

Metathesis of Nitrogen Atoms within Triple Bonds Involving Carbon, Tungsten, and Molybdenum

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 $(Bu^iO)_3Mo\equiv N$ and $W_2(OBu^i)_6(M\equiv M)$ react in hydrocarbons to form $Mo_2(OBu^i)_6(M\equiv M)$ and $(Bu^iO)_3W\equiv N$ via the reactive intermediate $MoW(OBu^i)_6(M\equiv M)$. $(Bu^iO)_3W\equiv N$ and $CH_3C\equiv N^{15}$ react in tetrahydrofuran (THF) at room temperature to give an equilibrium mixture involving $(Bu^iO)_3W\equiv N^{15}$ and $CH_3C\equiv N$. The $(Bu^iO)_3W\equiv N$ compound is similarly shown to act as a catalyst for N^{15} -atom scrambling between $MeC^{13}\equiv N^{15}$ and $PhC\equiv N$ to give a mixture of $MeC^{13}\equiv N$ and $PhC\equiv N^{15}$. From studies of degenerate scrambling of N atoms involving $(Bu^iO)_3W\equiv N$ and $MeC^{13}\equiv N$ in N^{15} -atom spectroscopy, the reaction was found to be first order in acetonitrile and the activation parameters were estimated to be $\Delta H^{\sharp}=13.4(7)$ kcal/mol and $\Delta S^{\sharp}=-32(2)$ eu. A similar reaction is observed for $(Bu^iO)_3Mo\equiv N$ and N^{15} upon heating in N^{15} -atom scale in pyridine solutions and not observed for the dimeric N^{15} -atom scale in N^{15} -and N^{15} -atom scale in N^{15} -

Introduction

The evolution of olefin metathesis involving transition-metal alkylidenes is one of the most fascinating and preparatively useful stories in chemistry. Not long after the initial industrial discovery of olefin metathesis, Pennella et al. disclosed a seemingly even more miraculous tale of triple-bond metathesis involving alkynes. In 1975, Katz and McGinnis addressed the mechanism of olefin metathesis involving metal carbene intermediates and proposed that alkyne metathesis proceeded via a similar mechanism involving the 2 + 2 cycloaddition of C≡C and M≡C bonds to give a planar metallacyclobutadiene intermediate. This was soon thereafter demonstrated by the elegant work of Schrock

and co-workers⁴ involving the reactions between tungsten trialkoxyalkylidynes and -alkynes. This reaction too has entered into the tool kit of organic reagents.⁵ We describe here our findings of the metathesis reaction involving M-N and C-N triple bonds where the metals are either W or Mo. A preliminary report on this work appeared in 2003,⁶ and subsequently related work was reported by Johnson in 2005.^{7,8}

Results and Discussion

Mo≡N + **W≡W** → **Mo≡W** + **W≡N**. The reaction between $(Bu^tO)_3Mo≡N^9$ and $(Bu^tO)_3W≡W(OBu^t)_3^{10}$ proceeds quite rapidly in hydrocarbon solutions to give the

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mixed-metal MoW triply bonded complex MoW(OBut)6 with the formation of (Bu^tO)₃W≡N.¹¹ The compound Mo-W(OBut)6 can readily be identified in the mass spectrum by the presence of a molecular ion that shows the distinct isotopic distribution predicted for a MoW-containing species. It is also seen in the ¹H NMR spectrum where two Bu^t singlets are observed, as is expected for each of the three alkoxide ligands bound to each metal. However, it has not been possible to isolate this heteronuclear MM'-bonded compound because this compound reacts further with (Bu^tO)₃Mo≡N with the ultimate formation of Mo₂(OBu^t)₆ and (ButO)₃W≡N. The compounds Mo₂(OBut)₆, Mo-W(OBut)₆, and W₂(OBut)₆ show similar solubilities and almost certainly cocrystallize. The driving force for this reaction rests with the favorable redox reaction involving the two metals, $Mo^{VI} + W^{III} \rightarrow Mo^{III} + W^{VI}$, and almost certainly proceeds by way of an activated trinuclear cluster involving a triply bridging nitride. Structural precedent for this exists in the form of the tetranuclear cluster $Mo_4(\mu_3$ N)₂(OPrⁱ)₁₂¹² and in related M₃(μ_3 -O)(OR)₁₀, ¹³ MoW₂(μ_3 -O)(OR)₁₀, ¹³ and W₃(μ_3 -P)(OR)₉ clusters. ¹⁴

