

A New Route to Heterobimetallic C₃ Complexes via C≡C Metatheses of 1,3-Diynyl Ligands: Synthesis and Structure of (η⁵-C₅Me₅)Re(NO)(PPh₃)(C≡CC≡)W(O-*t*-Bu)₃

Roman Dembinski,^{1a,b} Slawomir Szafert,^{1a,c} Pierre Haquette,^{1a} Tadeusz Lis,^{1c} and J. A. Gladysz*,^{1a,d}

Department of Chemistry, University of Utah, Salt Lake City, Utah 84112,
Department of Chemistry, University of Wroclaw, F. Joliot-Curie 14, 50–383 Wroclaw, Poland,
and Institut für Organische Chemie, Friedrich-Alexander Universität Erlangen–Nürnberg,
Henkestrasse 42, 91054 Erlangen, Germany

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Summary: Reactions of (η⁵-C₅Me₅)Re(NO)(PPh₃)(C≡CC≡CR) (R = Me, H) and (t-BuO)₃W≡W(O-*t*-Bu)₃ give (t-BuO)₃W≡CR and the title complex (79–76%), which crystallizes from toluene as a solvated *tert*-butoxy-bridged dimer. The NMR, IR, and crystallographic properties establish dominant Re—C≡C—C≡W as opposed to ⁺Re=C=C=C=W character.

Compounds in which wire-like unsaturated elemental carbon chains span two metals, L_nMC_xM'L'_n (I-C_x), are attracting attention from numerous fundamental and applied viewpoints.^{2–6} To date, C₁, C₂, C₃, C₄, C₅, C₆, C₈, C₁₀, C₁₂, C₁₆, and C₂₀ adducts have been isolated. However, odd carbon chains, which possess unique electronic characteristics,^{3d} remain conspicuously underrepresented. We have described a family of cumulated C₃ complexes of the formula [(η⁵-C₅Me₅)Re(NO)(PPh₃)(C≡C=C≡)M(CO)_a(η⁵-C₅H₅Cl_{5-a}b)]⁺BF₄[–] (M/a/b/c = Mn/2/1/5, Mn/2/1/4, Mn/2/1/0, Fe/4/0/–) and one labile C₅ homolog.^{3d} Templeton has reported labile group 6 cumulated systems, such as 2K⁺[Tp'(CO)₂Mo

(=C=C=C=)W(CO)₂Tp']^{2–}, which can be oxidized to isolable alkynyl/carbyne species, such as [Tp'(O)₂Mo(C≡CC≡)W(CO)₂Tp'].⁵ All other examples feature C₁ bridges.⁴

One factor contributing to this *gerade/ungerade* imbalance is that chains assembled solely by coupling readily available C≡C building blocks must contain even numbers of carbons.^{3d} Odd carbon chains require more sophisticated synthetic strategies. As illustrated in Scheme 1 (reaction a), the above C₃ complexes were prepared from various two-carbon and one-carbon precursors. However, we wondered about the feasibility of degrading ligands with even numbers of sp carbons (reaction b). For example, routes to longer polyynyl complexes, L_nM(C=C)_mR (m = 2–6), are constantly improving.^{3e,7} At the same time, methodologies for C≡C bond metathesis⁸ are rapidly advancing.⁹ In a conceptually pioneering experiment (Scheme 1, reaction c), Selegue demonstrated that RuC≡CMe and RC≡W species could condense to give the ruthenium/tungsten C₁ complex 1.^{4b} Chisholm reported related reactions of platinum alkynyl complexes believed to proceed via PtC≡W intermediates.¹⁰ Accordingly, we set out to probe extensions to longer carbon chains.

As shown in Scheme 2, the chiral rhodium 1,3-pentadiynyl complex (η⁵-C₅Me₅)Re(NO)(PPh₃)(C≡CC≡CMe) (**2a**)^{3a} and ditungsten *tert*-butoxy complex (t-BuO)₃W≡W(O-*t*-Bu)₃ were combined in toluene (1:1 mol ratio). The latter compound is known to participate in a variety of stoichiometric and catalytic C≡C metatheses.^{8–10} After 0.5 h, workup gave a brown-orange air-sensitive powder, which showed properties (below) consistent with the target rhodium/tungsten C₃ complex (η⁵-C₅Me₅)Re(NO)(PPh₃)(C≡CC≡)W(O-*t*-Bu)₃ (**3**), in 79% yield. The butadiynyl complex (η⁵-C₅Me₅)Re(NO)(PPh₃)(C≡CC≡CH) (**2b**),^{3a} which contains a potentially reactive end group but is available in fewer steps than **2a**, was similarly reacted. Workup gave **3** in 76% yield.

