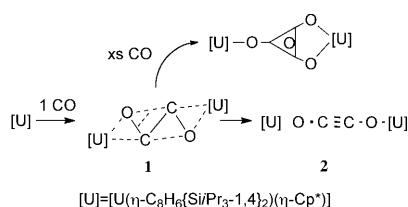


Facile Conversion of CO/H₂ into Methoxide at a Uranium(III) Center**

Alistair S. P. Frey, F. Geoffrey N. Cloke,* Martyn P. Coles, Laurent Maron, and Thomas Davin

The long-established Fischer–Tropsch process^[1] is employed on a very large scale to effect the conversion of synthesis gas (CO/H₂) to hydrocarbons and oxygenates, and continues to attract considerable interest.^[2] The C–C coupling reactions implicit in the latter have been extensively modeled using molecular organometallic systems,^[3] for example, the formation of enediolate complexes^[4] and ethene^[5] from reactions of early-transition-metal or f-block hydrides with CO. In 2006, we reported a novel CO-coupling reaction not previously observed in Fischer–Tropsch processes, namely the reductive cyclotrimerization of CO by the U^{III} complex [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)] to afford the deltate complex [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)]₂(μ-η¹:η²-C₃O₃).^[6] Subsequent computational studies indicated that this reaction proceeds through a proposed “zig-zag” C₂O₂ intermediate **1**, see Scheme 1: in the presence of excess (xs) CO the latter adds

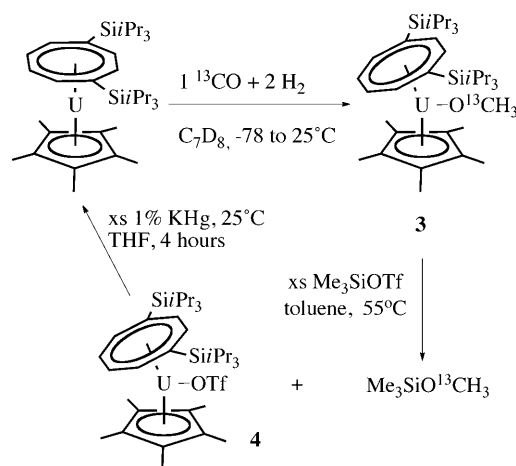


Scheme 1. Mechanism of the reductive cyclotrimerization of CO (xs = excess) by [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)].

a further molecule of CO to form a deltate structure, whereas in the absence of further CO the zig-zag intermediate slowly ($\Delta G^{\ddagger}_{\text{calc}} = 60 \text{ kJ mol}^{-1}$) transforms to the (isolated) linear yne diolate complex [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)]₂(μ-η¹:η¹-C₂O₂) **2**, see Scheme 1.^[7]

In the particular context of the Fischer–Tropsch conversion of CO/H₂ to oxygenates, such as ethylene glycol, we were interested to explore the reactivity of the C–C triple bond in **2** towards dihydrogen with a view to synthesizing the derived ethene or ethane diolate complexes. However, exposure of **2** in [D₈]toluene to excess dihydrogen (10 bar) did not result in

discernible reaction monitored by ¹H and ¹³C NMR spectroscopy, even after prolonged heating (60 °C, 3 days) and UV irradiation; a similar lack of reactivity towards dihydrogen has been noted for the related yne diolate complex [U(N-{SiMe₃})₂]₂(μ-η¹:η¹-C₂O₂).^[8] Instead, we turned our attention to the potential functionalization of the C₂ unit in **2** through reaction of [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)] with ¹³CO in the presence of H₂. Accordingly, [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)] in [D₈]toluene at –78 °C was treated with one equivalent of ¹³CO followed by two equivalents of H₂, with subsequent mixing and warming to room temperature. The ¹³C NMR spectrum of the resultant solution revealed the formation of an essentially sole ¹³C-containing product **3**, characterized by a single quartet resonance at $\delta = 319 \text{ ppm}$, with $J_{\text{CH}} = 137 \text{ Hz}$. Microanalytical, mass spectral and ¹H NMR data were consistent with the formulation of **3** as the U^{IV} methoxide complex [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)OMe], that is, the result of hydrogenation of CO at subambient to ambient temperatures and pressures (see Scheme 2).



Scheme 2. Complex **3** as result of hydrogenation of CO. Complex **4** is reduced back to the U^{III} starting complex with potassium amalgam.

