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Synthesis and reactivity of paramagnetic late transition metal complexes supported by tris(oxazolinyl)phenylborate

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Synthesis and reactivity of paramagnetic late transition metal complexes supported by tris(oxazolinyl)phenylborate

by

Regina Rose Reinig

A dissertation submitted to the graduate faculty
in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

Major: Inorganic Chemistry

Program of Study Committee:
Aaron D. Sadow, Major Professor
Levi Stanley
Gordon Miller
Javier Vela
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Iowa State University
Ames, IA
2018

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DEDICATION

To Dr. Chin for inspiring me to pursue research in chemistry.
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Our group previously developed the synthesis of main group Zn(II) and Mg(II) complexes supported by the $\text{To}^M$ (tris(4,4-dimethyl-2-oxazolinyl)phenylborate) ligand. The reactivity of these main group oxazolinyl based complexes in oxygenation, hydrosilylation, and hydroboration reactions provided motivation for studying the reactivity of late first-row transition metals. This thesis focuses on the synthesis, characterization, and reactivity of late transition metal complexes based on iron, nickel and cobalt containing $\text{To}^M$ as the ancillary as well as the synthesis and characterization of other paramagnetic molecules.

Salt metathesis reaction of Tl$\text{To}^M$ with CoCl$_2$•THF (THF = tetrahydrofuran), FeBr$_2$, or NiCl$_2$•DME (DME = dimethoxyethane) at room temperature in THF afforded $\text{To}^M$CoCl, $\text{To}^M$FeBr, and $\text{To}^M$NiCl, respectively. $\text{To}^M$CoCl, $\text{To}^M$FeBr, and $\text{To}^M$NiCl react with NaOAc to form the acetate complexes $\text{To}^M$CoOAc, $\text{To}^M$FeOAc, and $\text{To}^M$NiOAc, respectively. $\text{To}^M$CoOAc was studied in the oxidation of cyclohexane using meta-chloroperoxybenzoic acid ($m$CPBA) as the oxidant and found to selectively catalyze the formation of cyclohexanol over the over-oxidized products of cyclohexanone and ε-caprolactone.

Reaction of $\text{To}^M$CoCl, $\text{To}^M$FeBr, and $\text{To}^M$NiCl with alkyllithiums or potassium benzyl were attempted as a route to organometallic complexes. The nickel(II) alkyls were not identified under the reaction conditions attempted; however, $\text{To}^M$CoR (R = Me, Et, $n$Bu, CH$_2$SiMe$_3$, Bn, and Ph) and $\text{To}^M$FeR (R = Bn) compounds were found to be isolable.

$\text{To}^M$CoR (R = Me, Et, $n$Bu, CH$_2$SiMe$_3$, Bn, and Ph) complexes react with CO followed by O$_2$ to form the carboxylate compounds $\text{To}^M$CoO$_2$CR (R = Me, Et, $n$Bu, CH$_2$SiMe$_3$, Bn, and Ph). The direct insertion of CO$_2$ into the Co-R bond is determined to be much slower, requiring several days to weeks depending upon the R group. $\text{To}^M$FeBn reacts with CO to form
To $^3$Fe{C(O)Bn}(CO)$_2$, but sequential addition of O$_2$ does not provide the corresponding carboxylate compound suggesting the oxidative carbonylation reactivity is unique to cobalt. Lastly, the expertise gained from the synthesis and characterization of paramagnetic complexes provided guidance for the synthesis and characterization of deuterated biradicals to be used as polarizing agents in dynamic nuclear polarization (DNP).
CHAPTER 1: INTRODUCTION

General Introduction

Late first-row transition metals iron, cobalt, and nickel have the advantage of being relatively economical and environmentally benign. The utility of late first-row transition metals has been demonstrated in commercially and biologically relevant reactions including oxygenation and polymerization reactions.

Iron, cobalt, and nickel complexes supported by the tris(pyrazolyl)borate (Tp) ligand are well established. While TpM (M = Fe, Co or Ni) compounds have been very useful in catalytic transformations, a major general hindrance encountered in scorpionate chemistry is comproportionation reactions that result in, for example, Tp₂Fe.²⁻³ Tris(oxazolinyl)phenylborate (To⁻) has been reported to provide a more bowl-like steric environment suggesting it may support tetrahedral first-row transition metal complexes.⁴

A possible route to iron, cobalt, and nickel complexes supported by To⁻ is through salt metathesis reaction of TlTo⁻ with metal halides. The suitability of TlTo⁻ as a To⁻ transfer agent has been well-documented with examples including transfer of To⁻ from Tl to Zr, Zn, and Rh.⁵⁻⁷ The corresponding halide complexes, for example To⁻CoCl, could be expected to be used as a precursor to more reactive ligands. The general route of salt metathesis may also be applicable to replacement of the chloride ligand, although reactions involving To⁻ZrCl₃ have demonstrated that this route can be complicated by the hard basic nature of the To⁻ ligand that often favors transmetalation to Mg or Li.⁵ Replacement of the chloride ligand with acetate is desirable given acetate complexes have been demonstrated as effective catalysts in oxidation reactions and the acetate moiety is commonly encountered in biological systems. An oxidation reaction of
primary interest is the oxidation of hydrocarbons where catalysts based upon metals such as Co(II) and Cu(II) serve as catalysts on an industrial scale. A major application is the use of cobalt carboxylates as the catalysts for the oxidation of \( p \)-xylene to terephthalic acid or dimethyl terephthalate which represents the largest industrial use of homogeneous catalysts. Cyclohexane is another common substrate for such oxidation reactions (Figure 1). Oxidation of cyclohexane typically provides a mixture of cyclohexanol and cyclohexanone which commercially can be used as precursors to caprolactam and adipic acid which in turn are used to make nylon 6,6. However, the selective oxidation of cyclohexane is generally complicated by the over-oxidized products that so easily form.

![Figure 1](image1.png)

**Figure 1.** Oxidation of cyclohexane to cyclohexanol and cyclohexanone.

Iron, cobalt, and nickel complexes featuring an acetate coligand have been reported to serve as efficient catalysts for the oxidation of cyclohexane using \( m \)-CPBA as the terminal oxidant, thus demonstrating potential for further exploration of these late first-row transition metals.

Another reaction of interest related to cobalt is oxidative carbonylation. Traditionally, metals such as palladium, rhodium, and iridium are employed in these types of reactions. The utilization of more abundant late first-row transition metals in these processes is desirable. Precedence for cobalt as a suitable metal for such type of catalytic reactions can be found in nature where acetate formation is an important metabolic pathway in acetogens. The Wood-Ljungdahl pathway (Figure 2) proposes
reduction of CO$_2$ to CO followed by insertion of the CO into a Co-Me bond to form an acetyl group (Figure 2).$^{17-19}$

![Figure 2. Western and Eastern Branch of the Wood-Ljungdahl Pathway.](image)

Exploring the distinction between oxidative carbonylation, where CO$_2$ is the source of the carbonyl and O$_2$ is the oxidant, versus direct insertion of CO$_2$ to form carboxylates would provide insight into the synthesis of acetate and other carboxylate compounds. The synthesis of cobalt carboxylate compounds from cobalt alkyls could provide insight into rates and mechanisms for such processes.

This thesis contributes to the development of late first-row transition metals by demonstrating the reactivity of To$^M$CoOAc in the oxidation of cyclohexane and a series of cobalt alkyl compounds (To$^M$CoR; R = Me, Et, $^t$Bu, CH$_2$SiMe$_3$, Bn, and Ph) that are reactive in oxidative carbonylation. In addition, the reactivity of To$^M$FeBn towards CO, O$_2$, and CO$_2$ is reported as well as the synthesis and characterization of To$^M$NiCl and To$^M$NiOAc. Finally, in theme with the paramagnetic nature of these complexes, a series of deuterated biradical polarizing agents were synthesized and demonstrated to be effective for obtaining improved dynamic nuclear polarization (DNP) enhancements.
Dissertation Organization

This thesis contains seven chapters. Chapter 1 provides a general introduction to the synthesis and study of late first-row transition metal complexes. Chapters 2, 3, and 6 are published journal papers that have been modified to provide coherency while chapters 4 and 5 consist of work that is not yet published, and thus are manuscripts in preparation for publication.

Chapter 2 describes the synthesis of tris(4,4-dimethyl-2-oxazolinyl)phenylborato cobalt(II) chloride (To\textsuperscript{M}CoCl), which serves as a precursor to To\textsuperscript{M}CoO\textsubscript{t}Bu and To\textsuperscript{M}CoOAc. To\textsuperscript{M}CoOAc is demonstrated to selectively catalyze the oxidation of cyclohexane to cyclohexanol using meta-chloroperoxybenzoic acid (mCPBA) as the oxidant. Debabrata Mukherjee was the first member of our group to synthesize and obtain an X-ray quality crystal of To\textsuperscript{M}CoCl. Zachary B. Weinstein was responsible for the operation of the Chemspeed Technologies SWING-XL automated platform used for the oxidation catalysis as well as analysis of the oxidation catalysis results. Weiwei Xie was responsible for the X-ray powder diffraction analysis studies, Toshia Albright performed the EPR simulations, Sergey L. Bud’ko performed the magnetic susceptibility measurements, and previous experiments conducted by Benjamin Baird and Tristan S. Gray helped guide the synthesis reported in this chapter.

Chapter 3 presents the synthesis of To\textsuperscript{M}CoMe and detailed studies of the reactivity of To\textsuperscript{M}CoMe toward CO, O\textsubscript{2}, and CO\textsubscript{2}. The sequential addition of CO followed by O\textsubscript{2} to To\textsuperscript{M}CoMe affords To\textsuperscript{M}CoOAc. This oxidative carbonylation reactivity is explored in depth. Reaction of To\textsuperscript{M}CoMe with CO\textsubscript{2} also provides To\textsuperscript{M}CoOAc, but this reaction proceeds much more slowly than the two-step oxidative carbonylation pathway.
Ellie L. Fought carried out DFT, TDDFT, and DFT Hessian calculations of $^{M}\text{CoMe}$, $^{M}\text{CoMe}(\text{CO})$, $^{M}\text{Co}\{\text{C(O)Me}\}\text{CO}$, and $^{M}\text{CoOAc}$.

Chapter 4 describes the synthesis of $^{M}\text{CoR}$ ($R = \text{Me, Et, }^4\text{Bu, CH}_2\text{SiMe}_3, \text{Bn, and Ph}$) complexes to investigate the effect of the R group on the oxidative carbonylation reactivity observed for $^{M}\text{CoMe}$. The results suggest that both alkyl and aryl groups provide similar reactivity, and that oxidative carbonylation is kinetically favored over direct CO$_2$ insertion into the Co-C bond. Ellie L. Fought conducted the DFT, TDDFT, and DFT Hessian calculations included in this chapter.

Chapter 5 reports the synthesis of $^{M}\text{FeBr}$ and $^{M}\text{NiCl}$ compounds. $^{M}\text{FeBr}$ and $^{M}\text{NiCl}$ readily react with NaOAc to form $^{M}\text{FeOAc}$, and $^{M}\text{NiOAc}$. $^{M}\text{FeBr}$ also serves as a precursor to the organometallic complex $^{M}\text{FeBn}$. The reactivity of $^{M}\text{FeBn}$ is studied towards CO, O$_2$, and CO$_2$ and compared with the reactivity observed for $^{M}\text{CoBn}$.

Chapter 6 describes the synthesis and characterization of deuterated biradicals to be used as polarizing agents in dynamic nuclear polarization (DNP). This work was conducted in close collaboration with Frédéric A. Perras who was responsible for measuring the DNP enhancements for each of the biradicals studied.

Chapter 7 presents general conclusions. Dr. Arkady Ellern is credited for all of the X-ray data and solving of X-ray structures presented in this thesis.
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CHAPTER 2: SYNTHESIS AND OXIDATION CATALYSIS OF
TRI(OXAZOLINYL)BORATO COBALT(II) SCORPIONATES

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Regina R. Reinig, Debabrata Mukherjee, Zachary B. Weinstein, Weiwei Xie, Toshia Albright, Benjamin Baird, Tristan S. Gray, Arkady Ellern, Gordon J. Miller, Arthur H. Winter, Sergey L. Bud’ko, and Aaron D. Sadow*

**Abstract**

The reaction of CoCl\(_2\)·THF and thallium tris(4,4-dimethyl-2-oxazoliny1)phenylborate (TlTo\(^M\)) in tetrahydrofuran (THF) provides To\(^M\)CoCl (1) in 95 % yield; however, appropriate solvents and starting materials are required to favor 1 over two other readily formed side-products, (To\(^M\))\(_2\)Co (2) and {HTo\(^M\)}CoCl\(_2\) (3). ESR, NMR, FTIR, and UV/Vis spectroscopy were used to distinguish these cobalt(II) products and probe their electronic and structural properties. Even after the structures indicated by these methods were confirmed by X-ray crystallography, the spectroscopic identification of trace contaminants in the material was challenging. The recognition of possible contaminants in the synthesis of To\(^M\)CoCl in combination with the paramagnetic nature of these complexes provided impetus for the utilization of X-ray powder diffraction to measure the purity of the To\(^M\)CoCl bulk powder. The X-ray powder diffraction results provide support for the bulk-phase purity of To\(^M\)CoCl in preparations that avoid 2 and 3. Thus, 1 is a precursor for new tris(oxazoliny1)borato cobalt chemistry, as exemplified by
its reactions with KOrBu and NaOAc to give $\text{To}^\text{M}\text{CoO} \text{Bu}$ (4) and $\text{To}^\text{M}\text{CoOAc}$ (5), respectively. Compound 5 is a catalyst for the oxidation of cyclohexane with meta-chloroperoxybenzoic acid (mCPBA), and the rate constants and selectivity for cyclohexanol versus cyclohexanone and $\varepsilon$-caprolactone were assessed.

**Introduction**

Tridentate fac-coordinating monoanionic scorpionate-type ligands$^1$ support and stabilize first-row metal centers bonded to reactive moieties including hydrides,$^2$ alkyl groups containing $\beta$- hydrogen atoms,$^3$ imido and oxido ligands,$^4$ azides,$^5$ and oxidizing moieties such as peroxides.$^6$ In addition, 3d metal compounds coordinated by scorpionate ligands have served as models for metal sites in enzymes and provided motivation to study their spectroscopic and structural features in detail.$^7$ Typically, sterically encumbered scorpionates are required to support reactive species such as peroxides and superoxides.

Variation of the steric and electronic properties of the ancillary scorpionate donors can greatly influence the stability of reactive moieties and the reactivity of the complexes. For example, the steric encumbrance of tris(3-tert-butyl-5-methylpyrazolyl)borate ($\text{Tp}^{\text{Bu},\text{Me}}$), known as the tetrahedral enforcer, stabilizes reactive groups such as alkyl peroxides.$^8$ However, it also may limit the well-known catalytic oxidation chemistry of scorpionate peroxide complexes of first-row transition metals.$^6,^9$ The smaller parent tris(pyrazolyl)borate (Tp) or tris(3,5-dimethyl-pyrazolyl)borate (Tp*) ligands, which might provide more accessible metal centers, instead form octahedral $\{\kappa^3$-$\text{Tp}\}_2\text{M}$ compounds.$^10$ Even the bulkier tris(3-phenyl-5-methyl- pyrazolyl)borate
(Tp^{Bu,Me}) ligand forms octahedral (Tp^{Bu,Me})_2Co upon isomerization of a pyrazole ring.\textsuperscript{11} Thus, the steric properties of non-pyrazolyl-based scorpionate ligands, such as those involving nonplanar oxazoline donors, may sufficiently stabilize reactive moieties and also allow new metal-centered reactivity that involves switching between four- and five-coordinate species in a way not accessible with Tp^{Bu}-based compounds.

Recently, we discovered that tris(4,4-dimethyl-2-oxazolinyl)phenylborate (To^M) supports reactive main-group complexes of mononuclear and tetrahedral zinc(II) and magnesium(II). For example, a catalytically active mononuclear zinc hydride was readily synthesized by the reaction of To^MZnOtBu and PhSiH\textsubscript{3}.\textsuperscript{12} The To^MZnX system also supports and stabilizes alkylperoxides, such that To^MZnOOEt is thermally persistent even at temperatures above 100 °C.\textsuperscript{13} Moreover, To^MZnEt is sufficiently reactive to undergo selective oxidation upon treatment with O\textsubscript{2}. To^M gives a more open geometry and bowl-like steric profile, in contrast to the tetrahedral enforcer Tp^{Bu,Me} that typically stabilizes reactive moieties.

These examples provide motivation for studying the synthesis and reactivity of tetrahedral first-row transition-metal centers supported by To^M, in which redox-active metal centers could lead to new chemistry and catalysis. In particular, the electron-donating properties and bowl-like steric environment\textsuperscript{14} provided by the coordination pocket of To^M could stabilize reactive groups bonded to metal centers with accessible d electrons to facilitate catalytic chemistry. Moreover, a series of optically active tris(4-R-2-oxazolinyl)phenylborate ligands are also available for imposing chiral environments on tetrahedrally coordinated first-row transition metal sites.\textsuperscript{15}
The present study describes the synthesis and characterization of $\text{To}^M\text{CoCl}$ (1) as a precursor to reactive complexes. We also provide syntheses and characterization data of $(\text{To}^M)_2\text{Co}$ (2) and $\{\text{HTo}^M\}\text{CoCl}_2$ (3), which are side-products discovered during exploratory syntheses toward 1. The open steric profile of the $\text{To}^M$ ligand results in additional synthetic challenges compared with syntheses of sterically encumbered tris(pyrazolyl)borate based first-row metal compounds, in terms of controlling the formation of 1, 2, and 3. The chlorido ligand in 1 may be substituted through salt-metathesis reactions to give $\text{To}^M\text{CoO}t\text{Bu}$ (4) and $\text{To}^M\text{CoOAc}$ (5) without $\text{To}^M$ transmetalation or ligand redistribution demonstrating that 1 is a viable synthetic precursor for new cobalt(II) compounds. Complex 5 was studied as a catalyst for the oxidation of cyclohexane. Although the selective oxidation of cyclohexane remains a challenge, cobalt complexes have shown great promise in this area.\textsuperscript{9d}

**Results and Discussion**

The reaction of $\text{TITo}^M$ and $\text{CoCl}_2\cdot\text{THF}$ (THF = tetrahydrofuran) affords $\text{To}^M\text{CoCl}$ (1) as a bright blue solid in excellent yield (equation (1)). $\text{To}^M\text{CoCl}$ is paramagnetic, and its identity as a pseudotetrahedral $d^7$ center is supported by a host of characterization data including $^1\text{H}$ and $^{11}\text{B}$ NMR spectroscopy, IR spectroscopy, UV/Vis spectroscopy, ESR spectroscopy, X-ray diffraction studies, elemental analysis, and magnetic measurements that included both magnetometry and the Evans method. Many of these tools were needed initially to interpret the spectroscopic data, as the identities of the paramagnetic products were challenging to establish, and later to demonstrate the purities of the isolated materials. As outlined below, the ratio of $\text{TITo}^M/\text{CoCl}_2$, the use of $\text{CoCl}_2\cdot\text{THF}$,
and the choice of solvent are crucial to the high-yielding synthesis of 1 without contamination with the readily formed side-products (To\textsuperscript{M})\textsubscript{2}Co (2) and {HTo\textsuperscript{M}}CoCl\textsubscript{2} (3).

\[
\text{Ph} - \text{B} \quad \text{N} \quad \text{To} \quad \text{N} \quad \text{N} \quad \text{Tl} \quad + \quad \text{CoCl}_2 \cdot \text{THF} \quad \xrightarrow{\text{THF}} \quad \text{Ph} - \text{B} \quad \text{N} \quad \text{N} \quad \text{To} \quad \text{N} \quad \text{N} \quad \text{To} + \text{CoCl}_2 \cdot \text{THF} \quad \text{r.t., overnight} \quad \text{TlCl} \quad 1 \quad 95\%
\]

The signals in the \textsuperscript{1}H NMR spectrum of To\textsuperscript{M}CoCl dissolved in benzene-\textit{d}_6 were broad and dispersed over a large chemical shift range, as expected for a paramagnetic compound. Despite these spectroscopic effects, the number of \textsuperscript{1}H NMR signals and their integrated ratio provided characteristic data associated with a pseudo-C\textsubscript{3v}-symmetric oxazolinyloborate species. Broad resonances at \(\delta = 8.38\) (18 H) and 24.88 ppm (6 H) were assigned to the methyl and methylene groups, respectively, of equivalent oxazoline rings in the To\textsuperscript{M} ligand. The equivalence of the rings suggested the tridentate coordination of the To\textsuperscript{M} ligand to the cobalt center. A \textsuperscript{11}B NMR signal at \(\delta = -29\) ppm was shifted significantly from the isotropic value observed for the To\textsuperscript{M} ligand in diamagnetic environments (e.g., the \textsuperscript{11}B NMR chemical shift of TlTo\textsuperscript{M} is \(\delta = -16\) ppm). This paramagnetically shifted \textsuperscript{11}B NMR resonance further indicated the successful conversion of diamagnetic TlTo\textsuperscript{M} to a new species, and we note that the \textsuperscript{11}B NMR spectrum is useful for counting the number of To\textsuperscript{M} species in the reaction mixture.\textsuperscript{16}

The infrared spectrum contained a band at 1598 cm\textsuperscript{-1} (KBr), which was assigned to the C=N stretching mode (\(\nu\text{CN}\)) of the oxazoline groups. Typically, higher-energy \(\nu\text{CN}\) bands (> 1615 cm\textsuperscript{-1}) are observed for dissociated oxazoline moieties in oxazolinyloborate ligands; therefore, this data further supports tridentate ligand coordination. In addition, the observation of only a single peak from the symmetric mode (the asymmetric \(\nu\text{CN}\) band
had low intensity) in the solid-state and solution IR spectra (1586 cm\(^{-1}\)) suggested a similar configuration in the two phases. The tetrahedral geometry of I was also supported by the electronic absorption spectrum (Figure 1), which contained a band from \(\lambda \approx 550\) to 700 nm with \(\lambda_{\text{max}}\) at 568 (\(\varepsilon = 362\) M\(^{-1}\)cm\(^{-1}\)) and 635 nm (\(\varepsilon = 641\) M\(^{-1}\)cm\(^{-1}\)). The larger peak was sandwiched by shoulders at \(\lambda \approx 600\) and 660 nm. The spectra of high-spin pseudotetrahedral cobalt(II) complexes show peaks in this range, and these signals are likely related to the many-featured \(v_3\) band [{\(^4T_1(P) \leftarrow ^4A_2(F)\)}] observed for [CoCl\(_4\)]\(^{2-}\).\(^{17}\)

The To\(^M\)CoCl spectrum is also similar to the spectra of related tris(pyrazolyl)borate (Tp) cobalt(II) complexes such as Tp\(^{^tBu,Me}\)CoCl with maxima at \(\lambda = 526, 602, 636,\) and 660 nm\(^{18}\) or Tp\(^{^{iPr}_2CoEt,\) which exhibited maxima at \(\lambda = 580, 610,\) and 690 nm.\(^{19}\) For tetrahedral Co\(^{II}\) complexes, two lower-energy transitions [{\(^4T_1(F) \leftarrow ^4A_2(F)\)} and \(^4T_2(F) \leftarrow ^4A_2(F)\)] are also expected. Indeed, a weak absorption was also observed at \(\lambda \approx 1000\) nm.

**Figure 1.** UV/Vis spectra of To\(^M\)CoCl (1), (To\(^M\))\(_2-Co\) (2), {HTo\(^M\)}CoCl\(_2\) (3), and To\(^M\)CoOtBu (4).
The paramagnetic nature of \( \textbf{I} \) was further investigated by magnetic susceptibility measurements and ESR spectroscopy. The Evans method revealed a solution magnetic moment of 4.5(2) \( \mu_B \). This data is consistent with a high-spin cobalt(II) complex \((S = 3/2, \quad \text{for which spin-only } \mu_{\text{eff}} = 3.9 \ \mu_B)\) with the \( ^4 \text{A}_2 \) ground state, as discussed above. This electronic configuration was maintained at low temperature according to magnetometry measurements, which showed \( \mu_{\text{eff}} = 4.3(1) \ \mu_B \) at 10.3 K. The ESR spectrum (X-band, 10 K) of a neat point sample revealed the low, rhombic site symmetry. The ESR spectrum was simulated for an \( S = 3/2 \) spin system with anisotropic \( g \) values of \( g_x = 5.96(2), \ g_y = 3.50(2), \) and \( g_z = 2.10(2) \). Similar \( g \) values have been reported for other pseudotetrahedral cobalt(II) complexes featuring rhombic symmetry. \( ^2,^2^2 \) Hyperfine coupling to \( ^{59} \text{Co} \ (I = 7/2) \) was not detected under these conditions. \( ^2^1 \)

Blue crystals of \( \textbf{I} \), obtained from a saturated toluene solution cooled to \(-38^\circ\text{C}\), were determined to be \( \text{To}^\text{M} \text{CoCl} \) through an X-ray diffraction study (Figure 2). The X-ray crystal structure revealed a pseudotetrahedral cobalt center coordinated by the tridentate tris(oxazolinyl)borate ligand. Compound \( \textbf{I} \) and \( \text{To}^\text{M} \text{ZnCl} \) have similar structural features. For example, the \( \text{Co}–\text{N} \) and \( \text{Zn}–\text{N} \) bond lengths range from 2.0091(9) to 2.040(1) Å; the zinc complex has both the highest and the lowest values, and the cobalt compound has intermediate bond lengths. The \( \text{N}–\text{M}–\text{Cl} \) angles range from 115.33(4) to 130.82(4)° for the two compounds, and the cobalt compound has the extreme angles. In addition, the \( \text{B}–\text{M}–\text{Cl} \) angles of 170.99(3) and 174.27(2)° for the cobalt and zinc complexes, respectively, show a slight displacement of the Cl atom from the anticipated position in a pseudo-\( C_{3v} \) structure, and this distortion is slightly larger for the cobalt complex. The structural similarity of diamagnetic zinc(II) and paramagnetic cobalt(II)
compounds suggests that these distortions are sterically controlled rather than the result of electronic influences. The Co1–Cl1 bond length of 2.2025(4) Å in 1 is similar to that observed in other four-coordinate cobalt(II) scorpionate complexes (e.g., Tp^{Ph,Me}CoCl, 2.2004(9) Å; Tp^{tBu,H}CoCl, 2.216(2) Å; and Tp^{tBu,Me}CoCl, 2.2204(9) Å).

Although the reaction of TiToM and CoCl$_2$·THF is reproducible, the choice of CoCl$_2$·THF is critical to a successful and high yielding synthesis. In micromolar-scale reactions, equimolar amounts of TiToM and CoCl$_2$·THF provide 1 quantitatively. Excess CoCl$_2$·THF (1.5 equiv.) is needed in larger-scale preparations to optimize the To$^M$CoCl yield (calculated with respect to TiTo$^M$), because the separation of To$^M$CoCl from

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**Figure 2.** Thermal ellipsoid plot of To$^M$CoCl (1). Ellipsoids are plotted at 50 % probability. H atoms are omitted for clarity. Selected bond lengths (Å) and angles (°): Co1–Cl1 2.2026(5), Co1–N1 2.011(1), Co1–N2 2.030(1), Co1–N3 2.026(1); Cl1–Co1–N1 115.31(4), Cl1–Co1–N2 130.80(4), Cl1–Co1–N3 121.73(4), B1–Co1–Cl1 170.99(3).
CoCl$_2$·THF is easier than that from (To$^M$)$_2$Co (2). In contrast, the reaction of anhydrous CoCl$_2$ and TlTo$^M$ in THF provides a purple mixture of 1 and 2. During our initial synthetic studies, the appearance of two sets of signals in the $^1$H NMR spectra and the formation of purple material in these reactions made the isolation and characterization of pure To$^M$CoCl challenging. The $^1$H NMR spectrum of the purple reaction mixture contained signals at $\delta = 15.1$ and 12.5 ppm as well as the resonances later assigned to 1. The formation of two paramagnetic To$^M$Co species was also suggested by the two $^{11}$B NMR signals at $\delta = -29$ and 42 ppm; the former was assigned to To$^M$CoCl, and the latter was assigned to (To$^M$)$_2$Co on the basis of a single-crystal X-ray diffraction study and its independent synthesis, which is described below. Repeated recrystallizations from saturated toluene solutions at $-38$ °C yielded purple X-ray-quality crystals, and a single-crystal diffraction study revealed that the substance was (To$^M$)$_2$Co (2; Figure 3).

**Figure 3.** Thermal ellipsoid plot of (To$^M$)$_2$Co (2) with ellipsoids at 35% probability. H atoms and two cocrystallized toluene molecules are omitted for clarity. Selected bond lengths (Å) and angles (°): Co1–N1 2.010(4), Co1–N3 2.003(4), Co1–N4 2.013(4), Co1–N5 2.000(4); N1–Co1–N3 97.2(2), N4–Co1–N5 97.8(2), N1–Co1–N4 114.2(2), N3–Co1–N4 116.4(2), N5–Co1–N3 116.4(2), N5–Co1–N1 116.0(2).
The cobalt center in 2, as in 1, has a distorted tetrahedral geometry. Each of the tris(oxazolinyl)borate ligands in 2 is bidentate with one non-coordinated oxazoline ring, and the structure is similar to the solid-state structures of (ToM)₂Mg and (ToM)₂Zn.¹³⁻²⁵ Although the (κ²-ToM)₂Co structure forms readily, the ToM ligand apparently does not support six-coordinate pseudo-sandwich-type first-row metal compounds, in contrast to (κ³-Tp*)₂M compounds.²⁶ Moreover, we have not observed disproportionation of ToMCoCl into (ToM)₂Co and CoCl₂; therefore, ToMCoCl may be a suitable precursor for the synthesis of new inorganic compounds through chloride substitution. The crystallographic data show that the metal centers are sterically protected by the oxazoline methyl groups, and this likely prevents the formation of the six-coordinate structure.

Compound 2 has shorter Co–N interatomic distances than 1, and all of the N–Co–N angles in 2 are larger than those in 1. The interatomic distances and angles of 2 fall between those of (ToM)₂Mg and (ToM)₂Zn and bear greater resemblance to those of the latter. For example, the Co–N bonds in 2 are shorter than those in (ToM)₂Mg by ca. 0.05 Å but are equivalent within error to the Zn–N bonds encountered in (ToM)₂Zn.

Unfortunately, purification by crystallization is not effective for preparative-scale separations of mixtures of (ToM)₂Co and ToMCoCl. Moreover, the ¹H NMR spectrum of the reaction mixture did not contain features that easily identified inequivalent oxazoline rings in the bidentate coordinated structure. We prepared 2 independently to confirm that the unidentified signals in the ¹H NMR spectrum of the mixture of cobalt species corresponded to 2. The reaction of CoCl₂ and 2 equiv. of TlToM affords purple (ToM)₂Co in excellent yield (equation (2)). (ToM)₂Co and ToMCoCl are similarly soluble in
benzene, toluene, diethyl ether, THF, and dichloromethane, and this similarity likely causes the difficulties with the separation of mixtures of the two compounds.

The $^1$H and $^{11}$B NMR spectra of isolated 2 contained signals that matched those detected in the crude mixture with 1; therefore, (To$^\text{M}$)$_2$Co is indeed the second product obtained when anhydrous CoCl$_2$ is the starting material. However, these NMR spectroscopic data did not allow the direct quantitative estimation of the purity and offered few further structural insights. In contrast, the $\nu$$_{\text{CN}}$ bands at 1603 and 1554 (KBr) or 1609 and 1555 cm$^{-1}$ (CH$_2$Cl$_2$) in the infrared spectrum of 2 were structurally informative. These signals were assigned to the $\nu$$_{\text{CN}}$ modes of non-coordinated and coordinated oxazoline groups, respectively, consistent with the bidentate bonding mode determined by X-ray crystallography.

