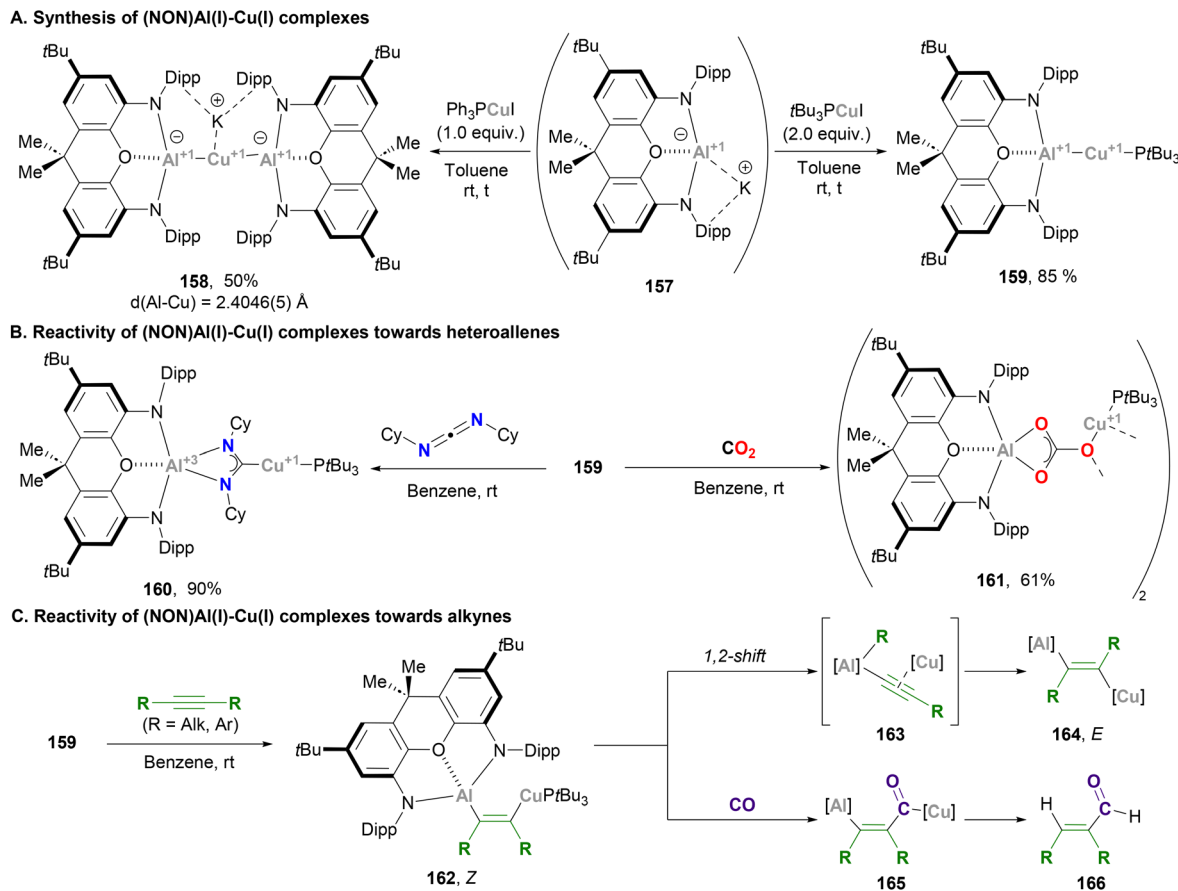
Scheme 33 Synthesis and reactivity of Cu–Al species reported by Mohan, McMullin and Hill.

and ligand exchange (Scheme 34A, left). The solid-state structure of **158** features a Cu–Al bond average distance of 2.4046(5) Å, which is shorter than the sum of their covalent radii (2.53 Å) and in agreement with previously reported Cu(I)–Al(I) species (*vide supra*). With the goal of synthesising 1 : 1 bimetallic systems with Cu salts that allow retention of the Cu–P linkage, phosphines with stronger σ -donor abilities were explored. Indeed, when the reaction was performed with 2.0 equivalents of *t*Bu₃PCuI, species **159** was obtained (Scheme 34A, right). Albeit there was no SC-XRD of this complex, its structure was unambiguously assigned by means of spectroscopic experiments and comparison with Ag and Ga analogues. After structural characterisation studies, the reactivity of Cu–Al bonds was interrogated with heteroallenes, including CO₂. Thus, reaction of complex **159** with *N,N'*-diisopropylcarbodiimide led to the formation of species **160** (Scheme 34B, pathway 1), whereas stirring **159** under a CO₂ atmosphere led to carbonate **161** (Scheme 34B, pathway 2). Mechanistic

studies unveiled that, similarly to previous examples,²⁰⁰ these products arise from an initial nucleophilic attack of the σ (Cu–Al) bond on the heteroallene, which yields species featuring μ - κ^1 (C): κ^2 (E,E') bridging units and Cu–C/Al–E bonds. Formation of carbonate species **161** was rationalised *via* a similar mechanism compared with species **151** (Scheme 33), which involves a rate-limiting CO extrusion and the intermediacy of a reactive μ -oxide Cu–O–Al species.

A follow-up study from Aldridge and co-workers focused on the reactivity of xanthene-based heterobimetallic Cu–Al systems towards C–C triple bonds (Scheme 34C).²⁰⁷ Contrary to the Cu–Al system reported by Hill and McMullin,²⁰⁶ when alkynes are reacted with **159**, spectroscopic features of the product suggest the formation of *Z*-alkene **162**. Indeed, SC-XRD analysis of the product with R = Et shows addition of the Cu–Al unit across the alkyne triple bond in *syn* fashion, which represents the first example of an (aluminumalkenyl) copper complex and the first structurally characterised 1,2-





Scheme 34 Synthesis and reactivity of Cu–Al species reported by Frenking, Goicoechea and Aldridge.

hetero-dimetallation by insertion of an alkyne into a heterobimetallic M–M' bond. Interestingly, if the reaction times are extended complete consumption of **162** is observed, and spectroscopic analysis of the crude reaction indicates formation of the corresponding *anti* dimetallated alkene **164**. The reaction mechanism of this isomerisation from *Z* to *E* is reminiscent of a recent report by Yamashita and co-workers describing the chemistry of B–Au species.²⁰⁸ Based on this mechanism, isomerisation from **162** is proposed to occur through Cu-alkynyl intermediate **163**, which can produce formal *anti* or *syn* dimetallation products after a 1,2-shift of the R group. Evidence of this mechanism was obtained by isolating the corresponding (NON)Al–Et complex, whose formation unambiguously demonstrates that C-to-Al migration is viable. Computational studies from Sorbelli, Belpassi and Belanzoni provided further insight into this mechanism and supported the idea of a cooperative radical-like reactivity of the Cu–Al bond.²⁰⁹ Reactivity of **162**, which contains a Cu–C and an Al–C bond, was also explored in the presence of CO. This resulted in an unprecedented copper acyl complex (**165**) after insertion of CO into the Cu–C bond, while the Al–C bond remained intact. It is important to note that this reaction was observed exclusively with the *Z*-(aluminyl-alkenyl)copper complex, which highlights the role played by Al, which offers additional stabilisation of the acyl unit in **165**

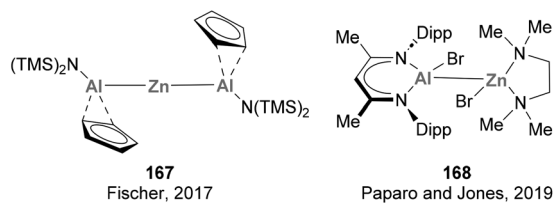
via an O → Al interaction. In addition, hydrolysis of such carbonylated species results in the formation of α,β -unsaturated aldehydes **166** *via* the net hydroformylation of an alkyne.

