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Facile C=O Bond Splitting of Carbon Dioxide Induced by Metal–Ligand Cooperativity in a Phosphinine Iron(0) Complex

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Abstract: New iron complexes $[\text{Cp}^*\text{FeL}]^-$ (**1-σ** and **1-π**, $\text{Cp}^* = \text{C}_5\text{Me}_5$) containing the chelating phosphinine ligand 2-(2'-pyridyl)-4,6-diphenylphosphinine (**L**) have been prepared, and found to undergo facile reaction with CO₂ under ambient conditions. The outcome of this reaction depends on the coordination mode of the versatile ligand **L**. Interaction of CO₂ with the isomer **1-π**, in which **L** binds to Fe through the phosphinine moiety in an η^5 fashion, leads to the formation of **3-π**, in which CO₂ has undergone electrophilic addition to the phosphinine group. In contrast, interaction with **1-σ**—in which **L** acts as a σ -chelating [P,N] ligand—leads to product **3-σ** in which one C=O bond has been completely broken. Such CO₂ cleavage reactions are extremely rare for late 3d metals, and this represents the first such example mediated by a single Fe centre.

The transformation of carbon dioxide into novel and useful chemicals remains one of the most important research challenges of modern chemistry. Application of CO₂ as a C₁ building block is highly attractive due to its relative abundance and, in particular, as a method to address long-established concerns about its role as an atmospheric greenhouse gas.^[1] One potential method is the transformation of CO₂ into carbon monoxide, CO, which is a common feedstock for large-scale industrial processes (e.g. Fischer–Tropsch alkane synthesis).^[2] This has led to great interest in the identification and study of well-defined molecular systems capable of directly cleaving the C=O bond in CO₂. Such transformations are highly challenging due to the high strength of the C=O bond (bond enthalpy 532 kJ mol⁻¹).^[3]

Nevertheless, these reactions have become relatively established for highly reducing, low-valent early 3d metals^[4] and a number of examples based on precious^[5] and f-block metals^[6] have also been reported. In comparison, the use of earth-abundant late 3d metals remains surprisingly underexplored, particularly given the role that such metals play in biological CO₂ reduction to CO, mediated by Ni, Fe CO dehydrogenase.^[7]

Currently, only a handful of examples of CO₂ cleavage based on 3d metals are known (Figure 1).^[8] In 2005, Sadighi et al. reported abstraction of an oxygen atom from CO₂ by

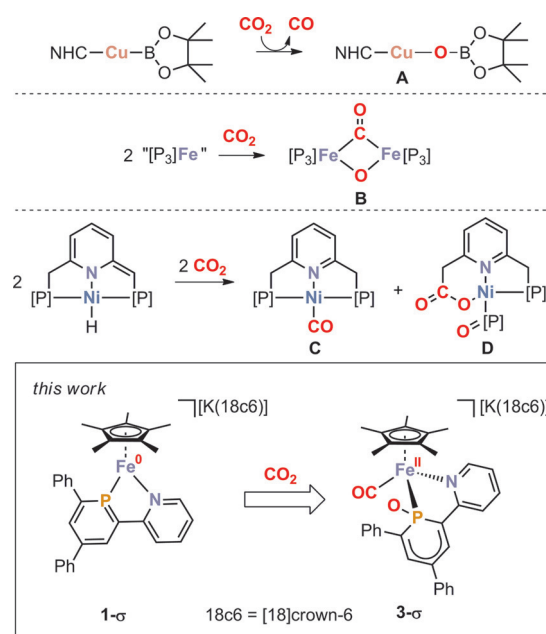


Figure 1. C=O cleavage in CO₂ mediated by well-defined late 3d metal complexes. NHC = N-heterocyclic carbene. $[\text{P}_3] = \text{PhB}(\text{CH}_2\text{P}(\text{CH}_2\text{Cy})_2)_3$. $[\text{P}] = \text{P}^i\text{Pr}_2$.