Furthermore, the d–d transitions in the UV/Vis spectrum of 2 again appeared in the expected region for a high-spin tetrahedral Co$^{II}$ complex at $\lambda_{\text{max}} = 562$ ($\varepsilon = 539$ M$^{-1}$ cm$^{-1}$) and 576 nm ($\varepsilon = 552$ M$^{-1}$ cm$^{-1}$). These signals were similar to those observed for the four-coordinate $C_2$-symmetric Bp$_2$Co species (Bp = H$_2$B(C$_3$N$_2$H$_3$)$_2$).$^{10a}$ The center of gravity of this band was blueshifted with respect to that for the related transition in 1 (see the absorption spectra in Figure 1). Note that a similar relationship was described between Bp$_2$Co and TpCoCl, in terms of a blueshifted absorption for the $^4$T$_1$(P)$\leftrightarrow^4$A$_2$(F) transition (for the tetrahedral site).$^{18}$ A second, lower-energy band was observed at $\lambda \approx$...
1040 nm ($\varepsilon = 120 \text{ M}^{-1} \text{ cm}^{-1}$), and this band was redshifted and more intense than the band for the transition in To$^M$CoCl. The high-spin nature of 2 was further supported by the solution magnetic moment of 4.2(2) $\mu_B$ (measured by the Evans method), which was slightly smaller than that of Bp$_2$Co but significantly reduced in comparison with that of six-coordinate Tp$_2$Co.$^{10a}$ The high-spin state of 2 at low temperature was confirmed by magnetometry experiments, which revealed $\mu_{\text{eff}} = 3.9(1) \mu_B$ at 12.5 K. An ESR spectrum of 2 acquired at 5 K as a point sample or as a glassed solution in 2-methyltetrahydrofuran contained only weak signals. One of the signals for the point sample showed evidence of the hyperfine coupling of the cobalt center in the form of a seven-line pattern. The discrepancy from the expected eight-line pattern for $I = 7/2$ was presumably caused by the broadness at the onset of the peak. In addition, the spectrum was consistent with a low-symmetry species.

Compound 1 and TlTo$^M$ react overnight to give 2, and this conversion occurs more rapidly than the reaction of TlTo$^M$ and CoCl$_2$ under equivalent conditions. In the attempted synthesis of 1 from anhydrous CoCl$_2$, the poor selectivity is hypothesized to come from the relative rates of transmetalation for CoCl$_2$ versus 1, and these rates are likely influenced by the heterogeneous nature of the reaction mixture. This contrasts with the reactivity of bulky tris(pyrazolyl)borato cobalt compounds. For example, Tp$^{\text{Ph,Me}}$CoCl and Tp$^{\text{Ph,Me}}$Tl do not afford (Tp$^{\text{Ph,Me}}$)$_2$Co.$^{11}$

In the course of our studies of the selective preparation of 1, polar solvents were tested to give a monophasic reaction (i.e., to improve the kinetics). In fact, the reaction of TlTo$^M$ and CoCl$_2$ in methanol gives a blue material, which was initially attributed to To$^M$CoCl on the basis of its color and $^1$H NMR spectrum in benzene-$d_6$, as a result of the
presence of 1 in the mixture. Moreover, the electronic absorption spectra of 1 and the
substance obtained from the reaction in methanol are similar (see Figure 1). However,
this material is less soluble in benzene than 1, and the XRD powder pattern of this
material did not match that expected for 1 (see Figure 5 below). The recrystallization of
this material from dichloromethane afforded blue X-ray-quality crystals, and a diffraction
study proved that this product is the protonated oxazoline derivative of 1, namely,
{HTo\textsuperscript{M}}CoCl\textsubscript{2} (3, Figure 4).

\begin{figure}
\centering
\includegraphics[width=\textwidth]{tca3}
\caption{Thermal ellipsoid plot of \{HTo\textsuperscript{M}}CoCl\textsubscript{2} (3) plotted at 50 % probability. H
atoms, with the exception of H1n, are omitted for clarity. Selected bond lengths (Å) and
angles (°): Co1–N1 2.006(3), Co1–N2 2.005(4), Co1–Cl1 2.237(3), Co1–Cl2 2.261(2);
N1–Co1–N2 96.2(1), N2–Co1–Cl1 109.77(9), N1–Co1–Cl1 115.18(9), N2–Co1–Cl2
111.68(9), N1–Co1–Cl2 110.6(1), Cl1–Co1–Cl2 112.40(9).
}
\end{figure}

The crystal structure of 3 shows a tetrahedral cobalt center featuring two chlorido
ligands and a bidentate To\textsuperscript{M}–cobalt interaction. The third oxazoline ring is protonated,
and HTo\textsuperscript{M} may be viewed as an overall charge-neutral ligand coordinated to CoCl\textsubscript{2}. Thus,
{HTo\textsuperscript{M}}CoCl\textsubscript{2} differs from 1 and 2 by its zwitterionic nature. Related iridium and rhodium compounds of protonated or methylated tris(oxazolinyl)borate complexes have been reported, as have zwitterionic cobalt(II) complexes stabilized by bulky tris(carbene)borate ligands.\textsuperscript{16,27} Compared with 1, compound 3 features slightly longer Co–Cl bonds (by more than 0.034 Å) and shorter Co–N bonds (by more than 0.025 Å).

The reaction of 1 and (HOEt\textsubscript{2})Cl provides an independent synthesis of 3 (equation (3)). Alternatively, LiTo\textsuperscript{M} and CoCl\textsubscript{2} react in wet dichloromethane to provide crystals of 3 directly from the reaction mixture.

$$\text{Ph-B} \begin{array}{c} \text{N} \\ \text{O} \end{array} \text{Co-Cl} + \text{HCl} \xrightarrow{\text{THF, r.t., 0.5 h}} \begin{array}{c} \text{H} \\ \text{N} \\ \text{B} \\ \text{O} \end{array} \text{O} \begin{array}{c} \text{N} \\ \text{O} \end{array} \text{Co-Cl} (3)$$

Compound 3 is highly soluble in dichloromethane and methanol but poorly soluble in benzene, toluene, tetrahydrofuran, and diethyl ether. The poor solubility of 3 and the high solubility of 1 in benzene-\textsubscript{d\textsubscript{6}} can result in erroneous interpretation of the purity of To\textsuperscript{M}CoCl on the basis of the NMR spectra of crude reactions mixtures. Thus, dry dichloromethane-\textsubscript{d\textsubscript{2}} is the best solvent for unambiguous NMR analysis of the purity of these compounds. The $^1$H NMR spectrum of 3 in dichloromethane showed paramagnetically shifted $^1$H NMR signals that were unassignable yet characteristic. The $^{11}$B NMR signal appeared at $\delta = -7.5$ ppm, and this peak was much less shifted from the diamagnetic range than those of 1 and 2. The infrared spectrum was consistent with the crystallographically determined structure and provided structural insights. The bands at
1588 and 1598 cm\(^{-1}\) were assigned to \(\nu_{CN}\) of cobalt-coordinated oxazoline and protonated oxazoline, respectively. Compound 3 was also a high-spin Co\(^{II}\) species, on the basis of magnetometry experiments that indicated a \(\mu_{\text{eff}}\) of 4.2(1) \(\mu_B\) at 5 K.

As all three compounds were formed from the same starting materials and \(^1\)H NMR spectroscopy provided ambiguous results during the course of these synthetic studies, we instead utilized powder XRD measurements to characterize the bulk compositions of the samples. These experiments, along with solution and solid-state IR spectroscopy, provide a connection between the solid-state and molecular structures that usually relies on solution-phase NMR spectroscopy (which was structurally uninformative for these (oxazolylborato)cobalt(II) compounds). Powder XRD is an appealing alternative characterization technique, because it is not complicated by the electronic structure or unpaired electrons and probes the composition of the crystalline component of the bulk powder.

The experimental powder XRD patterns matched the corresponding XRD patterns calculated from single-crystal data for samples of 1, 2, and 3. These results provided additional confidence that the single-crystal diffraction experiments corresponded to structures that describe the bulk samples. For example, the powder XRD experiments indicated that the preparation from CoCl\(_2\)·THF gives a reproducible and high-yielding route to 1 (Figure 5).
Figure 5. X-ray powder diffraction pattern of $\text{To}^\text{M}\text{CoCl}$ (1) at 293 K with Cu-$K\alpha$ radiation. The observed pattern (A) is shown in black, and the calculated intensity pattern (B) is indicated by red solid lines.

The viability of this method for the measurement of the purity was further tested with analytically pure $3/1$ in a 50:50 ratio. The XRD pattern of this mixture matched the expected powder pattern and verified that the presence of the two products could be identified by the diffraction method.

We also attempted to synthesize $\text{To}^\text{M}\text{CoX}$ compounds using cobalt acetate as the starting material. However, the reaction of cobalt acetate and $\text{TlTo}^\text{M}$ generated a mixture of 2 and another species, which was later identified as $\text{To}^\text{M}\text{CoOAc}$ (see below). Thus, $\text{CoCl}_2\cdot\text{THF}$ is the preferred starting material for the entry into the cobalt(II) chemistry of tris(oxazolinyl)borate ligands.
With isolable, fully characterized, and spectroscopically and analytically pure \( {\text{To}^M}\text{CoCl} \) in hand, we tested its reactivity in halide substitutions to prepare \( {\text{To}^M}\text{CoOrBu} \) (4) and \( {\text{To}^M}\text{CoOAc} \) (5). The reaction of \( {\text{To}^M}\text{CoCl} \) and KOrBu in tetrahydrofuran readily provides \( {\text{To}^M}\text{CoOrBu} \) (equation (4)).

\[
\begin{align*}
\text{Ph-} & \text{B} \quad \text{Co-} \begin{array}{c}
\text{Cl} \\
\text{O} \\
\text{O}
\end{array} + \text{KOrBu} \quad \xrightarrow{\text{THF}} \quad \text{Ph-} \text{B} \quad \text{Co-} \begin{array}{c}
\text{O} \\
\text{O}
\end{array} \\
\text{r.t., 12 h} & \quad 4, 87\%
\end{align*}
\]

In the \(^1\text{H} \) NMR spectrum of 4, all of the signals were readily assigned by integration, including a new signal at \( \delta = 11.16 \) ppm that integrated to 9 H relative to the \( {\text{To}^M} \) signals. This resonance was assigned to the OrBu group. Moreover, a new \(^{11}\text{B} \) NMR signal was detected at \( \delta = 73 \) ppm, which was further downfield than those of the other cobalt complexes. The paramagnetically shifted \(^{11}\text{B} \) NMR spectrum rules out the transmetalation of the \( {\text{To}^M} \) ligand to K.

Unlike blue \( {\text{To}^M}\text{CoCl} \), compound 4 is purple. Accordingly, the UV/Vis absorption spectrum of 4 contained a broad peak with a blueshifted onset edge at \( \lambda \approx 500 \) nm, as might be expected owing to the replacement of the weak-field chlorido ligand with a tert-butoxide ligand. The tridentate coordination of the tris(oxazolinyl)borate ligand to the cobalt center was suggested by a single strong IR band at 1590 cm\(^{-1} \), which was assigned to the oxazoline \( \nu_{\text{CN}} \) mode. Compound 4 is a high-spin cobalt(II) complex from room temperature to 36 K, as determined by magnetometry measurements (\( \mu_{\text{eff}} = 4.6(1) \mu_B \) at 36 K). Unlike compounds 1–3, \( {\text{To}^M}\text{CoOrBu} \) undergoes a broad, thermally induced spin transition, and \( \mu_{\text{eff}} \) is 1.3(1) \( \mu_B \) at 2.5 K, consistent with a low-spin state (\( S = 1/2 \)).
The recrystallization of 4 from toluene at –35 °C provided X-ray-quality crystals, and a single-crystal X-ray diffraction study confirmed the identity of 4 as To$^M$CoOrBu (Figure 6). The Co–N bonds in 4 are longer than those in 1. In addition, the B1–Co1–O4 angle in 4 is smaller than the B1–Co1–Cl1 angle in 1. The Co1–O4–C22 angle is 134.2(2)°, and this angle wedges the tBu group between the methyl groups of the N2 and N3 oxazolines.

**Figure 6.** Thermal ellipsoid plot of To$^M$CoOrBu (4). Ellipsoids are plotted at 50% probability. H atoms are omitted for clarity. Selected bond lengths (Å) and angles (°): Co1–O4 1.821(2), Co1–N1 2.049(3), Co1–N2 2.055(3), Co1–N3 2.026(2); O4–Co1–N1 111.2(1), O4–Co1–N2 128.0(1), O4–Co1–N3 131.0(1), B1–Co1–O4 166.2(1), Co1–O4–C22 134.2(2).

The reaction of To$^M$CoCl and NaOAc in THF provides To$^M$CoOAc (5) as a light purple solid in high isolated yield (97%; Equation (5)). Compound 5 was characterized by $^1$H and $^{11}$B NMR spectroscopy. The $^1$H NMR spectrum contained new tris(oxazolinyl)phenylborate signals, and the acetate signal appeared far downfield ($\delta =$
171.25 ppm). The $^{11}$B NMR signal of 5 appeared at 95 ppm, which is downfield relative to the signals for 1 and 4.

In the IR spectrum, a single band corresponding to the oxazoline $\nu_{\text{CN}}$ stretching mode was observed at 1591 cm$^{-1}$. Compound 5 is high spin ($\mu_{\text{eff}} = 4.8(2) \mu_B$), as determined by the Evans method. X-ray-quality crystals were obtained by recrystallization from pentane at $-35 \, ^\circ\text{C}$ (Figure 7). The Co–O bonds (2.098(2) and 2.089(2) Å) are longer than the Co–O bond in 4 (1.821(2) Å), and the Co–N bonds are also longer than those in 1 and 4.

Figure 7. Thermal ellipsoid plot of $\text{To}^\text{M} \text{CoOAc}$ (5). Ellipsoids are plotted at 50% probability. H atoms are omitted for clarity. Selected bond lengths (Å) and angles (°): Co1–O4 2.098(2), Co1–O5 2.089(1), Co1–N1 2.065(2), Co1–N2 2.064(2), Co1–N3 2.026(2); O5–Co1–N1 154.5(1), O5–Co1–N2 100.2(1), O5–Co1–N3 112.4(1), O5–Co1–O4 61.8(1).
Cobalt(II) acetate is used as an industrial catalyst in the oxidation of \( p \)-xylene to terephthalic acid,\(^{28} \) and nitrogen-ligand-based cobalt compounds have also been studied in cyclohexane oxidation to cyclohexanol.\(^{9d} \) In this oxidation catalysis, selectivity for a single oxidation to cyclohexanol, rather than over-oxidation to cyclohexanone or \( \varepsilon \)-caprolactone, is desired. Despite the important role of cobalt in oxidation chemistry, TpCoX-based compounds are reported as not effective for cyclohexane oxidation,\(^{29} \) so a study of To\(^{\text{M}}\)CoOAc as an oxidation catalyst provides a comparison with the pyrazolylborate analogues.

Complex 5 catalyzes the oxidation of cyclohexane to cyclohexanol with \textit{meta}-chloroperoxybenzoic acid (mCPBA) as the terminal oxidant and affords 536 equiv. of cyclohexanol per 1 equiv. of catalyst after 7 h. At the early stages of the reaction (3 h), the product ratio for cyclohexanol/cyclohexanone/\( \varepsilon \)-caprolactone is 20:1.5:1 with 337 equiv. of cyclohexanol formed per 1 equiv. of 5. This selectivity decreases as the reaction proceeds, and \( \varepsilon \)-caprolactone is formed from the reaction of mCPBA and cyclohexanone. Cumene hydroperoxide and \textit{tert}-butyl hydroperoxide were tested as oxidants for this catalytic transformation, but only mCPBA afforded the desired conversion. A control experiment, in which cyclohexane and mCPBA were mixed at room temperature, did not provide detectable quantities of cyclohexanol, cyclohexanone, or \( \varepsilon \)-caprolactone. Furthermore, anhydrous CoCl\(_2\) or anhydrous Co(OAc)\(_2\) (as Co\(_4\)(OAc)\(_7\)OH) are poorly soluble under these reaction conditions, and this limits comparisons with 5. However, the partially soluble suspended materials give lower activities and poorer selectivities than 5 under comparable conditions.
Figure 8. Concentration versus time plots for the products of cyclohexane oxidation with mCPBA. The experiment was performed four times.

To better identify the parameters associated with the oxidation process, the reaction mixture was monitored by GC to determine the concentrations of cyclohexanol, cyclohexanone, and ε-caprolactone. In a reaction with 0.16 mM 5, the concentration of cyclohexanol reaches a steady state after ca. 300 min. The plot of [cyclohexanol] versus time in Figure 8 can be fitted with a nonlinear least-squares regression ($R = 0.98$) to Equation (6).\textsuperscript{30}

From this analysis, the observed rate constant for the oxidation of cyclohexane to cyclohexanol ($k_1$) is $1.69(8) \times 10^{-4}$s\textsuperscript{-1}, and that for the oxidation of cyclohexanol is $3.9(1) \times 10^{-3}$s\textsuperscript{-1} (average of four experiments, see the Supporting Information below for individual curves). Thus, the selectivity for cyclohexanol production comes largely from the high initial [$C_6H_{12}$]. On the basis of these results, we are currently studying reaction
conditions (temperature, solvent polarity, and reagent concentrations) and catalyst structures that favor higher $k_1$ values and smaller $k_2$ values.

**Conclusion**

The synthesis of 1 from TlTo$^M$ and CoCl$_2$·THF provides an entry point into tris(oxazolinyl)borato cobalt chemistry. Like zinc and magnesium compounds, four-coordinate pseudotetrahedral cobalt complexes are supported by the tris(oxazolinyl)borate ligand, despite its reduced steric profile with respect to that of the highly bulky tert-butyl-substituted tris(pyrazolyl)borate scorpionates. The starting materials and reaction conditions are critical for obtaining 1, rather than 2 or 3. However, these other two compounds can be prepared by using 2 equiv. of TlTo$^M$ or polar protic solvents, respectively. Once the paramagnetic compounds were fully characterized by X-ray diffraction as well as IR, NMR, and UV/Vis spectroscopy, the characteristic spectroscopic signatures in the $^{11}$B NMR spectra, $^1$H NMR spectra, and powder XRD patterns are useful for establishing the synthetic reproducibility and product identities as well as the purities of the materials.

To$^M$CoCl is reactive with KOrBu and NaOAc in salt-metathesis reactions. In this context, we are currently working to synthesize, characterize, and study the reactivity of alkylcobalt(II) compounds supported by tris(oxazolinyl)borate ligands. To$^M$CoOAc is a catalyst for the selective oxidation of cyclohexane, in contrast to tris(pyrazolyl)borato cobalt compounds. We are currently preparing optically active analogues to study enantioselective C–H bond functionalization.
Experimental Section

General Experimental Methods: All reactions were performed by standard Schlenk techniques under dry argon. Tetrahydrofuran, diethyl ether, and toluene were dried and deoxygenated with an IT PureSolv system. Benzene-$d_6$ was heated under reflux over Na/K alloy and vacuum-transferred. CoCl$_2$ (Strem) was used to prepare CoCl$_2$·THF by Soxhlet extraction of CoCl$_2$ with THF. TlTo$^M$ was synthesized according to reported procedures. Caution! Thallium salts are highly toxic, and thallium-containing compounds and waste should be handled appropriately. KOrBu was purified by sublimation. mCPBA was purified by washing with K$_2$HPO$_4$/KH$_2$PO$_4$ buffer solution (pH 7.5). The $^1$H and $^{11}$B NMR spectra were recorded with a Bruker Avance III 600 spectrometer. The $^{11}$B NMR spectra were referenced to an external sample of BF$_3$·Et$_2$O. The infrared spectra were measured with a Bruker Vertex 80 FTIR spectrometer. The electron paramagnetic resonance (EPR) spectra were obtained with an X-band Elexsys 580 FT-EPR spectrometer in continuous-wave mode, and the spectra were simulated with XSophe. The direct current (DC) magnetization was measured with a Quantum Design MPMS-5 superconducting quantum interference device (SQUID) magnetometer. The UV/Vis spectra were recorded with an Agilent 8453 UV/Vis spectrophotometer with the analyte (2 mM) in dichloromethane. Elemental analyses were performed with a Perkin–Elmer 2400 Series II CHN/S analyzer. The catalytic cyclohexane oxidation experiments and the kinetic studies were performed with a Chemspeed Technologies SWING-XL automated platform. GC–MS was performed with Agilent 7890A GC and Agilent 5975C MS instruments equipped with an HP-5MS column. The powder XRD patterns were collected with a STOE WinXPOW powder diffractometer. The single-crystal X-ray
diffraction data were collected with an APEX II diffractometer. CCDC 1433255 (for 1), 1433256 (for 2), 1433257 (for 3), 1433258 (for 4), and 1457506 (for 5) contain the supplementary crystallographic data. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre.

**To**\textsuperscript{M}CoCl (1): TlTo\textsuperscript{M} (2.015 g, 3.43 mmol) was dissolved in THF (75 mL), and the solution was added dropwise to a stirred suspension of CoCl\textsubscript{2}·THF (1.01 g, 5.11 mmol) in THF (75 mL). The reaction mixture was stirred overnight, and then the solvent was evaporated to give a blue residue. The product was extracted with benzene (100 mL) and dried under vacuum to afford To\textsuperscript{M}CoCl (1) as a bright blue solid (1.55 g, 3.25 mmol, 94.8 % based on TlTo\textsuperscript{M}). Recrystallization from toluene gave analytically pure To\textsuperscript{M}CoCl. \textsuperscript{1}H NMR (benzene-\textit{d}\textsubscript{6}, 600 MHz): \(\delta = 24.88\) (s, 6 H, CNCMe\textsubscript{2}CH\textsubscript{2}O), 8.38 (s, 18 H, CNCMe\textsubscript{2}CH\textsubscript{2}O), 4.77 (s, 2 H, C\textsubscript{6}H\textsubscript{5}), 4.25 (s, 1 H, p-C\textsubscript{6}H\textsubscript{5}), –0.41 (s, 2 H, C\textsubscript{6}H\textsubscript{5}) ppm. \textsuperscript{11}B NMR (benzene-\textit{d}\textsubscript{6}, 128 MHz): \(\delta = –29.6\) ppm. IR (KBr): \(\nu = 2969\) (m), 2930 (m), 2898 (m), 2871 (m), 1588 (s, \(\nu_{\text{CN}}\)), 1461 (m), 1388 (m), 1369 (m), 1351 (m), 1273 (m), 1193 (m) 1162 (m), 1107 (m), 988 (s) cm\textsuperscript{-1}. UV/Vis (CH\textsubscript{2}Cl\textsubscript{2}): \(\lambda_{\text{max}} (\varepsilon) = 568\) (362), 635 (641), 940 (92 M\textsuperscript{-1}cm\textsuperscript{-1}) nm. Evans method: \(\mu_{\text{eff}} (C_{6}D_{6}) = 4.5\) \(\mu\text{B}\). C\textsubscript{21}H\textsubscript{29}BClCoN\textsubscript{3}O\textsubscript{3} (476.7): calcd. C 52.91, H 6.13, N 8.82; found C 53.26, H 6.20, N 9.03. M.p. 194–196 °C (dec.).

**To**\textsuperscript{M}\textsubscript{2}Co (2): TlTo\textsuperscript{M} (0.102 g, 0.173 mmol) and CoCl\textsubscript{2}·THF (0.0169 g, 0.0855 mmol) were stirred in THF (5 mL) for 1 d. The reaction mixture changed gradually from blue to purple. The reaction mixture was filtered, and the solvent was evaporated to afford (To\textsuperscript{M}\textsubscript{2}Co (2) as a purple solid (0.0689 g, 0.0837 mmol, 97.8 %). Recrystallization from toluene afforded analytically pure X-ray-quality single crystals of (To\textsuperscript{M}\textsubscript{2}Co. \textsuperscript{1}H NMR (benzene-\textit{d}\textsubscript{6}, 600 MHz): \(\delta = 15.04, 12.41, 2.63\) ppm. \textsuperscript{11}B NMR (benzene-\textit{d}\textsubscript{6}, 128 MHz): \(\delta = 15.04, 12.41, 2.63\) ppm.
= 42.3 ppm. IR (KBr): ν 2965 (m), 2928 (m), 2888 (m), 1602 (m, νCN), 1554 (s, νCN), 1463 (m), 1369 (m), 1279 (m), 1261 (m), 1198 (m) 1153 (m), 1100 (m), 1002 (m), 969 (m). UV/Vis (CH2Cl2): λmax (ε) = 562 (539), 576 (552), 1042 (120 M−1 cm−1) nm. Evans method: μeff (C6D6) = 4.2(2) μB. C42H58B2CoN6O6 (823.5): calcd. C 61.26, H 7.10, N 10.21; found C 61.21, H 7.20, N 9.78. M.p. 121–123 °C (dec).

{HToM}CoCl2 (3): Complex 1 (0.199 g, 0.417 mmol) was dissolved in THF (30 mL), and the solution was added dropwise to a solution of HCl in diethyl ether (210 µL, 0.420 mmol). The blue solution was stirred overnight, and the volatiles were removed in vacuo to afford {HToM}CoCl2 (3) as a bright blue solid (0.144 g, 0.281 mmol, 67.4%). Recrystallization from dichloromethane afforded analytically pure {HToM}CoCl2. 1H NMR (dichloromethane-d2, 600 MHz): δ = 17.3, 14.3, 9.9, 8.6, 7.0, 6.6, –21.4, –23.3 ppm. 11B NMR (dichloromethane-d2, 128 MHz): δ = –7.5 ppm. IR (KBr): ν 3270 (m, νNH), 2976 (m), 2932 (m), 2984 (m), 1598 (s, νCN), 1588 (s, νCN), 1462 (m), 1424 (m), 1371 (m), 1306 (m), 1274 (m), 1194 (m) 1171 (m), 965 (m), 937 (m) cm−1. UV/Vis (CH2Cl2): λmax (ε) = 567 (360), 614 (479), 635 (528), 999 (130 M−1 cm−1) nm. μeff (CD2Cl2) = 4.0(2) μB. C21H38BCl2CoN3O3 (513.1): calcd. C 49.16, H 5.89, N 8.19; found C 49.35, H 6.16, N 7.72. M.p. 218–221 °C (dec.).

ToMCoOrBu (4): Complex 1 (0.202 g, 0.423 mmol) was dissolved in THF (10 mL), and the solution was added to KOrBu (0.053 g, 0.472 mmol) to instantly afford a purple solution. The reaction mixture was stirred overnight, and then KCl was removed by filtration. The volatiles were evaporated in vacuo to afford a bright purple solid. The purple solid was washed with pentane (3 × 5 mL) and then dried under vacuum to yield ToMCoOrBu (0.189 g, 0.367 mmol, 86.8%). X-ray-quality crystals were obtained from
pentane at –40 °C. \(^1\)H NMR (benzene-\(d_6\), 600 MHz): \(\delta = 15.54\) (s, 6 H, CNCMe\(_2\)CH\(_2\)O), 11.33 (s, 2 H, C\(_6\)H\(_3\)), 11.16 (s, 9 H, CoOCMe\(_2\)), 9.51 (s, 2 H, C\(_6\)H\(_5\)), 8.00 (s, 1 H, \(p\)-C\(_6\)H\(_5\)), –6.26 (s, 18 H, CNCMe\(_2\)CH\(_2\)O) ppm. \(^{11}\)B NMR (benzene-\(d_6\), 128 MHz): \(\delta = 72.9\) ppm. IR (KBr): \(\nu = 2966\) (m), 2930 (m), 2889 (m), 1590 (s, \(\nu \text{CN}\)), 1465 (m), 1433 (m), 1386 (m), 1364 (m), 1278 (m), 1251 (m), 1193 (s), 1158 (m), 1002 (m), 966 (m), 928 (m) cm\(^{-1}\). UV/Vis (CH\(_2\)Cl\(_2\)): \(\lambda_{\text{max}}(\varepsilon) = 524\) (245), 575 (327), 624 (179), 657 (148 M\(^{-1}\)cm\(^{-1}\)) nm. C\(_{25}\)H\(_{38}\)BCoN\(_3\)O\(_4\) (514.3): calcd. C 58.38, H 7.45, N 8.17; found C 58.04, H 7.48, N 8.29. M.p. 233–235 °C (dec.).

**To\(^\text{M}\)CoOAc (5):** Complex 1 (0.101 g, 0.211 mmol) was dissolved in THF (5 mL), and the solution was added to NaOAc (0.017 g, 0.212 mmol). The reaction mixture was stirred for 24 h to afford a purple solution and a white precipitate. The NaCl precipitate was removed by filtration, and the volatiles were evaporated in vacuo to afford a light purple solid. The resulting purple solid was washed with pentane (3\(\times\) 5 mL) and dried under vacuum to yield To\(^\text{M}\)CoOAc (0.102 g, 0.204 mmol, 96.7 %). X-ray-quality crystals were obtained from pentane at –40 °C. \(^1\)H NMR (benzene-\(d_6\), 600 MHz): \(\delta = 171.25\) (s, 3 H, O\(_2\)CMe), 33.36 (s, 2 H, C\(_6\)H\(_3\)), 17.86 (s, 2 H, C\(_6\)H\(_5\)), 15.06 (s, 1 H, \(p\)-C\(_6\)H\(_5\)), 12.40 (s, 6 H, CNCMe\(_2\)CH\(_2\)O), 42.84 (s, 18 H, CNCMe\(_2\)CH\(_2\)O) ppm. \(^{11}\)B NMR (benzene-\(d_6\), 128 MHz): \(\delta = 95.3\) ppm. IR (KBr): \(\nu = 2964\) (m), 2930 (m), 2894 (m), 1590 (s, \(\nu \text{CN}\)), 1463 (m), 1442 (m), 1368 (m), 1351 (m), 1275 (m), 1197 (m), 1162 (m), 1098 (m), 1024 (m) 960 (m) cm\(^{-1}\). UV/Vis (CH\(_2\)Cl\(_2\)) \(\lambda_{\text{max}}(\varepsilon) = 486\) (51), 585 (104 M\(^{-1}\)cm\(^{-1}\)) nm. Evans method: \(\mu_{\text{eff}}\) (C\(_6\)D\(_6\)) = 4.8(2) \(\mu\). C\(_{23}\)H\(_{32}\)BCo\(_3\)NO\(_5\) (500.3): calcd. C 55.22, H 6.45, N 8.40; found C 55.21, H 6.58, N 7.79. M.p. 161–164 °C (dec.).
**ToMCoOAc-Catalyzed Oxidation of Cyclohexane:** The oxidation of cyclohexane was performed under conditions similar to those previously reported by Hikichi et al.\(^{[9d]}\) A reaction flask was charged with cyclohexane (1.6 mL, 15 mmol, 2.45 M) and ToMCoOAc (1 µmol, 0.16 mM) dissolved in dichloromethane/acetonitrile (95 %, 1 mL). Upon the addition of 1,2-dichloroethane/acetonitrile (3.5 mL, 97.5 %) containing mCPBA (345 mg, 2.0 mmol) and nitrobenzene (30 µL, 300 µmol), the reaction was initiated and maintained at 25 °C. At each time point, an aliquot (0.2 mL) was removed and quenched with triphenylphosphine (10 mg, 38.1 µmol). The organic products were identified and quantified by GC–MS through the integration of the peak areas with respect to a known amount of the nitrobenzene standard. In this analysis, the quenched reaction product (10 µL) was added to dichloromethane (1.5 mL) and analyzed by GC–MS. Each chromatogram was obtained under the following conditions: split, 25:1; inlet temperature, 250 °C; initial oven temperature, 45 °C; temperature ramp, 15 °C min\(^{-1}\) to 150 °C. From this data, the concentration of cyclohexanol versus time was analyzed with a nonlinear least-squares regression.

