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## Article

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# CO Coupling Chemistry of a Terminal Mo Carbide: Sequential Addition of Proton, Hydride, and CO Releases Ethenone

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**ABSTRACT:** The mechanism originally proposed by Fischer and Tropsch for carbon monoxide (CO) hydrogenative catenation involves C–C coupling from a carbide-derived surface methylidene. A single molecular system capable of capturing these complex chemical steps is hitherto unknown. Herein, we demonstrate the sequential addition of proton and hydride to a terminal Mo carbide derived from CO. The resulting anionic methylidene couples with CO (1 atm.) at low-temperature (-78 °C) to release ethenone. Importantly, the synchronized delivery of two reducing equivalents and an electrophile, in the form of a hydride (H<sup>-</sup> = 2e<sup>-</sup> + H<sup>+</sup>), promotes alkylidene formation from the carbyne precursor and enables coupling chemistry, under conditions milder than those previously described with strong one-electron reductants and electrophiles. Thermodynamic measurements bracket the hydricity and acidity requirements for promoting methylidene formation from carbide as energetically viable upon formal heterolysis of H<sub>2</sub>. Methylidene formation prior to C–C coupling proves critical for organic product release, as evidenced by direct carbide carbonylation experiments. Spectroscopic studies, a monosilylated model system, and Quantum Mechanics computations provide insight into the mechanistic details of this reaction sequence, which serves as a rare model of the initial stages of the Fischer Tropsch synthesis.

#### INTRODUCTION

The Fischer-Tropsch (FT) synthesis is a well-established industrial-scale process that uses synthesis gas, a mixture of CO and H<sub>2</sub>, as a feedstock for the generation of a broad range of catenated products.<sup>1-5</sup> While the variation in outputs can be versatile, it is often cost prohibitive in regards to petroleum substitution.<sup>1-2</sup> As efforts increase to synthesize biodiesels from renewably sourced synthesis gas,<sup>6-7</sup> shifting FT catalysis toward specific desirable products becomes ever more important.<sup>7</sup> Understanding the operative mechanism(s) of FT provides potential for rational catalyst design and, in turn, improved selectivity.<sup>1-2, 8</sup>

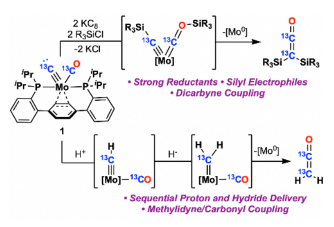
While there is little consensus as to the operative mechanism(s) of FT,9-10 all proposals share two fundamental commonalities—CO reduction (via C-H bond formation and/or C-O cleavage) and C-C coupling. The elementary step of C-C coupling between CO and surface hydrocarbons has been studied extensively in model systems.<sup>1,11-12,13-14</sup> In contrast, the investigation of carbide/CO coupling remains largely underexplored. In seminal studies of carbonyl-bridged metal carbide ensembles,<sup>15</sup> more exposed carbides demonstrated enhanced reactivity, including C/CO coupling in the Fe<sub>3</sub> cluster.<sup>16</sup> Additional examples of C≡O cleavage and coupling have been reported, likely via carbide intermediacy, with oxophilic metals.<sup>17-19</sup> To our knowledge, CO coupling has yet to be demonstrated from a terminal carbide precursor, likely due to the paucity of reactive yet well-characterized examples of such motifs.20-24

Previous work in our group has focused on reductive C–O catenation chemistry employing one electron reductants (KC<sub>8</sub>) and silyl electrophiles, demonstrating the deoxygenation of bound CO to a terminal carbide (**1**, Scheme 1).<sup>25</sup> This reactivity models the early steps proposed for FT, achieving the six-electron cleavage of CO using reducing equivalents stored in a redox non-innocent supporting ligand. C–C coupling was achieved via an isolated silyl alkylidyne complex, which, upon addition of two equiv. of strong reductant (KC<sub>8</sub> or naphthalenide) followed by strong electrophiles (silyl chlorides), afforded disilylketene (Scheme 1, top).<sup>25-26</sup>

In contrast to these relatively harsh reducing conditions, several molecular FT models have employed proton and hydride equivalents as the products of heterolytic cleavage of H<sub>2</sub>.<sup>27-29</sup> Herein, we revisit CO coupling chemistry from terminal carbide 1, the first reported example of a terminal group VI carbide bearing d-electrons. While C-C bond formation from this species is facile, it ultimately results in deleterious ligand functionalization. Sequential proton and hydride delivery convert complex **1** to a terminal methylidene via an isolated terminal methylidyne (Scheme 1, bottom). Importantly, thermochemical experiments with NaBHPh<sub>3</sub> and [Ar<sub>2</sub>PhPMe]Cl (Ar = 2,4,6-trimethoxyphenyl) provide a formal demonstration that thermodynamically H<sub>2</sub> can promote reduction and protonation of complex 1, illustrating the feasibility of using proton and hydride as H<sub>2</sub> surrogates in carbide conversion chemistry. Addition of exogenous CO to the formed methylidene releases ketene, demonstrating the sequential formation of two C-H bonds and C/CO coupling, to yield a silicon-free, metal-free  $C_2O_1$  organic.

