A heterobimetallic complex featuring a Ti–Co multiple bond and its application to the reductive coupling of ketones to alkenes†

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To explore metal–metal multiple bonds between first row transition metals, Ti/Co complexes supported by two phosphinoamide ligands have been synthesized and characterized. The Ti metalloidigand \( \text{Cl}_2\text{Ti}(\text{XylNPPr}_2) \) (1) was treated with \( \text{CoI}_2 \) under reducing conditions, permitting isolation of the Ti/Co complex \( [\mu-\text{Cl}(\text{XylNPPr}_2)\text{Co}]_2 \) (2). One electron reduction of complex 2 affords \( \text{Cl}(\text{XylNPPPr}_2)\text{CoPMe}_3 \) (3), which features a metal–metal triple bond and an unprecedentedly short Ti–Co distance of 2.0236(9) Å. This complex is shown to promote the McMurry coupling reaction of aryl ketones into alkenes, with concomitant formation of the tetranuclear complex \( [\text{Ti}(\text{OCMe}_2\text{CH}_2\text{PPh}_2)\text{Rh}] \) (4). A cooperative mechanism involving bimetallic C=O bond activation and a cobalt carbene intermediate is proposed.

Introduction

Metal–metal interactions have received considerable attention owing to their fundamentally interesting bonding properties and their potential use in multi-electron transfer processes. In particular, interactions between early and late metals are of interest due to their relevance to observed enhanced reactivity phenomena in the area of surface catalysis, e.g. the Fischer–Tropsch process. The formation of “strong metal-support interactions” in heterogeneous catalytic systems involving early metal-oxide supported late metal catalysts has been shown to promote reactivity, with the reduction of the early metal and formation of direct early/late metal–metal interactions being observed. To emulate this behavior in homogeneous systems, some metal–metal bonded early/late heterobimetallic complexes have been reported over the past several decades, and their reactivity towards small molecule substrates has been explored.

Wolczanski’s report of a metal–metal triple bond in the Ti/Rh complex \( \text{Ti}[\text{OCMe}_2\text{CH}_2\text{PPh}_2] \text{Rh} \) represents an important breakthrough in the area of early/late heterobimetallic complexes. Our group has studied a similar series of \( \text{C}_3 \)-symmetric tri(phosphinoamide) Zr–Co heterobimetallic complexes featuring Zr–Co multiple bonds and their remarkable ability to undergo a wide array of one, two, and four-electron transformations with small molecules such as hydrazines, alkyl halides, CO\(_2\), diarylketones, and organic azides. Since the dissociation of one of the phosphine ligands from Co and its coordination to Zr in an \( \eta^3 \) fashion are involved in many of our examples of reactivity, we posited that coordinatively unsaturated heterobimetallic complexes linked by just two phosphinoamide ligands may lead to more reactive and catalytically competent compounds (Scheme 1). Such a strategy was employed in the “A frame” heterobimetallic bis(alkoxyporphine)-linked Zr/M (M = Rh, Pt) complexes reported by Wolczanski et al., permitting the observation of intermetallic alkyl group exchange. However, other interesting cooperative reactivity was likely hindered in these systems by the coordinative saturation provided by a \( \text{Cp}^* \) ligand on Zr.

Furthermore, in light of the wide use of TiO\(_2\) as a catalyst support in heterogeneous catalysis and titanium’s ability to undergo one-electron redox processes, we chose to extend our studies of heterobimetallic chemistry to Ti/M complexes.

Scheme 1 Justification for exploring a bis(phosphinoamide) ligand framework for heterobimetallic complexes.
Nagashima and co-workers have reported a series of bis(phosphinoamide) Ti\textsuperscript{IV}/M (M = Ni, Pd, Pt, and Ru) heterobimetallic complexes, however, the reported reactivity of these complexes remains somewhat limited.\textsuperscript{20-22} Herein, we describe a synthetic method to construct metal–metal multiple bonds between Ti and a redox active first row transition metal, Co. Further, we explore the reactivity of a Ti/Co complex with aryl ketones, uncovering that the two metal centers in this heterobimetallic complex mediate the reductive coupling of ketones to alkenes (the McMurry reaction). Other reported examples of the deoxygenation of ketones by an early/late heterobimetallic complex include Gade’s report of the transfer of an oxygen atom from cyclopropenone to a bound carbonyl ligand by HC(Me\textsubscript{2}SiN(2,3,4-F\textsubscript{3}C\textsubscript{6}H\textsubscript{2}))\textsubscript{3}Zr–Fe(CO)\textsubscript{3}Cl\textsuperscript{32,24} and our own report of cleavage of the C–O bond of benzophe-none by (THF)Zr(MesNPiPr\textsubscript{2})\textsubscript{3}CoCl\textsuperscript{33,34} (the ratio of the metal–metal interatomic distance to the metal multiple bonds between Ti and Co (Fig. 1). Taking this metal interaction between Ti and Co (Fig. 1). Taking this metal interaction into consideration, the Ti center adopts a trigonal bipyramidal geometry and the geometry about Co is distorted tetrahedral.