**References**


**Supporting Information**

*Figure S1.* Concentration vs. time plots for the oxidation of cyclohexane by $\text{To}^\text{M} \text{CoOAc}$ (5) (0.16 mM) with mCBPA performed on a Chemspeed SwingXL Catalysis Platform. Figures S1-4 show reproducibility of the oxidation experiments. Species were quantified using GC-MS based on calibration to a nitrobenzene standard.
Figure S2. Concentration vs. time plots for the oxidation of cyclohexane by $\text{To}^M\text{CoOAc}$ (5) (0.16 mM) with mCBPA performed on a Chemspeed SwingXL Catalysis Platform. Figures S1- S4 show reproducibility of the oxidation experiments. Species were quantified using GC-MS based on calibration to a nitrobenzene standard.
Figure S3. Concentration vs. time plots for the oxidation of cyclohexane by $\text{To}^\text{M}\text{CoOAc}$ (5) (0.16 mM) with mCBPA performed on a Chemspeed SwingXL Catalysis Platform. Figures S1- S4 show reproducibility of the oxidation experiments. Species were quantified using GC-MS based on calibration to a nitrobenzene standard.
Figure S4. Concentration vs. time plots for the oxidation of cyclohexane by $^{\text{M}}\text{CoOAc}$ (5) (0.16 mM) with mCBPA performed on a Chemspeed SwingXL Catalysis Platform. Figures S1- S4 show reproducibility of the oxidation experiments. Species were quantified using GC-MS based on calibration to a nitrobenzene standard.
CHAPTER 3: RAPID AND ORDERED CARBONYLATION AND OXYGENATION OF A COBALT(II) METHYL

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Abstract

The oxidative carbonylation of $\text{To}^\text{M}\text{CoMe} \ (1; \text{To}^\text{M} = \text{tris}(4,4\text{-dimethyl}-2\text{-oxazolinyl})\text{phenylborate})$ involves its rapid, reversible reaction with CO to form $\text{To}^\text{M}\text{Co}\{\text{C(O)Me}\}\text{CO} \ (2)$ followed by rapid reaction with $\text{O}_2$ yielding $\text{To}^\text{M}\text{CoOAc} \ (3)$, in contrast to the slow direct carboxylation of $\text{To}^\text{M}\text{CoMe}$ by $\text{CO}_2$.

Other Author’s contributions

Ellie L. Fought: Responsible for DFT calculations, TDDFT calculations, and DFT Hessian calculations of $\text{To}^\text{M}\text{CoMe}$, $\text{To}^\text{M}\text{CoMe(CO)}$, $\text{To}^\text{M}\text{Co}\{\text{C(O)Me}\}\text{CO}$, and $\text{To}^\text{M}\text{CoOAc}$. 
Introduction

Catalytic oxidative carbonylation reactions are generally proposed to involve metal-based oxidations.\textsuperscript{1,2} Consider, for example, catalytic production of benzoic acid from benzene, CO, and an oxidant. Palladium(II) mediates the combination of the arene, CO and water to afford benzoic acid, with palladium(0) generated as a byproduct. Then, the suggested catalytic cycle is completed by metal-based oxidation, rather than oxygenation of an acylpalladium intermediate. Interestingly, the proposed biosynthesis of the energy-carriers acetate or acetyl-CoA follows a conceptually similar outline, involving insertion of CO into a metal-methyl bond followed by metal-based oxidation and thiolysis (reductive elimination of the C–S bond) to form acetyl-CoA.\textsuperscript{3,4}

Oxygenation of metal-carbon bonds (or the reductive coupling followed by oxidation pathway) is key to many hydrocarbon functionalization schemes. On the other hand, acetate may be derived directly from CO\textsubscript{2} and methyl transition metal compounds. Although CO\textsubscript{2} is a substrate for acetogenesis, it is reduced by carbon monoxide dehydrogenase (CODH) to CO prior to its interaction with the metal-methyl.\textsuperscript{4} Likewise, oxidative carbonylation yields are sometimes improved under CO\textsubscript{2}, yet CO is the ultimate source of carbonyl in the transformation. The distinction between CO/[O] in oxidative carbonylation versus CO\textsubscript{2} in carboxylation, in terms of their kinetic and thermodynamic parameters, is critical for the development of sustainable processes that utilize C\textsubscript{1} starting materials, such as methane, carbon monoxide, or carbon dioxide, because the processes must balance atom and energy economy with rate, yield, and selectivity to be viable.

Four-coordinate organocobalt(II) species are expected to be reactive toward CO, O\textsubscript{2} and CO\textsubscript{2} based on studies of bulky Tp\textsuperscript{Bu,Me}, Tp\textsuperscript{Pr}, Tp\textsuperscript{Pr2}, and [PhTt\textsuperscript{Bu}] (Tp\textsuperscript{R,R'} =
These reactions give either reduction or carbonylation products, oxidation or oxygenation, or carboxylation, respectively. CO affects one-electron reductions to give Tp\(^{Bu}\)CoCO or Tp\(^{Pr,Me}\)Co(CO), while carbonylation to cobalt(II) acyl species is also reported. Oxygenation of organocobalt(II) to a cobalt alkyl peroxide could follow our observations for related zinc compounds, where kinetics of zinc alkyl oxygenation to alkylperoxides or alkoxides are consistent with a turnover-limiting bimolecular substitution of zinc alkyl with alkylperoxy radical (S\(_{12}\)). This oxygenation pathway avoids metal-centered oxidation. Indeed, alkylperoxy and acylperoxy metal compounds are intermediates in oxygenation reactions. In the present work, a comparison of oxygenation, carbonylation and oxygenation, and carboxylation of To\(^{M}\)CoMe (1; To\(^{M}\) = tris(4,4-dimethyl-2-oxazolinyl)phenylborate) reveals that the kinetically favored pathway selectively produces acetate through an ordered, multistep sequence.

**Results and Discussion**

The reaction of To\(^{M}\)CoCl and MeLi at room temperature affords To\(^{M}\)CoMe (1) as a deep aquamarine solid (equation (1)).

\[ \text{To}^{M}\text{CoCl} + \text{Mel} \xrightarrow{\text{benzene, r.t., 1.5 h}} \text{To}^{M}\text{CoMe} \]

(1)

The \(^1\)H NMR spectrum of 1 showed signals at 15.44 and –12.05 ppm assigned to the oxazoline’s methylene and methyl groups, respectively, on the basis of integration.
These signals were shifted downfield compared to To^MCoCl (24.88 and 8.38 ppm, respectively). A resonance for the methyl ligand was not detected. The ^11^B signal at 100 ppm was significantly shifted in comparison to the peak of To^MCoCl (~29 ppm). A single ν_{CN} band at 1594 cm\(^{-1}\) in the IR spectrum suggested tridentate To^MCo coordination. The UV-Vis spectrum of 1 (in Et\(_2\)O) revealed intense absorptions at 346 (ε: 1412 M\(^{-1}\)cm\(^{-1}\)) and 697 (ε: 1078 M\(^{-1}\)cm\(^{-1}\)) assigned to charge transfer transitions associated with the alkyl ligand on the basis of their large molar absorbivities (>1000 M\(^{-1}\)cm\(^{-1}\)) and the lack of similar signals in To^MCoCl. Weaker absorptions at 581 (ε: 388 M\(^{-1}\)cm\(^{-1}\)) and 617 (ε: 424 M\(^{-1}\)cm\(^{-1}\)) were attributed to d←d transitions. The effective magnetic moment of 1 (4.2(2) μ\(_B\)), determined by Evans method, is consistent with a high-spin cobalt(II) (S = 3/2). The EPR spectrum of 1, acquired at 5 K in glassed toluene, showed a rhombic spectrum with hyperfine coupling to the ^59^Co center (I = 7/2) in a characteristic eight-line pattern.

The identity of 1 as the cobalt(II) methyl is unambiguously established by X-ray diffraction (Figure 1). The compound is slightly distorted from the C\(_{3v}\) symmetry suggested by solution-phase spectroscopy, with the C22 (methyl) displaced from the B–Co vector (∠B–Co–Me: 172.83°). The Co1–C22 distance in To^MCoMe (1.994(2) Å) is within the range of similar pseudotetrahedral methylcobalt(II) species, which vary from 1.9 to 2.1 Å.⁵,⁶,⁶d,¹⁶,¹⁰,¹⁷
**Figure 1.** Rendered thermal ellipsoid diagram of $\text{To}^\text{M}\text{CoMe}$ (1) with ellipsoids plotted at 50% probability. H atoms are omitted for clarity.

DFT calculations show that the quartet state ($S = 3/2$) is lowest energy and 49 kcal/mol lower than the doublet state. A TDDFT-calculated electronic transition at 310 nm (using implicit solvation) allows the experimental band at 346 nm to be classified as LMCT. This transition involves occupied orbitals mostly on the methyl carbon to unoccupied orbitals delocalized over Co, B, and the unsaturated C in the $\text{To}^\text{M}$ ligand. Additional peaks at 602 nm, 740 nm and 743 nm, found both in $\text{To}^\text{M}\text{CoMe}$ and $\text{To}^\text{M}\text{CoCl}$ calculations, support the assignment of the weaker signals in the experimental spectra as $d\rightarrow d$ transitions. The strong, experimentally observed band at 697 nm was not evident, which may be due to the single configuration approach of TDDFT.
To\textsuperscript{M}CoMe and CO (1 atm) rapidly react in benzene-$d_6$ or THF at room temperature, as evidenced by an immediate color change from blue to orange. The possible products of To\textsuperscript{M}CoMe and CO include reduced To\textsuperscript{M}CoCO or To\textsuperscript{M}Co(CO)\textsubscript{2} species, To\textsuperscript{M}Co(Me)CO or its inserted isomer To\textsuperscript{M}Co(C(O)Me), To\textsuperscript{M}Co{C(O)Me}CO, or To\textsuperscript{M}Co{C(O)Me}(CO)\textsubscript{2}. A single $^{11}\text{B}$ NMR signal at $-4$ ppm, shifted upfield by 104 ppm from the value for 1, suggested the formation of a single To\textsuperscript{M}Co-containing product (2). The $^1\text{H}$ and $^{11}\text{B}$ NMR spectra did not vary from room temperature to $-80\,^\circ\text{C}$.

Evaporation of a solution of 2 to dryness overnight gives 1 as the only detectable To\textsuperscript{M}Co-containing species, indicating that the reaction of 1 and CO is reversible. This reversibility hampers the isolation of 2. Compound 2 persists in partially degassed solutions, while evaporation of all solvent and immediate redissolution affords a mixture of 1 and 2. The reversible interaction of 1 and CO rules out formation of To\textsuperscript{M}CoCO because the byproducts of 1 e$^-$ reduction of 1 are unlikely to persist in a form that could re-methylate 2.

An in situ IR spectrum of the orange THF solution, collected on a ZnSe ATR crystal, revealed bands at 1984, 1886, 1687, and 1655 cm$^{-1}$. The two higher energy peaks were assigned to coordinated CO ligands, with the higher energy peak at 1984 cm$^{-1}$ significantly more intense. The lower energy absorptions were assigned to rotamers of the cobalt acyl group, and the signal at 1655 cm$^{-1}$ was notably non-Gaussian shaped with a shoulder tailing toward the red. On the basis of DFT calculations (see below), we assign this shoulder to the $\nu_{\text{CN}}$ of a weakly or non-coordinated oxazoline, which typically appears at 1630 cm$^{-1}$.$^{15}$ In addition, a peak at 1590 cm$^{-1}$, corresponding to the $\nu_{\text{CN}}$ of cobalt-coordinated oxazoline, was red-shifted by $\sim$4 cm$^{-1}$ in comparison to the To\textsuperscript{M}CoMe
starting material. For comparison, the IR spectrum of isolated Tp\(^{Pr_2}\)Co\{C(O)Et\}CO contained a single CO band at 1999 cm\(^{-1}\) and an acyl peak at 1636 cm\(^{-1}\).\(^7\) The IR spectrum of PhTt\(^{tBu}\)Co(C(O)Me)CO contained carbonyl \(v\)\(_{\text{CO}}\) at 1993 cm\(^{-1}\) and acyl \(v\)\(_{\text{CO}}\) at 1684 and 1663 cm\(^{-1}\) (assigned to rotomers).\(^10\) Thus, the interaction of To\(^M\)CoMe and CO affords a mixture of To\(^M\)Co{C(O)Me}CO (2; major, 1984, 1687, and 1655 cm\(^{-1}\)) and To\(^M\)Co(Me)CO ( minor, 1886 cm\(^{-1}\); Scheme 1).

Scheme 1. Reversible reaction of To\(^M\)CoMe (1) and CO.

The effective magnetic moment of 2 is 2.7(1) \(\mu_B\) (determined by Evans method). This value is reduced with respect to high spin To\(^M\)CoMe and is distinct from that of low-spin acyls PhTt\(^{tBu}\)Co{C(O)R}CO (R = Me, Et, Ph; \(\mu_{\text{eff}}\) = 1.9 – 2.1 \(\mu_B\)).\(^10\) The result for 2 does not fit the spin-only \(\mu_{\text{eff}}\) for low spin Co(II) (\(S = \frac{1}{2}\) is 1.73 \(\mu_B\)). While high spin Co(I) (\(S = 1\)) would give a spin-only value of 2.83 \(\mu_B\) (e.g., for Tp\(^{Np}\)CoCO and Tp\(^{Pr,Me}\)CoCO, \(\mu_{\text{eff}} = 3.1(1) \mu_B\)), this type of product is ruled out above.\(^11,18\) Instead, the effective magnetic moment of 2 is rationalized by a square pyramidal structure with a long axial Co–N interaction on the basis of the typical moments for square planar cobalt(II) complexes.\(^19-21\) While a trigonal bipyramidal structure is also consistent with the magnetic moment, DFT calculations (described below) are more consistent with the square pyramidal geometry. A room temperature isotropic signal (\(g_{\text{iso}} \approx 2.1\)) in the EPR spectrum of 2 further supported the low spin assignment.\(^22,23\) The UV-vis spectrum of 2
was distinct from 1 and contained a broad, weak band from 760 to 1100 nm with a $\lambda_{\text{max}}$ at 885 nm ($\varepsilon$: 305 M$^{-1}$ cm$^{-1}$). In addition, a strong absorption tails from 200 to 600 nm.

DFT calculations of $^{\text{Mo}}\text{Co}\{\text{C(O)Me}\}\text{CO}$ suggest a square pyramidal geometry for the optimized structure. The low spin state is 33 kcal/mol lower in energy than the high spin state. A TDDFT calculation with implicit solvation finds strong electronic transitions at 448 and 462 nm, with multiple smaller contributing transitions between 300 nm and 400 nm, consistent with experimental findings. In a DFT Hessian calculation, the frequency corresponding to the acyl carbonyl stretching mode is found at 1731 cm$^{-1}$ and the terminal carbonyl stretching is located at 2117 cm$^{-1}$. In addition, the calculated $\nu_{\text{CN}}$ stretches occur at 1636, 1667, and 1693 cm$^{-1}$ with the highest energy $\nu_{\text{CN}}$ being associated with the non-coordinated oxazoline. DFT calculations also suggest a square pyramidal geometry for the optimized structure of $^{\text{Mo}}\text{Co}(\text{Me})\text{CO}$. The low spin state is found to be 30 kcal/mol lower in energy than the high spin state. TDDFT calculations identify electronic transitions at 386 nm and 395 nm, which could be obscured by the multiple excitations found with the $^{\text{Mo}}\text{Co}\{\text{C(O)Me}\}\text{CO}$ species. The calculated frequency for the CO stretch at 2030 cm$^{-1}$ is lower energy than in $^{\text{Mo}}\text{Co}\{\text{C(O)Me}\}\text{CO}$, providing support for the assignment of the experimental spectrum.

The orange carbonylated cobalt 2 rapidly reacts with O$_2$ to give purple $^{\text{Mo}}\text{CoOAc}$ (3, equation (2)), identified by comparison with an authentic sample’s electronic spectrum ($\lambda_{\text{max}}$ = 486 and 585 nm), $^1$H and $^{11}$B NMR spectra, and X-ray diffraction pattern.\textsuperscript{15} This reaction is sufficiently rapid and selective (and $^{\text{Mo}}\text{CoOAc}$ is easily crystallized) that this product is typically isolated from attempted crystallizations of the monocarbonyl acyl complex at −80 °C due to trace O$_2$ impurities.
generated 2 reacts with O₂ to afford 3 under a range of conditions, including 1 atm of O₂ at room temperature (analyzed by ¹H and ¹¹B NMR spectroscopy) or addition of O₂ as a THF solution (5 – 20 equiv. O₂, with or without excess CO, analyzed by UV-vis spectroscopy) at room temperature. Also, addition of a solution of O₂ (10-20 equiv.) to 2 at −100 °C provides an intermediate (λ_max = 511, 550, and 585 nm) that converts into 3 upon warming to room temperature. That is, the transformation of equation (2) is fairly robust.

The reverse order of reaction of T⁰CoMe, oxidation followed by carbonylation, is not nearly so robust, selective or efficient. Reactions of 1 and O₂ at room temperature in THF or toluene-d₈ give a mixture of unidentified species that are unreactive toward CO. At −100 °C, a new, single ¹¹B NMR signal at −25 ppm suggested one species is formed from addition of T⁰CoMe and O₂ (1 atm). This species persists (based on unchanged ¹¹B NMR and EPR spectra) and is unreactive toward CO until the mixture is warmed to −20 °C. At that point, the solution becomes purple and T⁰CoOAc forms as a minor component in a mixture of T⁰Co-containing species.

The direct carboxylation reaction of 1 with CO₂ (1 atm or 85 psi) to form 3 requires 2 weeks (equation (3)), as monitored by ¹H NMR spectroscopy and a gradual blue to purple color change. The rate of this conversion is significantly slower than the rapid carbonylation/oxidation chemistry described above. A related reaction of 2 and CO₂
also sluggishly provides $\text{To}^M\text{CoOAc}$ at room temperature over ~2 weeks. Thus, while CO$_2$ insertion is possible, carbonylation and oxygenation by O$_2$ is the kinetically favored pathway.

These three sets of experiments provide considerable insight into the oxidative carbonylation sequence. First, a pathway from 2 to 3 involving 1 as an intermediate via initial decarbonylation steps is unlikely because decarbonylation appears to be slow as indicated by the long drying times needed to fully convert 2 to 1. Second, catalytic oxidation of CO to CO$_2$ by $\text{To}^M\text{Co}^{II}$/O$_2$ followed by carboxylation is ruled out as kinetically unfeasible by the experiment of equation (3).

**Conclusion**

We conclude that the sequence involving carbonylation of 1 to produce 2 followed by oxygenation to form 3 is kinetically favored. These results suggest that strategies for utilization of CO$_2$ involving carboxylation of organic compounds could benefit from an enzyme-inspired approach involving initial reduction to CO, CO migratory insertion, and finally oxidation rather than a direct, one-step carboxylation. At least in this case, a multistep compulsory-ordered pathway is considerably faster (and more selective) than random addition or direct insertion. Because acetate is important as a privileged ligand in oxidation catalysis and in CH activation pathways, the oxidative carbonylation studied here may be useful as part of selective hydrocarbon
functionalization schemes. We are currently investigating related cobalt(II) alkyl and aryl compounds in pursuit of catalytic conversions.

**Experimental Section**

**General Procedures.** All reactions were performed using standard Schlenk techniques under an atmosphere of dry argon. Benzene and diethyl ether were dried and deoxygenated using an IT PureSolv system. Benzene-$d_6$ was degased with freeze-pump-thaw cycles, heated to reflux over a Na/K alloy, and then vacuum transferred. $T^{M}$CoCl and $T^{M}$CoOAc (3) were synthesized following the reported procedure. $^1$H and $^{11}$B NMR spectra were collected on a Bruker Avance III 600 spectrometer. $^{11}$B NMR spectra were referenced to an external sample of BF$_3$·Et$_2$O. Infrared spectra were measured on a Bruker Vertex 80 FTIR spectrometer. EPR were obtained on an X-band Elexsys 580 FT-EPR spectrometer in continuous wave mode. UV-Vis spectra were recorded on an Agilent 8453 UV-vis spectrophotometer. The electronic spectrum of $T^{M}$CoOAc was not previously disclosed and is reported here. Elemental analyses were performed using a Perkin-Elmer 2400 Series II CHN/S. Single crystal X-ray diffraction data was collected on an APEX II.

$T^{M}$CoMe (1). Methylithium (1.6 M in diethyl ether, 1.00 mL, 1.6 mmol) was added to a solution of $T^{M}$CoCl (0.510 g, 1.07 mmol) in benzene (50 mL). The green reaction mixture was stirred for 1.5 h, the salt byproduct was removed by filtration, and the solvent was evaporated to afford $T^{M}$CoMe as a dark aquamarine solid (0.353 g, 0.773 mmol, 72%). X-ray quality crystals were obtained from pentane at −40 °C. $^1$H NMR (benzene-$d_6$, 600 MHz): δ 15.44 (s, 6 H, CNCMe$_2$CH$_2$O), 14.18 (s, 2 H, C$_6$H$_5$), 10.61 (s, 2 H, C$_6$H$_5$), 8.88 (s, 1 H, $p$-C$_6$H$_5$), −12.05 (s, 18 H, CNCMe$_2$CH$_2$O). $^{11}$B NMR (benzene-
$d_6$, 128 MHz): $\delta$ 100.3. IR (KBr, cm$^{-1}$): $\nu$ 2967 (m), 2925 (m), 2897 (m), 2861 (m), 1594 (s, $\nu$CN), 1462 (m), 1386 (m), 1352 (m), 1269 (m), 1194 (m), 1160 (m), 960 (m). UV-vis (Et$_2$O) $\lambda_{max} = 346$ ($\varepsilon$: 1412 M$^{-1}$cm$^{-1}$), 581 ($\varepsilon$: 388 M$^{-1}$cm$^{-1}$), 617 ($\varepsilon$: 424 M$^{-1}$cm$^{-1}$), 697 ($\varepsilon$: 1078 M$^{-1}$cm$^{-1}$). $\mu_{eff}(C_6D_6) = 4.2(2)$ $\mu_B$ as determined by the Evans method. Anal. Calcd. for C$_{22}$H$_{32}$BCoN$_3$O$_3$: C, 57.91; H, 7.07; N, 9.21 Found: C, 57.93; H, 7.18; N, 9.20. Mp. 235-237 °C, dec.

References


CHAPTER 4: CARBOXYLATES FROM REVERSIBLE CARBONYLATION FOLLOWED BY OXYGENATION OF A SERIES OF ORGANOCOBALT COMPOUNDS

Modified from a paper to be submitted to a journal

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Abstract

The organocobalt scorpionate compounds To\textsuperscript{M}CoR (To\textsuperscript{M} = tris(4,4-dimethyl-2-oxazoliny1)phenylborate; R = Bn, 1; CH\textsubscript{2}SiMe\textsubscript{3}, 2; Ph, 3; Et, 4; \textsuperscript{6}Bu, 5; Me, 6) are synthesized to investigate the impact of the organometallic moiety in carbonylation, oxidation, and carboxylation reactions. The compounds are prepared by reaction of To\textsuperscript{M}CoCl with the corresponding organolithium or organopotassium reagents. Compounds 1 – 6 were characterized by 8-line hyperfine coupling to cobalt in EPR spectra and solution phase magnetic measurements (\(\mu_{\text{eff}} = 4 - 5 \mu_B\)) as containing a high-spin cobalt(II) center. The UV-Vis spectra revealed a diagnostic band at ca. 700 nm (\(\varepsilon > 1000 \text{ M}^{-1}\text{cm}^{-1}\)) associated with the tetrahedral organocobalt(II) center and assigned to a \(\text{d} \leftarrow \text{d}\) transition on the basis of configuration interaction (CI) calculations. Complexes 1 – 6 react rapidly with CO to form equilibrating mixtures of the low spin organocobalt carbonyl To\textsuperscript{M}Co(R)CO, acyl To\textsuperscript{M}CoC(=O)R, and acyl carbonyl To\textsuperscript{M}Co{C(O)R}CO. The \(^1\text{H}\) and \(^{11}\text{B}\) NMR spectra contained only one set of signals for the CO-treated solutions, whereas the solution-phase IR spectra contained up to two \(\nu_{\text{CO}}\) and three \(\nu_{\text{C(=O)R}}\) signals with intensities varying depending on the R group (R = Bn, 7; CH\textsubscript{2}SiMe\textsubscript{3}, 8; Ph, 9; Et, 10; \textsuperscript{6}Bu, 11; Me, 12). Single crystal X-ray diffraction of To\textsuperscript{M}Co{C(O)Et}CO (10) supported its assignment as a square pyramidal cobalt(II) acyl carbonyl complex. Upon evaporation
of volatiles, solutions of 8 – 12 revert to the CO-free organocobalt starting materials 2 – 6, whereas attempts to isolate benzyl-derived 7 affords an unusual alkoxy ketone species, characterized by single crystal X-ray diffraction. Despite these differences observed in the carbonylation of 1 – 6 as a result of varying the R group, compounds 7 – 12 all react rapidly with O₂ through an oxygenation pathway to afford the corresponding carboxylate compounds ToMCoO₂CR (R = Bn, 13; CH₂SiMe₃, 14; Ph, 15; Et, 16; "Bu, 17; Me, 18). In contrast, the insertion of CO₂ into the Co–C bond in 7 – 12 requires several days to weeks.

Introduction

Oxidative carbonylation, an organotransition metal-mediated route to carboxylates, typically proceeds by a sequence in which a metal hydrocarbyl reacts with CO to form an acyl, followed by hydrolysis and reductive elimination. Under catalytic conditions, oxidation and metalation completes the cycle to generate a new metal hydrocarbyl. This kind of pathway has been proposed for palladium-catalyzed oxidative carbonylation of arenes as well as catalytic carboxylations of amides to give carbamates and ureas.¹ ² Remarkably, the biological synthesis of acetate also involves carbonylation of an organometallic nickel methyl to give an acetyl group that is transferred to acetyl Co-A to form a thioacetate and subsequently hydrolyzed.³ ⁶ The fact that carbon dioxide, which serves as the source of both carbon atoms in acetate, is not incorporated into acetate by direct insertion into the metal-methyl is perhaps even more remarkable. Instead, CO₂ is reduced both to the methyl and to CO by CO dehydrogenase.⁴ ⁵ Similarly in synthetic chemistry, the synthesis of acetate or acetic acid via the Monsanto process involves CO and oxidation rather than direct insertion of CO₂. These oxidative
carbonylations result in oxygenation of an acyl to carboxylate, but the pathway invokes hydrolysis followed by oxidation at the metal center rather than by direct oxygenation of the metal acyl species. The distinction between oxidation catalysis (including reactions mediated by oxidases) and oxygenation catalysis (catalyzed by oxygenases) affects the choice of reagent as the oxygen source and oxidant, as well as the conditions and occasions for their use. Nonetheless, acyl species are proposed as likely intermediates in multiple pathways, and identifying the conditions by which acyl metal compounds form and their subsequent reaction pathways are key to developing new transformations.

Routes to acyl compounds involve an insertive combination of CO and organometallic compounds. For tetrahedral organometallic compounds, however, a number of species and pathways can result from interactions with CO. The coordination of CO to the metal center gives a metal hydrocarbyl carbonyl adduct, which can undergo insertion to form an isomeric acyl species. Further coordination of one or two CO ligands is likely influenced by the steric properties of the ancillary ligand or acyl group and by the electronic configuration of the metal center. Alternatively, reactions of organometallics with CO can result in 1 e⁻ reduction to form carbonyl adducts. Reduction is typically observed with bulky ancillary ligands, such as in the reactions of CO and TpBuMeCoMe or PhTpBuFeMe that form cobalt(I) or iron(I) carbonyls respectively. Homolysis of tetrahedral cobalt alkyls is also proposed as the first step in the rearrangement of TpPhMeCoBu to TpPhMeCoCH2CHMe2. Interestingly, TpRCoEt (TpR = HB(3,5-iPr2-N2C3H), HB(3,4,5-Me3N2C3)3) and CO provides first the acylcobalt carbonyl species, which forms TpRCoCO upon removal of volatiles and this is suggested to occur by homolysis of TpRCoEt(CO). In contrast, allyl and benzyl derivatives
Tp^R Co\{C(O)R\}CO (R = CH$_2$C$_6$H$_4$OMe, C$_3$H$_5$) are isolable. The fate of the organometallic ligand in these reductive pathways has not been identified.

Conceivably, a metal acyl species in equilibrium with the metal alkyl could undergo ligand rearrangement to form a ketone that then inserts into a cobalt acyl species to form an OC–CO moiety. It is worth noting that such C–C bond formation to form OC–CO is extremely unusual, and to the best of our knowledge, is not known to occur under carbonylation conditions. Instead, complexes that contain coordination of an OC–CO moiety to the metal center are typically synthesized by ligand substitution reactions with an organic compound that already contains this structural feature.$^{11-13}$

Previously, we reported the synthesis of To$^M$CoMe by reaction of To$^M$CoCl with MeLi confirming that salt metathesis is a viable route to cobalt(II) organometallic complexes despite the hard basic nature of the To$^M$ ancillary ligand that can at times instead result in transmetalation to Mg or Li rather than the desired alkylation.$^{14,15}$ To$^M$CoMe reacts readily with CO to form To$^M$Co\{C(O)Me\}CO followed by rapid reaction with O$_2$ to produce To$^M$CoOAc. Direct insertion of CO$_2$ into To$^M$CoMe also affords To$^M$CoOAc, but the reaction requires several weeks. The observation that the multistep pathway is, at least in this case, significantly faster than direct insertion, motivated us to expand our study to other R groups to determine if this reactivity is general to other alkyl and aryl groups.

Herein, we prepare a series of alkyl, aryl, and benzyl cobalt(II) compounds supported by a tris(oxazoliny1)borate ligand. The spectroscopic, electronic, and structural features are compared within the series of organocobalt(II) compounds and to halide and pseudo-halide analogues, to identify signatures of classes of these paramagnetic
compounds and discover distinguishing features that might parallel reactivity differences within the series of compounds and to related four-coordinate cobalt alkyls. The products of carbonylation are characteristic of the hydrocarbyl ligand, as identified by signals observed in the infrared spectra in carbonyl and acyl C=O stretching regions. These carbonylation products are oxygenated by reaction with O$_2$, which occurs rapidly, in contrast to sluggish reactions of the organocobalt compounds and carbon dioxide.

**Results and Discussion**

**Synthesis and characterization of To$^M$CoR.** The organometallic cobalt(II) complexes To$^M$CoR (R = Bn (1), CH$_2$SiMe$_3$ (2), Ph (3), Et (4), $^n$Bu (5), Me (6); To$^M$ = tris(4,4-dimethyl-2-oxazolinyl)phenylborate) are prepared by salt metathesis reactions involving excess (1.4 – 1.7 equiv.) organopotassium (PhCH$_2$K) or organolithium (Me$_3$SiCH$_2$Li, PhLi, EtLi, nBuLi, MeLi) reagents and To$^M$CoCl (Scheme 1).$^{16}$ To$^M$CoMe and To$^M$CoBn are the most straightforward to prepare and form in good yields at room temperature under dilute conditions. For example, To$^M$CoBn (1, 0.129 g, 0.242 mmol, 78.7%) is synthesized from To$^M$CoCl (0.308 mmol, 0.031 M) and 1.7 equiv. of KBN in THF at room temperature.

Dilute conditions (~0.02 M) are also effective on a ~0.03 mmol scale for synthesizing organocobalt(II) complexes 2 – 5 in good yield (>50%). In contrast, preparative scale reactions for 2 – 5 (>0.06 mmol) under these dilute conditions consistently give less than 30% yield. Instead, 2 – 5 require more concentrated conditions (~0.1 M) and low temperature. Using 0.1 M To$^M$CoCl, 1.4 equiv. of alkylolithium, and mixtures cooled to −78 °C, To$^M$CoCH$_2$SiMe$_3$ (2), To$^M$CoPh (3), To$^M$CoEt (4) and To$^M$Co$^n$Bu (5) are reproducibly synthesized in greater than 0.20 mmol quantities and
>60% yields. These conditions provide spectroscopically pure and analytically pure

$\text{To}^M\text{CoR}$. Signals for $\text{To}^M\text{CoCl}$ in NMR and UV-vis spectra, even as a trace impurity, were not detected for these samples.

Scheme 1. Synthesis of $\text{To}^M\text{CoBn}$ (1), $\text{To}^M\text{CoCH}_2\text{SiMe}_3$ (2), $\text{To}^M\text{CoPh}$ (3), $\text{To}^M\text{CoEt}$ (4), $\text{To}^M\text{Co}^n\text{Bu}$ (5), and $\text{To}^M\text{CoMe}$ (6).