Slow cooling of a toluene solution from **3** to –50 °C gave red-brown crystals suitable for single-crystal X-ray diffraction studies, and the structure is shown in Figure 1, together with selected bond lengths and angles.^[9]

The structure shows the anticipated bent sandwich unit, with a terminal methoxide group. The distances between the metal and the ring centroids in the U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*) fragment (U–M1 2.4887(2) and U–M2 1.95590(2) Å) are identical within the estimated standard deviations (esds) to those for other U^{IV} complexes incorporating this fragment, for example, [U(η-C₈H₆{SiPr₃-1,4}₂)(η-Cp*)]₂(μ-η¹:η²-C₃O₃) (U–M1 2.480(8) and U–M2 1.950(8) Å), although the M1–U–

[*] Dr. A. S. P. Frey, Prof. Dr. F. G. N. Cloke, Dr. M. P. Coles
Division of Chemistry, School of Life Sciences
University of Sussex, Brighton BN1 9QJ (UK)
Fax: (+44) 1273-876-687
E-mail: f.g.cloke@sussex.ac.uk
Prof. Dr. L. Maron, Dr. T. Davin
LPCNO, INSA Toulouse, 31077 Toulouse (France)

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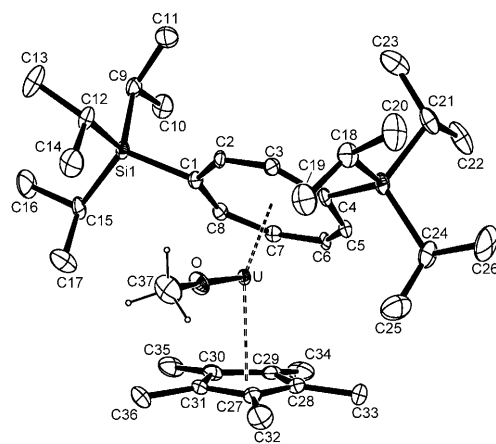


Figure 1. X-ray structure of **3** (thermal ellipsoids at 30%, hydrogens of the OMe group shown, all others removed for clarity). Selected bond lengths and angles: U–M1 2.4887(2), U–M2 1.95590(2), U–O 2.058(4) Å; C37–O–U 178.3(5), M1–U–M2 135.809(9)°. M1 and M2 are the centroids of the five- and eight-membered rings, respectively.

M2 angle in **3** (135.809(9)°) is slightly more acute than that in the latter (141.8(2)°).^[6] The U–OMe linkage in **3** is essentially linear (178.3(5)°) with a U–O distance of 2.058(4) Å, and the structural features are comparable to those found in other (rare) examples of U^{IV} terminal methoxide complexes.^[10]

To gain insight into the bonding situation in **3**, computational studies were carried out at the DFT (B3PW91/SDD-(U,Si)-6-31G(d,p)(C,O,H)) level. The optimized structure is in excellent agreement with the experimental one (see the Supporting Information) indicating that the method is suitable for the description of **3**. In particular, the metal-centroid distances are nicely reproduced (U–M1 2.486 and U–M2 1.958 Å) as well as the M1–U–M2 angle (136.3°). The U–OMe distance is also perfectly reproduced (2.055 Å) as well as the linearity (U–O–C angle of 178.9°). The bonding in complex **3** has been studied using MO and NBO analyses. Both methods indicate a double bond between U and O (see Figure 2 for the occupied MOs), strongly polarized towards



Figure 2. Bonding situation in **3**. Occupied molecular orbitals of the U–OMe bond.

oxygen. At the NBO level, the polarization is highlighted since the two bonds are only defined at the second-order donor–acceptor level (donation from a σ lone-pair orbital of the oxygen into an empty d orbital, 123 kcal mol^{−1}, as well as a donation from a π lone-pair orbital of the oxygen into an empty d orbital, 69 kcal mol^{−1}). The Wiberg bond index of 1.35 is also consistent with some double-bond character. This situation is rather unique since, for example, the U–O bond in

Cp''₂UO (Cp'' = C₅H₂tBu₃) was reported to only exhibit single-bond character.^[11] The Gibbs free-energy of formation of complex **3** is computed to be −76.6 kcal mol^{−1} with respect to the CO adduct (i.e. 2[U]–CO + 3H₂ → 2[U]–OMe, where [U] = [U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)]), in excellent agreement with the experimental observation.

Whilst we have been unsuccessful in liberating methanol from **3** (e.g. by treatment with stoichiometric HCl or HOTf), the OMe group may be smoothly converted into Me₃SiOMe by treatment of **3** with Me₃SiOTf with concomitant formation of the U^{IV} triflate complex [U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)OTf] **4** (see Scheme 2). Complex **4** can be reduced back to the U^{III} starting complex [U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)] with potassium amalgam in THF at a conversion of > 60% as determined by ¹H NMR spectroscopy—the final step in a hypothetical U^{III}-mediated cycle which converts CO + H₂ + Me₃SiOTf to Me₃SiOMe (see Scheme 2).