Overall, the recent eruption of heterobimetallic species featuring a Cu–Al bond has the potential to revolutionise the field of small molecule activation, as they present unprecedented reactivity modes. Based on the examples reported in this section, together with the breadth of application of boryl-based Cu species in organic synthesis,⁴³ the field of aluminyl–Cu species is expected to grow in the future, providing new and unexplored reactivity avenues for the most abundant, benign and inexpensive coinage metal.

3.10 Zinc

Heterobimetallic complexes bearing a covalent Zn–Al bond were reported a few years ago by Fischer and co-workers while studying the reactivity of 1st-row transition metals complexes bearing btsa ligands towards ECp* (E = Al, Ga) units.⁷² Insertion of AlCp* into the Zn–N bond of Zn(btsa)₂ resulted in complex **167**, with concomitant reduction of the Zn centre and formation of the first example of a heterobimetallic complex featuring a Zn–Al bond. Small molecule activation pathways of this early example, together with a recent Zn–Al species



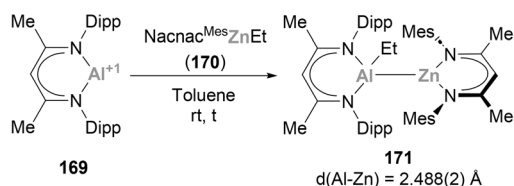


Scheme 35 Structurally characterised Zn–Al complexes.

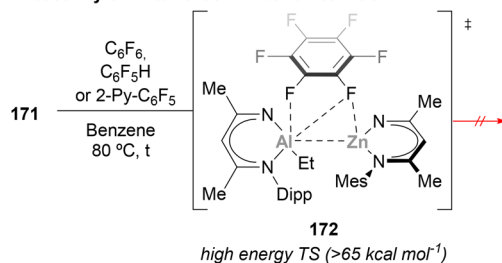
reported by Paparò and Jones (**168**), were not further investigated (Scheme 35).

3.10.1 Small molecule activation. The reactivity of heterobimetallic complexes featuring a Zn–Al bond remained unexplored until 2018, when Crimmin and co-workers synthesised a collection of heterobimetallic complexes that included an aluminium–base-metal complex featuring an L-type Zn–Al interaction.²¹⁰ To do so, neutral Al(i) precursor Nacnac^{Dipp}Al (**169**) was treated with Nacnac^{Mes}ZnEt (**170**), affording complex **171** (Scheme 36A) after a formal insertion of the Al(i) reagent into a Zn–C bond. SC-XRD analysis of such species revealed a Zn–Al bond length of 2.4877(10) Å, which was found to be similar to the Zn–Al distances observed by Fischer (**167**, 2.448 (2) Å).⁷² The reactivity of these complexes was tested for C–F bond activation. Contrary to heterobimetallic species featuring a Zn–Mg interaction, which proved to be active in C–F bond cleavage, complex **171** remained unreacted when mixed with a variety of fluorinated (hetero)arenes (Scheme 36B). This inactivity towards C–F bond activation was attributed to two main factors: the tetrahedral geometry of the Al centre, which diminished its Lewis acidity; and the steric hindrance of both Nacnac ligands, which disfavours the required disposition in the C–F bond activation TS (**172**). Additionally, computational studies are in agreement with the experimental results, revealing a large energetic barrier for this transformation with a value of $\Delta G^\ddagger = 65.1 \text{ kcal mol}^{-1}$.

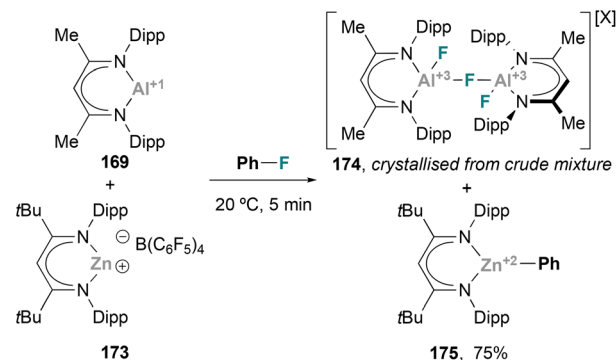
A. Synthesis of Zn–Al complex based on MesNacNac ligand



B. Reactivity of xx towards C–F bond activation



Scheme 36 Synthesis and attempted C–F bond activation with Zn–Al species.

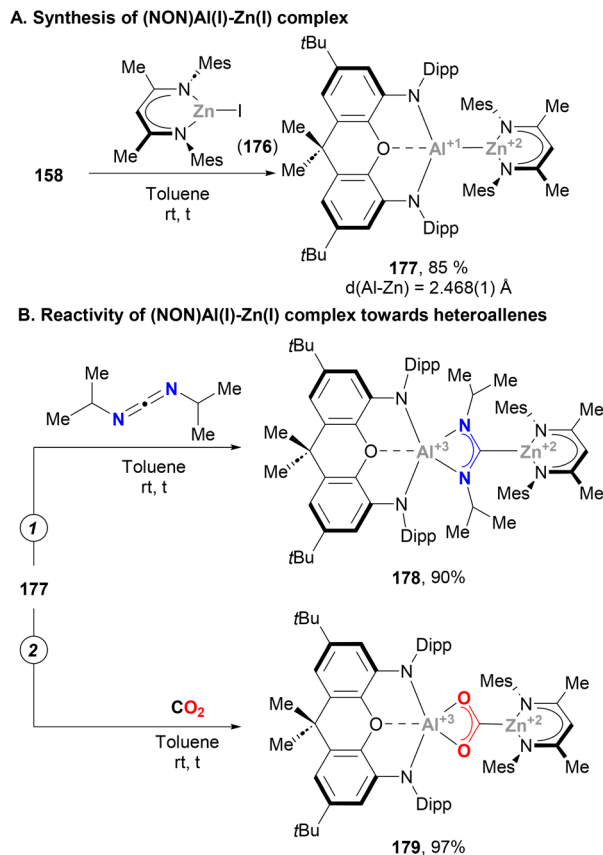


Scheme 37 Successful C–F bond activation with Zn–Al species via FLP-type mechanism.