NHC copper boryl complexes (NHC = N-heterocyclic carbene), resulting in formal O atom insertion into the Cu–B bond to give compound **A**.^[9] Subsequently, in 2007, Peters and co-workers reported the formation of a dinuclear Fe^{II} complex **B** containing bridging CO and O²⁻ ligands, which was prepared by reaction of CO₂ with an Fe^I precursor (similar results were also published by Holland et al. shortly thereafter).^[4d,10] Recently, the group of Milstein has described a more complex CO₂ activation sequence mediated by a nickel pincer complex, involving the cleavage of a ligand

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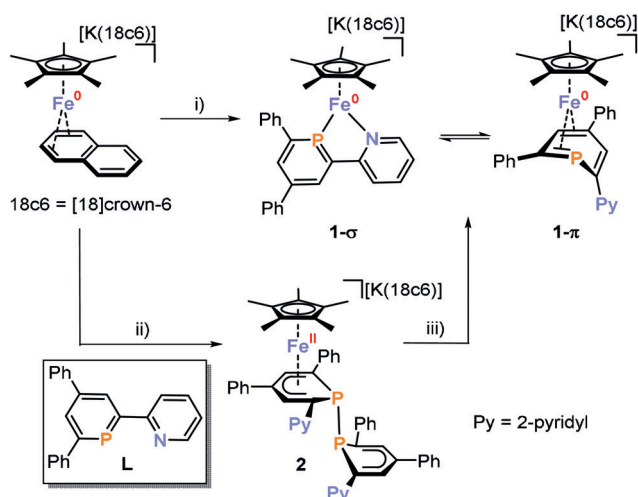
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C–P bond as well as one CO₂ C=O bond to give the products **C** and **D**.^[11]

The complexes **A–D** all form through some manner of cooperative CO₂ activation. In each case, the small-molecule substrate is cleaved as a result of combined interaction with either more than one metal centre (**B**, **C/D**), or with both a metal centre and a “non-innocent” ligand (**A**, **C/D**). On this basis, we reasoned that the use of other ligands capable of metal–ligand cooperation (MLC)^[12] should allow us to expand the range of structural motifs available for mediating challenging CO₂ cleavage reactions at late 3d metals.

In this context, we have recently become interested in the use of phosphinines as ligands in late 3d transition metal chemistry.^[13] Previous work has shown that these compounds exhibit a rich coordination chemistry, and are capable of displaying diverse coordination modes.^[13,14] Furthermore, we have shown that Cp*Fe complexes of the simple phosphinine TPP (TPP = 2,4,6-triphenylphosphinine) show considerable non-innocent behaviour, with the TPP ligand capable of displaying either nucleophilic or electrophilic character towards external reagents.^[15] We anticipated that the coordination chemistry (and, hence, onward reactivity) of these complexes might be further diversified through incorporation of the phosphinine moiety into a chelating ligand scaffold. We therefore decided to target the synthesis of an analogue of our previously reported complex [K([18]crown-6)][Cp*Fe(η⁴-TPP)],^[16] in which TPP is replaced by the known ligand 2-(2'-pyridyl)-4,6-diphenylphosphinine (**L**).^[17]

Pursuing an analogous method to that used previously to prepare [K([18]crown-6)][Cp*Fe(η⁴-TPP)], treatment of [K([18]crown-6)][Cp*Fe(η⁴-C₁₀H₈)] with **L** at –35 °C in 1,2-dimethoxyethane (DME), which was then warmed to room temperature (RT) and stirred overnight, gave a deep purple reaction mixture. Dark purple crystals of a new product **1** were obtained in good yield (68 %) by layering this reaction mixture with *n*-hexane and subsequent filtration (Scheme 1,



Scheme 1. Reaction of [K([18]crown-6)][Cp*Fe(C₁₀H₈)] with 2-(2'-pyridyl)-4,6-diphenylphosphinine (**L**) forming **1-σ** and **1-π**; i) **L**, DME, –35 °C to RT, –naphthalene; ii) 1 equiv. [K([18]crown-6)][Cp*Fe(C₁₀H₈)], 2 equiv. **L**, toluene/THF, –35 °C to RT; iii) 1 equiv. [K([18]crown-6)][Cp*Fe(C₁₀H₈)], THF.