The conversion of CO/H₂ to methanol (and higher alcohols) is an industrially important process carried out on a Cu/ZnO heterogeneous catalyst and has been extensively investigated.^[12] The reactions of CO with organometallic hydride complexes have also been studied, as potential models for key steps in the heterogeneous reaction. Early work by Bercaw and co-workers showed that the hydrogenation of a metal-bound CO ligand in [Zr(η-Cp*)₂(CO)₂] at 110°C to afford the methoxide complex [Zr(η-Cp*)₂–(OMe)(H)] proceeds through the hydride complex [Zr(η-Cp*)₂(H)₂CO].^[13] More recently, Andersen and co-workers have shown that [Ce(η-C₅H₂tBu₃)₂H] will effect the conversion of CO/H₂ to methoxide, forming [Ce(η-C₅H₂tBu₃)₂OMe], under relatively mild conditions, through a formyl intermediate.^[14] This immediately raises the question as to whether the formation of **3** proceeds through a U^{IV} hydride species, that is, U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)H. However, exposure of a [D₈]toluene solution of [U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)] to excess dihydrogen at 1 bar did not reveal evidence for formation of a hydride and there was no change in the ¹H NMR spectrum of the starting material. Complex [U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)] in [D₈]toluene was then reacted with one equivalent of ¹³CO at −78°C, allowed to warm briefly (1 min) to 20°C and then recooled to −78°C; at this point there is only a trace amount of yne diolate **2** monitored by ¹³C NMR spectroscopy, and under these conditions the proposed, relatively long-lived zig-zag intermediate **1** (see Scheme 1) is likely to be the dominant species in solution.^[7] Exposure of this solution to dihydrogen also results in the formation of the methoxide **3** as essentially the only ¹³CO-derived product. Thus, we suggest that **3** may arise from hydrogenation of the zig-zag intermediate **1**, as opposed to classical hydride reduction of bound CO. Detailed experimental and computational mechanistic studies on the formation of **3** are underway and will be reported in due course.

Experimental Section

Synthesis of 3: Complex [U(η-C₈H₆{SiPr₃-1,4})₂(η-Cp*)THF] (500 mg, 0.574 mmol) was heated under vacuum (45 min, 100°C, 1 × 10^{−6} mbar) to remove coordinated tetrahydrofuran (THF). The desolvated solid was dissolved in toluene (1 mL) to give a black

solution, and the ampoule was cooled to -78°C and attached to a Toepler pump equipped with a gas-addition line. To the cold degassed solution was added ^{13}CO (0.574 mmol) followed by H_2 (1.15 mmol, two equivalents per U); the reaction flask was then sealed and allowed to warm to room temperature overnight. The resulting red-brown solution was then stripped of solvent to provide the title compound as a brown powder (405 mg, 86%). An analytically pure sample, and crystals suitable for X-ray diffraction, were obtained by slow cooling of a pentane or toluene solution of **3** to -50°C . ^{13}C NMR (100 MHz, $[\text{D}_8]\text{toluene}$, 303 K, selected data): $\delta = 319$ ppm, O^{13}CH_3 (quartet, $J_{\text{C-H}} = 137$ Hz). ^1H NMR (400 MHz, $[\text{D}_8]\text{toluene}$, 303 K): $\delta = 142$ (d, 3H, O^{13}CH_3 , $J_{\text{H-C}} = 137$ Hz), 113 (s, 2H, COT ring-CH), -5.54 (br d, 18H, $i\text{Pr-CH}_3$, $J_{\text{H-H}} = 4.6$ Hz), -6.23 (s, 15H, $\text{Cp}^*\text{-CH}_3$), -14.9 (br d, 18H, $i\text{Pr-CH}_3$, $J_{\text{H-H}} = 4.6$ Hz), -18.0 (br m, 6H, $i\text{Pr-CH}$), -40.6 (s, 2H, COT ring-CH), -87.8 ppm (s, 2H, COT ring-CH); elemental analysis calcd (%) for $^{13}\text{C}_{36}\text{H}_{66}\text{OSi}_2\text{U}$: ^{13}C 54.18, H 8.09; found: ^{13}C 54.32, H 8.18; EIMS: m/z (%): 821 (25, M^+).

Reaction of **3** with Me_3SiOTf : To **3** (100 mg, 0.122 mmol) dissolved in $[\text{D}_8]\text{toluene}$ (0.5 mL) was added Me_3SiOTf (0.122 mmol, 27 mg, 22 μL) through a microsyringe, and the mixture was heated at 55°C overnight. The volatile components were transferred under vacuum to afford a red-brown solid residue and a $[\text{D}_8]\text{toluene}$ solution of $\text{Me}_3\text{SiO}^{13}\text{CH}_3$, identified by its NMR spectra. The ^1H , ^{19}F NMR and EI mass spectroscopic data for the solid residue showed it to be $\text{U}(\eta\text{-C}_8\text{H}_6\{\text{Si}(\text{Pr}_3\text{-}1,4)_2\})(\eta\text{-Cp}^*)\text{OTf}$, **4**. ^1H NMR (400 MHz, $[\text{D}_8]\text{toluene}$, 303 K): $\delta = 104$ (s, 2H, COT ring-CH), 13.6 (s, 15H, $\text{Cp}^*\text{-CH}_3$), -4.47 (br d, 18H, $i\text{Pr-CH}_3$), -6.15 (br m, 6H, $i\text{Pr-CH}$), -9.25 (br d, 18H, $i\text{Pr-CH}_3$), -112 (s, 2H, COT ring-CH), -115 ppm (s, 2H, COT ring-CH). ^{19}F NMR (376 MHz, $[\text{D}_8]\text{toluene}$, 303 K): $\delta = -94.4$ ppm (s); EIMS: m/z (%): 938 (3%, M^+).

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