In 2021, Harder and co-workers successfully achieved C–F bond activation using a heterobimetallic Zn–Al system.²¹¹ In this work, the synthesis of Zn–Al complexes from **169** and **173** was attempted using fluorobenzene as solvent (Scheme 37). Instead of the corresponding heterobimetallic species bearing a Zn–Al bond, the reaction resulted in mixture of a variety of species, from which Zn and Al complexes **174** and **175** were identified. These species are the result of a fast C–F bond activation of fluorobenzene, an unprecedented result with heterobimetallic Zn–Al systems that does not occur with Zn/Ga, Mg/Al or Ca/Al metal pairs.^{59,212–215} The authors speculated about the possibility of an intermediate species featuring a Zn–Al bond as they were able to isolate and characterise the corresponding Zn–Ga analogue. Nonetheless, computational studies indicate that C–F bond activation does not proceed through a previously formed heterobimetallic complex. Instead, the transformation is proposed to proceed *via* an FLP-type S_NAr reaction in which Al(i) species **169** acts as nucleophile while Zn cation **173** activates fluorobenzene, making it more electrophilic.

The synthesis of an unambiguously characterised low valent Zn(II)–Al(i) complex was recently reported by Aldridge, Goicoechea and co-workers taking advantage of the practicality of the salt elimination approach with anionic Al(i) species **158**.²¹⁶ Indeed, the reaction between **158** and Nacnac^{Mes}ZnI (**176**) results in the formation of a well-defined complex featuring a Zn–Al bond in 85% yield (**177**, Scheme 38A), the structure of which was unambiguously confirmed by SC-XRD analysis. Furthermore, such analysis revealed a Zn–Al bond length of 2.468(1) Å, which falls within the range of previously reported distances for heterobimetallic species with a Zn–Al bond (*vide supra*). Complementary computational studies suggest a high degree of Zn–Al covalency, which was further tested by means of reactivity towards heteroallenes such as diisopropylcarbodiimide and CO₂ (Scheme 38B, pathways 1 and 2, respectively). Although in both cases addition into the Zn–Al bond was observed, furnishing species **178** and **179**, only the CO₂ adduct could be isolated and fully characterised. As shown in Scheme 38B, insertion of CO₂ generates new Zn–C and Al–O bonds, unveiling the nucleophilic character of the Zn

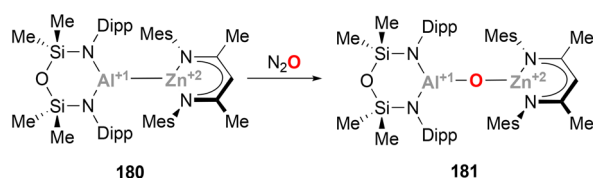




Scheme 38 Synthesis and reactivity of (NON)Al–Zn species towards heteroallenes.

centre, the electrophilic role played by the Al atom and the ambiphilic character of the heterobimetallic complex.

A similar heterobimetallic Zn(II)–Al(I) complex (**180**) featuring a SIONOSi backbone has been recently reported by McMullin and Coles (Scheme 39).²¹⁷ Complex **180** presents a Zn–Al bond length of 2.4860(5) Å, almost identical to species **177**, and shows the same reactivity pattern with CO₂. Interestingly, McMullin and Coles have been able to isolate and fully characterise the corresponding μ -oxo bridge Zn–O–Al species (**181**), which has been previously proposed to participate as intermediate in the conversion of CO₂ to carbonate in bimetallic systems.²⁰¹ Indeed, when species **181** was treated with CO₂, a carbonate species analogous to **162** was obtained, demonstrating the intermediacy of μ -oxo M–O–Al species.



Scheme 39 Reactivity of (SiNONSi)Al–Zn towards nitrous oxide.

4. Conclusion and prospect

In summary, heterobimetallic complexes pairing abundant base metals with aluminium are emerging as promising alternatives to noble metals in small molecule activation, mainly due to the synergy of aluminium with 3d centres. The studies showcased in this review demonstrate the potential of aluminium-based ligands to confer nobility to base metals and their potential to unveil cooperative catalytic methods for the functionalisation of organic molecules. Paralleling recent applications with noble metals (*vide supra*), implementation of heterobimetallic Al–base-metal complexes in C–X cleavage (X = hydrogen, halogen, ether, amine, *etc.*) and small molecule activation has gained significant attention, particularly with the development of X-type aluminyl anions. Such species have proved to be exceptional ligands not only for transition metal centres but also for various elements within the main group.^{203,216,218} We believe that this field will see further advancements combining forces through a multi-disciplinary approach, where the expertise of synthetic and computational chemists will be key to enhance the understanding of fundamental organometallic cooperative events involving heterobimetallic Al–BM complexes. Additionally, efforts should be directed towards expanding the catalytic applications beyond hydrogenation, exploring the integration of heterobimetallic complexes with popular sustainable technologies such as photochemical,²¹⁹ electrochemical and mechanochemical transformations. Furthermore, it is essential to extend the scope of heterobimetallic sustainable chemistry to encompass the main group elements and explore the potential applications of Al–MG (MG = main group) species in small molecule activation. This expansion would open new avenues for utilizing abundant resources in catalysis and contribute to the development of more sustainable chemical processes. By pursuing these directions, it will be possible to unlock the full potential of aluminium-based heterobimetallic complexes and drive innovation towards a greener and more efficient future in synthetic and coordination chemistry.

Author contributions

All authors participated in the writing, reviewing, and editing of the manuscript. O. P. supervised the project. Additionally, all authors have given their approval for the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Financial support for this work was provided by Queen Mary University of London. This project has received funding from



the Royal Society (Research Grant RGS/R1/221326) and the Engineering and Physical Sciences Research Council (EP/X019306/1). S. Fernández thanks the “Fundación Ramón Areces” for a Postdoctoral Fellowship (BEVP34A6835). We thank Dr S. Arseniyadis for insightful discussions and generous support.