NMR spectroscopy provided an initial assay for alkyl- or arylation of $\text{To}^M\text{CoCl}$. Despite the paramagnetic nature of these cobalt(II) complexes, both $^1\text{H}$ and $^{11}\text{B}$ NMR spectroscopy clearly distinguished $\text{To}^M\text{CoCl}$ from the organocobalt(II) complexes by their chemical shifts (Table 1). The $\text{To}^M$-based chemical shifts were consistent with $C_{3v}$-symmetric species and appeared in similar regions for compounds 1 – 6. For example, the signals attributed to the oxazoline methyl groups ranged from –9.6 to –14.5 ppm, which was more than 15 ppm upfield compared to corresponding signals in $\text{To}^M\text{CoCl}$ at 8.38 ppm. The oxazoline methylene peaks’ range was even smaller, from 14.8 – 16.7 ppm, whereas the corresponding signal in $\text{To}^M\text{CoCl}$ was observed at 24.88 ppm.
While the chemical shifts for the $\text{To}^M$ ligand were consistent across the organometallic complexes, the detectable signals for the alkyl and aryl ligands were wide ranging. For example, the benzyl ligand resonances were observed at 34, –77, and –89 ppm. $^1\text{H}$ NMR peaks that might be attributed to hydrogen on the $\alpha$-carbon were not detected in any of the alkyl compounds, which is typical for organic groups in close proximity to paramagnetic centers.

The $^{11}\text{B}$ NMR spectra of these complexes each contained one peak, the chemical shift of which ranged from 86.6 to 116.7 ppm. These signals were far downfield compared to the resonances observed for the chloride (~29 ppm) as well as diamagnetic species resulting from transmetalation of $\text{To}^M$ (ca. –17 ppm). Overall, the $^1\text{H}$ and $^{11}\text{B}$ NMR spectra associated with the $\text{To}^M$ ligand in the series of organometallic species are comparable, whereas the chemical shifts for the oxazolinylborate ligand in $\text{To}^M\text{CoX}$ (e.g. $X = \text{Cl, O}^{\prime}\text{Bu, OAc}$) complexes vary considerably. These data suggest that the organometallic compounds’ electronic structures, which are responsible for the paramagnetic chemical shifts, are similar between simple alkyl, $\beta$-H containing alkyl, trimethylsilyl-substituted alkyl, aryl, and benzyl ligands.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$^1\text{H}$ NMR (ppm)</th>
<th>$^{11}\text{B}$ NMR (ppm)</th>
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</thead>
<tbody>
<tr>
<td>$\text{To}^M\text{CoBn}$ (1)</td>
<td>$^{1}\text{H}$</td>
<td>$^{11}\text{B}$</td>
</tr>
<tr>
<td>$\text{To}^M\text{(CH}_2\text{)}$</td>
<td>16.33</td>
<td>34.47, –89.01, –77.45</td>
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<td>$\text{To}^M\text{(CH}_3\text{)}$</td>
<td>–12.46</td>
<td>–100.4</td>
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<td>$\text{To}^M\text{(C}_6\text{H}_5\text{)}$</td>
<td>14.90, 10.96, 9.14</td>
<td>12.72, 10.08, 8.54</td>
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<tr>
<td>$\text{To}^M\text{CH}_2\text{SiMe}_3$ (2)</td>
<td>15.73</td>
<td>8.54</td>
</tr>
<tr>
<td>$\text{To}^M\text{CoPh}$ (3)</td>
<td>16.73</td>
<td>8.54</td>
</tr>
<tr>
<td>$\text{To}^M\text{CoEt}$ (4)</td>
<td>14.85</td>
<td>12.72, 10.08, 8.54</td>
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<tr>
<td>$\text{To}^M\text{Co}^{\prime}\text{Bu}$ (5)</td>
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<td>8.54</td>
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<td>$\text{To}^M\text{CoMe}$ (6)</td>
<td>15.44</td>
<td>8.54</td>
</tr>
<tr>
<td>$\text{To}^M\text{CoCl}$</td>
<td>24.88</td>
<td>8.54</td>
</tr>
</tbody>
</table>

N.D. indicates not detected. $^a$See reference $^{15}$, $^b$See reference $^{16}$.
A single $\nu_{\text{CN}}$ band at $\sim 1590 \text{ cm}^{-1}$ in the IR spectra of 1 – 6 provided additional support for tridentate coordination of the $\text{To}^\text{M}$ ligand to cobalt. Signals associated with C=N stretching modes of non-coordinated oxazolinyl groups ($\sim 1630 \text{ cm}^{-1}$) were not detected. In the series, $\text{To}^\text{M}\text{CoPh}$ displayed the lowest energy $\nu_{\text{CN}}$ band at 1582 cm$^{-1}$, $\text{To}^\text{M}\text{CoEt}$ the highest at 1592 cm$^{-1}$, while $\text{To}^\text{M}\text{CoBn}$ was equivalent to that observed for $\text{To}^\text{M}\text{CoCl}$ (1588 cm$^{-1}$). The IR spectrum of 1 provided additional evidence of benzylolation, revealing a new aromatic $\nu_{\text{C-H}}$ mode at 3012 cm$^{-1}$ in addition to the aromatic $\text{To}^\text{M}$ bands at ca. 3070 and 3050 cm$^{-1}$. The aromatic $\nu_{\text{C-H}}$ bands of $\text{To}^\text{M}\text{CoPh}$ were more intense than in the spectra of 3, 4, or 5, but lacked additional bands. We attributed these observations to overlapping modes for CoPh and BPh.

The UV-vis spectra (Figure 1, Table 2) of compounds 1 – 6 contained intense absorptions at ca. 350 ($\varepsilon$: 1400-2200 M$^{-1}$cm$^{-1}$) and 700 nm ($\varepsilon$: 1200-1500 M$^{-1}$cm$^{-1}$) and two weaker bands at ca. 570 ($\varepsilon$: 300-400 M$^{-1}$cm$^{-1}$) and 620 nm ($\varepsilon$: 200-450 M$^{-1}$cm$^{-1}$). The former features are characteristic of these organometallic complexes and are attributed to transitions associated with the alkyl or aryl ligand on the basis of the lack of equivalent bands in $\text{To}^\text{M}\text{CoCl}$. The latter bands in the region of 500-650 nm were attributed to d$\leftarrow$d transitions and were similar to the bands observed for $\text{To}^\text{M}\text{CoCl}$ at 568 ($\varepsilon = 362 \text{ M}^{-1} \text{ cm}^{-1}$) and 635 nm ($\varepsilon = 641 \text{ M}^{-1} \text{ cm}^{-1}$). The observed d$\leftarrow$d transitions are related to the $^{4}\text{T}_1(P)\leftarrow^{4}\text{A}_2(F)$ transition in $[\text{CoCl}_4]^{2-}$ that split in lower symmetry. Across the organometallic compounds, the wavelengths of these d$\leftarrow$d bands do not vary very much, further supporting the idea, from the NMR chemical shift analysis discussed above, that their electronic structures are similar. It is worth noting that the benzyl compound 1, which also showed unique reactivity (see below), contained an additional
strong absorption at 439 nm ($\varepsilon$: 1964 M$^{-1}$cm$^{-1}$; see Figure 1) as well as the absorptions at 339 ($\varepsilon$: 3205 M$^{-1}$cm$^{-1}$), 589 nm ($\varepsilon$: 332 M$^{-1}$cm$^{-1}$), 628 ($\varepsilon$: 365 M$^{-1}$cm$^{-1}$), and 704 nm ($\varepsilon$: 1444 M$^{-1}$cm$^{-1}$).

The spectroscopic features assigned to transitions associated with the organometallic moiety on the basis of their absence in halide, alkoxide, and carboxylate analogues, are similar for compounds 1 – 6, irrespective of methyl, ethyl, butyl, phenyl, trimethylsilylmethyl, or benzyl groups that might be expected to have variable bonding and orbital energies. A possible interpretation of these observations is that the transfers are delocalized to include wavefunctions from the To$^M$ ligand (which is common to the series of compounds), and the allowedness of the transfer is enhanced by the organometallic ligand. This idea, at least in part, is supported by comparison with the spectra of reported for Tp$^{ipr2}$CoR complexes (Tp$^{ipr2}$ = tris(3,5-diisopropylpyrazolyl)borate). For example, the spectrum of Tp$^{ipr2}$CoEt, which had maxima at $\lambda = 388$ (1030 M$^{-1}$cm$^{-1}$), 580, 610, and 690 nm (810 M$^{-1}$cm$^{-1}$), contained similar features as To$^M$CoEt.$^{10}$ The absorption at 705 nm for To$^M$CoEt was 15 nm lower energy than the corresponding band in Tp$^{ipr2}$CoEt at 690 nm, whereas the d$\leftrightarrow$d bands appeared at similar energies. From these observations, we suggest that the wavefunctions contributed by the To$^M$ and Tp$^{ipr2}$ affect the energy of the transition. Tp$^{ipr2}$Co-$p$-tol also contains an intense absorption at 697 nm (1304 M$^{-1}$cm$^{-1}$).$^{17}$ However, the intense absorption at ca. 700 nm is not a universal features of cobalt(II) alky complexes. For example, [PhTt$^{t-Bu}$]CoCl contains absorptions at 664 (880 M$^{-1}$cm$^{-1}$) and 695 nm (740 M$^{-1}$cm$^{-1}$), which are more intense than any of the absorptions present in the corresponding methyl phenyl, or benzyl complex (e.g. [PhTt$^{t-Bu}$]CoMe 632 (570 M$^{-1}$cm$^{-1}$) and 725 nm
(574 M⁻¹cm⁻¹); [PhTr't-Bu]CoPh (643 (249 M⁻¹cm⁻¹), 719 (359 M⁻¹cm⁻¹), and 743 nm (578 M⁻¹cm⁻¹); [PhTr't-Bu]CoBn (511 (682 M⁻¹cm⁻¹), 655 (346 M⁻¹cm⁻¹), 711 (258 M⁻¹cm⁻¹), and 761 nm (430 M⁻¹cm⁻¹)). In addition, even when a strong absorption is observed at ca. 700 nm, the molar absorptivity value is typically less than 1,000 making it much weaker than the absorption at ca. 700 nm observed for this series of ToMCoR complexes (Tp't-Bu,MeCoMe 580 (235 M⁻¹cm⁻¹), 614 (293 M⁻¹cm⁻¹), and 685 nm (499 M⁻¹cm⁻¹); Tp't-BuCoMe 577 (277 M⁻¹cm⁻¹), 611 (374 M⁻¹cm⁻¹), and 688 nm (839 M⁻¹cm⁻¹); Tp't-Bu,MeCoEt 581 (220 M⁻¹cm⁻¹), 617 (275 M⁻¹cm⁻¹), and 688 nm (510 M⁻¹cm⁻¹)).

Figure 1. UV-vis spectra of ToMCoBn (1), ToMCoCH₂SiMe₃ (2), ToMCoPh (3), ToMCoEt (4), ToMCo't-Bu (5), and ToMCoMe (6) measured in diethyl ether. The d→d transitions are ~1 order of magnitude weaker intensity than those observed at ca. 300 and 700 nm.
Table 2. UV-vis spectroscopic data for compounds 1 – 6.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Intense absorptions (ε &gt; 1000), nm (ε, M⁻¹cm⁻¹)</th>
<th>d→d bands, nm (ε, M⁻¹cm⁻¹)</th>
<th>µ_{eff} (µB)</th>
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<td>339 (3205) 439 (1964) 704 (1444) 589 (332) 628 (365)</td>
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<td>Tonomy^3CoCH₂SiMe₃ (2)</td>
<td>355 (2228) 703 (1495) 591 (374) 628 (448)</td>
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<tr>
<td>Tonomy^3CoPh (3)</td>
<td>330 (1245) 708 (1333) 587 (176) 625 (252)</td>
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<tr>
<td>Tonomy^3CoEt (4)</td>
<td>382 (1935) 705 (1255) 579 (268) 615 (286)</td>
<td>4.1(6)</td>
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<tr>
<td>Tonomy^3CoBu (5)</td>
<td>382 (1679) 705 (1291) 576 (406) 613 (422)</td>
<td>4.5(2)</td>
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<tr>
<td>Tonomy^3CoMe (6)</td>
<td>346 (1412) 697 (1078) 581 (388) 617 (424)</td>
<td>4.2(2)</td>
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<td>Tonomy^3CoCl</td>
<td>n.a.</td>
<td>568 (362) 635 (641)</td>
<td>4.5(2)</td>
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</table>

Solution-phase magnetic moments of compounds 1 – 6 (Table 2) at room temperature, measured using Evans method, were consistent with high-spin cobalt(II) (S = 3/2, spin-only µ_{eff} = 3.87 µ_B). With the exception of Tonomy^3CoPh, the effective magnetic moments are within the range reported for other tetrahedral organocobalt(II) complexes. In high spin, tetrahedral cobalt(II) with an e⁹t₂³ configuration, orbital contributions to the magnetic moment are expected to be quenched in the ground state, but low-lying excited state mixing results in µ_{eff} ranging from 4 – 5 µ_B. EPR spectra for 1 – 6 all contain striking eight-line patterns resulting from hyperfine coupling (54 G) to the ^{59}Co center (I = 7/2). Other four-coordinate {κ^3-Tonomy^3}Co^II species, namely Tonomy^3CoCl and Tonomy^3CoO'Bu, produced rhombic signals that were devoid of hyperfine coupling.

Ultimately, the spectroscopic assigned structures of compounds 1 – 6 were validated by X-ray diffraction studies of single crystals obtained from saturated pentane solutions cooled at –40 °C (Figures 2-6, Table 3). The molecular structures are similar to previously reported tris(oxazolinyl)borate magnesium (d⁰) and zinc (d^{10}) organometallic
compounds. In each cobalt complex, the tris(oxazolinyl)borate ligand is coordinated in a tridentate motif, and the $\text{To}^M$ and alkyl or aryl ligands provide a distorted tetrahedral geometry (the $\tau_4$ scale for accessing distortions of four-coordinate compounds, is defined as $\tau_4 = 1$ for a $T_d$ geometry, 0.85 for a trigonal pyramid ($C_{3v}$), and 0 for a square planar geometry) for the cobalt centers ($\tau_4 = 0.75 - 0.82$), which is similar to $\text{To}^M\text{MgMe}$ ($\tau_4 = 0.75$) and $\text{To}^M\text{ZnMe}$ ($\tau_4 = 0.76$). In this series of cobalt compounds, the $\angle \text{N-Co-N}$ angles range over $91 - 93^\circ$, whereas $\angle \text{N-Co-C}$ angles vary from 116 to 131°. For comparison, the ranges of angles for $\text{To}^M\text{MgMe}$ and $\text{To}^M\text{ZnMe}$ are 122 – 130° and 120 – 131°, respectively. The $\tau_4$ scale, the $\angle \text{N-Co-C}$ angles, and the B–M–Me angles describe the distortion of the alkyl ligand away from the $C_3$ axis in the $\{\kappa^3-\text{To}^M\}M$ motif. The $\angle \text{B-M-Me}$ in $\text{To}^M\text{MgMe}$ (172.89°), $\text{To}^M\text{ZnMe}$ (174.74) and $\text{To}^M\text{CoMe}$ (172.83°) are similar. That is, the three unpaired d electrons in the high spin cobalt(II) compounds appear to have little consequence on the coordination geometry. The Co–C interatomic distances in compounds 1 – 6 are similar and vary only from 1.994(2) ($\text{To}^M\text{CoMe}$) to 2.023(2) Å for $\text{To}^M\text{CoBn}$ (see Table 3). The Co–N interatomic distances are also similar across the series, varying from 2.019(2) to 2.062(3) Å. Thus, the similar electronic features identified by UV-vis, EPR, and NMR spectroscopies are also reflected in similar structural parameters. The M–Me distances follow the expected periodic trend based on ionic radii (Mg (2.108(1) Å) > Co (1.994(2) Å) > Zn (1.972(1) Å)). A comparison of $\text{To}^M\text{CoBn}$ with $\text{To}^M\text{MgBn}$ reveals that the Co–C interatomic distance in $\text{To}^M\text{CoBn}$ (2.023(2) Å) is significantly shorter (by 0.12 Å) than in $\text{To}^M\text{MgBn}$ (2.143(2) Å). The Co-C distance of $\text{To}^M\text{CoCH}_2\text{SiMe}_3$ (1.999(4) Å) was similar to that reported for $\text{Tp}^{\text{Ph,Me}}\text{CoCH}_2\text{SiMe}_3$ (2.017(2) Å).
Table 3. Selected interatomic distances (Å) and angles (°) for compounds 1-6.

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Figure 2. Thermal ellipsoid diagram of To$^\text{N}$CoBn (1) with ellipsoids plotted at 50% probability. H atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1-C22 2.023(2), Co1-N1 2.040(1), Co1-N2 2.055(1), Co1-N3 2.041(1); N1-Co1-C22 122.94(6), N2-Co1-C22 131.33(6), N3-Co1-C22 116.37(6), N1-Co1-N2 91.68(5), N1-Co1-N3 94.00(4), N2-Co1-N3 91.00(5).
Figure 3. Thermal ellipsoid plot of $\text{To}^\text{M}\text{CoCH}_2\text{SiMe}_3$ (2) with ellipsoids at 50% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1-C41 1.999(4), Co1-N1 2.044(3), Co1-N2 2.062(3), Co1-N3 2.059(3); N1-Co1-C22 126.3(2), N2-Co1-C22 118.0(1), N3-Co1-C22 125.7(2), N1-Co1-N2 91.5(1), N1-Co1-N3 94.3(1), N2-Co1-N3 91.7(1).

Figure 4. Thermal ellipsoid plot of $\text{To}^\text{M}\text{CoPh}$ (3) with ellipsoids at 50% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1-C16 1.996(2), Co1-N1 2.041(2), Co1-N2 2.019(2), Co1-N3 2.048(2); N1-Co1-C16 121.41(9), N2-Co1-C16 121.50(9), N3-Co1-C16 128.00(1), N1-Co1-N2 92.29(9), N1-Co1-N3 91.37(9), N2-Co1-N3 92.95(9).
Figure 5. Thermal ellipsoid plot of $\text{To}^\text{M}\text{CoEt} \ (4)$ with ellipsoids at 50% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): $\text{Co1-C22} \ 1.980(3)$, $\text{Co1-N1} \ 2.046(2)$, $\text{Co1-N2} \ 2.045(2)$, $\text{Co1-N3} \ 2.046(2)$; $\text{N1-Co1-C22} \ 119.7(1)$, $\text{N2-Co1-C22} \ 126.9(1)$, $\text{N3-Co1-C22} \ 125.2(1)$, $\text{N1-Co1-N2} \ 91.07(9)$, $\text{N1-Co1-N3} \ 92.51(9)$, $\text{N2-Co1-N3} \ 91.8(1)$.

Figure 6. Thermal ellipsoid plot of $\text{To}^\text{M}\text{Co}^\text{Bu} \ (5)$ with ellipsoids at 50% probability. H atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): $\text{Co1-C22} \ 2.010(3)$, $\text{Co1-N1} \ 2.046(3)$, $\text{Co1-N2} \ 2.047(3)$, $\text{Co1-N3} \ 2.054(3)$; $\text{N1-Co1-C22} \ 121.34(2)$, $\text{N2-Co1-C22} \ 124.37(2)$, $\text{N3-Co1-C22} \ 126.29(2)$, $\text{N1-Co1-N2} \ 91.23(1)$, $\text{N1-Co1-N3} \ 91.52(1)$, $\text{N2-Co1-N3} \ 92.39(1)$. 
The absorption spectra for the series of organocobalt $\text{To}^M\text{CoR}$ ($1-6$) and heteroatom-bonded species $\text{To}^M\text{CoX}$ ($X = \text{Cl, OtBu, OAc}$) were further studied with representative electronic structure calculations. Gas-phase models for $\text{To}^M\text{CoBn}$ ($1\text{-calc}$), $\text{To}^M\text{CoMe}$ ($6\text{-calc}$) and $\text{To}^M\text{CoCl}$ ($\text{To}^M\text{CoCl\text{-calc}}$) were optimized using the coordinates from X-ray diffraction as initial geometries, to confirm tetrahedral geometries. The vibrational calculations for $6\text{-calc}$ and $\text{To}^M\text{CoCl\text{-calc}}$ contained $\nu_{CN}$ at 1667 and 1677 cm$^{-1}$, respectively. The high spin state for $1\text{-calc}$ and $6\text{-calc}$ are calculated to be 46 and 48 kcal/mol lower in energy than the low spin, respectively.

TDDFT calculations on $\text{To}^M\text{CoCl}$ and $\text{To}^M\text{CoMe}$ revealed transitions at 602, 740 and 743 nm for both species. However, the characteristic and intense low-energy experimental band at 697 nm is not calculated by this TDDFT model. We previously postulated that the single configuration TDDFT approach is insufficient to correctly model the electronic features of these compounds. In contrast, configuration interaction (CI) singles calculations show that the ground state wave functions of $\text{To}^M\text{CoBn}$, $\text{To}^M\text{CoMe}$ and $\text{To}^M\text{CoCl}$ are inequivalent. The ground state electronic structure of $\text{To}^M\text{CoCl}$ is best described by a single-reference wavefunction. In contrast, the $\text{To}^M\text{CoBn}$ and $\text{To}^M\text{CoMe}$ ground state wave functions contain multireference character (i.e., the ground state has more than one contributing electronic configuration). The additional contributing electronic configuration contains an excitation from a doubly occupied anti-bonding orbital to a singly occupied anti-bonding orbital both centered around the phenyl ring on the $\text{To}^M$ ligand. Multireference character in the ground state wave function of the $\text{To}^M\text{CoMe}$ helps to explain why the excitation at 697 nm from the experimental UV-Vis was not found in earlier TDDFT calculations. The CI singles calculation of $\text{To}^M\text{CoMe}$
contained an excited state (3\textsuperscript{rd} excited state) with an energy difference from the ground state corresponding to an excitation at 661 nm. The excitation is from a doubly occupied bonding orbital to a singly occupied anti-bonding orbital centered on the cobalt. The orbitals involved in this excitation are mostly, but not purely d orbitals and included contributions from the oxazolines and the methyl group; however, the d orbitals had the largest changes in electron density. The conclusions from CI calculations on To\textsuperscript{M}CoBn are consistent with those from the To\textsuperscript{M}CoMe study.

The complexes 1 – 6 persist in solution at elevated temperatures (in the absence of air and moisture). For example, ethyl and butyl compounds 4 and 5 do not eliminate detectable quantities of ethylene or butene, respectively, after thermal treatment at 120 °C, despite the possibility for β-hydrogen elimination. Similar resistance toward β-hydrogen elimination in a bulkier tris(pyrazolyl)borato cobalt ethyl Tp\textsuperscript{iPr}CoEt was attributed to its high spin electronic configuration.\textsuperscript{20} Products of Co–C bond homolysis, such as ethane or butane, are also not formed.

Unexpectedly, reactions of 2 – 5 and CO or CO\textsubscript{2} to form To\textsuperscript{M}Co\{C(R)O\}CO or To\textsuperscript{M}CoO\textsubscript{2}CR, respectively (see below), gave product mixtures that contained small signals associated with To\textsuperscript{M}CoCl (<10\%). Although the \textsuperscript{1}H and \textsuperscript{11}B NMR signals for 1 – 6 were paramagnetically shifted (see below), the peaks were characteristic and distinct from To\textsuperscript{M}CoCl. Moreover, samples of To\textsuperscript{M}CoR, apparently pure and To\textsuperscript{M}CoCl-free as determined by \textsuperscript{1}H NMR, \textsuperscript{11}B NMR and UV-vis spectroscopy, revealed To\textsuperscript{M}CoCl after addition of CO. Strangely, reactions of either 1 or 6 and CO or CO\textsubscript{2} provide acyl or carboxylate products that are free of To\textsuperscript{M}CoCl, implying that the source of To\textsuperscript{M}CoCl is not present in either the cobalt benzyl or methyl samples. The syntheses of 1 – 6 were
also attempted by reaction of $^{M}_{\text{Co}}\text{CoCl}$ with Grignard reagents, but transmetalation occurred to form $^{M}_{\text{Co}}\text{MgBr}$ as the dominant product rather than the desired alkylation of cobalt. Ultimately, recrystallization of organocobalt(II) species 2 – 5 afforded isolable organometallic species that did not generate detectable $^{M}_{\text{Co}}\text{CoCl}$ in subsequent reactions.

**Carbonylation of $^{M}_{\text{Co}}\text{CoR}$.**

Complexes 1 – 6 rapidly react with CO (1 atm) in benzene-$d_6$ at room temperature (Scheme 2), as evidenced by an immediate color change from blue/green to orange to give compounds labeled as 7 – 12 (7, Bn; 8, CH$_2$SiMe$_3$; 9, Ph; 10, Et; 11, nBu; 12, Me). Reactions of organometallics with CO are among the most studied, and the interaction of CO with four-coordinate iron, cobalt, and nickel alkyls can be complicated by the many pathways that lead to a wide range of products including coordination to Co(II) to give carbonyl adducts of cobalt alkyls, insertion to form the acyl isomer, and coordination of one or two carbon monoxide ligands to the cobalt acyl. In addition, CO could substitute one or more of the oxazoline groups to give bidentate coordinated $^{M}_{\text{Co}}$. Finally, 1 e$^-\text{ reduction to form Co(I) monocarbonyl or dicarbonyl adducts is presumed to involve M–C bond homolysis and occurs with bulky tris(pyrazolyl)borate iron or cobalt alkyls upon heating under CO.}\)\(^7,28\) Alkyl group-derived byproducts, presumably resulting from further reactions of alkyl radical species, are not described in those reactions.

The results may be summarized as follows: the reactions of $^{M}_{\text{Co}}\text{CoR}$ and CO give mixtures containing organocobalt carbonyl $^{M}_{\text{CoR}}$(CO) (7a – 12a), acyl $^{M}_{\text{Co}}$(C=O)R (7b – 12b), and acyl carbonyl $^{M}_{\text{Co}}$(C(O)R)CO (7c – 12c), and these species interconvert by insertion and CO coordination, or CO dissociation and decarbonylation (Scheme 2). $^1\text{H}$ and $^{11}\text{B}$ NMR spectroscopy suggest only a single species
is formed, whereas multiple $v_{\text{CO}}$ and $v_{(\text{C}=\text{O})R}$ in the IR spectra indicate that several species are present in each reaction mixture. Together, NMR and IR spectroscopy suggest that organocobalt carbonyl $a$, acyl $b$, and acyl carbonyl $c$ species are interconverted through a process that is faster than the NMR timescale and slower than the IR timescale. An additional interesting feature is that the apparent distribution of products depends on the $R$ group (Scheme 2, eqn (1) – (4)).

Scheme 2. Carbonylation of 1 – 6 to form interconverting mixtures of organocobalt(II) carbonyl ($a$), acyl ($b$), and acyl carbonyl ($c$) complexes. Species observed by IR spectroscopy are enclosed in boxes, and equilibrium arrows indicate favored and disfavored compounds. $[\text{Co}] = \kappa^3$-To$^M$Co.

Benzene-$d_6$ solutions of 7 – 12 were first assayed by $^{11}$B NMR spectroscopy in the presence of excess CO (1 atm). The $^{11}$B NMR resonance for 8 – 12 appeared at ca. –4 ppm, which was distinct from the downfield chemical shift range for the organocobalt(II)
1 – 6 from 87 to 116 ppm. Unexpectedly, the $^{11}$B NMR chemical shift for the carbonylated benzyl derivative 7 at 88 ppm was located in the region associated with four-coordinate cobalt alkyl/aryl or five-coordinate cobalt carboxylate compounds (e.g., $\text{To}^\text{M}\text{CoO}_2\text{CCH}_2\text{Ph}$: 84 ppm). Still, the $^{11}$B NMR spectrum provided evidence of carbonylation, giving a distinct signal from its precursor 1 (100 ppm). The benzyl compound 7 is also distinguished from the other tris(oxazolinyl)borate organocobalt compounds by its $^{1}$H NMR spectrum, its IR spectrum and its reactivity (see below).

Compounds 8 – 12 gave similar $^{1}$H NMR spectra that contained five, broad signals from 10 to −7 ppm, which were distinct from the spectra for the organocobalt(II) starting materials. Four of the $^{1}$H NMR signals were attributed to the $\text{To}^\text{M}$ ligand on the basis of their consistent chemical shifts across complexes 8 – 12 (9.6, 8.1, 7.7, and −1.2 ppm); however, additional assignments based on integration were impeded by their broad appearance. The fifth was assigned to the R group on the basis of its varying chemical shift among the complexes (e.g., −0.09 ppm for the $\text{CH}_2\text{SiMe}_3$ group in 8, and −7.7 ppm for the ethyl group in 10). The $^{1}$H NMR spectrum of cobalt benzyl-derived 7 contained many (>15) signals from 82 to −28 ppm that were not readily assigned to specific moieties in the complex.

The infrared spectra of 7 – 12 contained signals that were assigned to terminal carbonyl ($\nu_{\text{C=O}}$, 1886 and 1973 to 1986 cm$^{-1}$), acyl ($\nu_{\text{C(=O)R}}$, 1650 to 1718 cm$^{-1}$), and the oxazoline regions ($\nu_{\text{CN}}$: 1582 to 1594 cm$^{-1}$; see Table 4). Due to reactivity of the compounds, these infrared spectra were recorded in THF solutions saturated with CO in an air-free ATR configuration. The spectra of all of the compounds 7 – 12 contained a $\nu_{\text{C=O}}$ band at ca. 1886 cm$^{-1}$ (see Figure S36 – S40). This signal was the only $\nu_{\text{C=O}}$ in
benzyl 7, the dominant ν_{C≡O} in phenyl-derived 9, a significant signal in 8 and 11 from trimethylsilyl- and n-butylcobalt species, and the minor signal in ethyl 10 and methyl 12. A higher energy ν_{C≡O} band at approximately 1980 cm⁻¹ was observed for complexes 8 – 12 and appeared at the expense of the 1886 cm⁻¹ signal. These peaks were assigned to organocobalt carbonyl To^MCo(R)CO a and acylcobalt To^MCo{C(O)R}CO c, on the basis of the expected increased π-backdonation from organocobalt compared to acylcobalt species. This idea was supported by DFT calculations of To^MCo(Me)CO (12a) and To^MCo{C(O)Me}CO (12c), which reproduced the trend in ν_{C≡O} with 12a (2030 cm⁻¹) < 12c (2117 cm⁻¹). The acyl (ν_{C(=O)R}) signals were less intense than ν_{C≡O} and appeared as one (R = CH₂SiMe₃ or Ph) or two signals (R = Et, nBu, or Me) in the region from 1650 to 1690 cm⁻¹, but were absent for R = Bn (7). A ν_{C(=O)R} at 1717 cm⁻¹ was also observed for R = Bn, CH₂SiMe₃, Ph, or nBu and was assigned to an intermediate acyl species (see Table 4).

Table 4. Infrared spectroscopic data of 7 – 12 collected in CO-saturated THF using a ZnSe crystal in ATR mode. Not detected (N.D.) in the IR spectrum.
In the $\nu_{CN}$ region, complexes 7 – 12 contain a single $\nu_{CN}$ band in the region from 1590 to 1600 cm$^{-1}$ that is associated with coordinated oxazolines. In compound 12, a shoulder at 1630 cm$^{-1}$ was assigned to a non-coordinated oxazoline. This experimental result is consistent with the results of DFT calculations on 12. The optimized geometry of 12 (PBE0) contained a long apical Co–N interatomic distance in a square pyramidal structure (see X-ray discussion below). Computed $\nu_{CN}$ for 12c were found at 1611 cm$^{-1}$, 1641 cm$^{-1}$, and 1679 cm$^{-1}$, with the highest energy stretch corresponding to the weakly-coordinated oxazoline. In contrast the IR spectra of compounds 7 – 11 did not contain peaks that might be associated with non-coordinated oxazoline groups.