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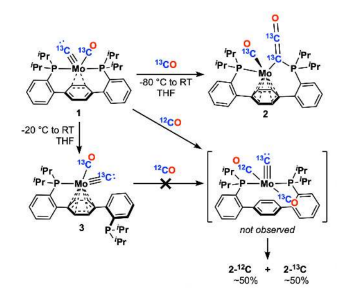
Scheme 1. Mechanistically Distinct Modes of C/CO Coupling to *Free* Ketene Fragments <sup>30</sup>



#### **RESULTS AND DISCUSSION**

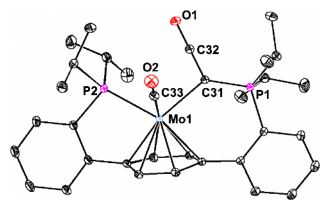
#### Carbide/CO Coupling Reactivity

Direct carbide carbonylation represents an early bifurcation in the mechanistically complicated FT process; surface C<sub>2</sub> formation via carbide/CO coupling has been proposed for syngas conversion on iron foils.<sup>31</sup> To explore carbide/CO coupling in a well-defined system, complex **1** was treated with CO (Scheme 2).<sup>30,32</sup> Admission of one atmosphere (1 atm) of <sup>13</sup>CO to a THF solution of 1 at -80 °C showed no effect; however, upon warming to 25 °C, new spectral features were observed. A peculiar high-field doublet of doublets (-32.14 ppm,  ${}^{1}/(P,C) = 118$  and  ${}^{1}/(C,C) = 103$  Hz) was present in the <sup>13</sup>C{<sup>1</sup>H} NMR spectrum, in addition to two downfield resonances. Concomitantly, two new features were observed in the <sup>31</sup>P{<sup>1</sup>H} NMR spectrum, including a doublet of doublet of doublets (37.64 ppm, <sup>1</sup>/(P,C) = 118.1 Hz, <sup>2</sup>/(P,C) = 33.2 Hz, & <sup>3</sup>/(P,P) = 4.2 Hz) displaying strong scalar coupling to the high-field carbon. The large P–C and C–C spin-spin coupling constants, in addition to the broken symmetry of the terphenyl diphosphine (P2) ligand, are most consistent with formation of a Mo-bound phosphoranylideneketene, 2. Scheme 2. CO Coupling Chemistry of Terminal Molybdenum Carbides<sup>a</sup>



<sup>*a*</sup> Carbide **1** was synthesized cleanly *in situ* via deprotonation of P2Mo(CH)(CO)(Cl) **4** with benzyl potassium. For more details, see the Supporting Information.

A single-crystal X-ray diffraction (XRD) study of **2** confirmed the proposed assignment (Figure 1). The short C31– C32 bond length and wide C31–C32-O1 angle (1.2885(11) Å and 179.01°, respectively) support assignment of the  $\eta^1$ -C(P)=C=O bonding motif.<sup>33</sup> Mo engages the central arene in an  $\eta^6$  fashion, to stabilize the low-valent metal center resulting from C–C coupling. Transition metal-bound phosphoranylidene ketenes, relatively rare motifs, are generally prepared via reaction of a metal precursor with preformed ketene, R<sub>3</sub>PCCO.<sup>34</sup> In one example, the P–C linkage was formed at the metal center, from cleavage of carbon suboxide.<sup>35</sup> Complex **2** therefore represents the first example of such a motif in which both the P–C and C–C bonds were formed at the metal center.



**Figure 1.** Solid-state structure of **2**. Thermal anisotropic displacement ellipsoids are shown at the 50% probability level. Hatoms are omitted. Selected bond distances [Å] and angles [°]: P1-C31 1.7144(11), C31-C32 1.2885(11), C32-O1 1.1865(10), Mo1-C31 2.2899(16), Mo1-C33 1.9563(12), C33-O2 1.1693(10), C31-C32-O1 179.01(8), P1-C31-Mo1 130.73(5).

We hypothesized that CO binding could induce C-C bond formation<sup>36-37</sup> in **1** via a putative dicarbonyl carbide intermediate (Scheme 2). Labeling experiments were conducted to explore this process. Addition of <sup>12</sup>CO to **1** results in a 50:50 mixture of isotopes at the phosphoranylideneketene  $\alpha$ -carbon (Figure 2). Likewise, the <sup>31</sup>P{<sup>1</sup>H} NMR spectrum displays a broad resonance for the Mo-bound phosphine arm comprised of an overlapping doublet of doublets and doublet, the former split by partial (ca. 50%) <sup>13</sup>C enrichment at the carbonyl carbon. Taken together, these data are consistent with an intramolecular coupling pathway from a proposed dicarbonyl carbide complex. This process contrasts the ostensibly similar cleavage and coupling of CO by Ta(silox)<sub>3</sub>,<sup>38</sup> which, through a series of detailed mechanistic studies, was shown to form the C-C bond prior to the first deoxygenation event, en route to a ditantalum dicarbide ((silox)<sub>3</sub>Ta=CC=Ta(silox)<sub>3</sub>).<sup>39</sup> Related transformations starting from alkylidyne carbonyl complexes, with addition of phosphines leading to the assembly of *neutral* PC(R)=C=O ligands, have also been reported.<sup>40-43</sup> Combined, these molecular systems represent complementary precedents for potential elementary steps in the early stages of the FT synthesis.