(1) (a) University of Utah. (b) Oakland University, Rochester, Michigan (current address). (c) University of Wroclaw (current address). (d) Universität Erlangen–Nürnberg (new permanent address).

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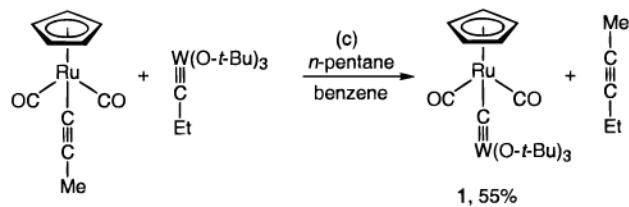
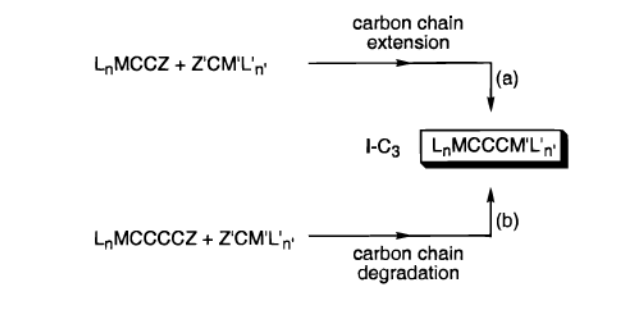
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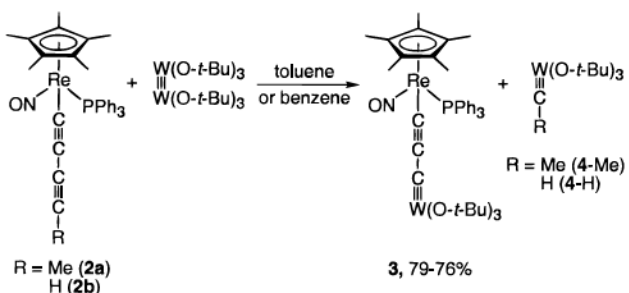
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Scheme 1. (Top) Some Synthetic Approaches to C_3 Complexes; (Bottom) $C\equiv C$ Metathesis Approach to C_1 Complexes



Scheme 2. Syntheses of a Rhenium/Tungsten C_3 Complex



Reactions were monitored by 1H , ^{13}C , and ^{31}P NMR in C_6D_6 . No intermediates or byproducts were detected, and the ^{13}C NMR signals of the tungsten coproduct (t -BuO) $_3$ W=CMe (**4**-Me) closely matched literature values.^{8b}

The 1H and ^{13}C NMR spectra supported the structural assignment **3**. First, the C_5Me_5 and t -Bu 1H signals gave the correct relative integration (15:27). Second, the ReC ^{13}C signal showed a chemical shift (94 ppm) and J_{CP} value (17.8 Hz) characteristic of an alkynyl ligand.^{3e,7} Distinctive changes occur with even slight $^{+}Re=C$ character.³ Third, the CW ^{13}C signal exhibited a chemical shift (236 ppm, s) characteristic of a carbyne ligand, similar to those of **4**-Me and Templeton's MC=CC=M' systems (254–246 ppm).⁵ The central carbon gave an intermediate chemical shift (146 ppm, s). The IR ν_{NO} value, which is also a function of $^{+}Re=C$ character, was essentially identical to those of **2a,b** (1643–1647 vs 1644–1651 cm^{-1}).^{3a,e} A strong $\nu_{C=C}$ band was also observed (1960–1973 cm^{-1}). However, efforts to obtain a correct microanalysis or a mass spectral parent ion were unsuccessful. NMR spectra showed no solvate molecules or impurities.

Deep red prisms were grown by slow evaporation of toluene solutions of **3**. A crystal structure was determined as described in the Supporting Information. As shown in Figure 1, a solvate of the *tert*-butoxy-bridged dimer **3** was obtained. The rhenium atoms are of opposite configurations and related by an inversion