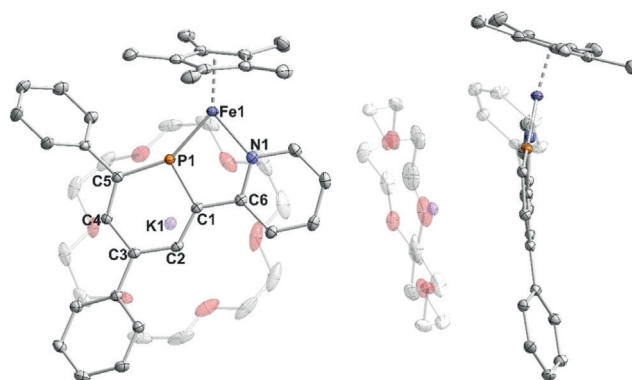


Figure 2. Solid-state molecular structure of **1-σ** (left);^[24] ellipsoids are drawn at the 40 % probability level; H atoms are omitted and the [K([18]crown-6)]⁺ cation is transparent for clarity; selected bond lengths [Å] and bond angles [°]: Fe1–P1: 2.0811(4), Fe1–N1: 1.9428(12), P1–C1: 1.7733(13), C1–C2: 1.3953(19), C2–C3: 1.3947(18), C3–C4: 1.4343(18), C4–C5: 1.3756(18), C5–P1: 1.7757(13), P1–Fe1–N1: 82.32(4), Fe1–P1–C1: 107.62(4), Fe1–N1–C6: 124.18(9), Cp*–Fe1–N1: 135.513(8), P1–Fe1–Cp*: 140.545(9). Side view of solid-state molecular structure of **1-σ** (right); for clarity, one additional phenyl group is also transparent.

top). X-ray analysis confirmed the expected formulation of **1** as [K([18]crown-6)][Cp*Fe**L**] (**1-σ**), in which the ligand **L** has displaced naphthalene from the anionic Cp*Fe moiety (Figure 2). The crystal structure reveals that, in contrast to the η⁴-TPP binding mode in [K([18]crown-6)][Cp*Fe(η⁴-TPP)], the phosphinine ligand acts as a σ-coordinated [P,N] chelate in [K([18]crown-6)][Cp*Fe**L**], with the Cp* and **L** ligands coordinating the iron centre in a trigonal-planar fashion.

Further structural assignment of **1-σ** in the solid state is supported by ³¹P CP MAS NMR spectroscopy, which indicates the presence of a single species with a signal at δ_{iso} = 121.1 ppm, in the range expected for this coordination mode of **L**.^[18] In contrast, solution-phase ³¹P{¹H} NMR analysis of crystalline **1** dissolved in suitable solvents^[19] shows two distinct resonances. For example, in [D₈]THF a sharp singlet at 130.7 ppm (or a triplet with ³J_{P-H} = 13 Hz in the coupled ³¹P NMR spectrum) is assigned to **1-σ**, but is accompanied by a second, very broad resonance (τ_{1/2} = 1000 Hz) at –46.2 ppm. Attempts to selectively crystallise the species corresponding to this resonance have been unsuccessful, with XRD and ³¹P CP MAS measurements consistently showing the presence of only **1-σ** in the solid state. Nevertheless, the broadness and chemical shift of this signal are very similar to those observed previously for the complex [K([18]crown-6)][Cp*Fe(η⁴-TPP)]. In this compound the TPP ligand is rapidly equilibrating between different η⁴ π-coordination modes.^[16] We therefore assign this signal to an isomeric structure of **1** (**1-π**) where the ligand **L** coordinates in an η⁴-fashion as shown in Scheme 1, and which exists in equilibrium with **1-σ** in solution. This conclusion is supported by DFT calculations. A minimum-energy structure corresponding to **1-π** is only 0.9 kcal mol^{–1} higher in energy than the optimized structure for **1-σ** (which reproduces well the structure observed by crystallography; see the Supporting Information). Furthermore, conversion of **1-π** to **1-σ** is calculated to proceed with a barrier of 27.0 kcal mol^{–1}, consistent with an equilibrium at room

temperature. $^{31}\text{P}\{^1\text{H}\}$ NMR integration indicates an approximately 2:1 ratio of **1-σ** to **1-π** at room temperature, which does not change significantly at lower temperature (-30°C ; at elevated temperatures other reactivity is observed, see the Supporting Information).