References

- R. H. Crabtree, *The Organometallic Chemistry of the Transition Metals*, John Wiley & Sons, Hoboken, NJ, 2005.
- J. F. Hartwig, *Organotransition Metal Chemistry: From Bonding to Catalysis*, University Science Books, Mill Valley, CA, 2010.
- A. de Meijere and F. Diederich, *Metal-Catalyzed Cross-Coupling Reactions*, Wiley-VCH Verlag GmbH & Co. KGaA, Mörtenbach, Germany, 2004.
- P. Nuss and M. J. Eckelman, *PLoS One*, 2014, **9**, e101298.
- R. M. Bullock, J. G. Chen, L. Gagliardi, P. J. Chirik, O. K. Farha, C. H. Hendon, C. W. Jones, J. A. Keith, J. Klosin, S. D. Minter, R. H. Morris, A. T. Radosevich, T. B. Rauchfuss, N. A. Strotman, A. Vojvodic, T. R. Ward, J. Y. Yang and Y. Surendranath, *Science*, 2020, **369**, eabc3183.
- P. J. Chirik and R. Morris, *Acc. Chem. Res.*, 2015, **48**, 2495–2495.
- Y. Feng, S. Long, X. Tang, Y. Sun, R. Luque, X. Zeng and L. Lin, *Chem. Soc. Rev.*, 2021, **50**, 6042–6093.
- T. Irrgang and R. Kempe, *Chem. Rev.*, 2019, **119**, 2524–2549.
- L. Rocard, D. Chen, A. Stadler, H. Zhang, R. Gil, S. Bezzenine and J. Hannedouche, *Catalysts*, 2021, **11**, 674.
- R. Arevalo and P. J. Chirik, *J. Am. Chem. Soc.*, 2019, **141**, 9106–9123.
- T. Zell and R. Langer, *ChemCatChem*, 2018, **10**, 1930–1940.
- P. L. Holland, *Acc. Chem. Res.*, 2015, **48**, 1696–1702.
- P. J. Chirik, K. M. Engle, E. M. Simmons and S. R. Wisniewski, *Org. Process Res. Dev.*, 2023, **27**(7), 1160–1184.
- B. Su, Z.-C. Cao and Z.-J. Shi, *Acc. Chem. Res.*, 2015, **48**, 886–896.
- C. B. Larsen and O. S. Wenger, *Chem. – Eur. J.*, 2018, **24**, 2039–2058.
- J. E. Zweig, D. E. Kim and T. R. Newhouse, *Chem. Rev.*, 2017, **117**, 11680–11752.
- B. Su, Z.-C. Cao and Z.-J. Shi, *Acc. Chem. Res.*, 2015, **48**, 886–896.
- R. J. M. K. Gebbink and M.-E. Moret, *Non-Noble Metal Catalysis: Molecular Approaches and Reactions*, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, Germany, 2019.
- O. R. Luca and R. H. Crabtree, *Chem. Soc. Rev.*, 2013, **42**, 1440–1459.
- A. Nakada, T. Matsumoto and H.-C. Chang, *Coord. Chem. Rev.*, 2022, **473**, 214804.
- J. I. van der Vlugt, *Chem. – Eur. J.*, 2019, **25**, 2651–2662.
- M. R. Elsby and R. T. Baker, *Chem. Soc. Rev.*, 2020, **49**, 8933–8987.
- J. I. van der Vlugt, *Eur. J. Inorg. Chem.*, 2012, **2012**, 363–375.
- M. D. Wodrich and X. Hu, *Nat. Rev. Chem.*, 2017, **2**, 1–7.
- D. G. A. Verhoeven and M.-E. Moret, *Dalton Trans.*, 2016, **45**, 15762–15778.
- K.-S. Feichtner and V. H. Gessner, *Chem. Commun.*, 2018, **54**, 6540–6553.
- A. Singh and D. Gelman, *ACS Catal.*, 2020, **10**, 1246–1255.
- D. V. Gutsulyak, W. E. Piers, J. Borau-Garcia and M. Parvez, *J. Am. Chem. Soc.*, 2013, **135**, 11776–11779.
- R. A. Manzano and R. D. Young, *Coord. Chem. Rev.*, 2021, **449**, 214215.
- A. Singh and D. Gelman, *ACS Catal.*, 2020, **10**, 1246–1255.
- P. M. Keil, T. Szilvási and T. J. Hadlington, *Chem. Sci.*, 2021, **12**, 5582–5590.
- P. M. Keil and T. J. Hadlington, *Angew. Chem., Int. Ed.*, 2022, **61**, e202114143.
- P. M. Keil, A. Soyemi, K. Weisser, T. Szilvási, C. Limberg and T. J. Hadlington, *Angew. Chem., Int. Ed.*, 2023, **62**, e202218141.
- J. Kim, Y.-E. Kim, K. Park and Y. Lee, *Inorg. Chem.*, 2019, **58**, 11534–11545.
- J. Kim, *Bull. Korean Chem. Soc.*, 2022, **43**, 538–548.
- J. A. Cabeza and P. García-Álvarez, *Chem. – Eur. J.*, 2023, **29**, e202203096.
- S. González-Gallardo, T. Bollermann, R. A. Fischer and R. Murugavel, *Chem. Rev.*, 2012, **112**, 3136–3170.
- C. Gemel, T. Steinke, M. Cokoja, A. Kempter and R. A. Fischer, *Eur. J. Inorg. Chem.*, 2004, **2004**, 4161–4176.
- G. Linti and H. Schnöckel, *Coord. Chem. Rev.*, 2000, **206**, 285–319.
- M. Asay, C. Jones and M. Driess, *Chem. Rev.*, 2011, **111**, 354–396.
- Y. Segawa, M. Yamashita and K. Nozaki, *Science*, 2006, **314**, 113–115.
- Y. Segawa, M. Yamashita and K. Nozaki, *Organometallics*, 2009, **28**, 6234–6242.
- D. Hemming, R. Fritzscheier, S. A. Westcott, W. L. Santos and P. G. Steel, *Chem. Soc. Rev.*, 2018, **47**, 7477–7494.
- F. Kong, P. Ríos, C. Hauck, F. J. Fernández-de-Córdova, D. A. Dickie, L. G. Habgood, A. Rodríguez and T. B. Gunnoe, *J. Am. Chem. Soc.*, 2023, **145**, 179–193.
- T.-P. Lin and J. C. Peters, *J. Am. Chem. Soc.*, 2014, **136**, 13672–13683.
- P. Ríos, A. Rodríguez and J. López-Serrano, *ACS Catal.*, 2016, **6**, 5715–5723.
- F. W. Seidel and K. Nozaki, *Angew. Chem., Int. Ed.*, 2022, **61**, e202111691.
- J. Hicks, P. Vasko, J. M. Goicoechea and S. Aldridge, *Nature*, 2018, **557**, 92–95.
- J. Hicks, P. Vasko, J. M. Goicoechea and S. Aldridge, *Angew. Chem., Int. Ed.*, 2021, **60**, 1702–1713.
- M. P. Coles and M. J. Evans, *Chem. Commun.*, 2023, **59**, 503–519.