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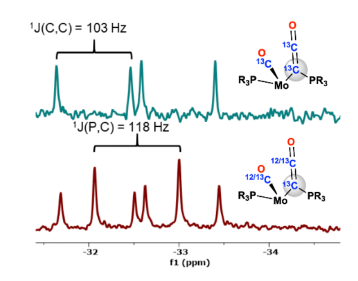
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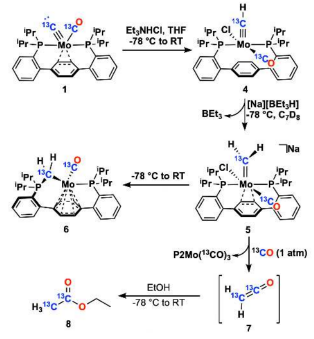
**Figure 2.** High-field partial <sup>13</sup>C{<sup>1</sup>H} NMR spectra of **2** generated under <sup>13</sup>CO (top) and <sup>12</sup>CO (bottom) showing splittings for the phosphoranylidene <sup>13</sup>C nucleus circled in grey.

Recognizing that carbide **1** need not bind an additional molecule of CO to form a ketenylidene, it was slowly warmed to RT. While this results in a composite mixture of largely intractable species,<sup>25</sup> one of these complexes proved highly soluble. Extraction of the crude residue with hexamethyldisiloxane afforded a deep purple solution, which upon standing at -35 °C, precipitated 3 as a purple powder. While single crystals of 3 have proven elusive, the structure can be confidently inferred from multinuclear NMR spectroscopy. A diagnostic carbidic <sup>13</sup>C{<sup>1</sup>H} resonance is observed at 569.86 ppm, further downfield shifted from that of **1**. Two distinct features in the <sup>31</sup>P{<sup>1</sup>H} NMR spectrum at 91.44 and -4.08 ppm support an arm-on arm-off binding mode for the P2 ligand, and four upfield aromatic <sup>1</sup>H shifts corroborate an n<sup>6</sup> metal-arene interaction. Addition of 1 atm of CO to solutions of **3** likewise did not afford the coupled product **2**, however. Instead, an intractable mixture is formed, together with free P2, reflecting additional, deleterious pathways that are enabled by dissociation of the second phosphine arm.

#### Carbide Protonation, Hydride Transfer, and Methylidene/CO Coupling

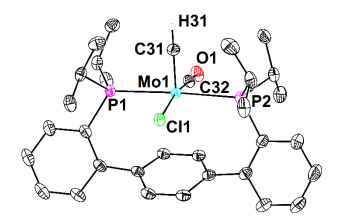
41 Toward modeling the proposed role of surface hydrides in 42 FT, and ultimately forming C–H bonds, carbide protonation 43 was targeted.<sup>21, 44</sup> Addition of Et<sub>3</sub>NHCl to a -78 °C THF solu-44 tion of **1** results in formation of a new compound upon 45 warming to room temperature. A single resonance between 46 3.5 and 6.5 ppm is observed in the <sup>1</sup>H NMR spectrum, a doublet of triplets centered at 5.06 ppm (<sup>1</sup>/(C,H) = 147.8 Hz, 47  $^{3}$ /(P,H) = 3.1 Hz), with a relative integration of one. This sug-48 gests both a terminal methylidyne<sup>20, 24, 45-51</sup> and a five-coor-49 dinate *pseudo*-square pyramidal structure, as in **4**, without 50 metal-arene interactions (vide infra; Scheme 3). The <sup>31</sup>P{<sup>1</sup>H} 51 and 13C{1H} NMR data further corroborate the proposed as-52 signment of **4**, with a doublet of doublets at  $\delta_P$  40.21 ppm 53  $(^{2}J(C,P) = 18.6 \text{ and } 10.2 \text{ Hz})$  and a characteristic methyli-54 dyne resonance at  $\delta_{C}$  281.19 ppm, respectively.<sup>45, 51</sup> 55

Scheme 3. Sequential Proton and Hydride Addition *en route* to Ketene Formation <sup>a</sup>



<sup>*a*</sup> Carbide **1** was synthesized cleanly *in situ* via deprotonation of P2Mo(CH)(CO)(Cl) **4** with benzyl potassium. For more details, see the Supporting Information.

Gratifyingly, **4** is thermally stable, facilitating both its use for reaction chemistry and permitting growth of single crystals. A single-crystal XRD study corroborates the structure inferred from spectroscopy (Figure 3). The Mo center sits above the central arene ring at a distance of 2.749 Å, suggesting negligible metal arene interaction. The Mo≡CH distance is appropriately short (1.764(2) Å) but slightly longer than that in a structurally characterized Mo(VI) methylidyne (1.702(5) Å),<sup>44</sup> likely a manifestation of the more-reduced metal center.