The orange carbonylated compounds 7 – 12 have distinct UV-vis spectra from their blue or green starting materials. The dominating bands at 350 nm and 700 nm in the starting materials are absent in the carbonylated product. Instead 7 – 12 are characterized by weak signals from 700 to 500 nm ($\varepsilon$: 100–170 M$^{-1}$ cm$^{-1}$) and a strong, tailing absorption from 200 to 500 nm. In addition, compounds 7 – 12 are low spin ($\mu_{\text{eff}} = 2.4 – 3.4(2)$ $\mu_B$) as determined by the Evans method. The optimized geometries of both $\text{To}^M\text{CoMe(CO)}$ (12a) and $\text{To}^M\text{Co}\{\text{CO}\}\text{Me}\{\text{CO}\}$ (12c) were found to be square-based pyramidal, with the low spin states being 30 and 33 kcal/mol lower than the high spin for 12a and 12c respectively. The magnetic moments are higher than the spin-only $\mu_{\text{eff}}$ value (1.73 $\mu_B$) for low spin Co(II) ($S = \frac{1}{2}$). A square pyramidal geometry with long axial Co–N interactions could be used to explain the higher than expected magnetic moments as square planar Co(II) complexes typically have magnetic moments in this range (2.1 – 2.9 $\mu_B$). Alternatively, contributions from the acyl species 7b – 12b (which are expected to be high spin) could also result in higher-than-expected magnetic moments. This idea
suggests that methyl (12) and ethyl (10) derivatives would have the lowest $\mu_{\text{eff}}$ values, because their IR spectra are dominated by the CO-coordinated species. Instead, experimental $\mu_{\text{eff}}$ values for 10 and 12 are in the middle of the series (2.7 $\mu_B$).

A pentane solution of 10, cooled to $-30^\circ$C provides X-ray quality crystals, and a single crystal diffraction study reveals the five-coordinate, acyl carbonyl form $\text{To}^\text{M} \text{Co}\{\text{C(O)Et}}\text{CO} (10\text{c})$ as a square pyramidal complex (Figure 7). Although the oxazolines are disposed trans to either CO, an acyl, or an open coordinate site, the Co1–N1 (2.037(3) Å), Co1–N2 (2.067(2) Å) and Co1–N3 (2.050(2) Å) distances are similar to each other and to the four-coordinate tris(oxazolinyl)borato organocobalt(II) compounds described above. Related [PhTt$^{\text{Bu}}$]Co(CO)(C(O)R) (R = Me, Et, or Ph) are also square pyramidal, with similar Co–S distances for basal and apical groups.\(^\text{19}\)

**Figure 7.** Thermal ellipsoid plot of $\text{To}^\text{M} \text{Co}\{\text{C(O)Et}}\text{CO} (10)$ plotted at 35% probability. H atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1–C22 1.973(4), Co1–C25 1.751(4), Co1–N1 2.037(3), Co1–N2 2.066(3), Co1–N3 2.050(2), C22–O4 1.207(5), C25–O5 1.156(5), N1–Co1–C22 89.7(1), N2–Co1–C22 174.0(1), N3–Co1–C22 97.1(1), N1–Co1–N2 87.0(1), N1–Co1–N3 92.6(1), N2–Co1–N3 88.1(1), C22–Co1–C25 85.5(2).
Comparison of the structure of ethyl 10, determined by X-ray diffraction, with minimum energy structures for $^{10}\text{MCo}\{\text{C(=O)Et}\}\text{CO}$ and $^{10}\text{MCo}\{\text{C(=O)Me}\}\text{CO}$ calculated using density functional theory (DFT) reveals that although all three species are square pyramidal, the geometry of the calculated energy minimized structure of $^{10}\text{MCo}\{\text{C(O)Me}\}\text{CO}$ contains a long apical Co–N bond (2.19 Å). The calculated Co–N distances of 13 in the basal plane (2.04 and 2.06 Å) are significantly shorter than the apical Co–N interatomic distance. In comparison, the apical Co–N bond in the DFT optimized geometry of 10 is only 2.12 Å, while calculated Co-N distances in the basal plane are 2.03 and 2.07 Å.

We have not been able to obtain X-ray quality crystals of any of the other species in the mixture of 10, nor have acyl or carbonyl containing compounds of 7 – 9, 11, or 12 been isolated. Attempts to crystallize 12, for example, provide either $^{10}\text{MCoMe}$ (6) or $^{10}\text{MCoOAc}$ (18, see below). It is likely that X-ray quality crystals of 10c are obtained because that is the most crystalline species in the reaction mixture. Additionally, the reversibility of the carbonylation of the organocobalt(II) alkyls posed challenges to the compounds’ isolation. Thus, evaporation and exhaustive drying of $^{10}\text{MCo}\{\text{C(O)Me}\}\text{CO}$ provides 6 after 24 h under dynamic vacuum, whereas complexes 9 – 11 show only partial conversion to 3 - 5 under these condition as assessed by $^{11}\text{B}$ NMR spectroscopy. Complex 8 is fully consumed upon evaporation, with $^{10}\text{MCoCH}_{2}\text{SiMe}_{3}$ as one of several other products that remain unidentified, as determined by $^{1}\text{H}$ and $^{11}\text{B}$ NMR spectroscopy.

An entirely unique outcome occurs when 7 is exposed to vacuum (Scheme 3). Unlike the evaporation of 8 – 12, which produces the corresponding organocobalt compounds 2 – 6, evaporation of solutions containing 7 does not produce $^{10}\text{MCoBn}$.
Instead, a mixture was obtained that gives two signals in the $^{11}$B NMR spectrum at 86 and 87 ppm. The former is assigned to 7, while the second signal and a corresponding dominant peak in the $^1$H NMR spectrum at −18 ppm were attributed to an alkoxy ketone cobalt(II) species (7d). The evidence for that structure is provided by X-ray diffraction studies of crystals grown from degassed solutions of $^M$CoBn and CO (Figure 8). In that compound, one $^M$Co, two CO, and three benzyl groups are combined to form $^M$Co{$O, O$-κ$^2$-O–C(Bn)$_2$–C(=O)Bn}.

**Scheme 3.** Carbonylation of $^M$CoBn (1) followed by removal of volatile materials provides 7d as confirmed by X-ray crystallography and a species tentatively assigned as $^M$CoCO (7e) as determined by IR spectroscopy.

The structure of 7d displays a distorted trigonal bipyramidal geometry ($\tau_5 = 0.5$) composed of N1-Co1-N2 88.8(1), N2–Co1–O4 129.4(1), and N1–Co1–O4 141.4(1) angles that sum to 359.6° and a nearly linear N3–Co1–O5 angle of 172.8(1)°. The interatomic distance between cobalt and the coordinated ketone oxygen (Co1–O5, 2.212(2) Å) is longer than the interatomic distance between cobalt and the alkoxy
oxygen (Co1–O4, 1.972(2) Å) while the ketone carbon-oxygen distance is shorter (O5–C37, 1.231(4) Å) than in the alkoxide (O4–C22, 1.386(4) Å). The Co–N interatomic distances (Co1-N1 2.098(3), Co1-N2 2.086(3), Co1-N3 2.136(3) Å; Co–N_{average} = 2.11 Å) are all longer than other structurally characterized Co(II) complexes supported by To^M (e.g. To^MCoBn Co–N_{average} = 2.055 Å).

**Figure 8.** Thermal ellipsoid plot of 7d with ellipsoids at 35% probability. H atoms and a pentane molecule in the unit cell are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1-O4 1.927(2), Co1-O5 2.212(2), Co1-N1 2.098(3), Co1-N2 2.086(3), Co1-N3 2.136(3); N1-Co1-N2 88.8(1), N2–Co1–O4 129.4(1), N1–Co1–O4 141.4(1), N3–Co1–O5 172.8(1), O4-Co1-O5 77.52(9).
The latter alkoxy ketone species 7d is attributed to the $^{11}$B NMR signal at 87 ppm that appears upon evaporation of 7 under vacuum overnight. IR spectroscopy (KBr) of the crystals contained a $\nu_{C=O}$ band (1943 cm$^{-1}$) and $\nu_{C(=O)R}$ band (1671 cm$^{-1}$). The $\nu_{C(=O)R}$ band is assigned to the alkoxy ketone species 7d while the $\nu_{C=O}$ band is tentatively assigned to a reduced To$^M$CoCO (7e) species based upon its similar frequency to signals in Tp’CoCO complexes (e.g., Tp$^{iPr^2}$CoCO, $\nu_{C=O} \sim 1950$ cm$^{-1}$; Tp$^{iPr,Me}$CoCO, $\nu_{C=O} = 1946$ cm$^{-1}$ and; Tp$^{Np}$CoCO, $\nu_{C=O} = 1950$ cm$^{-1}$, Tp$^{Np} = \text{tris}(3\text{-neopentylpyrazolyl})$borate; Tp$^{iPr,Me} = \text{tris}(3\text{-isopropyl-5-methylpyrazolyl})$borate).$^{10,28,30}$ In these Co(I) compounds, the $\nu_{C=O}$ appears at lower energy than in the cobalt(II) acyl carbonyls (e.g., Tp$^{iPr^2}$Co{C(O)Et}CO, $\nu_{C=O} = 1999$ cm$^{-1}$). Unfortunately, X-ray quality crystals of To$^M$CoCO were not obtained to support this assignment. When the crystals were dissolved in C$_6$D$_6$ and CO was re-introduced, the diagnostic $^1$H NMR signal at –18 ppm assigned to the alkoxy ketone species disappeared and the remaining signals match those that have been assigned to To$^M$Co{C(O)Bn}CO suggesting that its formation is reversible.

A possible mechanism is suggested (Scheme 4). The first step is formation of a ketone species that is proposed to occur by combination of the cobalt(II) acyl with cobalt(II) benzyl forming dibenzylketone and a reduced Co–CO species. The combination of an acyl and alkyl species to form a ketone has been proposed in the Fischer Tropsch reaction.$^{31}$ Then, species 7d could form by insertion of dibenzylketone into the cobalt-carbon bond of the cobalt acyl species (7b). Nucleophilic acylation of ketones to form alpha hydroxy ketones has been proposed using acyllithium reagents.$^{32}$
A) Ketone formation

\[
\begin{align*}
\text{[Co]} & \text{Ph} \quad \text{CO} \rightarrow \quad \text{[Co]} \quad \text{7b} \\
\text{[Co]} & \text{Ph} \quad \text{Ph} \quad \text{O} \quad \text{[Co]} \quad \text{7c}
\end{align*}
\]

\[
\begin{align*}
\text{[Co]} & \text{Ph} \quad \text{Ph} \quad \text{Ph} \quad \text{O} \\
\text{[Co]} & \text{Ph} \quad \text{Ph} \quad \text{Ph} \quad \text{O}
\end{align*}
\]

B) C-C bond formation

\[
\begin{align*}
\text{[Co]} & \text{Ph} \quad \text{Ph} \quad \text{O} \quad \text{[Co]} \quad \text{7d} \\
\text{[Co]} & \text{Ph} \quad \text{Ph} \quad \text{O} \quad \text{[Co]} \quad \text{7e}
\end{align*}
\]

Scheme 4. Proposed formation of 7d and 7e by reaction of To\textsuperscript{M}CoBn (1) and CO.

Oxidation of To\textsuperscript{M}Co{C(O)R}CO and Carboxylation of To\textsuperscript{M}CoR.

Complexes 7 – 12 rapidly react with O\textsubscript{2} to form the corresponding carboxylate complexes (To\textsuperscript{M}CoO\textsubscript{2}CBn (13), To\textsuperscript{M}CoO\textsubscript{2}CCH\textsubscript{2}SiMe\textsubscript{3} (14), To\textsuperscript{M}CoO\textsubscript{2}CPh (15),
To\textsuperscript{M}CoO\textsubscript{2}CEt (16), To\textsuperscript{M}CoO\textsubscript{2}C"Bu (17), and To\textsuperscript{M}CoO\textsubscript{2}CMc (18)) (eqn (5)).
The orange mixtures of carbonyl and acyl immediately change to purple solutions upon addition of \( \text{O}_2 \) at room temperature. The \(^1\text{H}\) NMR spectra of the products were characteristic, with assignable signals for methyl (18 H) at an unusually upfield chemical shift of ca. \(-45\) ppm and methylene (6 H) groups from the oxazoline. For comparison, the oxazoline methyl signals in the organocobalt starting materials appear at \(-12\) ppm. The \(^{11}\text{B}\) NMR spectra of 13 – 18 displayed a single signal in the region from 86 to 112 ppm, which is also distinct from that observed for the carbonylated species (c.a. \(-4\) ppm) but similar to the region observed for \(\text{To}^\text{M}\text{CoR}\).

A second route, involving reaction of \(\text{To}^\text{M}\text{CoCl}\) and \(\text{MO}_2\text{CR}\) (\(M = \text{Na or K}; R = \text{Bn, Ph, Et, Me}\)) provides 13, 15, 16, and 18. Unfortunately, we were unable to isolate the carboxylate products from this route; however, the \(^1\text{H}\) and \(^{11}\text{B}\) NMR spectra, UV-Vis spectra (as THF solutions), and IR spectra (as KBr pellets) of these samples matched the spectra obtained from sequential carbonylation and oxygenation.
Additionally, the reaction of $\text{To}^\text{M} \text{CoCH}_2\text{SiMe}_3$ (2) and $\text{CO}_2$ provides $\text{To}^\text{M} \text{CoO}_2\text{CCH}_2\text{SiMe}_3$ (14), and $\text{To}^\text{M} \text{Co}''\text{Bu}$ (5) and $\text{CO}_2$ provides $\text{To}^\text{M} \text{CoO}_2\text{C}''\text{Bu}$ (17). These species were not accessible through salt metathesis. Likewise, compounds 3, 4, and 6 react with $\text{CO}_2$ to provide the corresponding carboxylates. Again, the $^1\text{H}$ and $^{11}\text{B}$ NMR spectra, UV-vis spectra, and IR spectra matched the material obtained from oxygenation. Notably, direct insertion of $\text{CO}_2$ into the Co-C bond required longer reaction times of three days for $R = \text{Ph}$ and over two weeks for $R = \text{Me, Et, }''\text{Bu, or CH}_2\text{SiMe}_3$. Strangely, the reaction of $\text{To}^\text{M} \text{CoBn}$ (1) and $\text{CO}_2$ provides a compound with an $^{11}\text{B}$ NMR signal at 108 ppm that does not match the $^{11}\text{B}$ NMR signal of $\text{To}^\text{M} \text{CoO}_2\text{CBn}$ (13) (86 ppm).

Finally, the structural assignments of 13 and 15 – 18 were confirmed by X-ray diffraction studies of X-ray quality crystals grown from saturated pentane solutions at $-40 \, ^\circ\text{C}$ (Figures 9-11). The crystals of 13 were obtained from reaction of $\text{To}^\text{M} \text{CoCl}$ with KO$_2\text{Bn}$, 16 – 18 by reaction of $\text{To}^\text{M} \text{CoR}$ ($R = \text{Et, }''\text{Bu, and Me}$) with CO followed by O$_2$, and 15 by reaction of $\text{To}^\text{M} \text{CoPh}$ with $\text{CO}_2$. The cobalt-oxygen interatomic distances associated with the carboxylate moiety are inequivalent in each of these structures. For example, in 13 the Co–O4 and Co–O5 interatomic distances are 2.040(2) and Co1–O5 2.14(2) Å, respectively. The Co–O4 and Co–O5 interatomic distance in each of the carboxylate complexes characterized by X-ray diffraction are less than or similar to the covalent radii of Co–O (2.2 Å). As comparison, the Co–O interatomic distance in $\text{To}^\text{M} \text{CoO}''\text{Bu}$ is 1.821(2) Å.
Figure 9. Thermal ellipsoid plot of $\text{To}^M\text{CoO}_2\text{CBn}$ (13) with ellipsoids at 50% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1-O4 2.040(2), Co1-O5 2.214(2), Co1-N1 2.066(3), Co1-N2 2.023(3), Co1-N3 2.059(3); N1-Co1-N2 92.2(1), N1-Co1-N3 88.6(1), N2-Co1-N3 89.9(1), N1-Co1-O4 98.4(1), N2-Co1-O4 118.2(1), N3-Co1-O4 150.6(1), O4-Co1-O5 61.3(1).

Figure 10. Thermal ellipsoid plot of $\text{To}^M\text{CoO}_2\text{CPh}$ (15) with ellipsoids at 20% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Co1-O4 2.150(3), Co1-O5 2.085(3), Co1-N1 2.024(3), Co1-N2 2.057(3), Co1-N3 2.062(3); N1-Co1-N2 91.3(1), N1-Co1-N3 90.5(1), N2-Co1-N3 89.4(1), N1-Co1-O4 119.3(1), N2-Co1-O4 146.5(2), N3-Co1-O4 102.7(1), O4-Co1-O5 58.7(1).
Figure 11. Thermal ellipsoid plot of $\text{To}^M\text{CoO}_2\text{C}^\text{Et}$ (16) with ellipsoids at 50% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): $\text{Co1-O4 2.006(4), Co1-O5 2.185(5), Co1-N1 2.102(4), Co1-N2 2.080(4), Co1-N3 2.030(4); N1-Co1-N2 102.4(2), N1-Co1-N3 89.9(2), N2-Co1-N3 87.7(2), N1-Co1-O4 90.6(2), N2-Co1-O4 160.0(2), N3-Co1-O4 107.7(2), O4-Co1-O5 67.0(2)}$.

Figure 12. Thermal ellipsoid plot of $\text{To}^M\text{CoO}_2\text{C}^\text{Bu}$ (17) with ellipsoids at 50% probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): $\text{Co1-O4 2.054(2), Co1-O5 2.173(2), Co1-N1 2.073(2), Co1-N2 2.068(2), Co1-N3 2.023(2); N1-Co1-N2 87.79(6), N1-Co1-N3 90.70(6), N2-Co1-N3 90.76(6), N1-Co1-O4 98.47(6), N2-Co1-O4 154.95(8), N3-Co1-O4 113.25(8), O4-Co1-O5 60.94(6)}$. 
The possibility of adventitious H$_2$O serving as the source of oxygen was ruled out by the independent reaction of 12 with over 10 equivalents of H$_2$O (eqn (6)). $^1$H NMR spectrum analysis revealed only 12 remained after 30 minutes. Upon introduction of O$_2$ to this same sample, within seconds complete conversion to 18 was observed.

When complex 12 was generated using isotopically labeled $^{13}$CO and then reacted with O$_2$, a new signal at 124 ppm assigned to $^{13}$CO$_2$ was detected in the $^{13}$C{$^1$H} NMR spectrum of complex 18. The $^{13}$C{$^1$H} NMR spectrum of 18 also contained an isotopically enriched signal at 82 ppm that was assigned to the O$_2$-$^{13}$CMe moiety. The latter observation supports the formation of the carboxylic group as occurring by O$_2$ oxygenation of the acetyl moiety. Together, these studies suggest that the oxidation of the acyl carbonyl complex 12 involves oxidation of both the acyl moiety to form the carboxylate derivative as well as oxidation of the terminal carbonyl to form CO$_2$ as a byproduct. A potential intermediate is an acylperoxo species formed from O$_2$ insertion into the cobalt acyl bond. Such a species is proposed by Akita and coworkers in an analogous reaction involving a cobalt(II) $p$-tolyl complex, CO, and O$_2$ to form a carboxylate species. Similarly, oxidations of hydrocarbons to form alcohols involving...
$m$CPBA as the oxidant are proposed to involve binding of $m$CPBA to the metal center to form an acylperoxy species that undergoes O atom transfer to the hydrocarbon and release of a benzoic derivative. To$^M$CoOAc has been reported to be reactive in in oxidations of hydrocarbons using mCPBA. Conceptually, under the conditions presented here, a similar reaction could occur with oxidation of a acyl ligand rather than a hydrocarbon substrate.\textsuperscript{34}

**Conclusion**

The series of organocobalt(II) complexes 1–6 share similar spectroscopic features by $^1$H and $^{11}$B NMR, IR, and UV-Vis spectroscopy. Particularly, characteristic intense absorptions at ca. 350 and 700 nm in the UV-Vis spectra are a general feature of tetrahedral alkyl or aryl To$^M$Co(II) complexes and distinguishes them from other four-coordinate compounds lacking a Co–C bond such as To$^M$CoOAc, To$^M$CoOtBu or To$^M$CoCl. Reactions of 1–6 with CO afford mixtures containing organocobalt carbonyl, cobalt acyl, and/or cobalt acyl carbonyl species that interconvert. The speciation of the mixture varies, depending upon the R group, with only the benzyl (7) favoring organocobalt carbonyl and cobalt acyl, whereas ethyl (10) and methyl (12) favor cobalt acyl and its carbonyl adduct. The reactivity of the mixtures appears to correlate with this speciation. Evaporation of solutions of 8–12 provides the starting alkyl or aryl compounds via decarbonylation; however, evaporation of solutions of 7 (R = Bn) results in combination of three benzyl groups, two CO and one To$^M$Co-moiety to form an alkoxy ketone species. The CO is formally oxidized in this process, and we propose that two of the three benzyl groups are derived from To$^M$CoBn species in which Co is reduced to Co(I). Reaction of O$_2$ and in situ-generated carbonylated species provides the
carboxylate complexes. The rapid multi-step oxidative carbonylation route to form the carboxylate derivatives sharply contrasts the timescale of the alternative direct insertion of \( \text{CO}_2 \) into the cobalt carbon bond, which requires several days to weeks. In addition, oxidative carbonylation selectively yields the carboxylates making it superior to the independent salt metathesis reaction of \( \text{To}^\text{MCoCl} \) with the \( \text{KOO}_2\text{CR} \) salts, which yields unidentified byproducts. Importantly, if C–H activation of a hydrocarbon could complete the cycle to form the starting cobalt alkyl and the carboxylate, this reactivity could be incorporated into new catalytic cycles for making carboxylate acids. The general oxidative carbonylation reactivity observed for \( R = \text{Bn}, \text{CH}_2\text{SiMe}_3, \text{Ph}, \text{Et}, \text{^nBu}, \text{and Me} \) demonstrates the versatility of this reaction that could prove useful in synthesizing an array of carboxylates. Further studies will explore such catalytic application of this oxidative carbonylation reactivity as well comparison with other late transition metals such as iron and nickel in place of cobalt.

**Experimental Section**

**General Procedures.** All reactions were performed using standard Schlenk techniques under an atmosphere of dry argon. Benzene and diethyl ether were dried and deoxygenated using an IT PureSolv system. Benzene-\( d_6 \) was degassed with freeze-pump-thaw cycles, heated to reflux over a Na/K alloy, and then vacuum transferred. \( \text{To}^\text{MCoCl}, \text{To}^\text{MCoMe}, \text{and To}^\text{MCoOAc} \) were synthesized following the reported procedures.\(^1,3\) \( ^1\text{H} \) and \( ^{11}\text{B} \) NMR spectra were collected on a Bruker Avance III 600 spectrometer. \( ^{11}\text{B} \) NMR spectra were referenced to an external sample of \( \text{BF}_3\cdot\text{Et}_2\text{O} \). Infrared spectra were measured on a Bruker Vertex 80 FTIR spectrometer. EPR were obtained on an X-band Elexsys 580 FT- EPR spectrometer in continuous wave mode. UV-Vis spectra were
recorded on an Agilent 8453 UV-vis spectrophotometer. Elemental analyses were
performed using a Perkin-Elmer 2400 Series II CHN/S. Single crystal X-ray diffraction
data was collected on an APEX II.

**Computational methods.** Density Functional Theory (DFT) optimizations were
performed using NWChem\(^{35}\) on \(6, 10c, 12a, 12c\), and \(\text{To}^M\text{CoCl}\) using the PBE0 hybrid
functional,\(^{25}\) the 6-311+G* basis\(^{26}\) for first and second row elements, and the Stuttgart
RSC 1997\(^{27}\) effective core potential for the cobalt. Single-point energy calculations were
also performed on both the high-spin and low-spin configurations to determine the lowest
energy spin state for each structure. DFT Hessians were performed on \(6, 12a,\) and \(12c\) to
compute vibrational frequencies. A Time-Dependent DFT (TDDFT) calculation using
implicit solvation\(^{36}\) was initially performed to calculate excited states for \(6\), but it was
determined that a more robust method allowing for multiple electronic configurations
was needed due to missing excitations in the results. Gas-phase configuration interaction
(CI) singles calculations were performed using the GAMESS quantum chemistry
package\(^ {37}\) on \(6\) and \(\text{To}^M\text{CoCl}\) to compute the first 25 excited states for each structure.

**\(\text{To}^M\text{CoBn (1)}\).** \(\text{To}^M\text{CoCl} (0.147 \text{ g}, 0.308 \text{ mmol})\) and \(\text{KBN} (0.067\text{g}, 0.515 \text{ mmol})\) were
stirred in tetrahydrofuran (10 mL) for 2 h. The green solution was dried under vacuum,
the residue was extracted with benzene (10 mL), and the extracts were dried *in vacuo* to
afford \(\text{To}^M\text{CoBn}\) as a green solid (0.129 g, 0.242 mmol, 78.7%). X-ray quality crystals
were obtained from a saturated pentane solution at –40 °C. \(^1\)H NMR (benzene-\(d_6\), 600
MHz): \(\delta\) 34.47 (s, 2 H), 16.33 (s, 6 H, CNCMe\(_2\)CH\(_2\)O), 14.90 (s, 2 H), 10.96 (s, 2 H),
9.14 (s, 1 H), –12.46 (s, 18 H, CNCMe\(_2\)CH\(_2\)O), –77.45 (s, 1 H), -89.01 (s, 1 H). \(^{11}\)B NMR
(benzene-\(d_6\), 128 MHz): \(\delta\) 100.4. IR (KBr, cm\(^{-1}\)): ν 3072 (w), 3046 (w), 3012 (w), 2966
(m), 2926 (m), 2897 (m), 2869 (m), 1701 (w), 1588 (s, $\nu_{CN}$), 1482 (m), 1459 (m), 1384 (m), 1366 (m), 1351 (m), 1271 (m), 1194 (m) 1160 (m), 1027 (m), 1008 (m), 963 (m).

UV-vis (Et$_2$O) $\lambda_{max} = 339$ ($\varepsilon$: 3205 M$^{-1}$cm$^{-1}$), 439 ($\varepsilon$: 1964 M$^{-1}$cm$^{-1}$) 589 ($\varepsilon$: 332 M$^{-1}$cm$^{-1}$), 628 ($\varepsilon$: 365 M$^{-1}$cm$^{-1}$), 704 ($\varepsilon$: 1444 M$^{-1}$cm$^{-1}$). $\mu_{eff}$(C$_6$D$_6$) = 4.2(7) $\mu_B$ as determined by the Evans method. Anal. Calcd. for C$_{42}$H$_{58}$B$_2$CoN$_6$O$_6$: C, 63.17; H, 6.82; N, 7.89. Mp. 151-153°C, dec.

To$^M$CoCH$_2$SiMe$_3$ (2). LiCH$_2$SiMe$_3$ (0.041 g, 0.44 mmol) dissolved in pentane (1 mL) was added dropwise to a solution of To$^M$CoCl (0.150 g, 0.31 mmol) in tetrahydrofuran (3 mL) cooled to −78 °C. The purple reaction mixture was stirred for 15 min. at −78 °C, warmed to room temperature, and then stirred for an additional 45 min. The solvent was evaporated, and the residue was extracted with benzene. Evaporation of benzene afforded To$^M$CoCH$_2$SiMe$_3$ as a blue solid (0.161 g, 0.30 mmol, 97% yield). X-ray quality crystals were obtained from a saturated pentane solution at −40 °C. $^1$H NMR (benzene-$d_6$, 600 MHz): $\delta$ 15.73 (s, 6 H, CNCMe$_2$CH$_2$O), 12.72 (s, 2 H, C$_6$H$_5$), 10.08 (s, 2 H, C$_6$H$_5$), 8.54 and 8.48 (s, 10 H, CH$_2$SiMe$_3$, p-C$_6$H$_5$), −9.58 (s, 18 H, CNCMe$_2$CH$_2$O). $^{11}$B NMR (benzene-$d_6$, 128 MHz): $\delta$ 86.6. IR (KBr, cm$^{-1}$): v 2966 (m), 2892 (m), 1586 (s, $\nu_{CN}$), 1490 (m), 1462 (m), 1434 (m), 1386 (m), 1366 (m), 1275 (m), 1251 (m) 1195 (m), 1159 (m), 1020 (m), 1001 (m), 966 (m). UV-vis (Et$_2$O) $\lambda_{max} = 355$ ($\varepsilon$: 2228 M$^{-1}$cm$^{-1}$), 591 ($\varepsilon$: 374 M$^{-1}$cm$^{-1}$), 628 ($\varepsilon$: 448 M$^{-1}$cm$^{-1}$), 703 ($\varepsilon$: 1495 M$^{-1}$cm$^{-1}$). $\mu_{eff}$(C$_6$D$_6$) = 4.9(3) $\mu_B$ as determined by the Evans method. Anal. Calcd. for C$_{25}$H$_{40}$BCoN$_3$O$_3$Si: C, 56.82; H, 7.63; N, 7.95 Found: C, 56.25; H, 7.72; N, 7.54. Mp. 234-235 °C, dec.

To$^M$CoPh (3). Phenyllithium (1.8 M in dibutyl ether, 0.250 mL, 0.45 mmol) was added dropwise to a solution of To$^M$CoCl (0.151 g, 0.317 mmol) in THF (3 mL) cooled to −78
°C. The dark blue reaction mixture was stirred for 15 min. at −78 °C, warmed to room temperature, and stirred for an additional 45 min. The solvent was evaporated, and the residue was extracted with benzene. Evaporation of benzene provided ToMCoPh as a blue solid (0.119 g, 0.230 mmol, 72.6% yield). X-ray quality crystals were obtained from saturated pentane solutions cooled to −40 °C. $^1$H NMR (benzene-$d_6$, 600 MHz): δ 73.95 (s, 1 H), 16.73 (s, 6 H, CNCMe$_2$CH$_2$O), 15.74 (s, 2 H, C$_6$H$_5$), 11.30 (s, 2 H, C$_6$H$_5$), 10.61 (s, 1 H, p-C$_6$H$_5$), 9.42 (s, 1 H, p-C$_6$H$_5$), −13.68 (s, 18 H, CNCMe$_2$CH$_2$O). $^{11}$B NMR (benzene-$d_6$, 128 MHz): δ 107.7. IR (KBr, cm$^{-1}$): ν 3042 (m), 2967 (m), 2927 (m), 2890 (m), 2869 (m), 1582 (s, νCN), 1494 (m), 1461 (m), 1433 (m), 1388 (m), 1368 (m), 1352 (m), 1274 (m), 1196 (m), 1159 (m), 964 (m), 951 (m). UV-vis ($(Et_2O)$: $\lambda_{max}$ = 330 (ε: 1245 M$^{-1}$cm$^{-1}$), 587 (ε: 176 M$^{-1}$cm$^{-1}$), 625 (ε: 252 M$^{-1}$cm$^{-1}$), 708 (ε: 1333 M$^{-1}$cm$^{-1}$). $\mu_{eff}$ (C$_6$D$_6$) = 4.0(1) $\mu_B$ as determined by the Evans method. Anal. Calcd. for C$_{27}$H$_{34}$BCoN$_3$O$_3$: C, 62.57; H, 6.61; N, 8.11 Found: C, 63.05; H, 6.81; N, 7.75. Mp. 194-197 °C, dec.