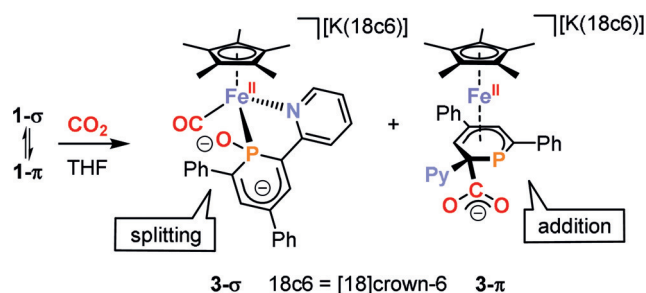
In the hope of gaining more insight into the formation of both isomers of **1**, the reaction of $[\text{K}([18]\text{crown-6})][\text{Cp}^*\text{Fe}(\eta^4\text{-C}_{10}\text{H}_8)]$ with **L** was monitored by $^{31}\text{P}\{^1\text{H}\}$ NMR spectroscopy at 273 K. The initial spectrum showed immediate, full consumption of ligand **L**, and formation of two major new groups of resonances at ca. -25 and -42 ppm.^[20] Only after approximately 30 minutes did signals attributable to **1-σ** and **1-π** become readily discernible, growing in steadily over the course of several hours with concomitant loss of the intermediate resonances.

Close inspection of the observed multiplets suggested that they could be assigned to a pair of AB spin systems with very similar chemical shifts and coupling constants. Notably, the chemical shifts at approximately -42 ppm are almost identical to those of the P–P bonded dimeric species $[\text{Cp}^*\text{Fe}(\eta^5\text{-TPP})]_2$.^[16] Indeed, when the reaction between **L** and $[\text{K}([18]\text{crown-6})][\text{Cp}^*\text{Fe}(\eta^4\text{-C}_{10}\text{H}_8)]$ was repeated with a 2:1 molar ratio, these resonances were seen to form selectively. Only upon addition of a further equivalent of $[\text{K}([18]\text{crown-6})][\text{Cp}^*\text{Fe}(\eta^4\text{-C}_{10}\text{H}_8)]$ were the isomers of **1** observed (Scheme 1, bottom). Conversely, addition of 1 equiv. of **L** to an isolated sample of **1** regenerated the intermediate resonances. It was therefore proposed that these resonances might correspond to two isomers of the P–P dimer $[\text{K}([18]\text{crown-6})][\text{Cp}^*\text{Fe}(\eta^5\text{-L}_2)]$, **2**, which differ only in the relative orientations of the **L** pyridyl groups. This assignment was confirmed by X-ray analysis of single crystals grown by slow diffusion of *n*-hexane into a DME solution of **2**, which resolved to show the expected **L**₂ dimer. One half of the molecule is coordinated in an η^5 fashion to a Cp^*Fe moiety (Figure 3).^[21] The P1–P2 bond is longer than typical P–P single bonds (2.3149(14) vs. 2.04 Å)^[22] suggesting a relatively

weak interaction, consistent with facile cleavage upon addition of a second equivalent of $[\text{K}([18]\text{crown-6})][\text{Cp}^*\text{Fe}(\eta^4\text{-C}_{10}\text{H}_8)]$.

Complex **2** likely arises through rapid addition of initially-formed **1** to a second equivalent of **L**. When **L** is consumed, further coordination of “ Cp^*Fe^- ” leads to formation of free **1**. Transiently-formed **1** is believed to act as a nucleophile in this reaction, with the second equivalent of **L** behaving as a formal electrophile. This is consistent with previous observations of nucleophilic character for the P atom in $[\text{K}([18]\text{crown-6})][\text{Cp}^*\text{Fe}(\eta^4\text{-TPP})]$,^[15a] as well as with DFT calculations, which suggest that the phosphinine moiety not directly bound to Fe is best described as an anionic phosphahexacyclodienyl fragment (see the Supporting Information). These calculations also indicate that formation of **2** from **1-π** and **L** is favourable by 23.5 kcal mol^{−1}, and proceeds over a low energy barrier of 7.5 kcal mol^{−1}.