**Figure 3.** Solid-state structure of **4**. Thermal anisotropic displacement ellipsoids are shown at the 50% probability level. Hatoms are omitted, except for the methylidyne proton, which was refined freely. Selected bond distances [Å] and angles [°]: Mo1–C31 1.764(2), Mo1–C32 1.930(4), O1–C32 1.197(6).

Hydride addition to both terminal<sup>21</sup> and bridging<sup>52</sup> methylidyne ligands has precedent, and we sought to target this reactivity, considering proton and hydride as a crude surrogate for  $H_2$  (*vide infra*). NaBEt<sub>3</sub>H was added to a C<sub>7</sub>D<sub>8</sub>

solution of 4 at -78 °C. Mixing the reagents immediately resulted in a color change to dark brown, and VT NMR spectroscopy showed signals in the  ${}^{31}P{}^{1}H{}$  (s, 52.07 ppm), <sup>13</sup>C{<sup>1</sup>H} (br s, 296.91 and br s, 237.52 ppm), and <sup>1</sup>H (15.03, d, <sup>1</sup>/(C,H) = 114.90 Hz and 13.27, d, <sup>1</sup>/(C,H) = 142.7 Hz) spectra, in line with formation of methylidene 5 (Scheme 3).<sup>21</sup> Warming the sample to 0 °C, showed quantitative conversion to a new asymmetric complex with a triplet of doublets  $(84.68 \text{ ppm}, {}^{2}J(P,C_{C0}) = 15.5 \& {}^{2}J(P,C_{CH2}) = 6.0 \text{ Hz})$  and doublet of doublets (47.41 ppm, <sup>1</sup>/(P,C<sub>CH2</sub>) = 21.9 ppm &  $^{2}$ /(P,C<sub>co</sub>) = 17.0 Hz) in the  $^{31}$ P{ $^{1}$ H} NMR spectrum. The <sup>13</sup>C{<sup>1</sup>H} NMR spectrum likewise showed two signals, a broad doublet of triplets centered at 251.88 ppm (<sup>2</sup>/(C<sub>CO</sub>,P) = 15.3 &  ${}^{2}J(C_{CO},C_{CH2})$  = 2.2 Hz) and an extremely high-field resonance at -34.87 ppm (ddd,  ${}^{1}/(C_{CH2},P) = 21.8, {}^{2}/(C_{CH2},P) =$ 6.2 Hz &  ${}^{2}/(C_{CH2},C_{CO}) = 2.2$  Hz). The high-field chemical shift in the <sup>13</sup>C{<sup>1</sup>H} NMR spectrum<sup>49</sup> and the C<sub>1</sub> symmetry are consistent with P-C bond formation, giving 6,53-54 unexpectedly suggesting some electrophilic carbene character in electron-rich, anion **5**.<sup>53-55</sup> The small <sup>1</sup>*J*(P,C<sub>CH2</sub>) spin-spin coupling<sup>56</sup> observed for **6** is consistent with phosphonium ylide complexation to transition metal centers; ligation results in rehybridization— $C(sp^2)$  to  $C(sp^3)$ —that is reflected in the smaller one-bond scalar coupling constant.<sup>57</sup> The structure of **6** was confirmed in a single-crystal XRD study, which corroborated the assignment of the  $\eta^1$ -C-bound ylide moiety (see the SI).

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We hypothesized that similarly to the effect observed for carbide/CO coupling, methylidene/CO coupling may be favored by addition of exogenous CO, either via direct CO attack<sup>58</sup> or coordination to Mo. Placing a -78 °C solution of *in situ* generated **5** under a <sup>13</sup>CO atmosphere resulted in a gradual color change to bright orange, as the gas diffused down the sample (Scheme 3). The <sup>13</sup>C{<sup>1</sup>H} NMR spectrum at -78 °C, showed both free <sup>13</sup>CO and three resonances in the metal-bound region—the CO signals of previously characterized **P2Mo(<sup>13</sup>CO)**<sub>3</sub>.<sup>59</sup>

Anticipating that zero-valent Mo was indicative of a C-C coupling process, identification of ethenone by <sup>13</sup>C{<sup>1</sup>H} NMR spectroscopy was attempted. The diagnostic signals for free ketene ( ${}^{13}C\delta$  = 193.7 and 2.7 ppm) ${}^{60}$  were absent, but coupling doublets at 154.69 and 83.90 ppm (<sup>1</sup>/(C,C) = 76.1 Hz) were observed, attributed to a putative ethenone-triethyl borane adduct.61 While further efforts to detect free ethenone (7) were unsuccessful, ketene trapping with ethanol has precedent, giving stable and easily detectable ethyl acetate (EtOAc).<sup>60, 62</sup> Forming **5** at low temperature, adding <sup>13</sup>CO, and condensing EtOH into the reaction tube afforded EtOAc-<sup>13</sup>C<sub>2</sub>, **8**, unequivocally demonstrating the generation and ejection of parent ketene (Scheme 3). Alkylidene carbonylation has been demonstrated on a variety of molecular scaffolds.<sup>13</sup> but often requires CO overpressures.<sup>58, 63-65</sup> proceeds from activated Fischer carbene complexes, 13, 66-68 or occurs at bridging alkylidene ligands.<sup>69-70</sup> Rarely is this coupling achieved at monometallic complexes with carbene fragments derived from CO.67, 71-73