ToMCoEt (4). Ethyllithium (0.5 M in a mixture of benzene and cyclohexane, 0.880 mL, 0.44 mmol) was added dropwise to a solution of ToMCoCl (0.150 g, 0.31 mmol) in THF (3 mL) cooled to −78 °C. The purple reaction mixture was stirred for 15 min. at −78 °C, warmed to room temperature, and stirred for an additional 45 min. Evaporation of volatiles, extraction of the solid residue with benzene, and evaporation of the benzene afforded ToMCoEt as a green solid (0.114 g, 0.24 mmol, 77% yield). X-ray quality crystals were obtained from saturated pentane solutions at −40 °C. $^1$H NMR (benzene-$d_6$, 600 MHz): δ 15.95 (s, 2 H, C$_6$H$_5$), 14.85 (s, 6 H, CNCMe$_2$CH$_2$O), 11.33 (s, 2 H, C$_6$H$_5$), 9.42 (s, 1 H, p-C$_6$H$_5$), −14.45 (s, 18 H, CNCMe$_2$CH$_2$O), −31.31 (s, 1 H, CH$_2$CH$_3$). $^{11}$B
NMR (benzene-$d_6$, 128 MHz): δ 116.7. IR (KBr, cm$^{-1}$): ν 3078 (w), 3045 (w), 2967 (m), 2929 (m), 2895 (m), 2870 (m), 2837 (m), 1592 (s, ν$_{CN}$), 1495 (m), 1461 (m), 1434 (m), 1385 (m), 1366 (m), 1351 (m), 1268 (m), 1193 (m), 1159 (m), 1101 (w), 1017 (w), 961 (m). UV-vis ((Et$_2$O) λ$_{max}$ = 382 (ε: 1935 M$^{-1}$cm$^{-1}$), 579 (ε: 268 M$^{-1}$cm$^{-1}$), 615 (ε: 286 M$^{-1}$cm$^{-1}$), 705 (ε: 1255 M$^{-1}$cm$^{-1}$). µ$_{eff}$(C$_6$D$_6$) = 4.1(6) µ$_B$ as determined by the Evans method. Anal. Calcd. for C$_{23}$H$_{34}$BCoN$_3$O$_3$: C, 58.74; H, 7.29; N, 8.94 Found: C, 58.83; H, 7.07; N, 8.64. Mp. 179-181°C, dec.

**To$^M$Co$^n$Bu (5).** To$^M$CoCl (0.155 g, 0.33 mmol) was dissolved in THF (3 mL), cooled to −78 °C, and n-butyllithium (2.5 M in hexanes, 0.19 mL, 0.48 mmol) was added in a dropwise fashion. The dark purple reaction mixture was stirred at −78 °C for 15 min., warmed to room temperature, and stirred for an additional 45 min. THF was removed en vacuo, the residue was extracted with benzene, and benzene was evaporated to afford To$^M$Co$^n$Bu as a dark green solid (0.100 g, 0.20 mmol, 61% yield). X-ray quality crystals were obtained from saturated pentane solutions cooled to −40 °C. $^1$H NMR (benzene-$d_6$, 600 MHz): δ 15.91 (s, 2 H, C$_6$H$_5$), 14.88 (s, 6 H, CNCMe$_2$CH$_2$O), 14.21 (s, 2 H, (CH$_2$)$_3$CH$_3$), 11.34 (s, 2 H, C$_6$H$_5$), 9.47 (s, 1 H, p-C$_6$H$_5$), −2.67 (s, 3 H, (CH$_2$)$_3$CH$_3$), −14.31 (s, 18 H, CNCMe$_2$CH$_2$O). $^{11}$B NMR (benzene-$d_6$, 128 MHz): δ 115.0. IR (KBr, cm$^{-1}$): ν 3083 (w), 3047 (w), 2966 (m), 2928 (m), 2890 (m), 2871 (m), 1587 (s, ν$_{CN}$), 1495 (m), 1462 (m), 1434 (m), 1386 (m), 1368 (m), 1351 (m), 1271 (m), 1198 (m), 1159 (m), 1104 (w), 1025 (m), 995 (m). UV-vis (Et$_2$O): λ$_{max}$ = 382 (ε: 1679 M$^{-1}$cm$^{-1}$), 576 (ε: 406 M$^{-1}$cm$^{-1}$), 613 (ε: 422 M$^{-1}$cm$^{-1}$), 705 (ε: 1291 M$^{-1}$cm$^{-1}$). µ$_{eff}$(C$_6$D$_6$) = 4.5(2) µ$_B$ as determined by the Evans method. Anal. Calcd. for C$_{25}$H$_{38}$BCoN$_3$O$_3$: C, 60.26; H, 7.69; N, 8.43 Found: C, 60.25; H, 7.37; N, 8.32. Mp. 244-246°C, dec.
Preparation of Carbonylated Complexes. As a typical procedure, $\text{To}^M\text{CoEt}$ (0.01 g, 0.02 mmol) was dissolved in benzene-$d_6$ (0.5 mL), the headspace was removed in vacuo, and then CO (1 atm) was introduced. An immediate color change from green to orange was observed. Benzene-$d_6$ was evaporated in vacuo to afford $\text{To}^M\text{Co}\{\text{C(O)Et}\}_2\text{CO}$ as a yellow solid (0.01 g, 0.02 mmol, 95% yield). X-ray quality crystals were obtained from saturated pentane solutions cooled to –40 °C. The other carbonylated complexes 7 – 12 were synthesized by this same procedure.

$\text{To}^M\text{Co}\{\text{C(O)Bn}\}_2\text{CO}$ (7): $^1\text{H}$ NMR (benzene-$d_6$, 600 MHz): $\delta$ 82.15, 64.43, 42.01, 26.06, 14.98, 12.63, 10.91, 9.11, 5.63, 2.78, –2.29, –5.98, –16.04, –25.97, –28.18. $^{11}\text{B}$ NMR (benzene-$d_6$, 128 MHz): $\delta$ 87.0. IR (KBr, cm$^{-1}$): v 3076 (w), 3062 (w), 2965 (m), 2927 (m), 1982 (w), 1948 (m), 1888 (m), 1717 (w), 1669 (m), 1594 (s, $\nu_{\text{CN}}$), 1495 (m), 1460 (m), 1366 (m), 1351 (m), 1273 (m), 1195 (m), 1158 (m), 1125 (w), 1095 (w), 1069 (w), 1028 (m), 957 (m). $\mu_{\text{eff}}$ ($C_6D_6$) = 2.8(2) $\mu_B$ as determined by the Evans method.

$\text{To}^M\text{Co}\{\text{C(O)CH}_2\text{SiMe}_3\}_2\text{CO}$ (8): $^1\text{H}$ NMR (benzene-$d_6$, 600 MHz): $\delta$ 9.56, 8.16, 7.62, 6.89, –0.09, –0.98. $^{11}\text{B}$ NMR (benzene-$d_6$, 128 MHz): $\delta$ –4.7. IR (KBr, cm$^{-1}$): v 3077 (w), 3046 (w), 2967 (m), 2385 (m), 2055 (m, $\nu_{\text{CO}}$), 1982 (m, $\nu_{\text{CO}}$), 1938 (m, $\nu_{\text{CO}}$), 1887 (m, $\nu_{\text{CO}}$), 1590 (s, $\nu_{\text{CN}}$), 1573 (m, $\nu_{\text{CN}}$), 1556 (m, $\nu_{\text{CN}}$), 1494 (m), 1463 (m), 1433 (m), 1388 (m), 1368 (m), 1352 (m), 1275 (m), 1251 (m), 1197 (m), 1162 (m), 1024 (m), 962 (m). UV-vis (THF): $\lambda_{\text{max}}$ = 669 ($\epsilon$: 66 M$^{-1}$cm$^{-1}$) and 903 ($\epsilon$: 104 M$^{-1}$cm$^{-1}$) $\mu_{\text{eff}}$ ($C_6D_6$) = 2.4(2) $\mu_B$ as determined by the Evans method.

$\text{To}^M\text{Co}\{\text{C(O)Ph}\}_2\text{CO}$ (9): $^1\text{H}$ NMR (benzene-$d_6$, 600 MHz): $\delta$ 9.76, 8.22, 7.71, 7.46, 7.16, –1.49. $^{11}\text{B}$ NMR (benzene-$d_6$, 128 MHz): $\delta$ –5.7. IR (KBr, cm$^{-1}$): v 3077 (w), 3044 (w), 2965 (m), 2926 (m), 2892 (m), 2853 (m), 1985 (w, $\nu_{\text{CO}}$), 1986 (w, $\nu_{\text{CO}}$), 1591 (s, $\nu_{\text{CN}}$).
1557 (m), 1494 (m), 1462 (m), 1415 (m), 1388 (m), 1368 (m), 1353 (m), 1279 (m), 1198 (m), 1160 (m), 1053 (m), 1025 (m), 968 (m). UV-vis (THF): $\lambda_{\text{max}} = 909$ (\(\varepsilon\): 135 M$^{-1}$cm$^{-1}$) $\mu_{\text{eff}}(C_6D_6) = 2.4(2)$ \(\mu_B\) as determined by the Evans method.

$\text{To}^M\text{Co}\{\text{C(O)Et}\}CO$ (10). $^1$H NMR (benzene-$d_6$, 600 MHz): $\delta$ 9.61, 8.17, 7.62, -1.18, -7.69. $^{11}$B NMR (benzene-$d_6$, 128 MHz): $\delta$ -4.44. IR (KBr, cm$^{-1}$): v 3080 (w), 3046 (w), 2967 (m), 2931 (m), 2890 (m), 1989 (w), 1966 (w), 1942 (w), 1886 (m), 1594 (s, \(\nu_{\text{CN}}\)), 1565 (m), 1495 (m), 1465 (m), 1433 (m), 1387 (m), 1367 (m), 1352 (m), 1275 (m), 1196 (m), 1161 (m), 1075 (w), 1024 (m), 961 (m). UV-vis (THF): $\lambda_{\text{max}} = 669$ (\(\varepsilon\): 102 M$^{-1}$cm$^{-1}$) and 704 (\(\varepsilon\): 116 M$^{-1}$cm$^{-1}$) $\mu_{\text{eff}}(C_6D_6) = 2.7(2)$ \(\mu_B\) as determined by the Evans method.

$\text{To}^M\text{Co}\{\text{C(O)Bu}\}CO$ (11). $^1$H NMR (benzene-$d_6$, 600 MHz): $\delta$ 9.61, 8.17, 7.63, 0.94, -1.16. $^{11}$B NMR (benzene-$d_6$, 128 MHz): $\delta$ -4.9. IR (KBr, cm$^{-1}$): v 3077 (w), 3046 (w), 2965 (m), 2930 (m), 2893 (m), 2873 (m), 2055 (w, \(\nu_{\text{CO}}\)), 1980 (w, \(\nu_{\text{CO}}\)), 1941 (m, \(\nu_{\text{CO}}\)), 1887 (w, \(\nu_{\text{CO}}\)), 1594 (s, \(\nu_{\text{CN}}\)), 1556 (m), 1494 (m), 1463 (m), 1434 (m), 1367 (m), 1354 (m), 1276 (m), 1196 (m), 1161 (m), 1026 (m), 960 (m).

Preparation of Carboxylate Complexes. As a typical procedure, the carbonylated species 10 was generated in situ. The headspace was removed and O$_2$ (1 atm) was introduced. A color change from orange to purple was immediately visible. Benzene-$d_6$ was evaporated in vacuo to afford $\text{To}^M\text{CoO}_2\text{CEt}$ as a purple. X-ray quality crystals were obtained for complexes 13 and 15 – 18 from a saturated pentane solution cooled to $-40 \degree C$.

$\text{To}^M\text{Co}(\text{O}_2\text{CBn})$ (13). $^1$H NMR (benzene-$d_6$, 600 MHz): $\delta$ 49.61, 32.02, 28.01, 23.86, 23.64, 17.32, 13.17, 10.93, 9.68, $-40.84$. $^{11}$B NMR (benzene-$d_6$, 128 MHz): $\delta$ 86.4. IR (KBr, cm$^{-1}$): v 3080 (w), 3040 (w), 2964 (m), 2928 (m), 2891 (m), 1667 (m), 1591 (s,
\(\nu_{\text{CN}}\), 1517 (m), 1495 (m), 1462 (m), 1402 (m), 1367 (m), 1351 (m), 1275 (m), 1196 (m),
1160 (m), 1126 (m), 1095 (m), 1029 (m), 961 (m). UV-vis (THF): \(\lambda_{\text{max}} = 663 \, (\varepsilon: 200 \, \text{M}^{-1} \, \text{cm}^{-1})\). \(\mu_{\text{eff}}(\text{C}_6\text{D}_6) = 4.7(2) \, \mu_\text{B}\) as determined by the Evans method. Mp. 230-232 °C.

**\(\text{To}^\text{M} \text{Co(O}_2\text{CCH}_2\text{SiMe}_3\) (14).** \(^1\)H NMR (benzene-\(d_6\), 600 MHz): \(\delta\) 208.77 (s, 2 H, \(\text{O}_2\text{CCH}_2\text{SiMe}_3\)), 37.30 (s, 2 H, \(\text{C}_6\text{H}_5\)), 19.31 (s, 2 H, \(\text{C}_6\text{H}_5\)), 16.20 (s, 1 H, \(p\)-\(\text{C}_6\text{H}_5\)), 15.92 (s, 9 H, \(\text{O}_2\text{CCH}_2\text{SiMe}_3\)), 11.34 (s, 6 H, \(\text{CNCMe}_2\text{C}_2\text{H}_5\)), \(\delta\) -49.53 (s, 18 H, \(\text{CNCMe}_2\text{CH}_2\text{O}\)). \(^{11}\)B NMR (benzene-\(d_6\), 128 MHz): \(\delta\) 113.2. IR (KBr, cm\(^{-1}\)): \(\nu\) 3074 (w), 3045 (w), 2965 (m), 2932 (m), 2893 (m), 2874 (m), 1588 (s, \(\nu_{\text{CN}}\)), 1544 (m, \(\nu_{\text{CN}}\)), 1496 (m), 1462 (m), 1408 (m), 1368 (m), 1352 (m), 1273 (m), 1197 (m), 1159 (m), 1101 (m), 1025 (m), 962 (m), 949 (m). UV-vis (THF): \(\lambda_{\text{max}} = 588 \, (\varepsilon: 98 \, \text{M}^{-1} \, \text{cm}^{-1})\). \(\mu_{\text{eff}}(\text{C}_6\text{D}_6) = 4.7(2) \, \mu_\text{B}\) as determined by the Evans method. Mp. 186-188 °C.

**\(\text{To}^\text{M} \text{Co(O}_2\text{CPh}\) (15).** \(^1\)H NMR (benzene-\(d_6\), 600 MHz): \(\delta\) 48.80 (s, 2 H), 31.55 (s, 2 H), 23.34 (s, 2 H), 17.16 (s, 2 H), 14.48 (s, 1 H), 13.16 (s, 6 H, \(\text{CNCMe}_2\text{C}_2\text{H}_5\)), 12.83 (s, 1 H), \(-40.00\) (s, 18 H, \(\text{CNCMe}_2\text{CH}_2\text{O}\)). \(^{11}\)B NMR (benzene-\(d_6\), 128 MHz): \(\delta\) 83.6. IR (KBr, cm\(^{-1}\)): \(\nu\) 3074 (w), 3045 (w), 2965 (m), 2927 (m), 2893 (m), 2870 (m), 1589 (s, \(\nu_{\text{CN}}\)), 1536 (m, \(\nu_{\text{CN}}\)), 1496 (m), 1462 (m), 1408 (m), 1368 (m), 1352 (m), 1273 (m), 1197 (m), 1159 (m), 1101 (m), 1025 (m), 962 (m), 949 (m). UV-vis (THF): \(\lambda_{\text{max}} = 587 \, (\varepsilon: 157 \, \text{M}^{-1} \, \text{cm}^{-1})\). \(\mu_{\text{eff}}(\text{C}_6\text{D}_6) = 4.6(2) \, \mu_\text{B}\) as determined by the Evans method. Mp. 238-240 °C.

**\(\text{To}^\text{M} \text{Co(O}_2\text{CEt}\) (16).** \(^1\)H NMR (benzene-\(d_6\), 600 MHz): \(\delta\) 35.16 (s, 4 H), 33.81 (s, 2 H, \(\text{C}_6\text{H}_5\)), 18.05 (s, 2 H, \(\text{C}_6\text{H}_5\)), 15.20 (s, 1 H, \(\text{C}_6\text{H}_5\)), 12.32 (s, 6 H, \(\text{CNCMe}_2\text{CH}_2\text{O}\)), \(-43.62\) (s, 18 H, \(\text{CNCMe}_2\text{CH}_2\text{O}\)). \(^{11}\)B NMR (benzene-\(d_6\), 128 MHz): \(\delta\) 95.0. IR (KBr, cm\(^{-1}\)): \(\nu\) 3080 (w), 3045 (w), 2965 (m), 2932 (m), 2893 (m), 2874 (m), 1588 (s, \(\nu_{\text{CN}}\)), 1544 (m, \(\nu_{\text{CN}}\)).
ν\textsubscript{CN}, 1466 (m), 1437 (m), 1387 (w), 1369 (m), 1352 (m), 1276 (m), 1199 (m), 1160 (m), 1074 (w), 1027 (w), 964 (m), 949 (m). UV-vis (THF): \(\lambda_{\text{max}} = 584 (\varepsilon: 98 \text{ M}^{-1}\text{cm}^{-1})\). \(\mu_{\text{eff}}\) \((\text{C}_6\text{D}_6) = 4.8(2) \mu_B\) as determined by the Evans method. Mp. 174-176 °C.

\textbf{To}^{\text{M}}\text{Co(O}_2\text{C}^\text{Bu}) (17). \(^1\text{H} \text{NMR (benzene-}d_6, 600 \text{ MHz):} \delta 161.09 (s, 2 \text{ H}, \text{O}_2\text{CCH}_2\text{CH}_2\text{CH}_2\text{CH}_3), 34.53 (s, 2 \text{ H}, \text{O}_2\text{C(CH}_2)_3\text{CH}_3), 33.72 (s, 2 \text{ H}, \text{O}_2\text{C(CH}_2)_3\text{CH}_3), 19.24 (s, 2 \text{ H}, \text{C}_6\text{H}_3), 17.93 (s, 2 \text{ H}, \text{C}_6\text{H}_3), 15.08 (s, 1 \text{ H}, p-\text{C}_6\text{H}_3), 12.27 (s, 6 \text{ H}, \text{CNCMe}_2\text{CH}_2\text{O}), 10.83 (s, 3 \text{ H}, \text{O}_2\text{C(CH}_2)_3\text{CH}_3), -43.55 (s, 18 \text{ H}, \text{CNCMe}_2\text{CH}_2\text{O}). \(^{11}\text{B} \text{NMR (benzene-}d_6, 128 \text{ MHz):} \delta 95. \text{ IR (KBr, cm}^{-1}): \nu 3079 (w), 3055 (w), 2963 (m), 2930 (m), 2893 (m), 2871 (m), 1595 (s, \nu\text{CN}), 1540 (m, \nu\text{CN}), 1498 (w), 1461 (m), 1437 (m), 1385 (m), 1365 (m), 1316 (w), 1274 (m), 1107 (w), 1024 (w), 956 (s).

UV-vis (THF): \(\lambda_{\text{max}} = 584 (\varepsilon: 101 \text{ M}^{-1}\text{cm}^{-1})\). \(\mu_{\text{eff}}\) \((\text{C}_6\text{D}_6) = 4.3(2) \mu_B\) as determined by the Evans method. Mp. 197-199 °C.

\textbf{References}


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CHAPTER 5: SYNTHESIS, CHARACTERIZATION, AND REACTIVITY OF TRIS(OXAZOLINYL)BORATO IRON AND NICKEL COMPLEXES

Modified from a paper to be submitted to a journal

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Abstract

The reaction of FeBr$_2$ and thallium tris(4,4-dimethyl-2-oxazolinyl)phenylborate (TlToM) in tetrahydrofuran (THF) provides ToMFeBr (1). Alternatively, FeBr$_2$ and 2 equiv. of TlToM react to give (ToM)$_2$Fe (2); however, this species is not detected in one-to-one reactions of FeBr$_2$ and TlToM. (ToM)$_2$Fe and FeBr$_2$ react in THF overnight via a comproportionation process to give ToMFeBr. Salt metathesis of 1 and KBn affords ToMFeBn (3) in 89% yield. Complexes 1 – 3 contain a high-spin iron(II) center characterized by the effective magnetic moments ranging from 4.9 to 5.4 $\mu_B$ determined by Evans Method. A single $^1$H NMR signal assigned to the methyl groups of the ToM ligand and a single $\nu_{CN}$ band in the IR suggested tridentate coordination of the ToM ligand to iron in 1 and 3 while two $\nu_{CN}$ bands at 1604 and 1548 cm$^{-1}$ indicated bidentate coordination of ToM to iron in 2. The coordination mode is supported by X-ray crystallography studies of 1 – 3 that further identify these complexes as containing a tetrahedral iron center. ToMFeBn and CO (1 atm) react to afford isolable ToMFe{C(O)Bn}(CO)$_2$ (4) as a yellow solid. Complex 4 is diamagnetic ($S = 0$), and the three distinct methyl signals in the $^1$H NMR spectrum are consistent with a six-coordinate, $C_s$-symmetric species. This assignment is supported by its IR spectrum, which revealed intense bands at 2004 and 1935 cm$^{-1}$ (symmetric and asymmetric $\nu_{CO}$), at 1680 and 1662 cm$^{-1}$ (acetyl rotomers, $\nu_{CO}$), and at 1593 and 1553 cm$^{-1}$ ($\nu_{CN}$). A single-
crystal X-ray diffraction study of $\text{To}^\text{M}\text{Fe}\{\text{C} \text{(O)Bn}\}(\text{CO})_2$ confirmed the spectroscopic assignments. The synthesis of $\text{To}^\text{M}\text{NiCl}$ and $\text{To}^\text{M}\text{NiOAc}$ is also reported; however, these complexes did not lead to any organonickel(II) complexes.

**Introduction**

Iron-containing compounds have attracted interest as biological mimics in order to gain further understanding of iron based proteins. In particular, iron complexes stabilized by tris(pyrazolyl)borate serve as models for oxygenation reactions due to the pyrazolyl ligands which mimic the histidine ligands which are so prevalent at the iron coordination site.\(^1\)\(^-\)\(^4\) Many of these systems focus on the reactivity of the iron complexes towards $\text{O}_2$ in order to better understand the mechanism of iron oxygenases. While TpFe compounds have been very useful in studying biological systems, a major general hindrance encountered in scorpionate chemistry is comproportionation reactions that result in, for example, $\text{Tp}_2\text{Fe}$.\(^5\) Synthetic systems oftentimes prevent this undesirable reactivity by introducing steric bulk to the ancillary ligand while biological systems rely more upon second sphere coordination environments to stabilize the metal center. The problem with stabilization of the metal center by increasing the steric bulk of the ancillary ligand is that the metal center becomes less accessible to substrates, thus potentially limiting its reactivity. Previously, we reported the synthesis and isolation of coordinatively unsaturated $\text{To}^\text{M}\text{CoCl}$ and found that the open coordination environment provided by $\text{To}^\text{M}$ allows for the synthesis of $(\text{To}^\text{M})_2\text{Co}$ yet at the same time provides sufficient steric protection such that once $\text{To}^\text{M}\text{CoCl}$ is isolated, the chloride ligand can be readily exchanged with other potentially more reactive ligands without further
complication of $(\text{To}^\text{M})_2\text{Co}$. The results demonstrate $\text{To}^\text{M}$ as a suitable ligand for the study of late transition metal complexes.

Of related importance is the reactivity of $\text{To}^\text{M}\text{CoR}$ ($\text{R} = \text{Me, Et, CH}_2\text{SiMe}_3, \text{^6Bu, Ph, Bn}$), which undergo rapid oxidative carbonylation by reaction of CO followed by $\text{O}_2$ to form carboxylate compounds. The carboxylate compounds can also be formed by direct insertion of $\text{CO}_2$ into the Co-C bond; however, the reaction is kinetically less favorable requiring up to several weeks. Thus, the results parallel biological systems, which also favor a multi-step route to acetate synthesis over direct utilization of $\text{CO}_2$. The intriguing reactivity demonstrated by organocobalt(II) compounds stabilized by $\text{To}^\text{M}$ in oxidative carbonylation lead us to question if analogous iron(II) alkyl compounds might also be reactive towards oxidative carbonylation or if instead the reactivity is unique to cobalt. At the same time, given TpFe complexes have shown promising reactivity as models of enzymes in oxidation reactions, the synthesis of $\text{To}^\text{M}\text{Fe}$ compounds could provide entryway into similar explorative studies.

Herein we report the synthesis and characterization of $\text{To}^\text{M}\text{FeBr}$ (1) and compare it to the synthesis of $\text{To}^\text{M}\text{CoCl}$. The independent synthesis of $(\text{To}^\text{M})_2\text{Fe}$ (2), which was expected as a side product in the synthesis of $\text{To}^\text{M}\text{FeBr}$, is also presented and it is shown that $(\text{To}^\text{M})_2\text{Fe}$ can be converted to $\text{To}^\text{M}\text{FeBr}$ by reaction of 2 with excess $\text{FeBr}_2$. The latter reactivity explains the ease of synthesis of $\text{To}^\text{M}\text{FeBr}$, which contrasts with the synthetic challenges encountered in the synthesis of $\text{To}^\text{M}\text{CoCl}$. The utility of $\text{To}^\text{M}\text{FeBr}$ as a precursor to iron(II) alkyl compounds is demonstrated by the salt metathesis reaction of $\text{To}^\text{M}\text{FeBr}$ with KBn in THF yielding $\text{To}^\text{M}\text{FeBn}$ (3). The reactivity of 3 towards CO, $\text{O}_2$, and $\text{CO}_2$ is studied and presented in relation to the reactivity observed for $\text{To}^\text{M}\text{CoBn}$. 
Results and Discussion

The reaction of TlTo$^M$ and FeBr$_2$ in tetrahydrofuran at room temperature readily affords paramagnetic To$^M$FeBr (1) as a cream colored solid (equation (1)). To$^M$FeBr was characterized by a single signal in the $^{11}$B NMR spectrum at 74 ppm that was significantly displaced from the region in which diamagnetic borate complexes typically appear (e.g., $^{11}$B NMR signal for TlTo$^M$ is located at −16 ppm). The $^1$H NMR spectrum of 1 contained broad, yet diagnostic resonances from 24 to −30 ppm that were assigned to the methyl (−28.9 ppm, 18 H), methylene (20.9 ppm, 6 H) and phenyl group in the To$^M$ ligand on the basis of integration. The IR spectrum of 1 contained a single $\nu$CN band at 1586 cm$^{-1}$, which is consistent with tridentate coordination of the To$^M$ ligand to iron. The 5.1(2) $\mu_B$ effective magnetic moment of 1, measured by the Evans method, characterized 1 as high-spin iron(II). This value is lower than similar tris(pyrazolyl)borate-coordinated tetrahedral iron(II) halide complexes, such as Tp$^{\text{Bu,Me}}$FeBr (5.4(1) $\mu_B$) and Tp$^{\text{Bu}}$FeCl (5.7 $\mu_B$). 8-9

\begin{equation}
\text{TlTo}^M + \text{FeBr}_2 \xrightarrow{\text{tetrahydrofuran}} \text{To}^M\text{FeBr} (\text{1})
\end{equation}

An X-ray diffraction study confirmed tridentate coordination of the To$^M$ ligand to a pseudo-tetrahedral iron center (Figure 1). The average Fe-N interatomic distance is 2.067 Å and the Fe-Br interatomic distance is 2.3517(9) Å. The Fe-N and Fe-Br interatomic distances are longer than those observed in To$^M$CoCl (Co-N$_{\text{average}} = 2.055$ Å and Co-Cl = 2.206(5) Å) as would be expected on the basis of the greater ionic radii of Br.
compared to Cl, and similar to other scorpionate iron(II) halide complexes. For example, in TpBuMeFeBr the Fe-N_{average} interatomic distance is 2.090 Å and the Fe-Br bond interatomic distance is 2.3604(8) Å.  

![Figure 1](image.png)

**Figure 1.** Thermal ellipsoid plot of To^{M}FeBr (1) with ellipsoids at 20 % probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Fe1-Br1 2.3517(9), Fe1-N1 2.072(4), Fe1-N2 2.061(4), Fe1-N3 2.069(4); N1-Fe1-N2 91.4(2), N1-Fe1-N3 92.4(2), N2-Fe1-N3 90.4(2), N1-Fe1-Br1 120.4(2), N2-Fe1-Br1 129.4(1), N3-Fe1-Br1 122.8(1), B1-Fe1-Br1 174.4(2).

The synthesis of To^{M}FeBr is straightforward, and the possible side product To^{M}_{2}Fe is not detected under the conditions of equation 1. Previously, we reported the synthesis of To^{M}CoCl and To^{M}_{2}Co. In the cobalt system, To^{M}_{2}Co forms readily as a contaminant, and synthetic conditions to isolate To^{M}CoCl require CoCl_{2}·THF (rather than CoCl_{2}) used in excess. We sought to identify underlying chemical features responsible for the difference in selectivity for To^{M}MX vs. To^{M}_{2}M compounds. First, the independent synthesis of (To^{M})_{2}Fe was attempted. Interestingly, (To^{M})_{2}Fe forms as the
only detected product from the reaction of FeBr$_2$ and two equiv. of TlTo$^M$ in THF (equation (2)). (To$^M$)$_2$Fe (2) is isolated as an analytically pure white solid in excellent yield (93%).

![Equation 2]

The $^1$H NMR spectrum was rather unusual and contained peaks in the diamagnetic region as well as the expected paramagnetic signals. Given the selective synthesis of To$^M$FeBr above, we considered a possible interpretation of the apparent diamagnetic signals in the $^1$H NMR spectrum that compound 2 was in fact a mixture of To$^M$FeBr and TlTo$^M$. Several data eliminate that possibility. First, the chemical shifts of the diamagnetic signals and those of TlTo$^M$ were distinct. Moreover, mixtures of TlTo$^M$ and (To$^M$)$_2$Fe gave $^1$H NMR spectra that contained two sets of diamagnetic signals, with one set matching the shifts expected for TlTo$^M$ (1.03 (CNCMe$_2$CH$_2$O) and 3.41 (CNCMe$_2$CH$_2$O) ppm) and the other corresponding to the resonances observed in isolated (To$^M$)$_2$Fe (1.02 and 2.96 ppm). The $^{11}$B NMR spectrum of (To$^M$)$_2$Fe contained a characteristic signal at −122 ppm, and this signal was distinct from the downfield $^{11}$B NMR signal in samples of To$^M$FeBr and the diamagnetic signal of TlTo$^M$ ($^{11}$B NMR: −16 ppm). Nonetheless, the $^{11}$B NMR is clearly affected by the high spin Fe(II) center. Finally, the effective magnetic moment for 2 of 4.9(2) $\mu_B$, measured in solution by the Evans method, is consistent with high spin Fe(II). If the signals were in fact due to
TlTo\textsuperscript{M} then the effective magnetic moment would be expected to be much lower due to the lower than expected concentration of the paramagnetic To\textsuperscript{M}Fe species. The reason for the distinct diamagnetic signals intermixed with paramagnetic signals is currently not understood, but it can be concluded that paramagnetic (To\textsuperscript{M})\textsubscript{2}Fe is analytically and spectroscopically pure despite these diamagnetic signals. The infrared spectrum of 2 suggested bidentate coordination of the To\textsuperscript{M} ligands to iron with two \(\nu_{\text{CN}}\) bands appearing at 1604 and 1548 cm\(^{-1}\) similar to the stretching frequencies observed for (To\textsuperscript{M})\textsubscript{2}Co (\(\nu_{\text{CN}} = 1602\) and 1554 cm\(^{-1}\)).