Encouraged by the evidence of both coordinative flexibility and non-innocent reactivity already displayed by the phosphinine ligand **L** during the synthesis of compound **1**, we proceeded to investigate the reactivity of this complex towards CO₂. Thus, a THF solution of **1-σ** and **1-π** was charged with CO₂ (1 atm) at room temperature (Scheme 2).



Scheme 2. Reaction of **1-σ** and **1-π** with CO₂ (1 atm) in THF at room temperature.

Gratifyingly, an immediate colour change from deep purple to deep green was observed, and $^{31}\text{P}\{^1\text{H}\}$ NMR spectroscopy confirmed the complete consumption of **1** within 10 minutes with concomitant formation of two major new species, identified by signals at 97.0 and 116.0 ppm, with the former splitting into a triplet in the proton-coupled ^{31}P NMR spectrum ($^3J_{\text{P-H}} = 13$ Hz). The species corresponding to these two resonances could be separated by fractional crystallisation, and the identity of both was established by single-crystal X-ray diffraction studies (Figure 4).

Thus, the $^{31}\text{P}\{^1\text{H}\}$ resonance at 116.0 ppm can be assigned to the new compound **3-π**, in which an intact CO₂ moiety has attached to the phosphinine carbon “*ipso*” to the pyridyl group.^[23] The resulting carboxylate-substituted phosphacyclohexadienyl ligand coordinates to Fe in an η^5 fashion, and the carboxylate itself interacts in a κ^2 manner with the potassium counteranion. It is noteworthy that no reaction is observed between CO₂ and the free ligand **L** under analogous conditions.

A more dramatic outcome is revealed by the second major reaction product, **3-σ**. In this case, the molecular structure in

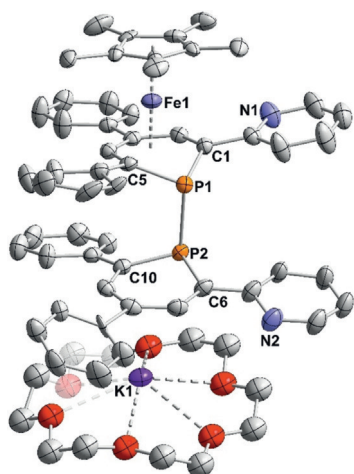


Figure 3. Solid-state molecular structure of **2**;^[24] ellipsoids are drawn at the 40% probability level; H atoms are omitted for clarity; selected bond lengths [Å] and bond angles [°]: P1–P2: 2.3149(14), C1–P1–C5: 92.99(17), C6–P2–C10: 99.58(19), fold angle C1–P1–C5: 32.64, fold angle C6–P2–C10: 14.79.