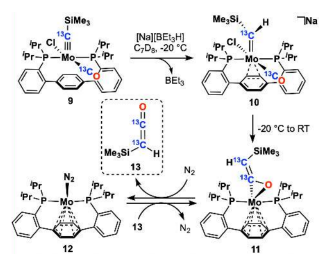
Isotopic labeling studies were conducted in which <sup>12</sup>CO was added to **5**. Mixed isotopolog **P2Mo(**<sup>12/13</sup>**CO)**<sup>3</sup> was formed, consistent with either i) rapid exchange of the <sup>13</sup>CO ligand in **5** with free <sup>12</sup>CO and on-metal C–C bond formation or ii) <sup>12</sup>CO binding and C–C coupling preceding from a

methylidene dicarbonyl complex. We favor the latter, given that coordination to Mo facilitates C–C coupling chemistry in the related carbide/CO chemistry described above.

#### **Reactivity of a Silylcarbyne Model System**

Expecting that similar C/CO coupling chemistry would be achieved from silvl carbyne 9, affording a more stable organic product, a related reactivity sequence was explored with this silvl model system (Scheme 4). Low-temperature addition of NaBEt<sub>3</sub>H to a C<sub>7</sub>D<sub>8</sub> solution of alkylidyne 9 and subsequent warming to -20 °C resulted in the gradual formation of a new species demarcated by a <sup>31</sup>P{<sup>1</sup>H} resonance at 48.53 ppm and low-field doublet in the <sup>1</sup>H NMR spectrum  $(\delta = 15.79 \text{ ppm}, {}^{1}/(C,H) = 121.7 \text{ Hz})$ . Such a signal could be consistent with either formyl generation,74-77 via CO insertion into an intermediate Mo hydride (a process that is facilitated by borane Lewis acids),<sup>29, 78-80</sup> or hydride attack at the carbyne to give a silyl alkylidene. The <sup>13</sup>C NMR spectrum was more informative, displaying a broad doublet at 315.09 ppm  $(^{1}/(C,H) = 121.7 Hz)$  and a broad singlet at 251.83 ppm (Figure S28). C-H coupling to the more downfield resonance is suggestive of a carbene/carbonyl isomer rather than the alternative carbyne/formyl complex. 2D <sup>1</sup>H/<sup>13</sup>C correlation experiments further disambiguated the structure (Figure S29), with strong HMBC cross peaks between the H-bound carbon atom and the SiMe<sub>3</sub> methyl protons. These data point to hydrogen transfer to the alkylidyne carbon,<sup>21,81-83</sup> affording silyl carbene **10**, in analogy to the reactivity observed for parent methylidyne 4 (vide supra).

#### Scheme 4 Hydride-Initiated C-C Coupling.



While **10** was the major species observed at -20 °C, it proved transient, slowly reacting further to give a mixture of two Mo complexes with a doublet and singlet in the <sup>31</sup>P NMR spectrum at 58.38 and 76.24 ppm, respectively. The latter corresponds to previously characterized Mo<sup>0</sup>-N<sub>2</sub> adduct **12**,<sup>59, 84</sup> demonstrating loss of the carbon-based ligands, presumably via C–C coupling. The isotopically enriched <sup>13</sup>C signals at 84.76 and 191.35 ppm showed strong scalar coupling (<sup>1</sup>/(C,C) = 69.7 Hz), the latter split into a doublet of triplets (<sup>2</sup>/(C,P) = 26.3 Hz) by the trans-spanning phosphines of the **P2** ancillary ligand. This spectral signature is consistent with C–C bond formation affording the trimethylsilyl ketene complex **11**. Interestingly, in this case, CO addition is not requisite for C–C coupling.

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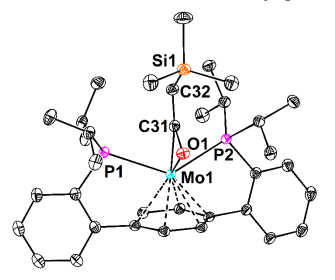
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The assignment of **11** was further supported via independent synthesis.<sup>85</sup> Anticipating that the observed mixture of **11** and **12** resulted from a ligand substitution equilibrium between the isoelectronic Mo<sup>0</sup> adducts, a C<sub>6</sub>D<sub>6</sub> solution of **12** was treated with trimethylsilyl ketene.<sup>86</sup> While only starting material was observed under an N<sub>2</sub> atmosphere, degassing the reaction mixture resulted in significant conversion to **11**, as evidenced by <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy. The N<sub>2</sub>/ketene exchange was borne out in the hydride-initiated C–C coupling chemistry; in addition to the coupling doublets of the bound ketene, a pair of doublets at 192.48 and 60.28 ppm (<sup>1</sup>/(C,C) = 73.5 Hz) were observed in the <sup>13</sup>C NMR spectrum, corresponding to the metal-free C<sub>2</sub> fragment.<sup>61</sup>

An XRD study of single crystals of **11** grown from slow evaporation of liquid butane confirmed the inferred  $\eta^2$ -CO ketene assignment (Figure 4), with comparable 01-C31 (1.2799(12) Å) and C31-C32 (1.3536(14) Å) distances.<sup>87-89</sup>  $\eta^2$ -CC ketene adducts are known,<sup>90-92</sup> but the O-bound form is often thermodynamically favored.<sup>90</sup> Contrasting the structure of **9**,<sup>25</sup> As in the structure of **2** above, the central arene is engaged in  $\eta^6$  coordination to Mo, consistent with the more reduced Mo center formed on C–C coupling.