Despite our previous successes with heterobimetallic Zr/Co complexes featuring the [MesNP\textsubscript{3}Pr\textsubscript{3}]\textsuperscript{-} ligand,\textsuperscript{16,17} our initial efforts targeting bimetallic Ti[MesNP\textsubscript{3}Pr\textsubscript{3}]Co complexes were largely unsuccessful as a result of the smaller ionic radius of Ti and the resulting inability to install more than one phosphinoamide ligand around the Ti center. We found, however, that reducing the steric hindrance by changing the N-aryl group from 2,4,6-trimethylphenyl to 3,5-dimethylphenyl permits 

Fig. 1  Displacement ellipsoid (50%) representation of 2. All hydrogen atoms have been omitted for clarity. Selected bond distances (\AA) and angles (\textdegree): Ti1–Co1, 2.0251(4); Ti1–Cl1, 2.3843(5); C1–Cl1–N1, 126.73(4); C1–Cl1–N2, 115.90(4); N1–Ti1–N2, 116.91(6); P1–Co1–I1, 119.458(14); P2–Co1–I1, 125.626(14); P1–Co1–P2, 111.923(17).

sum of the single bond atomic radii of the two metal ions, FSR\textsuperscript{\textdegree}) of only 0.89, suggesting a relatively strong metal–metal interaction between Ti and Co (Fig. 1). Taking this metal–metal bond into consideration, the Ti center adopts a trigonal bipyramidal geometry and the geometry about Co is distorted tetrahedral. The Ti–Cl bond trans to the Ti–Co bond is considerably longer than the one cis to the Ti–Co bond (2.5745(5) \AA vs. 2.3843(5) \AA), in line with previous reports of polar metal–metal bonds exerting trans influences on apical ligands in heterobimetallic complexes.\textsuperscript{16,35}

Formally, the oxidation states of the two metals in 2 are Ti\textsuperscript{III} and Co\textsuperscript{I}, and solution magnetic moment measurements suggest an S = 1 ground state (\mu\textsubscript{eff} = 3.11 \mu\textsubscript{B}). A computational investigation of 2 was carried out using density functional theory (DFT). Although the unmodified ligand set was used in calculation, the cobalt-bound iodide ligand was changed

Results and discussion

Despite our previous successes with heterobimetallic Zr/Co complexes featuring the [MesNP\textsubscript{3}Pr\textsubscript{3}]\textsuperscript{-} ligand,\textsuperscript{16,17} our initial efforts targeting bimetallic Ti[MesNP\textsubscript{3}Pr\textsubscript{3}]Co complexes were largely unsuccessful as a result of the smaller ionic radius of Ti and the resulting inability to install more than one phosphinoamide ligand around the Ti center. We found, however, that reducing the steric hindrance by changing the N-aryl group from 2,4,6-trimethylphenyl to 3,5-dimethylphenyl permits access to the desired precursor Cl\textsubscript{1}Ti[XylNP\textsubscript{3}Pr\textsubscript{3}]\textsubscript{2} (1) (Xyl = 3,5-dimethylphenyl) via treatment of TiCl\textsubscript{4} with two equivalents of Li[XylNP\textsubscript{3}Pr\textsubscript{3}] (see ESI†). In contrast to our reported synthesis of ClZr(MesNP\textsubscript{3}Pr\textsubscript{3})Co,\textsuperscript{18} successful installation of a second metal to metallocligand 1 required the addition of an external reductant. Treatment of Co\textsubscript{1} with 1 in the presence of excess Zn powder afforded a symmetric tetrametallic complex [([μ-Cl]Ti[XylNP\textsubscript{3}Pr\textsubscript{3}])CoCl\textsubscript{2}] (2) with two heterobimetallic Ti–Co fragments bridged by two Cl\textsuperscript{-} ligands between the two Ti centers (Scheme 2).