- 51 K. Hobson, C. J. Carmalt and C. Bakewell, *Chem. Sci.*, 2020, **11**, 6942–6956.
- 52 M. Zhong, S. Sinhababu and H. W. Roesky, *Dalton Trans.*, 2020, **49**, 1351–1364.
- 53 Y. Liu, J. Li, X. Ma, Z. Yang and H. W. Roesky, *Coord. Chem. Rev.*, 2018, **374**, 387–415.
- 54 N. Hara, K. Semba and Y. Nakao, *ACS Catal.*, 2022, **12**, 1626–1638.
- 55 A. L. Allred and E. G. Rochow, *J. Inorg. Nucl. Chem.*, 1958, **5**, 264–268.
- 56 J. Takaya and N. Iwasawa, *J. Am. Chem. Soc.*, 2017, **139**, 6074–6077.
- 57 S. Morisako, S. Watanabe, S. Ikemoto, S. Muratsugu, M. Tada and M. Yamashita, *Angew. Chem., Int. Ed.*, 2019, **58**, 15031–15035.
- 58 N. Hara, T. Saito, K. Semba, N. Kuriakose, H. Zheng, S. Sakaki and Y. Nakao, *J. Am. Chem. Soc.*, 2018, **140**, 7070–7073.
- 59 I. Fujii, K. Semba, Q.-Z. Li, S. Sakaki and Y. Nakao, *J. Am. Chem. Soc.*, 2020, **142**, 11647–11652.
- 60 N. Hara, K. Aso, Q.-Z. Li, S. Sakaki and Y. Nakao, *Tetrahedron*, 2021, **95**, 132339.
- 61 R. Seki, N. Hara, T. Saito and Y. Nakao, *J. Am. Chem. Soc.*, 2021, **143**, 6388–6394.
- 62 I. Fujii, K. Semba and Y. Nakao, *Org. Lett.*, 2022, **24**, 3075–3079.
- 63 S. Ogoshi, M. Ueta, T. Aral and H. Kurosawa, *J. Am. Chem. Soc.*, 2005, **127**, 12810–12811.
- 64 C.-S. Wang, S. Di Monaco, A. N. Thai, M. S. Rahman, B. P. Pang, C. Wang and N. Yoshikai, *J. Am. Chem. Soc.*, 2020, **142**, 12878–12889.
- 65 T. Zhang, Y.-X. Luan, N. Y. S. Lam, J.-F. Li, Y. Li, M. Ye and J.-Q. Yu, *Nat. Chem.*, 2021, **13**, 1207–1213.
- 66 R. A. Fischer and J. Weiß, *Angew. Chem., Int. Ed.*, 1999, **38**, 2830–2850.
- 67 C. Gemel, T. Steinke, M. Cokoja, A. Kempter and R. A. Fischer, *Eur. J. Inorg. Chem.*, 2004, **2004**, 4161–4176.
- 68 M. J. Butler and M. R. Crimmin, *Chem. Commun.*, 2017, **53**, 1348–1365.
- 69 J. P. Collman, *Acc. Chem. Res.*, 1975, **8**, 342–347.
- 70 A. Noor, G. Glatz, R. Müller, M. Kaupp, S. Demeshko and R. Kempe, *Nat. Chem.*, 2009, **1**, 322–325.
- 71 M. Molon, C. Gemel, P. Jerabek, L. Trombach, G. Frenking and R. A. Fischer, *Inorg. Chem.*, 2014, **53**, 10403–10411.
- 72 J. Weßing, C. Göbel, B. Weber, C. Gemel and R. A. Fischer, *Inorg. Chem.*, 2017, **56**, 3517–3525.
- 73 J. M. Burlitch, M. E. Leonowicz, R. B. Petersen and R. E. Hughes, *Inorg. Chem.*, 1979, **18**, 1097–1105.
- 74 K. Sugita and M. Yamashita, *Chem. – Eur. J.*, 2020, **26**, 4520–4523, and references within.
- 75 J. P. Collman, R. G. Finke, J. M. Cawse and J. I. Brauman, *J. Am. Chem. Soc.*, 1977, **99**, 2515–2526.
- 76 S. M. Rummelt, P. O. Peterson, H. Zhong and P. J. Chirik, *J. Am. Chem. Soc.*, 2021, **143**, 5928–5936.
- 77 G. Feng, K. L. Chan, Z. Lin and M. Yamashita, *J. Am. Chem. Soc.*, 2022, **144**, 22662–22668.
- 78 L. Pauling, *J. Am. Chem. Soc.*, 1947, **69**, 542–553.
- 79 K. Ziegler, E. Holzkamp, H. Breil and H. Martin, *Angew. Chem.*, 1955, **67**, 541–547.
- 80 G. Natta, *J. Polym. Sci.*, 1955, **16**, 143–154.
- 81 E. P. Beaumier, A. J. Pearce, X. Y. See and I. A. Tonks, *Nat. Rev. Chem.*, 2019, **3**, 15–34.
- 82 L. Tebben, C. Mück-Lichtenfeld, G. Fernández, S. Grimme and A. Studer, *Chem. – Eur. J.*, 2016, **23**, 5864–5873.
- 83 M. L. Cooper and J. B. Rose, *J. Chem. Soc.*, 1959, 795–802.
- 84 E. H. Adema, H. Bos and C. H. Vrinssen, *Recl. Trav. Chim. Pays-Bas*, 1960, **79**, 1282–1288.
- 85 P. E. M. Allen, J. K. Brown and R. M. S. Obaid, *Trans. Faraday Soc.*, 1963, **59**, 1808–1814.
- 86 G. Natta and F. Danusso, *Stereoregular Polymers and Stereospecific Polymerizations*, Pergamon Press Ltd, Oxford, United Kingdom, 1967.
- 87 A. J. Amass, J. N. Hay and J. C. Robb, *Br. Polym. J.*, 1969, **1**, 277–281.
- 88 J. N. Hay and R. M. S. Obaid, *Eur. Polym. J.*, 1978, **14**, 965–969.
- 89 J. Holton, M. F. Lappert, D. G. H. Ballard, R. Pearce, J. L. Atwood and W. E. Hunter, *J. Chem. Soc., Dalton Trans.*, 1979, 45–53.
- 90 B. M. Bulychev, S. E. Tokareva, G. L. Soloveichik and E. V. Evdokimova, *J. Organomet. Chem.*, 1979, **179**, 263–273.
- 91 V. E. Lvovsky, E. A. Fushman and F. S. Dyachkovsky, *J. Mol. Catal.*, 1980, **10**, 43–56.
- 92 B. M. Bulichev, E. V. Evdokimova, A. I. Sizov and G. L. Soloveichik, *J. Organomet. Chem.*, 1982, **239**, 313–320.
- 93 E. B. Lobkovskii, G. L. Soloveichik, A. I. Sisov, B. M. Bulychev, A. I. Gusev and N. I. Kirillova, *J. Organomet. Chem.*, 1984, **265**, 167–173.
- 94 E. B. Lobkovskii, G. L. Soloveichik, A. I. Sizov and B. M. Bulychev, *J. Organomet. Chem.*, 1985, **280**, 53–66.
- 95 J. Poláček, H. Antropiusová, V. Hanuš, L. Petrusová and K. Mach, *J. Mol. Catal.*, 1985, **29**, 165–180.
- 96 U. M. Dzhemilev and O. S. Vostrikova, *J. Organomet. Chem.*, 1985, **285**, 43–51.
- 97 E. B. Lobkovskii, A. I. Sizov, B. M. Bulychev, I. V. Sokolova and G. L. Soloveichik, *J. Organomet. Chem.*, 1987, **319**, 69–75.
- 98 A. I. Sizov, I. V. Molodnitskaya, B. M. Bulychev, E. V. Evdokimova, G. L. Soloveichik, A. I. Gusev, E. B. Chuklanova and V. I. Andrianov, *J. Organomet. Chem.*, 1987, **335**, 323–330.
- 99 A. I. Sizov, I. V. Molodnitskaya, B. M. Bulychev, V. K. Bel'skii and G. L. Soloveichik, *J. Organomet. Chem.*, 1988, **344**, 185–193.
- 100 T. Y. Sokolova, A. I. Sizov, B. M. Bulychev, E. A. Rozova, V. K. Belsky and G. L. Soloveichik, *J. Organomet. Chem.*, 1990, **388**, 11–19.
- 101 M. Bochmann, *J. Chem. Soc., Dalton Trans.*, 1996, 255–270.
- 102 D. G. Kelly, A. J. Toner, N. M. Walker, S. J. Coles and M. B. Hursthouse, *Polyhedron*, 1996, **15**, 4307–4310.