**Figure 4.** Solid-state structure of **11**. Thermal anisotropic displacement ellipsoids are shown at the 50% probability

level. H-atoms are omitted for clarity. Selected bond distances [Å] and angles [°]: Mo1–O1 2.1858(7), Mo1–C31 2.1629(10), Mo1–Carene (ave.) 2.2599(9), O1–C31 1.2799(12), C31–C32 1.3536(14),  $\angle$ O1-C31-C32 1.32.32(9).

#### **Quantum Mechanics Computations**

Akin to the sequential addition of potassium graphite and silvl electrophile to alkylidyne 9,25-26 NaBEt<sub>3</sub>H proved proficient for enacting the reductive coupling of silvl carbyne with CO. The latter reactivity manifold, preceding through a carbene intermediate (10) rather than a dicarbyne complex, is accessible under much milder conditions and shows equal efficacy in C–C coupling. Given the facility with which the catenation proceeds, Quantum Mechanics computation (using the M06-2X flavor of Density Functional Theory)<sup>93</sup> was employed to interrogate the electronic structure requirements in the critical bond-forming step. The optimized structure following hydride addition to 9 resembles silyl alkylidene **10**,<sup>94</sup> in agreement with experiment; however, a close contact between Na<sup>+</sup> and the Cl ligand is observed in addition to borane binding to the carbonyl (Figure S42). 95 The HOMO of this complex, with  $d_{xz}$  parentage, is dominated by a  $\pi$ -bonding interaction with CO (**10**•**BEt**<sub>3</sub>, Figure 5, inset). As the C-C distance contracts, the alkylidene ligand rotates, directing the Mo–CSi  $\pi$ -system towards CO (<sup>1</sup>X, Figure 5). This geometry has a  $d_{yz}$  HOMO stabilized by M-arene backbonding. With further C–C shortening, the ground state crosses to a triplet, resulting from electron promotion from the  $d_{vz}$  (Mo- arene  $\pi$ ) to the  $d_{xz}$  (Mo-CSi  $\pi^*$ ) and representing cleavage of the Mo–C  $\pi$ -bond (<sup>1</sup>X to <sup>3</sup>X, Figure 5, right insets). This alkylidene activation process, with significant electron density localization on the carbon, is facilitated by the proximal Na<sup>+</sup> ion, without which a low-energy pathway could not be identified. Bond formation is additionally assisted by borane coordination, stabilizing the oxygen lone pair as electron density is transferred to CO.95 The direct impact of the Lewis acids Na<sup>+</sup> and BH<sub>3</sub> on this C-C coupling process is particularly notable. While BEt<sub>3</sub> has no direct relevance to FT, there is precedent for both Lewis acidic promoters in FT catalysis<sup>10</sup> and selectivity perturbations resulting from alkali metal cation addition in electrochemical CO<sup>96-97</sup> reduction. Our computations highlight in a well-defined molecular system the necessity of these interactions for coupling chemistry resulting in ketene.

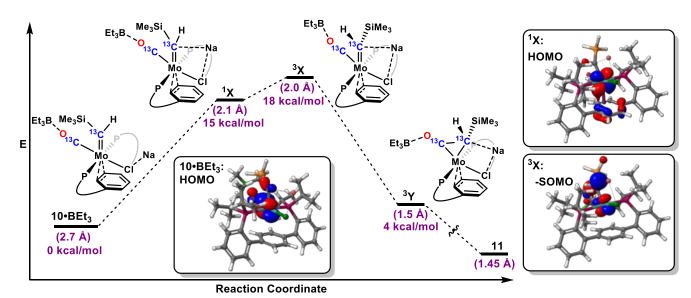


Figure 5. Potential Energy Landscape for C–C Bond Formation. Truncated transition state structures, computed C–C bond distances, and energies (relative to **10**•BEt<sub>3</sub>) are provided. The insets depict key molecular orbitals (0.05 eÅ<sup>-3</sup> isosurfaces) involved in bondbreaking and making steps.

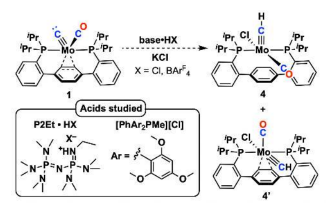
#### Thermochemistry of Proton and Hydride Transfer to Carbide 1

Thus far, a moderate acid (Et<sub>3</sub>NHCl,  $pK_a = 14.9$  in THF)<sup>98</sup> and a strong hydride donor (NaBHEt<sub>3</sub>,  $\Delta G^{\circ}_{H}$  = 26 kcal/mol in MeCN<sup>99</sup>) have been described in the reaction chemistry for the formation of ethenone from **1**. These reagents are only a crude surrogate for H<sub>2</sub>, as the free energy of hydrogen formation is favorable by ca. 25 kcal/mol, considering the reaction free energy of H<sub>2</sub> heterolysis in MeCN.<sup>100</sup> Given the basicity of related Mo(VI) and W(VI) carbide complexes  $(Mo(\equiv CH)(NRAr)_3, pK_a \sim 30 \text{ kcal/mol} \text{ in THF};^{20}$  $W(\equiv CH)(CO)_2(Tp^*)$ ,  $pK_a = 28.7$  kcal/mol in THF<sup>21</sup>), we anticipated that weaker acids could be employed to access methylidyne 4.