X-ray crystallography of single crystals of 2 reveals a Ti–Co interatomic distance of 2.2051(4) \AA with a “formal shortness ratio” (the ratio of the metal–metal interatomic distance to the
Population Analysis (NPA). As shown in Table 1, the more

ence in natural charges of the two atoms calculated

polarity of the Ti

contributing 75% and 20% of the electron density in the bond,

signi
cantly more localized on Co than on Ti, indicating that

these are best described as donor–acceptor interactions. The

polarity of the Ti–Co bond in 3 is also evident from the
difference in natural charges of the two atoms calculated via

Natural Population Analysis (NPA). As shown in Table 1, the more

negative charge on Co and more positive charge on Ti in

complex 3 is indicative of a polar bond. The difference in

natural charge between the two metal atoms is far smaller for

complex 3 (1.04) than for the previously described Zr/Co

complex (THF)Zr(MeSi(Me2SiN(4-CH3C6H4))3Ti (Ref. 39)

Based on these metrics, the Ti–Co bond is more covalent

than the metal–metal bond in the (THF)Zr(MeNSPPr2)2CoN2

system,9 likely owing to the better orbital overlap between

the two sets of 3d orbitals (rather than 3d and 4d) in the Ti–Co

complex. The enhanced orbital overlap between the two first

row metals in complex 3 is likely also responsible for its shorter

metal–metal distance compared to the isoelectronic complex

Ti(OCMc2CH2PPh3)2Rh.15

The formal oxidation states of the two metals in 3 could be
described as TiIII/Co0 or TiIV/Co−1. A recent study using X-ray

absorption near edge structure (XANES) spectroscopy showed

that the effective oxidation states in the similar complex (THF)

Zr(MeNSPPr2)2CoN2 are ZrIV/Co−1.22 However, given the polar
cova
ten nature of the metal–metal σ bond, a non-zwitterionic

TiIII/Co0 description may be favored in the case of 3 and further

spectroscopic study will be required to determine the effective

oxidation states in this molecule.

It is also worth noting that Gade’s Ti/Co complexes

MeSi(Me2SiN(4-CH3C6H4))3Ti–Co(CO)3L (L = CO, P(Tol3)) have
much weaker Ti-Co bonds than 2 or 3 based on their longer intermetallic distances (2.5542(10) Å and 2.471(4) Å, respectively). The metal-metal bonds in these molecules were also deemed polar covalent in nature. The calculated WBI of simplified models of Gade’s complexes are much smaller than those computed for 2 and 3, and the computed charge differences between Ti and Co are larger (Table 1). The weaker and more polar bonding between Ti and Co in MeSi[Me₂SiN(4-CH₃C₆H₄)]₂Ti-Co(CO)₃L may be a function of oxidation state differences: as shown in Table 1, the natural charge calculated for the Ti atom in the (NH₃)₂Ti-Co(CO)₃L complexes is greater than that of 3 by ~1, implicating a more reduced TiIII center in 2 and 3.

Owing to its open coordination geometry, the bis(phosphinoamide) Ti/Co platform is expected to be able to accommodate a wide range of small molecule substrates in the context of σ and π bond activation, and we chose to initially investigate the representative reactivity of 3 with diaryl ketones. Allowing a benzene solution of 3 to react with benzophenone at room temperature resulted in the loss of PMe₃ (detected by ¹³P NMR spectroscopy) and afforded a brown complex 4 that featured paramagnetically shifted resonances in its ¹H NMR spectrum (Scheme 3, Fig. S5†). Several attempts to crystallize this complex failed due to its thermal instability. However, addition of NaI to a solution of 4 generated a purple TiIVCoI complex [Ti(μ-O)(XylNPPr₂)₂Co]₂ (4) along with tetraphenylethylene, which was observed by GC-MS (m/z = 332). The structure of 4 was determined by a single-crystal X-ray diffraction study, revealing a tetrametallic structure in which a μ₁-oxo ligand bridges between Ti and Co as well as a second Ti center, giving a dimeric structure (Fig. 4). The tetrametallic complex 4 features an elongated Ti-Co distance of 2.4397(4) Å (FSR = 0.985), indicating a diminished metal-metal interaction between Ti and Co. The Co-O distance of 2.0342(14) Å in 4 is longer than that observed for (η²-MesNPPr₂)Zr(μ-O)(MesNPPr₂)₂Co=CH₂ (1.9710(16) Å) (5), in which the oxo ligands are only bridging between Zr and Co.