The independent synthesis of (To\textsuperscript{M})\textsubscript{2}Fe from reaction of TlTo\textsuperscript{M} and FeBr\textsubscript{2} in THF at room temperature indicates that (To\textsuperscript{M})\textsubscript{2}Fe is indeed a viable side product under the reaction conditions adopted for the synthesis of To\textsuperscript{M}FeBr. Thus, the results suggest that a thermodynamic difference may be playing a role in the facile synthesis of To\textsuperscript{M}FeBr. To explore this possibility, FeBr\textsubscript{2} and (To\textsuperscript{M})\textsubscript{2}Fe were allowed to react in THF. Upon stirring overnight, NMR analysis confirmed the complete conversion of (To\textsuperscript{M})\textsubscript{2}Fe to To\textsuperscript{M}FeBr. The reaction was found to occur in other solvents as well, such as toluene, but in these more nonpolar solvents the reaction requires heating at 60°C for several days. Thus, it appears that the reason FeBr\textsubscript{2} is an effective precursor for the synthesis of To\textsuperscript{M}FeBr is that while it allows for the formation of (To\textsuperscript{M})\textsubscript{2}Fe, (To\textsuperscript{M})\textsubscript{2}Fe in turn readily reacts with FeBr\textsubscript{2} in THF to form the desired To\textsuperscript{M}FeBr.
The observed transformation of $(\text{To}^M)_2\text{Fe}$ to $\text{To}^M\text{FeBr}$ is particularly unexpected given in the analogous cobalt system, $(\text{To}^M)_2\text{Co}$ occurs readily if conditions are not used that kinetically favor instead the synthesis of $\text{To}^M\text{CoCl}$. In comparison, the analogous $\text{Tp}_2\text{Fe}$ complexes are reported to be stable and in fact present a challenge when attempting to synthesize heteroleptic Fe compounds. For instance, the synthesis of $\text{TpFeR} (R = \text{alkyl or aryl})$ is complicated by the strong tendency of these latter compounds to disproportionate to form $\text{Tp}_2\text{Fe}$.\(^1\) Perhaps one reason for the increased favorability of $\text{Tp}_2\text{Fe}$ versus $(\text{To}^M)_2\text{Fe}$ is that $\text{Tp}_2\text{Fe}$ forms as an octahedral complex whereas $(\text{To}^M)_2\text{Fe}$ maintains a tetrahedral geometry with each $\text{To}^M$ ligand bound in a bidentate fashion. The coordinatively unsaturated $(\text{To}^M)_2\text{Fe}$ could explain its enhanced reactivity compared with $\text{Tp}_2\text{Fe}$. Importantly, this change in coordination environment indicates that the slightly more sterically hindered $\text{To}^M$ versus $\text{Tp}^{\text{Me}2}$ concluded from earlier studies of solid angles is significant enough to cause this change in preferential geometry.\(^{10}\)

Synthesis of $\text{To}^M\text{NiCl} (3)$ was similarly achieved by the reaction of $\text{TlTo}^M$ and $\text{NiCl}_2\cdot\text{DME} (\text{DME} = \text{dimethoxyethane})$ in THF. This reaction can be performed on a multi-gram scale, with respect to $\text{TlTo}^M$, affording $\text{To}^M\text{NiCl}$ as a pink crystalline solid in over 90% yield (equation (4)). An X-band EPR spectrum of 3 collected at 10K showed no signal. This result is in accord with literature reports of other nickel(II) complexes in a tetrahedral geometry which are EPR silent due to large zero-field splitting parameters.\(^{11}\) The UV-Vis spectrum of 1 displayed $\lambda_{\text{max}}$ at 310 ($\varepsilon$: 233 M$^{-1}$cm$^{-1}$), 478 ($\varepsilon$: 355 M$^{-1}$cm$^{-1}$), 817 ($\varepsilon$: 84 M$^{-1}$cm$^{-1}$), and 925 nm ($\varepsilon$: 115 M$^{-1}$cm$^{-1}$), which is similar to that reported for
tris(pyrazolyl)borate nickel(II) halide complexes (e.g. Ni\textsuperscript{II}(Cl)(Tp\textsubscript{Me2,Br}) in CH\textsubscript{2}Cl\textsubscript{2} $\lambda_{\text{max}} = 481$ ($\varepsilon$: 360 M\textsuperscript{-1} cm\textsuperscript{-1}), 801 ($\varepsilon$: 105 M\textsuperscript{-1} cm\textsuperscript{-1}), and 900 nm ($\varepsilon$: 120 M\textsuperscript{-1} cm\textsuperscript{-1})).\textsuperscript{12}

![Salt metathesis reaction of To\textsuperscript{M}NiCl with NaOAc proceeds facilely in THF at room temperature overnight. To\textsuperscript{M}NiOAc was isolated as a bright green solid (equation (5)). Single X-ray quality crystals of To\textsuperscript{M}NiOAc grown from a saturated pentane solution at $-40$ °C showed a tetrahedral geometry around the nickel center. However, despite attempts to form alkylated products To\textsuperscript{M}NiR (R = Me, CH\textsubscript{2}SiMe\textsubscript{3}, or Bn), no nickel(II) alkyls could be isolated and instead decomposition occurred.](image)

In contrast, To\textsuperscript{M}FeBr reacts with KBn to form To\textsuperscript{M}FeBn (5) as a yellow solid (equation (6)). Complex 5 was characterized by a single signal at $-225$ ppm in the $^{11}$B NMR spectrum which was the most downfield resonance observed for the iron(II) complexes reported here. As in To\textsuperscript{M}FeBr, the methyl and methylene signals of the oxazoline group in the To\textsuperscript{M} ligand were equivalent in the $^1$H NMR spectrum supporting tridentate coordination. The signal corresponding to the methyl resonances is shifted even further upfield ($-56.35$ ppm) relative to 1 ($-30$ ppm). The infrared spectrum (KBr) of 5 contained several notable features. Instead of a single $\nu_{\text{CN}}$ signal observed as expected for tridentate coordination of To\textsuperscript{M} to iron, two $\nu_{\text{CN}}$ signals were observed at
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1586 and 1552 cm\(^{-1}\). Given no \(\nu_{\text{CN}}\) signals appear at higher intensity than 1600 cm\(^{-1}\), the three oxazoline rings are clearly coordinated to iron. However, the presence of two rather than one \(\nu_{\text{CN}}\) band indicates that the oxazoline rings are not bound to iron in an equivalent manner. In addition, a distinct band at 1701 cm\(^{-1}\) was evident.

\[
\begin{align*}
\text{Ph-B}&\text{N}^\text{N-Fe-Br} + \text{KBN} & \text{THF} \\
\text{r.t., overnight} & -\text{KBr} & \text{Ph-B}&\text{N}^\text{N-Fe-Br} \\
\text{1} & \text{5} & 89\% \\
\end{align*}
\]

X-ray quality crystals of 5 revealed a tetrahedral iron center with the Fe-C interatomic distance of 2.067(3) Å only slightly longer than the analogous Co-C interatomic distance of 2.023(2) Å in To\(^{M}\)CoBn and similar to other organoiron compounds (e.g. Tp\(^{3Pr}\)Fe-CH\(_2\)C\(_6\)H\(_4\)Me-\(\text{Me-}\phi\)Fe-C 2.05(1) Å (Figure 2).\(^{13}\) The B-Fe-C angle was nearly linear (170°) but the benzyl ligand deviated further from linearity than in To\(^{M}\)FeBr where B-Fe-Br measures 174°.

**Figure 2.** Thermal ellipsoid plot of To\(^{\phi}\)FeBn (5) with ellipsoids at 20 % probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Fe1-C1 2.067(3), Fe1-N1 2.073(2), Fe1-N2 2.086(3), Fe1-N3 2.092(2); N1-Fe1-N2 90.0(1), N1-Fe1-N3 90.3(1), N2-Fe1-N3 88.6(1), N1-Fe1-C1 134.8(1), N2-Fe1-C1 121.6(1), N3-Fe1-C1 119.2(1), B1-Fe1-C1 170.0(1).
To\textsuperscript{M}FeBn reacts readily with CO to form To\textsuperscript{M}Fe\{C(O)Bn\}(CO)\textsubscript{2} (6); upon exposure of a solution of 5 to CO (atm) the solution maintains its yellow color but if allowed to stand a precipitate settles out (equation (7)). NMR spectroscopy analysis of the product revealed a new \textsuperscript{11}B NMR signal at –28 ppm and diamagnetic signals in the \textsuperscript{1}H NMR that can be assigned to an octahedral iron(II) configuration with three distinct signals visible for the methyl groups of the oxazoline ring. The absence of any paramagnetic signals in the \textsuperscript{1}H NMR spectrum suggests that species 6 is low-spin iron(II). The IR spectrum of 6 contained two equally intense signals at 2004 and 1935 cm\textsuperscript{-1} and two weaker signals at 1680 and 1662 cm\textsuperscript{-1}. The two higher energy bands are assigned as two terminal CO ligands coordinated to iron and the two lower energy bands suggest the presence of an acyl ligand that exists as two rotamers. In combination with the two coordinated CO ligands and acyl ligand, the IR spectrum indicates that all of the oxazoline rings of the To\textsuperscript{M} ligand are coordinated to iron with two \textnu_{\text{CN}} signals observed at 1593 and 1553 cm\textsuperscript{-1} as explained by To\textsuperscript{M} occupying a combination of equatorial and axial coordination sites.

Finally, X-ray crystallography confirmed the octahedral structural assignment of 6 (Figure 3). The reaction of To\textsuperscript{M}FeBn with CO to form To\textsuperscript{M}Fe\{C(O)Bn\}(CO)\textsubscript{2} is not unprecedented given similar carbonylated TpFe complexes have been isolated (e.g. HBpz\textsubscript{3}Fe\{C(O)Me\}(CO)\textsubscript{2} and \textit{t}-BuTp\textsuperscript{\textit{i}-Pr}Fe\{C(O)Me\}(CO)\textsubscript{2}).\textsuperscript{14-15} The Fe–C interatomic distances measure 1.753(7) and 1.753(6) Å for the carbonyl groups and 1.984(6) Å for
the acetyl moiety. These interatomic distances are similar to related compounds. For example, in HBpz₃Fe{C(O)Me}(CO)₂ the Fe–C interatomic distances are 1.758(6) and 1.771(5) Å for the carbonyl groups and 1.968(5) Å for the acetyl group. Similarly, as in HBpz₃Fe{C(O)Me}(CO)₂ where the longest Fe–N interatomic distance (2.082(4) Å) lies trans to the acetyl group, the Fe–N interatomic distance (2.097(4) Å) trans to the acetyl group is longer than the Fe–N interatomic distances (2.028(4) and 2.047(4) Å) which are trans to the carbonyls.

Figure 3. Thermal ellipsoid plot of To⁴Fe{C(=O)Bn}(CO)₂ (6) with ellipsoids at 20 % probability. Hydrogen atoms are omitted for clarity. Selected interatomic distances (Å) and angles (°): Fe1-C22 1.753(6), Fe1-C23 1.753(7), Fe1-C24 1.984(6), Fe1-N1 2.028(4), Fe1-N2 2.097(4), Fe1-N3 2.047(4), C22-O4 1.159(7), C23-O5 1.155(7), C24-O6 1.228(7); N2-Fe1-C22 97.6(2), C22-Fe1-C24 91.1(3), C24-Fe1-N3 86.8(2), N3-Fe1-N2 84.9(2), N1-Fe1-C23 174.7(2).
To\textsuperscript{M}Fe\{C(O)Bn\}(CO)\textsubscript{2} and O\textsubscript{2} reacts to form a species that has a \textsuperscript{11}B signal indistinguishable from 6 (–18 ppm) as well as similar signals in the \textsuperscript{1}H NMR at 1 and 3 ppm; however, an identifying feature in the \textsuperscript{1}H NMR at 10 ppm indicates a new product has formed. The \textsuperscript{1}H signal at 10 ppm typically signifies formation of HTo\textsuperscript{M}; however the \textsuperscript{1}H signals of the new species are shifted slightly upfield in comparison with the signals reported for HTo\textsuperscript{M}. In addition, if authentic HTo\textsuperscript{M} is added to a benzene-d\textsubscript{6} solution of the new species, the signals corresponding to the new species as well as a separate set of signals for HTo\textsuperscript{M} are observed further ruling out HTo\textsuperscript{M} as the product. The UV-Vis supports conversion of 6 to a new iron containing species with the characteristic broad band at 800 nm disappearing upon addition of O\textsubscript{2} to 6. The only absorption in the UV-Vis for the new species is a broad tailing absorption from 200 to 500 nm. Loss of the carbonyl and acetyl function groups in 4 is confirmed by IR spectroscopy where in the carbonyl and acetyl region only one weak band at 1700 cm\textsuperscript{-1} was observed for the new species. The direct reaction of To\textsuperscript{M}FeBn and O\textsubscript{2} forms the same product as the reaction of 4 with O\textsubscript{2} as evidenced by matching signals in the \textsuperscript{1}H NMR and \textsuperscript{11}B NMR spectra. The unexpected similarity between these two reactions suggests that the oxidative carbonylation to form carboxylate compounds, as found in the cobalt(II) alkyl system, does not occur in the analogous iron(II) alkyl complexes. Support for this conclusion was further gained by reaction of To\textsuperscript{M}FeBn with CO\textsubscript{2} to provide a product that was characterized by a new major signal at –24.27 ppm assigned to the methyl groups on the oxazoline ring on the basis of integration and a \textsuperscript{11}B signal at 35 ppm. The product is presumably To\textsuperscript{M}FeO\textsubscript{2}CBn and supports that this carboxylate compound is not forming in the reaction of To\textsuperscript{M}FeBn with CO followed by O\textsubscript{2}.
Conclusion

In synthesizing $\text{To}^\text{M} \text{FeBr}$, several striking differences with regards to cobalt were discovered. Primarily, solvent-free $\text{FeBr}_2$ readily provides spectroscopically and analytically pure $\text{To}^\text{M} \text{FeBr}$ whereas a solvent adduct of the cobalt chloride starting material, $\text{CoCl}_2 \cdot \text{THF}$ ($\text{THF} = \text{tetrahydrofuran}$), is required in order to effectively promote salt metathesis reaction with $\text{TITo}^\text{M}$ and to avoid formation of $(\text{To}^\text{M})_2 \text{Co}$. Interrelated to this first observation was the discovery that $(\text{To}^\text{M})_2 \text{Fe}$ also serves as precursor to $\text{To}^\text{M} \text{FeBr}$. Thus, reaction of $(\text{To}^\text{M})_2 \text{Fe}$ with $\text{FeBr}_2$ in THF at room temperature affords $1$ in THF. $\text{To}^\text{M} \text{NiCl}$ is also isolable and serves as precursor to $\text{To}^\text{M} \text{NiOAc}$; however, organonickel compounds could not be isolated. Complex $1$ readily undergoes salt metathesis reaction with KBN to provide $\text{To}^\text{M} \text{FeBn}$. Like similar TpFeBn complexes, $5$ reacts with CO to form a dicarbonyl acyl species. We are currently working to synthesize alternative iron(II) alkyl complexes supported by $\text{To}^\text{M}$ to more fully understand the reactivity of this class of organometallic compounds.

Experimental Section

**General Procedures.** All reactions were performed using standard Schlenk techniques under argon or in a glovebox under an inert atmosphere of nitrogen. Benzene and tetrahydrofuran were dried and deoxygenated using an IT PureSolv system. Benzene-$d_6$ was degased with freeze-pump-thaw cycles, heated to reflux over a Na/K alloy, and then vacuum transferred. $^1\text{H}$ and $^{11}\text{B}$ NMR spectra were collected on a Bruker Avance III 600 spectrometer. $^{11}\text{B}$ NMR spectra were referenced to an external sample of BF$_3 \cdot \text{Et}_2\text{O}$. Infrared spectra were measured on a Bruker Vertex 80 FTIR spectrometer. UV-Vis spectra were recorded on an Agilent 8453 UV-vis spectrophotometer. The electron
paramagnetic resonance (EPR) spectra were obtained with an X-band Elexsys 580 FT-EPR spectrometer in continuous-wave mode. Elemental analyses were performed using a Perkin-Elmer 2400 Series II CHN/S. Single crystal X-ray diffraction data was collected on an APEX II.

**ToMFeBr (1).** Tl[ToM] (0.200 g, 0.341 mmol) was dissolved in tetrahydrofuran (3 mL) and added to a suspension of FeBr₂ (0.081 g, 0.38 mmol) in tetrahydrofuran (10 mL). The cloudy white solution was stirred overnight and the solvent evaporated. The product was extracted using benzene and isolated *in vacuo* to afford ToMFeBr as a cream colored solid (0.113 g, 0.218 mmol, 63.9%). X-ray quality crystals were obtained from pentane at –40°C. ¹H NMR (benzene-d₆, 600 MHz): δ 24.36 (s, 2 H, C₆H₅), 20.09 (s, 6 H, CNCMe₂CH₂O), 11.97 (s, 1 H, p-C₆H₅), –28.93 (s, 18 H, CNCMe₂CH₂O). ¹¹B NMR (benzene-d₆, 128 MHz): δ 74. IR (KBr, cm⁻¹): ν 2965 (m), 2929 (w), 2897 (w), 2870 (w), 1586 (s, νCN), 1496 (w), 1461 (m), 1434 (w), 1387 (m), 1368 (m), 1351 (m), 1270 (s), 1191 (s), 1161 (m), 1099 (w), 1017 (w), 982 (w), 956 (s). UV-vis (THF) λₘₐₓ = 312 (ε: 597 M⁻¹cm⁻¹). µₑffective(C₆D₆) = 5.1(2) μB as determined by the Evans method. Anal. Calcd. for C₂₁H₂₉BBrFeN₃O₃: C, 48.69; H, 5.64; N, 8.11 Found: C, 48.42; H, 5.68; N, 8.00. Mp.238-241.

**(ToM)₂Fe (2).** Tl[ToM] (0.200 g, 0.341 mmol) was dissolved in tetrahydrofuran (3 mL) and added to a suspension of FeBr₂ (0.037 g, 0.17 mmol) in tetrahydrofuran (10 mL). The cloudy white solution was stirred overnight and the solvent evaporated. The product was extracted using benzene and isolated *in vacuo* to afford ToMFeBr as a white colored solid (0.130 g, 0.158 mmol, 93.2%). X-ray quality crystals were obtained from pentane at –40°C. ¹H NMR (benzene-d₆, 600 MHz): δ 20.16, 17.78, 16.59, 16.56, 8.05, 7.16, 7.05,
$^{11}$B NMR (benzene-$d_6$, 128 MHz): $\delta$ -122. IR (KBr, cm$^{-1}$): v 3069 (w), 3046 (w), 2966 (m), 2930 (m), 2881 (m), 1604 (m, $v_{CN}$), 1548 (s, $v_{CN}$), 1491 (w), 1463 (m), 1433 (w), 1370 (m), 1281 (m), 1250 (m), 1197 (m), 1151 (m), 1107 (w), 1027 (w), 1001 (m), 966 (m). UV-vis (Et$_2$O) $\lambda_{max} = 304$ ($\varepsilon$: 156 M$^{-1}$cm$^{-1}$), 581 ($\varepsilon$: 388 M$^{-1}$cm$^{-1}$), 617 ($\varepsilon$: 424 M$^{-1}$cm$^{-1}$), 697 ($\varepsilon$: 1078 M$^{-1}$cm$^{-1}$). $\mu_{eff}$ (C$_6$D$_6$) = 4.9(2) $\mu_B$ as determined by the Evans method. Anal. Calcd. for C$_{42}$H$_{58}$B$_2$FeN$_6$O$_6$: C, 61.49; H, 7.13; N, 10.24 Found: C, 61.35; H, 7.15; N, 10.05. Mp. 217–219°C, dec.

**To$^M$NiCl (3).** TiTo$^M$ (3.89 g, 6.64 mmol) dissolved in tetrahydrofuran (400 mL) was added to NiCl$_2$•DME (1.45 g, 6.60 mmol) and the mixture was stirred for 18 hours at room temperature. Tetrahydrofuran was removed in vacuo and the product extracted with methylene chloride (200 mL). Evaporation of methylene chloride afforded To$^M$NiCl as a pink solid (2.86 g, 6.00 mmol, 91%). Recrystallization from pentane gave analytically pure To$^M$NiCl. $^1$H NMR (benzene-$d_6$, 600 MHz): $\delta$ 16.94 (s, 6 H, CNCMe$_2$CH$_2$O), 7.59 and 7.49 (4 H, C$_6$H$_5$), 6.73 (s, 1 H, C$_6$H$_5$), -0.93 (s, 18 H, CNCMe$_2$CH$_2$O). $^{11}$B NMR (benzene-$d_6$, 128 MHz): $\delta$ -17. IR (KBr, cm$^{-1}$): v 2966 (m), 2933 (m), 2899 (m), 2861 (m), 1600 (s, $v_{CN}$), 1463 (m), 1388 (m), 1369 (m), 1353 (m), 1276 (m), 1195 (m) 1167 (m), 955 (m). UV-vis (THF): $\lambda_{max}$ 310 nm ($\varepsilon$: 233 M$^{-1}$cm$^{-1}$), 478 nm ($\varepsilon$: 355 M$^{-1}$cm$^{-1}$), 817 nm ($\varepsilon$: 84 M$^{-1}$cm$^{-1}$), 925 nm ($\varepsilon$: 115 M$^{-1}$cm$^{-1}$). Evans method: $\mu_{eff}$ (C$_6$D$_6$) = 3.3(1) $\mu_B$. Anal. Calcd. for C$_{23}$H$_{32}$BNiN$_3$O$_5$: C, 52.94; H, 6.14; N, 8.82 Found: C, 52.88; H, 6.11; N, 8.85. Mp. 267–267 °C.

**To$^M$NiOAc (4).** To$^M$NiCl (0.051 g, 0.107 mmol) dissolved in tetrahydrofuran (5 mL) was added to NaOAc (0.009 g, 0.110 mmol) and the mixture was stirred for 18 hours at room temperature. The reaction mixture was filtered and the solvent evaporated to afford
To\textsuperscript{M}NiOAc as a yellow solid (0.050 g, 0.100 mmol, 93%). Recrystallization from pentane gave analytically pure To\textsuperscript{M}NiOAc. \textsuperscript{1}H NMR (benzene-\textit{d}_6, 600 MHz): \(\delta\) 91.01 (s, 3 H, CoO\textsubscript{2}CCH\textsubscript{3}), 13.28 (s, 6 H, CNCMe\textsubscript{2}CH\textsubscript{2}O), 8.25 (s, 2 H, C\textsubscript{6}H\textsubscript{5}), 7.72 (s, 2 H, C\textsubscript{6}H\textsubscript{5}), 6.96 (s, 1 H, \(p\)-C\textsubscript{6}H\textsubscript{5}), and -1.51 (s, 18 H, CNCMe\textsubscript{2}CH\textsubscript{2}O).

\textsuperscript{11}B NMR (benzene-\textit{d}_6, 128 MHz): \(\delta\) -5.9. IR (KBr, cm\textsuperscript{-1}): \(\nu\) 2964 (m), 2928 (m), 2891 (m), 2853 (m), 1602 (s, \(\nu\)CN), 1535 (m), 1463 (m), 1387 (m), 1366 (m), 1353 (m), 1274 (m), 1198 (m) 1163 (m), 959 (m). UV-vis (CH\textsubscript{2}Cl\textsubscript{2}): \(\lambda_{\text{max}}\) 422 nm (\(\varepsilon\): 233 M\textsuperscript{-1}cm\textsuperscript{-1}), 704 nm (\(\varepsilon\): 75 M\textsuperscript{-1}cm\textsuperscript{-1}), 874 nm (\(\varepsilon\): 63 M\textsuperscript{-1}cm\textsuperscript{-1}). Evans method: \(\mu_{\text{eff}}\) (C\textsubscript{6}D\textsubscript{6}) = 3.1(3) \(\mu\)B. Anal. Calcd. for C\textsubscript{23}H\textsubscript{32}BNiN\textsubscript{3}O\textsubscript{5}: C, 55.25; H, 6.45; N, 8.40 Found: C, 55.03; H, 6.62; N, 8.34. Mp. 181-183 °C.

To\textsuperscript{M}FeBn (5). KBn (0.030 mg, 0.23 mmol) was dissolved in THF (3 mL) and added dropwise to a THF solution (10 mL) of To\textsuperscript{M}FeBr (0.100 g, 0.19 mmol). The yellow solution was stirred for 30 min. at room temperature. The solvent was removed \textit{in vacuo} and filtered with toluene to provide To\textsuperscript{M}FeBn as a yellow solid (0.090 g, 0.17 mmol, 89% yield). \textsuperscript{1}H NMR (benzene-\textit{d}_6, 600 MHz): \(\delta\) 41.70 (s, 2 H), 41.02 (s, 2 H), 21.42 (s, 2 H), -17.44 (s, 1 H), 13.60 (s, 7 H, CNCMe\textsubscript{2}CH\textsubscript{2}O), -17.61 (s, 1 H), -56.35 (s, 18 H, CNCMe\textsubscript{2}CH\textsubscript{2}O), -63.08 (s, 1 H).

\textsuperscript{11}B NMR (benzene-\textit{d}_6, 128 MHz): \(\delta\) -225. IR (KBr, cm\textsuperscript{-1}): \(\nu\) 3070 (w), 3048 (w), 2966 (m), 2928 (m), 2896 (w), 2871 (m), 1701 (m), 1586 (s, \(\nu\)CN), 1552 (m, \(\nu\)CN), 1493 (m), 1460 (m), 1433 (w), 1388 (m), 1367 (m), 1275 (m), 1198 (s), 1158 (m), 1027 (w), 1005 (m), 969 (s). UV-vis (THF) \(\lambda_{\text{max}}\) = 395 (\(\varepsilon\): 1739 M\textsuperscript{-1}cm\textsuperscript{-1}). \(\mu_{\text{eff}}\) (C\textsubscript{6}D\textsubscript{6}) = 5.4(2) \(\mu\)B as determined by the Evans method. Mp.218-220 °C.

To\textsuperscript{M}Fe\{C(O)Bn\}(CO)\textsubscript{2} (6). To\textsuperscript{M}FeBn (0.01 g, 0.02 mmol) was dissolved in benzene-\textit{d}_6 (0.5 mL), the headspace was removed \textit{in vacuo}, and then CO (1 atm) was introduced.
Benzene-$d_6$ was evaporated \textit{in vacuo} to afford $^{10}M\text{Fe}\{C(O)Bn\}(CO)_2$ as a yellow solid (0.01 g, 0.02 mmol, 91\% yield). X-ray quality crystals were obtained from pentane cooled to $-40$ °C. $^1\text{H}$ NMR (benzene-$d_6$, 600 MHz): $\delta$ 8.31, 8.30, 7.51, 7.33, 7.25, 7.22, 7.13, 7.11, 7.00, 4.48, 3.49, 3.41, 3.36, 1.15, 0.94, 0.87. $^{11}\text{B}$ NMR (benzene-$d_6$, 128 MHz): $\delta$ –18. IR (KBr, cm$^{-1}$): $\nu$ 3078 (w), 3041 (w), 3000 (m), 2969 (m), 2931 (m), 2881 (m), 2004 (s, $\nu_{\text{CO}}$), 1935 (s, $\nu_{\text{CO}}$), 1680 (m, $\nu_{\text{C}=\text{O}}$), 1662 (m, $\nu_{\text{C}=\text{O}}$), 1593 (s, $\nu_{\text{CN}}$), 1553 (m, $\nu_{\text{CN}}$), 1495 (w), 1460 (m), 1388 (w), 1369 (m), 1355 (m), 1284 (m), 1248 (m), 1199 (s), 1156 (m), 1070 (w), 1028 (w), 993 (m), 970 (s). UV-vis (THF) $\lambda_{\text{max}} = 776$ (e: 92 M$^{-1}$ cm$^{-1}$).

\textbf{References}


CHAPTER 6: EFFECTS OF BIRADICAL DEUTERATION ON THE PERFORMANCE OF DNP: TOWARDS BETTER PERFORMING POLARIZING AGENTS


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Abstract

We study the effects of the deuteration of biradical polarizing agents on the efficiency of dynamic nuclear polarization (DNP) via the cross-effect. To this end, we synthesized a series of bTbK and TOTAPol biradicals with systematically increased deuterium substitution. The deuteration increases the radical’s relaxation time, thus contributing to a higher saturation factor and larger DNP enhancement, and reduces the pool of protons within the so-called spin diffusion barrier. Notably, we report that full or partial deuteration leads to improved DNP enhancement factors in standard samples, but also slows down the build-up of hyperpolarization. Improvements in DNP enhancements factors of up to 70% and time savings of up to 38% are obtained upon full deuteration. It is foreseen that this approach may be applied to other DNP polarizing agents thus enabling further sensitivity improvements.
Frédéric A. Perras: Responsible for measuring the DNP enhancements for each of the biradicals studied.

Introduction

Dynamic nuclear polarization (DNP) is the most widely applicable hyperpolarization technique to enhance the sensitivity of nuclear magnetic resonance (NMR) experiments. In most DNP solid-state (SS)NMR applications, a sample is placed in contact with a source of unpaired electrons (usually an exogenous radical polarization agent) and is irradiated, at low temperature (<110 K), with high power microwaves near the electron Larmor frequency.\(^1\) Electron polarization can then be transferred to the nuclei by a series of mechanisms, the cross-effect generally being the most efficient. The cross-effect is a process involving three coupled spins: two electrons and one nucleus.\(^2-4\)

Large strides have been made in recent years towards the improvement of DNP for solids. Specifically, the development of stable high-power microwave sources (gyrotrons), and advances in probe technology for low temperature magic angle spinning (MAS) have enabled the application of DNP SSNMR to high magnetic field strengths (49.4 T).\(^5-12\) Another important line of inquiry, of relevance here, concerns the development of polarizing agents capable of generating larger NMR enhancement factors (ε) and operating at higher temperatures. A notable breakthrough was reached by Hu et al. who demonstrated that the cross-effect condition was easier to satisfy by tethering two nitrooxide radicals together in a single biradical molecule.\(^13,14\) Matsuki et al. then showed
that further improvement could be obtained by using a rigid linker, thus fixing the relative orientation of the two electrons’ g tensors.\textsuperscript{15,16} Most recently, Zagdoun \textit{et al.} demonstrated that even larger enhancements may be obtained by using a biradical with a high molecular weight.\textsuperscript{17-19} It was hypothesized that the lessened molecular motions in these compounds slow the electrons’ relaxation, thus enabling a higher electron saturation factor, and a larger DNP enhancement.\textsuperscript{17,20} These breakthroughs may be combined, thus multiplying the improvements available from each approach, as is the case for the TEKPol biradical.\textsuperscript{20} Note that such efforts may not be beneficial for DNP experiments performed at liquid helium temperatures, where the electron relaxation times can become much longer.\textsuperscript{21}

Instead of hindering the molecular motions that are responsible for the electrons’ relaxation, the size of the interactions contributing to relaxation may also be lessened. Given that dipolar coupling to protons, particularly those in rapidly rotating methyl groups, is a leading cause of electron relaxation under the DNP conditions, these interactions may be reduced by a factor of 6.5 by simply perdeuterating the polarizing agent.\textsuperscript{22,23} Perdeuteration also eliminates the $^1$H spins that are nearest to the radical. These nuclei possess short relaxation times and thus become a polarization sink. Since spin diffusion from these nuclei to the bulk is also slowed by the presence of pseudocontact shifts, they are said to be within the so-called spin diffusion barrier.\textsuperscript{24-26} As a result of deuteration, less electron polarization will be spent repolarizing these rapidly relaxing $^1$H spins. It is important to note that the use of deuterated solvent, or sample, is also known to lead to an increase in DNP enhancement factor.\textsuperscript{27-29}
Results and Discussion

We have prepared four versions of the benchmark organic soluble polarizing agent bTbK\textsuperscript{15} using reagents at natural abundance (bTbK-d\textsubscript{0}), a deuterated pentaerythritol linker (bTbK-d\textsubscript{8}), deuterated nitroxides (bTbK-d\textsubscript{32}), as well as perdeuterated reagents (bTbK-d\textsubscript{40}), see Figure 1.

![Biradical polarizing agents](image_url)

Figure 1. Biradical polarizing agents used in this study.
All isotopologues of bTbK were dissolved in a 96 : 4 tetrachloroethane (TCE) : methanol-\textit{d}_4 (CD\textsubscript{3}OD) solvent mixture to form a 16 mM solution, and 20 mL of this solution was pipetted into 3.2 mm sapphire rotors. The solutions were degassed by repeatedly inserting and ejecting the sample, as previously described,\textsuperscript{19,30} until the DNP enhancement reached a plateau. As shown in Figure 2 and Table 1, the maximum achievable DNP enhancements at 109 K, using a constant microwave power near 30 W, progressively increase with the level of deuteration, from 53 in the protonated bTbK to 91 in the fully perdeuterated version. This corresponds to a 70% increase in DNP enhancement by simply deuterating the polarizing agent.