35 Further protonation studies assessed the basicity of car-36 bide **1**, and supported a p $K_a$  (Cl) for methylidyne **4** of  $\geq 33.^{101}$ 37 Treatment of carbide **1** with the phosphazene base 38 P2Et•HCl (Scheme 5;  $pK_a$  26.6 in THF<sup>102</sup>) resulted in quanti-39 tative conversion to methylidyne 4, with observation of the expected conjugate base  $P_2Et$  at 15.1 ppm in the  ${}^{31}P{}^{1}H$ 40 NMR spectrum. Reaction with the weaker acid 41  $[Ar_2PhPMe]Cl$  (Ar = 2,4,6-trimethoxyphenyl; pK<sub>a</sub> 33.5 in 42 THF<sup>102</sup>) likewise furnished methylidyne **4**, along with a new 43 species, 4', whose NMR signature mirrored that of 4, sug-44 gesting a closely related isomer. Most diagnostic was a <sup>31</sup>P 45 chemical shift at 41.81 ppm, shifted only ca. 1.5 ppm down-46 field from **4**, and a diagnostic <sup>13</sup>C resonance at 267.8 ppm, 47 which correlated by HMQC analysis to a triplet in the <sup>1</sup>H 48 NMR spectrum at 2.64 ppm for the methylidyne proton 49  $(^{3}/(H,P)= 3.8 \text{ Hz})$ . Independent synthesis, together with a 50 single-crystal XRD study, confirmed 4' to be an isomer of 4 in which the methylidyne is oriented cis with respect to the central arene (Figure 6). As in the previously reported cis silylcarbyne isomer,<sup>26</sup> the *cis* methylidyne isomer adopts a *pseudo*-octahedral geometry with a clear  $\eta^2$ -arene interaction trans to CO. The Mo-C31 distances are similar between the two isomers (cis/trans 1.781 Å vs 1.764(2), respectively), and both are consistent with a Mo-C triple bond.

The conjugate ylide Ar<sub>2</sub>PhP=CH<sub>2</sub> was not detected, however ( $\delta_P$  7.00 ppm). Instead, a new peak in the <sup>31</sup>P NMR at 7.4 ppm was observed, together with free P2 ( $\delta_P$  -3.45 ppm), suggesting displacement of P2 by ylide in the formed methylidyne complex. Competing decomposition of the carbide complex by other pathways, however, cannot be strictly ruled out. Taken together, these protonation studies suggest a lower limit of 33 for the pK<sub>a</sub> (Cl) of methylidyne 4, roughly three orders of magnitude higher than the closest previously reported comparator,  $Mo(\equiv CH)(NRAr)_{3.20}$  The higher effective basicity of carbide 1 may reflect diminished  $\pi$ -donation from the carbide ligand in this more electronrich, formally Mo(IV), complex, and/or the favored energetics of chloride binding to Mo.

Scheme 5. Protonation studies of carbide 1.<sup>*a*, *b*</sup>



<sup>a</sup>pKa values of HCl salts in THF: P2Et•HCl, 26.6;98 [PhAr<sub>2</sub>PMe][Cl], 33.5.<sup>102</sup> <sup>b</sup> Carbide **1** was synthesized cleanly in situ via deprotonation of P2Mo(CH)(CO)(Cl) 4 with benzyl potassium. For more details, see the Supporting Information.

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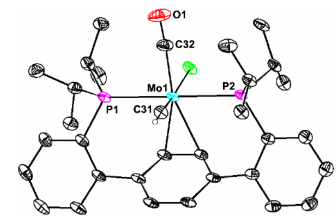
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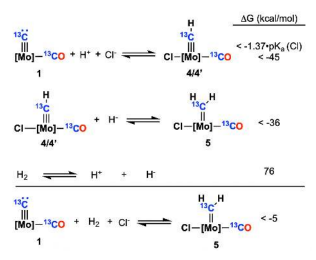
**Figure 6.** Solid-state structure of **4**'. Thermal anisotropic displacement ellipsoids are shown at the 50% probability level. H-atoms are omitted for clarity, except the methylidyne proton. Selected bond distances [Å] and angles [°]: Mo1-C31 1.781(2), Mo1-C32 1.972(2), O1-C32 1.155(2).