The reductive coupling of ketones (or aldehydes) to form alkenes as occurs upon addition of benzophenone to 3 is known as the McMurry reaction. A typical McMurry reaction involves the reduction of TiCl₄ or TiCl₃ with an external reductant such as Li, Na, K, Mg, Mg(Hg), Zn or LiAlH₄, followed by the reductive coupling of ketones to alkenes by the active low valent titanium species. Stoichiometric reagents and elevated temperatures are often required, and problems of reactivity and product selectivity (alkene vs. pinacol, E vs. Z) have limited the scope of this reaction. Furthermore, the mechanism of this reaction is poorly understood and operative mechanisms proceeding through carbenoid or pinacolate intermediates have been proposed. A number of ketone substrates were screened to investigate the substrate scope of the McMurry reaction promoted by complex 3. We were pleased to find that complex 3 is reactive towards a variety of ketone substrates, affording moderate to high yields of alkene products at room temperature with relatively short reaction times, provided that stoichiometric NaI is added (Table 2). In all cases, complex 4 is formed as a byproduct of the reaction, as confirmed by ¹H NMR spectroscopy. Asymmetric ketones gave a mixture of Z and E isomers in a ca. 2 : 1 ratio determined by GC-MS (entries 2–4, 7–8). It was also found that ketone substrates with more electron donating arene substituents (entries 6–9) require longer reaction times, and that only aryl-substituted ketones afforded alkene products. These observations suggest that, as with McMurry reactions in the literature, electron transfer from 3 to the ketone substrate is an essential step.

To determine the role of Ti and Co and the potential cooperative reactivity between the two metals, the reactivity of monometallic Ti and Co analogues with benzophenone was also screened. Neither 1 nor ICo(Ph₂PNCPr₃)₃ (ref. 16) shows similar reactivity with aryl ketones. A reduced monometallic titanium(III) complex, Ti(XylNPPr₂)₃, was also synthesized and shows no activity to promote the reductive coupling of ketones within 3 hours. Furthermore, neither Ti(XylNPPr₂)₃ nor ICo(Ph₂PNCPr₃)₃ afforded the McMurry coupling product from benzophenone when Zn powder or KC₈ were added as external reductants. These control reactions show that both Ti and Co play an essential role in the reaction.

The mechanism by which benzophenone and other aryl ketones are reductively coupled into alkenes is intriguing. Our
group previously reported that the reaction of benzophenone with heterobimetallic Zr/Co complex (THF)Zr(MesNPiPr2)3CoN2 (6) leads to a benzophenone radical coupling product 7 and thermolysis of 7 affords the \( \mu \)-oxo terminal carbene complex 5 (Scheme 4).\(^{23,24}\) Upon thermolysis at 110 °C, complex 5 decomposes, producing a mixture of tetraphenylethane, tetraphenylethylene, and diphenylmethane as the \( \text{CPh}_2 \) fragment is extruded. By analogy to the Zr/Co system, it is proposed that the intermediate complex 4′ obtained upon initial exposure of 3 to benzophenone is a \( \mu \)-oxo/carbene complex formed upon oxidative addition of the ketone’s C=O double bond (Scheme 5). Another possible structure of 4′ is a bridging \( \mu_2 \)-\( \eta^1 \),\( \eta^1 \)-ketone adduct (Scheme 5). In either case, upon addition of NaI, the \( \text{CPh}_2 \) fragment is released from the Co center to form the tetraphenylethylene coupled product. The resulting \( \mu \)-oxo complex dimerizes to form complex 4 (Scheme 5). McMurry coupling reactions have also been shown to proceed through pinacolate intermediates; however, several key experiments have ruled out the possibility that 4′ contains a bound pinacolate. First, two separate solutions of 3 were treated with benzophenone and 4,4′-dimethylbenzophenone to form 4′ and its \( p \)-tolyl-substituted analogue in situ. Combination of these two solutions in the presence of NaI affords a statistical distribution of alkene hetero- and homo-coupling products, namely Ph2C=CHPh2, Ph2C=CH(p-tolyl)2, and (p-tolyl)2C=CH(p-tolyl). The crossover product could not form if a pinacolate mechanism was operative. Furthermore, addition of stoichiometric \( \text{H}_2\text{O} \) to intermediate 4′ affords free ketone and a small amount of tetraphenylethylene, with no evidence for diol hydrolysis products that would form from a pinacolate complex.

### Conclusions

In summary, we have synthesized heterobimetallic Ti–Co complexes linked by two phosphinoamide ligands and the reduced low coordinate Ti–Co complex 3 features a strong metal–metal triple bond, with planar geometry at both Ti and Co. This low coordinate Ti–Co complex is shown to promote the reductive coupling of aryl ketones into alkenes. Future studies will focus on exploring the mechanism of this reaction and the reactivity of 3 towards other unsaturated substrates.
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Notes and references