- 103 J. P. Corden, W. Errington, P. Moore and M. G. H. Wallbridge, *Chem. Commun.*, 1999, 323–324.
- 104 P. Arndt, A. Spannenberg, W. Baumann, S. Becke and U. Rosenthal, *Eur. J. Inorg. Chem.*, 2001, **2001**, 2885–2890.
- 105 M. P. Coles and P. B. Hitchcock, *J. Chem. Soc., Dalton Trans.*, 2001, 1169–1171.
- 106 V. Taberero and T. Cuenca, *Eur. J. Inorg. Chem.*, 2005, **2005**, 338–346.
- 107 G. B. Nikiforov, H. W. Roesky, P. G. Jones, R. B. Oswald and M. Noltemeyer, *Dalton Trans.*, 2007, 4149–4159.
- 108 G. K. P. Dathara and D. S. Mainardi, *Mol. Simul.*, 2008, **34**, 201–210.
- 109 A. Hernán-Gómez, A. Martín, M. Mena and C. Santamaría, *Dalton Trans.*, 2013, **42**, 5076–5084.
- 110 T. Oswald, M. Diekmann, A. Frey, M. Schmidtman and R. Beckhaus, *Acta Crystallogr., Sect. E: Crystallogr. Commun.*, 2017, **73**, 691–693.
- 111 P. Corradini and A. Sirigu, *Inorg. Chem.*, 1967, **6**, 601–605.
- 112 K. Sugita and M. Yamashita, *Organometallics*, 2020, **39**, 2125–2129.
- 113 E. del Horno, J. Jover, M. Mena, A. Pérez-Redondo and C. Yélamos, *Chem. – Eur. J.*, 2022, **28**, e202103085.
- 114 H. J. de Liefde Meijer, J. W. G. van den Hurk and G. J. M. van der Kerk, *Recl. Trav. Chim. Pays-Bas*, 1966, **85**, 1025–1038.
- 115 A. G. Evans, J. C. Evans and E. H. Moon, *J. Chem. Soc., Dalton Trans.*, 1974, 2390–2395.
- 116 A. Anundskås and H. A. Øye, *J. Inorg. Nucl. Chem.*, 1975, **37**, 1609–1619.
- 117 H. Hagen, J. Boersma and G. van Koten, *Chem. Soc. Rev.*, 2002, **31**, 357–364.
- 118 I. E. Soshnikov, N. V. Semikolenova, A. A. Shubin, K. P. Bryliakov, V. A. Zakharov, C. Redshaw and E. P. Talsi, *Organometallics*, 2009, **28**, 6714–6720.
- 119 M. M. Schulte, E. Herdtweck, G. Raudaschl-Sieber and R. A. Fischer, *Angew. Chem., Int. Ed. Engl.*, 1996, **35**, 424–426.
- 120 R. A. Fischer, M. M. Schulte, J. Weiss, L. Zsolnai, A. Jacobi, G. Huttner, G. Frenking, C. Boehme and S. F. Vyboishchikov, *J. Am. Chem. Soc.*, 1998, **120**, 1237–1248.
- 121 H. Fölsing, O. Segnitz, U. Bossek, K. Merz, M. Winter and R. A. Fischer, *J. Organomet. Chem.*, 2000, **606**, 132–140.
- 122 Q. Yu, A. Purath, A. Donchev and H. Schnöckel, *J. Organomet. Chem.*, 1999, **584**, 94–97.
- 123 I. M. Riddlestone, S. Edmonds, P. A. Kaufman, J. Urbano, J. I. Bates, M. J. Kelly, A. L. Thompson, R. Taylor and S. Aldridge, *J. Am. Chem. Soc.*, 2012, **134**, 2551–2554.
- 124 J. A. B. Abdalla, I. M. Riddlestone, J. Turner, P. A. Kaufman, R. Tirfoin, N. Phillips and S. Aldridge, *Chem. – Eur. J.*, 2014, **20**, 17624–17634.
- 125 R. Y. Kong and M. R. Crimmin, *Dalton Trans.*, 2021, **50**, 7810–7817.
- 126 X. Zhang and L. L. Liu, *Angew. Chem., Int. Ed.*, 2021, **60**, 27062–27069.
- 127 X. Cong and X. Zeng, *Synlett*, 2021, **32**, 1343–1353.
- 128 X. Cong and X. Zeng, *Acc. Chem. Res.*, 2021, **54**, 2014–2026.
- 129 I. M. Riddlestone, J. Urbano, N. Phillips, M. J. Kelly, D. Vidovic, J. I. Bates, R. Taylor and S. Aldridge, *Dalton Trans.*, 2012, **42**, 249–258.
- 130 D. W. Agnew, C. E. Moore, A. L. Rheingold and J. S. Figueroa, *Dalton Trans.*, 2017, **46**, 6700–6707.
- 131 R. M. Philip, S. Radhika, C. M. A. Abdulla and G. Anilkumar, *Adv. Synth. Catal.*, 2021, **363**, 1272–1289.
- 132 W. Liu and L. Ackermann, *ACS Catal.*, 2016, **6**, 3743–3752.
- 133 W. Liu and J. T. Groves, *Acc. Chem. Res.*, 2015, **48**, 1727–1735.
- 134 R. Jamatia, A. Mondal and D. Srimani, *Adv. Synth. Catal.*, 2021, **363**, 2969–2995.
- 135 R. A. Fischer and T. Priermeier, *Organometallics*, 1994, **13**, 4306–4314.
- 136 H. Braunschweig, J. Müller and B. Ganter, *Inorg. Chem.*, 1996, **35**, 7443–7444.
- 137 J. Weiss, D. Stetzkamp, B. Nuber, R. A. Fischer, C. Boehme and G. Frenking, *Angew. Chem., Int. Ed. Engl.*, 1997, **36**, 70–72.
- 138 B. N. Anand, I. Krossing and H. Nöth, *Inorg. Chem.*, 1997, **36**, 1979–1981.
- 139 T. Steinke, M. Cokoja, C. Gemel, A. Kempter, A. Krapp, G. Frenking, U. Zenneck and R. A. Fischer, *Angew. Chem., Int. Ed.*, 2005, **44**, 2943–2946.
- 140 C. Jones, S. Aldridge, T. Gans-Eichler and A. Stasch, *Dalton Trans.*, 2006, 5357–5361.
- 141 B. Buchin, C. Gemel, A. Kempter, T. Cadenbach and R. A. Fischer, *Inorg. Chim. Acta*, 2006, **359**, 4833–4839.
- 142 P. A. Rudd, S. Liu, L. Gagliardi, V. G. Young Jr. and C. C. Lu, *J. Am. Chem. Soc.*, 2011, **133**, 20724–20727.
- 143 G. Tan, T. Szilvási, S. Inoue, B. Blom and M. Driess, *J. Am. Chem. Soc.*, 2014, **136**, 9732–9742.
- 144 T. Agou, T. Yanagisawa, T. Sasamori and N. Tokitoh, *Bull. Chem. Soc. Jpn.*, 2016, **89**, 1184–1186.
- 145 D. Dange, C. P. Sindlinger, S. Aldridge and C. Jones, *Chem. Commun.*, 2016, **53**, 149–152.
- 146 T. Yanagisawa, Y. Mizuhata and N. Tokitoh, *Heteroat. Chem.*, 2018, **29**, e21465.
- 147 J. Fajardo Jr. and J. C. Peters, *Inorg. Chem.*, 2021, **60**, 1220–1227.
- 148 K. Nakaya, A. Ishii and N. Nakata, *Mendeleev Commun.*, 2022, **32**, 71–73.
- 149 S. Sinhababu, M. R. Radzhabov, J. Telser and N. P. Mankad, *J. Am. Chem. Soc.*, 2022, **144**, 3210–3221.
- 150 N. Gorgas, A. J. P. White and M. R. Crimmin, *J. Am. Chem. Soc.*, 2022, **144**, 8770–8777.
- 151 S. Sinhababu and N. P. Mankad, *Organometallics*, 2022, **41**, 1917–1921.
- 152 J. Uddin and G. Frenking, *J. Am. Chem. Soc.*, 2001, **123**, 1683–1693.
- 153 R. Lehmann and M. Schlosser, *Tetrahedron Lett.*, 1984, **25**, 745–748.
- 154 R. B. Bates, L. M. Kroposki and D. E. Potter, *J. Org. Chem.*, 1972, **37**, 560–562.