\textbf{Figure 2.} (a) DNP enhancement of 16 mM 96:4 TCE:CD\textsubscript{3}OD solutions of bTbK-\textit{dn} radicals. The ‘MW off’ spectrum is shown on the bottom along with this same spectrum scaled by the enhancement factor listed in Table 1. The ‘MW on’ spectrum is shown on the top of the figure and dashed lines highlight the relative intensities of the spectra acquired using bTbK-\textit{d}0 and bTbK-\textit{d}40. A comparison of ‘MW off’ spectra of equal volume solutions of the non- (black) and perdeuterated (red) biradicals is shown in (b) highlighting the minute difference of depolarization for both radicals.
Surprisingly, the deuteration of the linker, containing only 8 hydrogen atoms, was particularly important, leading to a comparable improvement in enhancement per atom than the deuteration of the TEMPO moiety (1.4–2.1% vs. 1.5–1.7% higher enhancement per hydrogen atom). As shown in Table 1, this increase in DNP enhancement is also accompanied by an increase in the DNP buildup time ($T_{\text{DNP}}$) from 3.5 s in the nonlabeled compound to 7.8 s in the perdeuterated compound. Because the signal to noise increases linearly as a function of the square of the number of scans, the relative sensitivity per unit of time ($e^2/T_{\text{DNP}}$) is still 30% higher when using the perdeuterated radical. $^{13}$C and $^{29}$Si DNP-enhanced NMR experiments performed on 3-(3-phenylureido)propyl-functionalized mesoporous silica nanoparticles (PUP-MSNs) in fact demonstrate that larger enhancement factors can also be obtained on solid samples of interest, although in PUP-MSN the relative sensitivity per unit of time remains
approximately the same. As can be seen in Figure 2b and 3b, the extent of the signal loss, thought to be related to both blanking and MAS-induced depolarization, does not increase by a measurable amount for the deuterated biradicals.\textsuperscript{31}

To confirm that this is indeed a generally applicable strategy for improving the effectiveness of other DNP polarization agents we have also synthesized and tested deuterated versions of the hydrophilic TOTAPol biradical.\textsuperscript{1b} Natural abundance (TOTAPol-d0), perdeuterated (TOTAPol-d40), as well as a biradical with deuterium labeling on only the amine-terminated nitroxide (TOTAPol-d17) were prepared, see Figure 1. These radicals were then dissolved in 60:30:10 glycerol-$d_8$:D$_2$O:H$_2$O (the so-called ‘DNP juice’) at a concentration of 10 mM as well as in a deuterium-free 60 : 40 mixture of glycerol and H$_2$O.\textsuperscript{32} The DNP enhancement measurements on these solutions were performed following the protocol described for the bTbK radicals.

As shown in Figure 3a and Table 1, the DNP enhancement increases upon deuteration of TOTAPol from 67 to 85 for the d0 and d17 derivatives, respectively, when using DNP juice. The fully deuterated biradicals, TOTAPol-d40, yields the same enhancement as partly-deuterated TOTAPol-d17, within uncertainty in the estimates (5%). Since the DNP build-up time increases from 8.5 to 11.0 and 23.5 s upon deuteration, the sensitivity per time is affected accordingly; the reason for this will be expanded later in the text. As was expected, however, the enhancement increases steadily from 39 to 51 and 59 for the d0, d17, and d40 derivatives, respectively, when using the fully protonated solvent. The enhancement factor for TOTAPol then increases by 50% upon perdeuteration and the sensitivity per unit of time increases by 38%.
Figure 3. DNP enhancement of 10 mM 60:30:10 glycerol-d8: D2O: H2O (a) and 60:40 glycerol:H2O (c) solutions of various TOTAPol radicals. The ‘MW off’ spectrum is shown on the bottom along with this same spectrum scaled by the enhancement factor listed in Table 1. The ‘MW on’ spectrum scaled by the enhancement factor listed in Table 1. The ‘MW on’ spectrum is shown on the top of the figure and dashed lines highlight the relative intensities of the spectra acquired using TOTAPol-d0 and TOTAPol-d40. A comparison of ‘MW off’ spectra of equal-volume solutions of the non- (black) and per-deuterated (red) biradicals is shown in (b and d) highlighting the minute difference of depolarization for both radicals.

Using a model spin system consisting of two electrons and one nucleus, Thurber and Tycko have shown theoretically that long T1e values prevent the loss of magnetization between the MAS-induced frequency crossing events that lead to cross-effect polarization transfer. The subsequent investigation of a series of functionalized nitroxide biradicals by Zagdoun et al. confirmed that electron relaxation properties were highly correlated with the DNP enhancements. Thus, both theoretical and experimental studies of the cross effect under MAS demonstrate that longer T1e relaxation times of the electrons generally lead to higher enhancement factors; as was mentioned earlier,
this was the main motivation for the synthesis of the perdeuterated biradicals by the present authors. As can be seen from the plots shown in Figure 4, the $T_{1e}$ and $\varepsilon$ values indeed increase linearly as the biradical polarizing agents are progressively deuterated. The deuteration of the solvent, on the other hand, does not lead to an increase of $T_{1e}$.\textsuperscript{35} We also note that, for bTbK, the observed increase in DNP enhancement is in agreement with the numerical calculations of Mance et al. for the increase in $T_{1e}$ that we observed.\textsuperscript{33} However, for TOTAPol, the modest increase in $T_{1e}$ does not appear to fully explain the observed 50\% increase in DNP enhancement factor.\textsuperscript{33} This may be the result of the use of a much lower magnetic field for the EPR measurements (9 vs. 263 GHz). Another plausible mechanism contributing to an increase in $\varepsilon$ values would be that, as mentioned earlier, there is a diminished loss of polarization within the so-called spin diffusion barrier when perdeuterated radicals are used.\textsuperscript{24} The elimination of $^1$H spins within the spin diffusion barrier also, unfortunately, has the effect of increasing the $^1$H DNP build-up time due to the reduced efficiency of DNP-assisted spin diffusion (see Table 1).\textsuperscript{36} Hovav and coworkers demonstrated that the rate at which hyperpolarization is transferred to the bulk nuclei depends on the strength of the dipolar coupling between the core nuclei and the electron and depends very little on the dipolar coupling between the bulk nuclei. The deuteration of the radicals then reduces the dipolar coupling between the core nuclei and the electrons and essentially slows down the spin diffusion of hyperpolarization to the bulk. In extreme cases when deuterated solvents are used this can lead to particularly long $T_{\text{DNP}}$ values and a loss of sensitivity.
Figure 4. (a) The DNP enhancement factors of various bTbK and TOTAPol radicals are plotted as a function of the deuteration level of the molecule. The linear fits correspond to $e = 52.1 \pm 0.36$ D for bTbK and $e = 40.2 \pm 0.21$ D, where D corresponds to the % deuteration of the molecule. The correlation between the deuteration levels of a biradical and the $T_{1e}$ values are plotted in (b) and (c) for bTbK and TOTAPol, respectively. In (c) the black circles correspond to the use of fully protonated solvent whereas the red squares correspond to the data acquired when using a 90% deuterated solvent.

Conclusion

In conclusion, we have explored a simple approach for improving the DNP enhancement afforded by biradical polarizing agents. Indeed, using the bTbK and TOTAPol biradicals, an increase in DNP enhancement factors of up to 70%, and an
improved sensitivity per unit of time of up to 38% can be obtained by simply perdeuterating the polarizing agent. The deuteration of biradicals increases their $T_{1e}$ values, thus improving the saturation factors upon continuous wave microwave irradiation, and consequently contributing to higher enhancement factors. The use of deuterated polarizing agents also reduces the number of $^1H$ spins within the spin diffusion barrier, which has the effects of reducing the polarization loss, but also leads to slower DNP-assisted spin diffusion and longer build-up times.$^{36}$ The perdeuteration, or partial deuteration, of the current state-of-the-art polarizing agents (AMUPol$^{18}$ and TEKPol$^{19}$), which can already yield enhancement factors over 200 at 105 K, is likely to yield the next generation of biradicals giving enhancement factors well over 300; we are actively exploring this avenue. The results of this study may also help in understanding the mechanisms of the cross effect in DNP of solids, particularly with respect to the polarization of remote spins.

**Experimental Section**

**EPR Spectroscopy.** The $T_{1e}$ measurements were performed on an X-band Elexsys 580 FT- EPR spectrometer by using an inversion recovery method at a temperature of 100 K. The $\pi/2$ and $\pi$ pulse durations used for the relaxation measurements were 16 ns and 32 ns, respectively. As had been previously described,$^{37}$ the relaxation process in these samples is not monoexponential and is better represented by a stretched exponential function:

$$I(t) = I_0 + I_1 e^{-\left(\frac{t}{T_{1e}^*}\right)^\beta}.$$  

In the expression above, $I_0$ is the initial intensity, $I_1$ is a proportionality constant, $T_{1e}^*$ is a
decay constant, and $\beta$ is the stretching parameter. The $T_{1e}$ values mentioned in the text are in fact the mean $T_{1e}$ values, which correspond to the first moment of the stretched exponential decay. These are calculated as:

$$\langle T_{1e} \rangle = \frac{T_{1e}^*}{\beta} \Gamma\left(\frac{1}{\beta}\right).$$

The $T_{1e}^*$ and $\beta$ fitting parameters are listed in Table 2 below.

**Table 2.** Fitting parameters for the $T_{1e}$ measurements of the radicals described in the text.

<table>
<thead>
<tr>
<th>polarizing agent</th>
<th>solvent</th>
<th>$T_{1e}^*$ / µs</th>
<th>$\beta$</th>
<th>$\langle T_{1e} \rangle$ / µs</th>
</tr>
</thead>
<tbody>
<tr>
<td>bTbK</td>
<td>96:4</td>
<td>26.8</td>
<td>0.76</td>
<td>35</td>
</tr>
<tr>
<td>bTbK-d8</td>
<td>TCE : CD$_2$OD</td>
<td>40.8</td>
<td>0.66</td>
<td>62</td>
</tr>
<tr>
<td>bTbK-d32</td>
<td>46.8</td>
<td>0.54</td>
<td>87</td>
<td></td>
</tr>
<tr>
<td>bTbK-d40</td>
<td>69.4</td>
<td>0.57</td>
<td>121</td>
<td></td>
</tr>
<tr>
<td>TOTAPol-d0</td>
<td>60:30:10</td>
<td>71.8</td>
<td>0.60</td>
<td>120</td>
</tr>
<tr>
<td>TOTAPol-d17</td>
<td>glycerol-d8</td>
<td>90.7</td>
<td>0.69</td>
<td>131</td>
</tr>
<tr>
<td>TOTAPol-d40</td>
<td>D$_2$O : H$_2$O</td>
<td>111.3</td>
<td>0.73</td>
<td>153</td>
</tr>
<tr>
<td>TOTAPol-d0</td>
<td>60:40 glycerol</td>
<td>76.50</td>
<td>0.62</td>
<td>124</td>
</tr>
<tr>
<td>TOTAPol-d17</td>
<td>: H$_2$O</td>
<td>86.5</td>
<td>0.67</td>
<td>128</td>
</tr>
<tr>
<td>TOTAPol-d40</td>
<td>103.3</td>
<td>0.68</td>
<td>153</td>
<td></td>
</tr>
</tbody>
</table>

**General Procedure:** All deuterated chemicals were purchased from C/D/N Isotopes Inc. with the exception of pentaerythritol-d12, which was purchased from Cambridge Isotope Laboratories, Inc. Acetonitrile was dried by heating to reflux over calcium hydride followed by distillation. All other chemicals were used as received from suppliers. $^1$H and $^{13}$C{$^1$H} NMR spectra were collected on a Bruker AVII 600 spectrometer. Infrared spectra were measured on a Bruker Vertex 80 FTIR spectrometer. Accurate mass measurements were achieved using an Agilent QTOF 6540 mass spectrometer.

**Experimental Procedures:**

Preparation of the diamine 2,2,4,4,14,14,16,16-octamethyl-7,11,18,21-tetraoxa-3,15-diazatrispiro[5.2.2.5^12.2^9.2^6]henicosane followed a combination of the methods
previously reported by Matsuki and Griffin and later by Thankamony, Lafon, and Polshettiwar.\textsuperscript{38,39} Synthesis of the bTbK series followed the method reported by Matsuki and Griffin.\textsuperscript{38} Synthesis of the epoxide precursor to TOTAPOL followed the procedure reported by Zhang whereas the synthesis of TOTAPOL from the epoxide followed the procedure reported by Song and Griffin.\textsuperscript{40, 41} All of the syntheses were performed on a significantly smaller scale than what was previously reported. While the reactions proved to be scalable, the yields of the diamines were variable and this is attributed to the low scale of these reactions. The conditions for the deuterated syntheses were never optimized due to the cost of the isotopically labeled starting materials and the nature of the DNP experiments, which only require milligram quantities of the polarizing agent.

\textbf{Synthesis of } 2,2,4,4,14,14,16,16-\textit{octamethyl-7,11,18,21-tetraoxa-3,15-diazatrispiro-[5.2.2.5.12.2.9.2.6]henicosane.}

Pentaerythritol (0.237 g, 1.60 mmol), 2,2,6,6-\textit{tetramethyl-4-piperidone} (0.500 g, 3.22 mmol) and \textit{p}-toluenesulfonic acid (0.730 g, 3.84 mmol) were mixed together in toluene (15 mL) and heated to reflux for 12 hours in a Dean-Stark trap. The solution was cooled to room temperature and then washed with 2.0 M NaOH solution (2 \times 10 mL) and brine (1 \times 10 mL). The organic layer was dried over Na\textsubscript{2}SO\textsubscript{4} and filtered, and the solvent was removed under vacuum. Recrystallization from diethyl ether afforded white crystals (0.418 g, 1.02 mmol, 64\%). The \textsuperscript{1}H and \textsuperscript{13}C{\textsuperscript{1}H} NMR spectra match the reported literature values.\textsuperscript{39} \textsuperscript{1}H NMR (600 MHz, CDCl\textsubscript{3}): \textit{\delta} 3.76 (s, 8 H), 1.68 (s, 8 H), 1.21 (s, 24 H). \textsuperscript{13}C{\textsuperscript{1}H} NMR (150 MHz, CDCl\textsubscript{3}): \textit{\delta} 99.65, 63.56, 51.18, 43.08, 32.64, 32.24. IR (KBr, cm\textsuperscript{-1}): \textit{\nu} 3337, 2954, 2919, 2856, 1487, 1461, 1445, 1420, 1377, 1356, 1311, 1245,
Synthesis of 2,2,4,4,14,14,16-octamethyl-7,11,18,21-tetraoxa-3,15-diazatrispiro-
[5.2.2.5\textsuperscript{12},2.9.2.6]henicosane-8,8,10,10,19,19,20,20-d8 (diamine-d8).

Pentaerythritol-d\textsubscript{12} (0.237 g, 1.60 mmol), 2,2,6,6-tetramethyl-4-piperidone (0.500 g, 3.22 mmol) and \textit{p}-toluenesulfonic acid (0.730 g, 3.84 mmol) were mixed together in toluene (15 mL) and heated to reflux for 12 hours in a Dean-Stark apparatus with azeotropic removal of water. The solution was cooled to room temperature and then washed with 2.0 M NaOH solution (2 \times 10mL) and brine (1 \times 10 mL). The organic layer was dried over Na\textsubscript{2}SO\textsubscript{4} and filtered, and the solvent was removed under vacuum. Recrystallization from diethyl ether afforded white crystals (0.105 g, 0.25 mmol, 16%). \textsuperscript{1}H NMR (600 MHz, CDCl\textsubscript{3}): \( \delta \) 1.67 (s, 8 H), 1.21 (s, 24 H). \textsuperscript{13}C{\textsuperscript{1}H} NMR (150 MHz, CDCl\textsubscript{3}): \( \delta \) 99.56, 62.76 (br), 51.14, 43.11, 32.25, 31.95. IR (KBr, cm\textsuperscript{-1}): \( \nu \) 3553, 3499, 3021, 2927, 2864, 2229, 2095, 1652, 1488, 1448, 1424, 1375, 1362, 1314, 1240, 1200, 1155, 1086, 1034, 1018, 1001, 955. Mp. 131-133 °C. ESI-MS: m/z Calcd. for C\textsubscript{23}H\textsubscript{34}D\textsubscript{8}N\textsubscript{2}O\textsubscript{4} [M+H]\textsuperscript{+}: 419.37; Found: 420.3782.

Synthesis of 2,2,4,4,14,14,16-octakis(methyl-d\textsubscript{3})-7,11,18,21-tetraoxa-3,15-
diazatrispiro[5.2.2.5\textsuperscript{12},2.9.2.6]henicosane-1,1,5,5,13,13,17,17-d8 (diamine-d\textsubscript{32}).

Pentaerythritol (0.197 g, 1.45 mmol), 4-oxo-2,2,6,6-tetramethylpiperidine-d\textsubscript{17} (0.500 g, 2.90 mmol) and \textit{p}-toluenesulfonic acid (0.662 g, 3.48 mmol) were mixed together in toluene (15 mL) following the procedure described for the unlabeled or partially labeled diamine described above to afford 0.489 g (1.10 mmol, 76%) of diamine-d\textsubscript{32}. \textsuperscript{1}H NMR (600 MHz, CDCl\textsubscript{3}): \( \delta \) 3.76 (s, 8 H). \textsuperscript{13}C{\textsuperscript{1}H} NMR (150 MHz, CDCl\textsubscript{3}): \( \delta \) 99.62, 63.57,
50.62, 43.00, 42.60, 32.62, 31.15 (br). IR (KBr, cm\(^{-1}\)): \(\nu\) 3553, 3499, 2949, 2870, 2253, 2214, 2123, 2061, 1652, 1473, 1381, 1351, 1261, 1194, 1166, 1086, 1067, 907. Mp. 114-116 °C. ESI-MS: m/z Calcd. for \(\text{C}_{23}\text{H}_{10}\text{D}_{32}\text{N}_{2}\text{O}_{4}\) \([\text{M+H}]^+\): 442.79; Found: 439.4974. The experimentally measured molecular mass distribution indicates there is a small amount of proton exchange occurring and a fraction of the diamine-d32 contains a decreased level of deuteration. This proton exchange occurs at the acidic methylene position of the TEMPO moiety via the enolization of 4-oxo-2,2,6,6-tetramethylpiperidine-d17, and this position provides a small signal in the \(^1\)H NMR spectrum.

**Synthesis of 2,2,4,4,14,14,16,16-octakis(methyl-d3)-7,11,18,21-tetraoxa-3,15-diazatrispiro[5.2.2.5\(^{12}\).2\(^8\).2\(^6\)]henicosane-1,1,5,5,8,8,10,10,13,13,17,17,19,19,20,20-d16 (diamine-d40).**

Pentaerythritol-d12 (0.107 g, 0.72 mmol), 4-oxo-2,2,6,6-tetramethylpiperidine-d17 (0.250 g, 1.45 mmol), and \(p\)-toluenesulfonic acid (0.360 g, 1.89 mmol) were allowed to react following the procedure described above to provide tan colored crystals of diamine-d40 (0.127 g, 0.28 mmol, 39%). \(^{13}\)C\({}^{1}\)H\} NMR (150 MHz, CDCl\(_3\)): \(\delta\) 99.48, 62.69 (br), 50.59, 42.98, 42.58 (br), 31.89, 31.16 (br). IR (KBr, cm\(^{-1}\)): \(\nu\) 3551, 3497, 2961, 2927, 2850, 2214, 2096, 2070, 1654, 1472, 1454, 1349, 1316, 1286, 1263, 1238, 1196, 1175, 1154, 1103, 1071, 1040, 1024, 950. Mp. 112-115 °C. ESI-MS: m/z Calcd. for \(\text{C}_{23}\text{H}_{2}\text{D}_{40}\text{N}_{2}\text{O}_{4}\) \([\text{M+H}]^+\): 450.84; Found: 448.5544. As observed for diamine-d32, the experimentally measured molecular mass indicates there is proton exchange occurring resulting in a decreased level of deuteration.
**Synthesis of bTbK-d0.**

A mixture of the diamine-d0 (0.107 g, 0.261 mmol), sodium tungstate dehydrate (0.010 g, 0.030 mmol), and methanol (10 mL) was cooled to 0 °C. Hydrogen peroxide solution (30% in water, 128 µL, 1.28 mmol) was added in a dropwise fashion. The reaction mixture was allowed to warm to room temperature and was stirred for 20 hours. Potassium carbonate (~0.3 g) was added, and the mixture was filtered. The filtrate was dried over magnesium sulfate, the volatiles were evaporated, and the product was purified by silica gel chromatography (hexanes:ethyl acetate = 4:5) to afford bTbK as an orange colored solid (0.076 g, 0.172 mmol, 66%). IR (KBr, cm⁻¹): ν 2976, 2935, 2864, 1471, 1453, 1377, 1344, 1243, 1191, 1167, 1110, 1085, 1024, 1008. Mp. 167-169 °C. ESI-MS: m/z Calcd. for C₂₃H₄₀N₂O₆²⁺ [M+H]⁺: 441.58; Found: 441.2953.

**Synthesis of bTbK-d8.**

The procedure described for bTbK-d0 was used with a mixture of the diamine-d8 (0.058 g, 0.138 mmol), sodium tungstate dihydrate (0.023 mg, 0.070 mmol), and methanol (0.9 mL) to afford bTbK-d8 as an orange colored solid (0.031 g, 0.068 mmol, 57%). IR (KBr, cm⁻¹): ν 3566, 3318, 2935, 2870, 2235, 2099, 1728, 1633, 1476, 1363, 1346, 1239, 1195, 1094, 1034, 999. Mp. 174-176 °C. ESI-MS: m/z Calcd. for C₂₃H₃₂D₈N₂O₆²⁺ [M+H]⁺: 449.63; Found: 450.3528.

**Synthesis of bTbK-d32.**

The procedure described for bTbK-d0 was used with a mixture of the diamine-d32 (0.100 g, 0.225 mmol), sodium tungstate dihydrate (0.009 mg, 0.027 mmol), and methanol (10 mL) to afford bTbK-d32 as an orange colored solid (0.039 g, 0.082 mmol, 36%). IR (KBr, cm⁻¹): ν 2982, 2933, 2865, 2228, 1462, 1386, 1355, 1252, 1199, 1181, 1168, 1067,
Synthesis of bTbK-d40.

The procedure described for bTbK-d0 was used with a mixture of the diamine-d40 (0.050 g, 0.110 mmol), sodium tungstate dihydrate (0.018 mg, 0.055 mmol), and methanol (0.75 mL) to afford bTbK-d40 as an orange colored solid (0.031 g, 0.065 mmol, 59%). IR (KBr, cm\(^{-1}\)): \(\nu\) 3564, 3488, 3317, 2963, 2935, 2241, 2228, 1743, 1353, 1329, 1299, 1261, 1236, 1186, 1139, 1094, 1076, 1042, 1021. Mp. 158-161 °C. ESI-MS: m/z Calcd. for \(\text{C}_{23}\text{H}_{32}\text{D}_{40}\text{N}_{2}\text{O}_{6}\)\(^2\)\([\text{M+H}]^+\): 481.83; Found: 478.5280.

Synthesis of 4-(2,3-Epoxypropoxy)-2,2,6,6-tetramethyl-1-piperidin-1-oxyl (epoxide-d0).

Epichlorohydrin (2.25 mL, 28.7 mmol), 4-hydroxy-TEMPO (0.250 g, 1.45 mmol) and tetrabutylammonium hydrogen sulfate (0.029 g, 0.085 mmol) were added to a 90% w/w aqueous NaOH solution (0.232 g NaOH/0.263 mL H\(_2\)O). The mixture was stirred at 30 °C for 4 hours, at which point the mixture was poured into ice water and the product was extracted with diethyl ether (3 \(\times\) 10 mL). The organic layer was washed with brine solution, dried over Na\(_2\)SO\(_4\), and concentrated in vacuo. The product was purified by silica gel chromatography (hexanes:ethyl acetate = 4:5) to obtain a red oil (0.119 g, 0.521 mmol, 36%). The infrared spectrum of the product matched the literature report.\(^{40}\) IR (KBr, cm\(^{-1}\)): \(\nu\) 2976, 2938, 2869, 1635, 1465, 1378, 1363, 1245, 1179, 1096. ESI-MS: m/z Calcd. for \(\text{C}_{12}\text{H}_{22}\text{NO}_3\)\(^-\)\([\text{M+H}]^+\): 229.16; Found: 229.1676.
Synthesis of 1-(2,2,6,6-Tetramethyl-1-oxy-4-piperidinyl)oxy-3-(2,2,6,6-tetramethyl-1-oxy-4-piperidinyl)amino-propan-2-ol (TOTAPol-d0).

Under an inert atmosphere of nitrogen, the epoxide-d0 (0.100 g, 0.438 mmol) and LiClO₄ (0.047 g, 0.442 mmol) were dissolved in acetonitrile (0.8 mL). A solution of 4-amino-TEMPO (0.074 g, 0.432 mmol) dissolved in acetonitrile (0.2 mL) was added in a dropwise fashion. The solution was stirred overnight, concentrated, and purified by silica gel chromatography (hexanes:ethyl acetate = 4:5) to obtain a red solid (0.084 g, 0.210 mmol, 49%). The infrared spectrum of the product matched the literature report.¹⁰

IR (KBr, cm⁻¹): ν 3424, 2976, 2938, 2867, 1635, 1466, 1364, 1244, 1179, 1112, 1089. ESI-MS: m/z Calcd. for C₂₁H₄₁N₃O₄ • [M+H]⁺: 400.32; Found: 400.3169.

Synthesis of 4-(2,3-Epoxypropoxy)-2,2,6,6-tetramethyl-1-piperidin-1-oxyl (epoxide-d22).

Epichlorohydrin-d₅ (1.0 g, 10.3 mmol), 4-hydroxy-TEMPO-d₁₇ (0.122 g, 0.644 mmol) and tetrabutylammonium hydrogen sulfate (0.013 g, 0.038 mmol) were added to a 90% w/w aqueous NaOH solution (0.101 g NaOH/0.117 mL H₂O). The mixture was stirred at 30°C for 4 hours, at which point the mixture was poured into ice water and the product was extracted with diethyl ether (3 × 10 mL). The organic layer was washed with brine solution, dried over Na₂SO₄, and concentrated in vacuo. The product was purified by silica gel chromatography (hexanes:ethyl acetate = 4:5) to obtain a red oil (0.086 g, 0.343 mmol, 53%). ESI-MS: m/z Calcd. for C₁₂D₂₂NO₃⁺ [M+H]⁺: 251.30; Found: 251.3057.
Synthesis of 1-(2,2,6,6-Tetramethyl-1-oxy-4-piperidinyl)oxy-3-(2,2,6,6-tetramethyl-1-oxy-4-piperidinyl)amino-propan-2-ol (TOTAPol-d17).

Under an inert atmosphere of nitrogen, the epoxide-d0 (0.045 g, 0.197 mmol) and LiClO$_4$ (0.021 g, 0.197 mmol) were dissolved in acetonitrile (0.3 mL). A solution of 4-amino-TEMPO-d$_{17}$ (0.038 g, 0.202 mmol) dissolved in acetonitrile (0.1 mL) was added in a dropwise fashion. The solution was stirred overnight, concentrated, and purified by silica gel chromatography (hexanes:ethyl acetate = 4:5) to obtain a red solid (0.018 g, 0.043 mmol, 22%). IR (KBr, cm$^{-1}$): $\nu$ 3435, 2975, 2922, 2851, 2235, 1632, 1467, 1383, 1367, 1261, 1178, 1099, 1054. ESI-MS: m/z Calcd. for C$_{21}$H$_{24}$D$_{17}$N$_{3}$O$_{4}$• [M+H]$^+$: 417.42; Found: 417.4241.

Synthesis of 1-(2,2,6,6-Tetramethyl-1-oxy-4-piperidinyl)oxy-3-(2,2,6,6-tetramethyl-1-oxy-4-piperidinyl)amino-propan-2-ol (TOTAPol-d40).

Under an inert atmosphere of nitrogen, the epoxide-d22 (0.047 g, 0.188 mmol) and LiClO$_4$ (0.020 g, 0.188 mmol) were dissolved in acetonitrile (0.3 mL). A solution of 4-amino-TEMPO-d$_{17}$ (0.035 g, 0.186 mmol) dissolved in acetonitrile (0.5 mL) was added in a dropwise fashion. The solution was stirred overnight, concentrated, and purified by silica gel chromatography (hexanes:ethyl acetate = 4:5) to obtain a red solid (0.042 g, 0.095 mmol, 51%). IR (KBr, cm$^{-1}$): $\nu$ 3437, 2962, 2929, 2768, 2232, 2107, 1735, 1630, 1436, 1368, 1257, 1180, 1109, 1054. ESI-MS: m/z Calcd. for C$_{21}$H$_{24}$D$_{40}$N$_{3}$O$_{4}$• [M+H]$^+$: 440.82; Found: 439.5619.
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CHAPTER 7: CONCLUSION

General Conclusions

Research based on the iron triad consisting of iron, cobalt, and nickel is of particular interest given these metal’s usefulness in both commercial and biological applications. In addition, they have the benefit of being relatively abundant in the earth’s crust compared with other metals. This thesis demonstrates $TOM$ as a suitable ancillary ligand for isolation and study of iron, cobalt, and nickel complexes.

Synthesis of paramagnetic $TOM\text{MX} (\text{M,X} = \text{Fe,Br}; \text{Co,Cl}; \text{Ni,Cl})$ was achieved by reaction of $\text{TITOM}$ with the corresponding metal halide. The synthesis of $TOM\text{CoCl}$ was the most challenging requiring careful control of reaction conditions to exclude formation of the side products ($TOM$)$_2\text{Co}$ and $\text{HTOMCoCl}_2$. The synthesis of $TOM\text{FeBr}$ and $TOM\text{NiCl}$ was more straightforward and formed selectively under various reaction conditions. $(\text{TOM})_2\text{Ni}$ was not identified even in attempted independent syntheses. On the other hand, $(\text{TOM})_2\text{Fe}$ can be readily independently synthesized. Interestingly, $(\text{TOM})_2\text{Fe}$ reacts with $\text{FeBr}_2$ to form $TOM\text{FeBr}$.

$TOM\text{CoCl}$ reacts with NaOAc to provide $TOM\text{CoOAc}$. The catalytic activity of $TOM\text{CoOAc}$ in oxidation of hydrocarbons was studied using cyclohexane as the substrate and $m\text{CPBA}$ as the oxidant and found to be selective towards cyclohexanol over the over-oxidized products of cyclohexane and $\varepsilon$-caprolactone.

Organometallic cobalt(II) and iron(II) complexes were found to be accessible through salt metathesis reactions $TOM\text{CoCl}$ and $TOM\text{FeBr}$ and alkyl lithium or benzyl
potassium. Reactions involving Grignard reagents instead resulted in transmetalation. 

$\text{To}^M\text{CoR} \ (R = \text{Me, Et, } \text{^nBu, CH}_2\text{SiMe}_3, \text{Bn, and Ph})$ were found to react with CO followed by $O_2$ to form the corresponding carboxylate compounds. The oxidative carbonylation reactivity observed for these cobalt(II) alkyl complexes was found to be unique to cobalt, thus, similar reactivity was not observed for $\text{To}^M\text{FeBn}$. $\text{To}^M\text{FeBn}$ reacts with CO to form low-spin, diamagnetic $\text{To}^M\text{Fe}\{\text{C(O)Bn}\}(\text{CO})_2$ as a yellow solid.

Lastly, deuterated biradicals were synthesized and found to result in a larger DNP enhancement as a result of the radical’s longer relaxation time. Consequently, the results suggest that deuteration of biradicals is a promising approach to achieving larger DNP enhancements and thus increase the sensitivity of nuclear magnetic resonance (NMR) experiments.