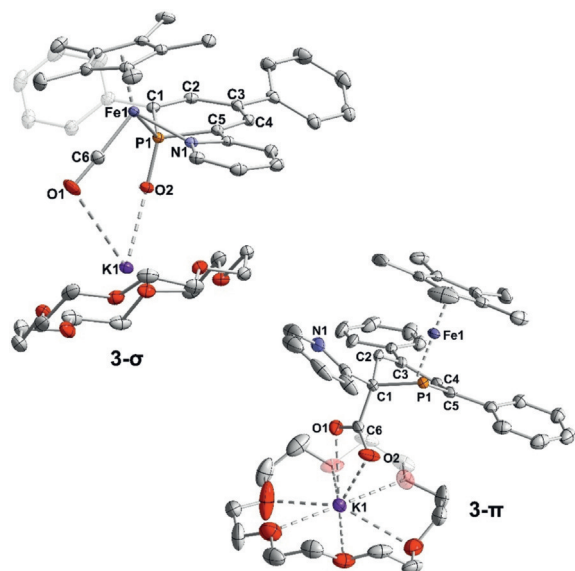


Figure 4. Solid-state molecular structures of **3-σ** and **3-π**;^[24] ellipsoids are drawn at the 40% probability level; H atoms are omitted for clarity; selected bond lengths [Å] and bond angles [°] of **3-σ**: Fe1-P1: 2.2434(4), Fe1-N1: 2.0060(14), Fe1-C6: 1.7272(18), C6-O1: 1.170(2), P1-O2: 1.5272(12), K1-O1: 3.3043(16), K2-O2: 2.558(17), O1-C6-Fe1 = 177.49(17), C6-Fe1-P1 = 88.41(6), Fe1-P1-O2 = 114.45(5), C6-Fe1-N1: 98.78(7); for **3-π**: C6-O1: 1.236(5), C6-O2: 1.229(5), C1-C6: 1.575(5), K1-O1: 2.723(3), K1-O2: 3.050(4), O1-C6-O2 = 127.3(4), P1-C1-C2 = 97.5(3), C1-P1-C5 = 98.36(19).

the crystal clearly shows complete cleavage of one C=O bond in the CO₂ substrate, with coordination of the resulting CO molecule to the Fe centre, and of the remaining O atom to the ligand P atom. Nonetheless, the coordination mode of the ligand **L** remains unchanged from **1-σ**, which may be attributed to the electronic flexibility of the phosphinine moiety. In contrast to Milstein's product **D** (as well as an Ir-based system described recently by Langer, Hamza and Pápai), O-atom transfer to P does not require cleavage of any bonds in the ligand backbone.^[5e,11] Again, this can be attributed to the use of an unsaturated phosphinine moiety in place of a more conventional phosphine donor. This reaction represents the first example of full CO₂ C=O cleavage mediated by a single well-defined Fe centre. That the activation is achieved under ambient conditions using only earth-abundant elements renders this system a highly attractive starting point for further investigations into productive CO₂ utilization.

Intuitively, the structures observed for **3-σ** and **3-π** suggest that they are formed through the reaction of CO₂ with isomers **1-σ** and **1-π** of **1**, respectively. This conclusion is further supported by ³¹P{¹H} spectra of the crude product mixture, which indicate the formation of **3-σ** and **3-π** in the same 2:1 ratio observed for the equilibrium between **1-σ** and **1-π** (vide supra). To provide more conclusive evidence, DFT studies were performed to establish the mechanisms of formation of both **3-σ** and **3-π** (see the Supporting Information). As expected, interaction of CO₂ with **1-σ** was calculated to lead to formation of **3-σ**, with only a very small energy barrier of 3.5 kcal mol⁻¹. Interaction with **1-π** was calculated to lead to

3-π over a similarly small barrier (5.5 kcal mol⁻¹). Notably, both barriers are significantly lower than the barrier to isomerisation between **1-σ** and **1-π** (vide supra).

In summary, we have prepared a new anionic Fe complex **1** incorporating a chelating phosphinine ligand. Complex **1** exists in solution as a pair of isomers in which the phosphinine adopts distinct coordination modes at the metal. Both isomers react with CO₂; for the π-coordinated isomer **1-π** addition of CO₂ to a carbon atom of the phosphinine moiety is observed, while full cleavage of one C=O bond occurs in the σ-coordinated isomer **1-σ** to give an Fe-coordinated CO moiety and P-bound O atom. This is the first reported example of C=O cleavage of a CO₂ molecule mediated by a single Fe centre, and clearly demonstrates the potential of coordinatively and electronically flexible phosphinine ligands to help mediate challenging bond-activation reactions. Research into the further applications of such complexes, the derivatisation of CO₂ cleavage product **3-σ**, and the use of these systems to address the crucial and topical challenge of efficient CO₂ transformation are currently underway in our laboratories.

Please note: Minor changes have been made to this manuscript since its publication in *Angewandte Chemie* Early View. The Editor.

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Conflict of interest

The authors declare no conflict of interest.

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- [24] CCDC 1942537, 1942538, 1942550 and 1942542 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre.

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