- 155 N. Gorgas, A. J. P. White and M. R. Crimmin, *Chem. Commun.*, 2022, **58**, 10849–10852.
- 156 B. Stadler, N. Gorgas, A. J. P. White and M. R. Crimmin, *Angew. Chem., Int. Ed.*, 2023, **62**, e202219212.
- 157 M.-E. Moret and J. C. Peters, *J. Am. Chem. Soc.*, 2011, **133**, 18118–18121.
- 158 M. Sircoglou, N. Saffon, K. Miqueu, G. Bouhadir and D. Bourissou, *Organometallics*, 2013, **32**, 6780–6784.
- 159 P. A. Rudd, N. Planas, E. Bill, L. Gagliardi and C. C. Lu, *Eur. J. Inorg. Chem.*, 2013, **2013**, 3898–3906.
- 160 J. P. Campbell and W. L. Gladfelter, *Inorg. Chem.*, 1997, **36**, 4094–4098.
- 161 R. F. Moreira, E. Y. Tshuva and S. J. Lippard, *Inorg. Chem.*, 2004, **43**, 4427–4434.
- 162 T. P. Hanusa, *Encyclopedia of Inorganic and Bioinorganic Chemistry*, John Wiley & Sons, Ltd, 2015, pp. 1–25.
- 163 I. Bauer and H.-J. Knölker, *Chem. Rev.*, 2015, **115**, 3170–3387.
- 164 K. E. Schwarzans and H. Steiger, *Angew. Chem., Int. Ed. Engl.*, 1972, **11**, 535–535.
- 165 J. J. Schneider, C. Krüger, M. Nolte, I. Abraham, T. S. Ertel and H. Bertagnolli, *Angew. Chem., Int. Ed. Engl.*, 1995, **33**, 2435–2437.
- 166 C. Üffing, A. Ecker, R. Köppe and H. Schnöckel, *Organometallics*, 1998, **17**, 2373–2375.
- 167 A. B. Thompson, D. R. Pahls, V. Bernales, L. C. Gallington, C. D. Malonzo, T. Webber, S. J. Tereniak, T. C. Wang, S. P. Desai, Z. Li, I. S. Kim, L. Gagliardi, R. L. Penn, K. W. Chapman, A. Stein, O. K. Farha, J. T. Hupp, A. B. F. Martinson and C. C. Lu, *Chem. Mater.*, 2016, **28**, 6753–6762.
- 168 P. J. Black, *Acta Metall.*, 1956, **4**, 172–179.
- 169 M. M. Clark, W. W. Brennessel and P. L. Holland, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2009, **65**, 391–391.
- 170 M. A. Laffey and P. Thornton, *J. Chem. Soc., Dalton Trans.*, 1982, 313–318.
- 171 L. Lukasevics, A. Cizikovs and L. Grigorjeva, *Chem. Commun.*, 2021, **57**, 10827–10841.
- 172 M. Moselage, J. Li and L. Ackermann, *ACS Catal.*, 2016, **6**, 498–525.
- 173 S. M. Ujwaldev, N. A. Harry, M. A. Divakar and G. Anilkumar, *Catal. Sci. Technol.*, 2018, **8**, 5983–6018.
- 174 K. Gao and N. Yoshikai, *Acc. Chem. Res.*, 2014, **47**, 1208–1219.
- 175 M. V. Vollmer, J. Ye, J. C. Linehan, B. J. Graziano, A. Preston, E. S. Wiedner and C. C. Lu, *ACS Catal.*, 2020, **10**, 2459–2470.
- 176 K. Semba, F. Shimoura and Y. Nakao, *Chem. Lett.*, 2022, **51**, 455–457.
- 177 Y. Cao, W.-C. Shih and O. V. Ozerov, *Organometallics*, 2019, **38**, 4076–4081.
- 178 L. Escomel, I. Del Rosal, L. Maron, E. Jeanneau, L. Veyre, C. Thieuleux and C. Camp, *J. Am. Chem. Soc.*, 2021, **143**, 4844–4856.
- 179 K. Fischer, K. Jonas, P. Misbach, R. Stabba and G. Wilke, *Angew. Chem., Int. Ed. Engl.*, 1973, **12**, 943–953.
- 180 K. Ziegler, E. Holzkamp, H. Breil and H. Martin, *Angew. Chem.*, 1955, **67**, 426–426.
- 181 J. J. Eisch, X. Ma, M. Singh and G. Wilke, *J. Organomet. Chem.*, 1997, **527**, 301–304.
- 182 J. J. Eisch, S. R. Sexsmith and K. C. Fichter, *J. Organomet. Chem.*, 1990, **382**, 273–293.
- 183 C. Dohmeier, H. Krautscheid and H. Schnöckel, *Angew. Chem., Int. Ed. Engl.*, 1995, **33**, 2482–2483.
- 184 T. Steinke, C. Gemel, M. Cokoja, M. Winter and R. A. Fischer, *Angew. Chem., Int. Ed.*, 2004, **43**, 2299–2302.
- 185 B. Buchin, T. Steinke, C. Gemel, T. Cadenbach and R. A. Fischer, *Z. Anorg. Allg. Chem.*, 2005, **631**, 2756–2762.
- 186 M. Molon, T. Bollermann, C. Gemel, J. Schaumann and R. A. Fischer, *Dalton Trans.*, 2011, **40**, 10769–10774.
- 187 Q. Lai, M. N. Cosio and O. V. Ozerov, *Chem. Commun.*, 2020, **56**, 14845–14848.
- 188 R. C. Cammarota and C. C. Lu, *J. Am. Chem. Soc.*, 2015, **137**, 12486–12489.
- 189 M. V. Vollmer, J. Xie, R. C. Cammarota, V. G. Young Jr., E. Bill, L. Gagliardi and C. C. Lu, *Angew. Chem., Int. Ed.*, 2018, **57**, 7815–7819.
- 190 M. V. Vollmer, R. C. Cammarota and C. C. Lu, *Eur. J. Inorg. Chem.*, 2019, **2019**, 2140–2145.
- 191 B. J. Graziano, M. V. Vollmer and C. C. Lu, *Angew. Chem., Int. Ed.*, 2021, **60**, 15087–15094.
- 192 B. Cordero, V. Gómez, A. E. Platero-Prats, M. Revés, J. Echeverría, E. Cremades, F. Barragán and S. Alvarez, *Dalton Trans.*, 2008, 2832–2838.
- 193 W. H. Harman and J. C. Peters, *J. Am. Chem. Soc.*, 2012, **134**, 5080–5082.
- 194 E. De Leon, F. Gonzalez, P. Bauskar, S. Gonzalez-Eymard, D. De Los Santos and M. M. Shoshani, *Organometallics*, 2023, **42**, 435–440.
- 195 Z. Weng, S. Teo, L. L. Koh and T. S. A. Hor, *Chem. Commun.*, 2006, 1319.
- 196 A. E. Nako, Q. W. Tan, A. J. P. White and M. R. Crimmin, *Organometallics*, 2014, **33**, 2685–2688.
- 197 K. L. Mears, C. R. Stennett, E. K. Taskinen, C. E. Knapp, C. J. Carmalt, H. M. Tuononen and P. P. Power, *J. Am. Chem. Soc.*, 2020, **142**, 19874–19878.
- 198 B. J. Graziano, T. R. Scott, M. V. Vollmer, M. J. Dorantes, V. G. Young, E. Bill, L. Gagliardi and C. C. Lu, *Chem. Sci.*, 2022, **13**, 6525–6531.
- 199 O. J. García-de-Jesus, A. Mondragón-Díaz, B. Donnadiou and M.-Á. Muñoz-Hernández, *Inorg. Chem.*, 2023, **62**, 2518–2529.
- 200 H.-Y. Liu, R. J. Schwamm, M. S. Hill, M. F. Mahon, C. L. McMullin and N. A. Rajabi, *Angew. Chem., Int. Ed.*, 2021, **60**, 14390–14393.
- 201 C. McManus, J. Hicks, X. Cui, L. Zhao, G. Frenking, J. M. Goicoechea and S. Aldridge, *Chem. Sci.*, 2021, **12**, 13458–13468.
- 202 C. Ganesamoorthy, J. Weßing, C. Kroll, R. W. Seidel, C. Gemel and R. A. Fischer, *Angew. Chem., Int. Ed.*, 2014, **53**, 7943–7947.