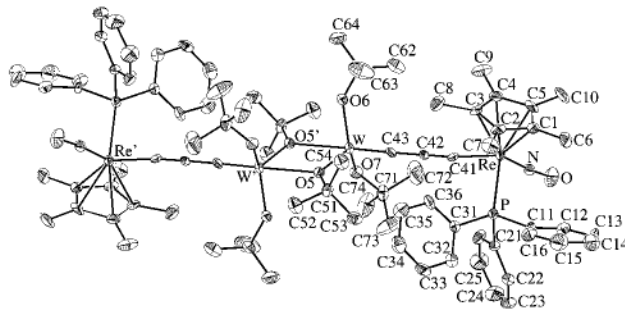


Figure 1. Structure of the dimeric portion of **3**·3·(toluene)₄. Selected interatomic distances and angles: Re–P, 2.369(2); Re–N, 1.753(6); Re–C41, 2.039(8); C41–C42, 1.227(10); C42–C43, 1.398(11); C43–W, 1.769(8); Re–W, 6.422(3); W–O5, 1.927(5); W–O5', 2.492(5); W–O6, 1.881(5); W–O7, 1.880(5). Key angles: N–Re–P, 88.5(2); P–Re–C41, 90.2(2); N–Re–C41, 101.9(3); Re–N–O, 173.0(6); Re–C41–C42, 173.8(6); C41–C42–C43, 179.0(8); W–C43–C42, 178.2(6); C43–W–O5, 102.0(3); C43–W–O5', 177.3(2); C43–W–O6, 102.4(3); C43–W–O7, 101.9(3); W–O5–W', 104.6(2); O5–W–O5', 75.4(2); O5–W–O6, 112.5(2); O5–W–O7, 113.7(2); O5'–W–O6, 78.4(2); O5'–W–O7, 79.8(2); O6–W–O7, 121.0(2).

center (meso isomer). 3D graphics show that the ditungsten core occupies about the same volume as each rhenium end group (32 non-hydrogen atoms each), with no van der Waals contacts. Other complexes with (*t*-BuO)₃W=C moieties have been structurally characterized. Some crystallize as dimers (**4**-Me, **4**-NMe₂),¹¹ whereas others do not (**1**, **4**-Ph).^{4b,12} The tungsten atoms in the dimers adopt distorted trigonal bipyramidal geometries, with axial bridging tungsten–oxygen bonds that are much longer than the three equatorial bridging and nonbridging tungsten–oxygen bonds (2.492(5) vs 1.927(5), 1.881(5), 1.880(5) \AA in **3**·3). Simple nitrogen donor ligands and **4**-R give similar trigonal bipyramidal adducts.^{8,13}

In accord with the solution NMR and IR data, **3**·3 exhibits ReC=C, C=C, and C=CC=C bond lengths (2.039(8), 1.227(10), 1.398(11) \AA) close to those in several (η^5 -C₅Me₅)Re(NO)(PPh₃)(C=CC=CX) species (1.998(12)–2.037(5), 1.202(7)–1.28(2), 1.35(2)–1.389(5) \AA).^{3b,7} The C=W bond length (1.769(8) \AA) is similar to those in **1**, **4**-Me, **4**-NMe₂, and **4**-Ph (1.75(2)–1.77(2) \AA) and related amine adducts.¹³ The ReC=C angle is slightly bent (173.8(6)°), but the C=CC and CC=C angles are nearly linear (179.0(8)°, 178.2(6)°). Hence, the Re–W distance (6.432 \AA) is very close to the sum of the bond lengths (6.432 \AA). Only one other C₃ complex has been structurally characterized to date.^{3d} It exhibits a $^{+}Re=C=C=C=Mn$ linkage and much different NMR and IR properties.^{8,13}

As a powder, **3** melted with decomposition at 121–123 °C. In contrast to $^{+}Re=C=C=C=Mn$ systems, UV-visible spectra showed only featureless tails arising from shoulders at 290/266 nm (benzene/CH₂Cl₂, ϵ 18 000/17 000 $M^{-1}cm^{-1}$). The absence of special metal–metal

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interactions, and the $\text{ReC}\equiv\text{CC}\equiv\text{W}$ bonding model, is further supported by density functional calculations.¹⁴ Finally, the silylalkyne $\text{EtC}\equiv\text{CSiMe}_3$ and $(t\text{-BuO})_3\text{W}\equiv\text{W}(\text{O-}t\text{-Bu})_3$ are known to undergo metathesis.⁸ However, the silylbutadiynyl complex $(\eta^5\text{-C}_5\text{Me}_5)\text{Re}(\text{NO})(\text{PPh}_3)\text{-}(\text{C}\equiv\text{C}\equiv\text{CSiMe}_3)$ ^{3a} gave no reaction over the course of 24 h in benzene at 60 °C, even with the reactive neopentylidyne complex **4**-*t*-Bu.⁸

In conclusion, we have demonstrated the viability of strategy b in Scheme 1 for accessing complexes of the type **I** with $>\text{C}_1$ odd-carbon chains. Furthermore, our approach gives species with $\text{C}\equiv\text{W}$ linkages that should

be metathesis active. These have possibilities as precursors to monometallic complexes with longer chains, as progenitors of other types of bimetallic adducts, and as polymerization catalysts. Efforts toward these ends will be reported in due course.

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Supporting Information Available: Experimental procedures and tables of crystallographic data. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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