To gain insight into the hydride affinity of methylidyne **4**, weaker hydride sources were employed. Hydride transfer was complete using NaBHPh<sub>3</sub> ( $\Delta G^{\circ}_{H^-} = 36$  in MeCN<sup>99</sup>), as judged by the quantitative formation of **6** upon warming the reaction to RT. However, in a variable temperature NMR study, methylidene **5** was not observed on maintaining the reaction at -78 °C. Instead, observation of **5** necessitated warming to *ca.* -30 °C, consistent with a higher activation energy of hydride transfer from the bulkier borohydride. Up to equal proportions of **5** and **4** were observed at -20 °C, suggesting that the hydride affinity of methylidyne **4** is at least comparable to that of NaBHPh<sub>3</sub>. Competing conversion of **5** to **6** occurred at these temperatures; however, preventing extraction of the exact thermodynamic parameters.

Addition of NaBHPh<sub>3</sub> at -78 °C, followed by reaction with CO (1 atm), likewise furnished **P2Mo(CO)**<sub>3</sub> on warming to RT, as expected based on the thermodynamic favorability of conversion from **5** described above. A small peak assigned to **5** was apparent only at intermediate temperatures (0 °C), again consistent with the higher activation barrier using NaBHPh<sub>3</sub>.

Reconsidering the relevance to  $H_2$  splitting, the p $K_a$  of [ $Ar_2PhPMe$ ]Cl and hydricity of NaBHPh<sub>3</sub> suggest that a reaction between carbide **1** and  $H_2$  is thermodynamically downhill by at least 5 kcal / mol (Scheme 6).<sup>103</sup> These experiments therefore constitute a thermochemical demonstration that  $H_2$  is a suitable source of protons and hydrides in the studied system. However, warming a THF solution of **1** under 1 atm  $H_2$  showed only slight conversion to **3**. Direct reaction of carbides with  $H_2$  therefore appears to be kinetically disfavored.<sup>104</sup> Cleavage of  $H_2$ —either heterolytically or on a surface—may be important to reaction with carbide and subsequent enabling of C–C coupling reactivity. This is in contrast to the reactivity of transition metal imides and carbenes, for which direct activation of R–H bonds (R = C, H) is kinetically allowed.<sup>105-107</sup>

Scheme 6. Thermochemical Cycle Evaluating Relevance to Syngas Chemistry

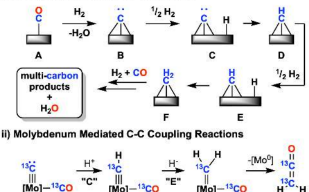


#### CONCLUSION

In summary, a terminal Mo carbide complex has been demonstrated to undergo direct carbonylation, in conjunction with ligand functionalization, affording a phosphoranylideneketene complex. Carbide protonation provides a thermally stable methylidyne carbonyl complex, which, upon sequential treatment with hydride and CO, induces C– C coupling. Experimental and Quantum Mechanics computational interrogation of reactions between the silyl alkylidyne congener and hydride suggest that this C–C bond formation differs from the mechanism established for stepwise reduction/electrophile addition, proceeding through carbene carbonyl intermediates rather than dicarbynes.

Notably, this reaction sequence models, with high fidelity, the proposed mechanism of FT: surrogates of H<sub>2</sub> react with a terminal carbide (1) to afford a reactive methylidene (5) (cf. B - F, Figure 7). This methylidene is the precursor to C-C bond formation, coupling with CO to give ketene. These steps provide valuable precedent for the reactivity proposed for the heterogenous FT process, mimicking the multi-electron C-C coupling reactivity postulated following C-O scission at a well-defined monometallic Mo carbide complex. Critically, the use of hydride as a FT relevant twoelectron reductant provides an efficient strategy to accomplish C-C coupling and C-H bond formation. The reaction of the **1** with weak acid and hydride sources (the products of formal H<sub>2</sub> heterolytic cleavage) to access a methylidene contrasts the lack of reactivity observed when treated with  $\ensuremath{H_2}$ directly, highlighting the need for pre-activation to achieve productive carbide hydrogenation in either the present molecular model system or via cleavage on the catalyst surface in FT.

i) Proposed Mechanism of FT Synthesis



**Figure 7. i)** Schematic representation of the early steps of Fischer-Tropsch synthesis. **ii)** Molecular Mo Complexes Modeling C/CO Coupling to Metal-Free Organics.

"F'

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#### ASSOCIATED CONTENT

"R'

#### Supporting Information

Detailed experimental procedures, full characterization, crystallographic details (CIF), and spectroscopic data. This material is available free of charge via the Internet at http://pubs.acs.org.

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#### Notes

The authors declare no competing financial interests.

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however; nor was it detected for bound ketene complex **11** generated in the presence or absence of BEt<sub>3</sub>. We conclude that O-borane interactions are likely important for stabilizing reaction intermediates during C-C coupling, but that they are not ubiquitously present in the ground-state structures.

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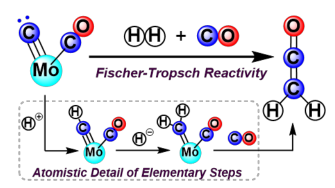
104. Soluble KCl is carried forward from the *in situ* synthesis of **1**, as evidenced by experiments in which **1** was treated with acids bearing  $BAr^{F_4}$  counterions, and *chloride*-bound methylidynes **4/4'** were generated as the primary products (see the SI). Even so, no reaction was observed between carbide **1** and H<sub>2</sub>.

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