- 203 R. J. Schwamm, M. P. Coles, M. S. Hill, M. F. Mahon, C. L. McMullin, N. A. Rajabi and A. S. S. Wilson, *Angew. Chem., Int. Ed.*, 2020, **59**, 3928–3932.
- 204 H.-Y. Liu, S. E. Neale, M. S. Hill, M. F. Mahon and C. L. McMullin, *Dalton Trans.*, 2022, **51**, 3913–3924.
- 205 X. Guo, T. Yang, Y. Zhang, F. K. Sheong and Z. Lin, *Inorg. Chem.*, 2022, **61**, 10255–10262.
- 206 H.-Y. Liu, S. E. Neale, M. S. Hill, M. F. Mahon and C. L. McMullin, *Chem. Sci.*, 2023, **14**, 2866–2876.
- 207 C. McManus, A. E. Crumpton and S. Aldridge, *Chem. Commun.*, 2022, **58**, 8274–8277.
- 208 A. Suzuki, L. Wu, Z. Lin and M. Yamashita, *Angew. Chem., Int. Ed.*, 2021, **60**, 21007–21013.
- 209 D. Sorbelli, L. Belpassi and P. Belanzoni, *Inorg. Chem.*, 2022, **61**, 21095–21106.
- 210 C. Bakewell, B. J. Ward, A. J. P. White and M. R. Crimmin, *Chem. Sci.*, 2018, **9**, 2348–2356.
- 211 A. Friedrich, J. Eysel, J. Langer, C. Färber and S. Harder, *Angew. Chem., Int. Ed.*, 2021, **60**, 16492–16499.
- 212 D. D. L. Jones, I. Douair, L. Maron and C. Jones, *Angew. Chem.*, 2021, **133**, 7163–7168.
- 213 T. X. Gentner, B. Rösch, G. Ballmann, J. Langer, H. Elsen and S. Harder, *Angew. Chem., Int. Ed.*, 2019, **58**, 607–611.
- 214 F. Rekhroukh, W. Chen, R. K. Brown, A. J. P. White and M. R. Crimmin, *Chem. Sci.*, 2020, **11**, 7842–7849.
- 215 A. S. S. Wilson, M. S. Hill, M. F. Mahon, C. Dinoi and L. Maron, *Tetrahedron*, 2021, **82**, 131931.
- 216 M. M. D. Roy, J. Hicks, P. Vasko, A. Heilmann, A.-M. Baston, J. M. Goicoechea and S. Aldridge, *Angew. Chem., Int. Ed.*, 2021, **60**, 22301–22306.
- 217 M. J. Evans, G. H. Iliffe, S. E. Neale, C. L. McMullin, J. R. Fulton, M. D. Anker and M. P. Coles, *Chem. Commun.*, 2022, **58**, 10091–10094.
- 218 J. T. Boronski, L. R. Thomas-Hargreaves, M. A. Ellwanger, A. E. Crumpton, J. Hicks, D. F. Bekiş, S. Aldridge and M. R. Buchner, *J. Am. Chem. Soc.*, 2023, **145**, 4408–4413.
- 219 J. T. Moore, M. J. Dorantes, Z. Pengmei, T. M. Schwartz, J. Schaffner, S. L. Apps, C. A. Gaggioli, U. Das, L. Gagliardi, D. A. Blank and C. C. Lu, *Angew. Chem., Int. Ed.*, 2022, **61**, e202205575.