Alkylidyne and Nitride Ligand Transfer between Molybdenum and Tungsten. Although the triple bonds between the metals Mo and W and between C and N are estimated to be $\sim 120-155$ kcal/mol, 15,16 they are kinetically labile and readily transferable between metal centers as shown in reactions 1–3, where L = Bu^tO.

$$L_3W \equiv {}^{13}CMe + L_3Mo \equiv CPh \rightleftharpoons L_3W \equiv CPh +$$

$$L_3Mo \equiv {}^{13}CMe (1)$$

$$L_3W \equiv^{13}CMe + L_3Mo \equiv N \rightleftharpoons L_3W \equiv N + L_3Mo \equiv^{13}CMe$$
(2)

$$L_3W = {}^{15}N + L_3Mo = N \Rightarrow L_3W = N + L_3Mo = {}^{15}N$$
 (3)

These reactions can be monitored by NMR spectroscopy in solvents such as toluene- d_8 . The presence of ¹⁸³W, $I={}^{1}/_{2}$, which occurs in \sim 15% natural abundance aids in the detection of ¹³C and ¹⁵N nuclei by the presence of ¹ J_{183} W satellites.

These observations led us to interrogate the metathesis reactions involving $M \equiv N$ and $C \equiv N$ described below.

$$W \equiv N^* + RC \equiv N \rightleftharpoons W \equiv N + RC \equiv N^*$$

The compound $(Bu^tO)_3W \equiv ^{15}N$ is readily prepared from the reaction between $W_2(OBu^t)_6$ and labeled acetonitrile $CH_3C \equiv ^{15}N$, and if this reaction is carried out in a hydro-

carbon solvent, the tungsten nitride is formed as a finely divided microcrystalline product. ¹⁷ The compound $(Bu^tO)_3$ -W \equiv N is only sparingly soluble in solvents such as toluene and in the solid state exists as an infinite polymeric chain involving connectivity (\rightarrow W \equiv N \rightarrow W \equiv N \rightarrow). ¹¹ Because of poor solubility in hydrocarbons, our studies of N-atom exchange were carried out in the more polar and coordinating solvent tetrahydrofuran (THF). The molybdenum analogue (Bu tO) $_3$ Mo \equiv N adopts an analogous structure in the solid state but is somewhat more soluble in hydrocarbon solvents because of the weaker donor strength of the less polar Mo \equiv N bond.

The reaction between MeC \equiv ¹⁵N and (Bu^tO)₃W \equiv N in THF- d_8 was followed by ¹⁵N NMR. The signal at $\delta \sim 248$ ppm decreased in intensity as the signal due to (Bu^tO)₃W \equiv ¹⁵N grew in intensity at $\delta \sim 732$ ppm. A similar reaction was carried out employing (Bu^tO)₃Mo \equiv N in THF- d_8 , but no reaction was observed at room temperature. However, heating the solution to 70 °C did bring about a similar reaction with the formation of (Bu^tO)₃Mo \equiv ¹⁵N.

The tungsten nitride complex was then used as a catalyst for the N-atom exchange as shown in reaction 4. The reactions were again monitored by ¹⁵N NMR spectroscopy.

$$MeC \equiv ^{15}N + RC \equiv N \stackrel{cat.}{\rightleftharpoons} MeC \equiv N + RC \equiv ^{15}N$$
 (4)

A broad range of nitriles was chosen both to test the generality of the N-atom exchange and to test the tolerance of $(Bu^iO)_3W\equiv N$ toward various functional groups. For the groups where R= ortho and $p\text{-}C_6H_4F$, $-C_6H_4Cl$, $(CH_2)_4Cl$, Me_3C , Ph, $PhCH_2$, Ph_2CH , Ph_3C , and $-CH\equiv CHCN$ scrambling of the label was observed. The reaction is quite tolerant to steric factors and carbon—halogen as well as $C\equiv C$ bonds. Rather interestingly, the o-fluorophenyl nitrile showed $^{15}N^{-19}F$ coupling, with its signal at $\delta=271.7$ ppm appearing as a doublet (see Figure 1). At this point, it is worth noting that Johnson has recently demonstrated that alkynes will react with fluorinated alkoxide molybdenum and tungsten nitrides to yield the respective alkylidynes and nitriles.

In order to establish that in reaction 4 the metathesis involved solely N-atom exchange and not a cyano group exchange, we also examined the reaction between Me¹³C=N, MeC≡¹⁵N, and PhC≡N. As shown in Figure 2, the exchange catalyzed by (Bu¹O)₃W≡N involved exclusively W≡N and C≡N metathesis.

The exchange reactions noted above do not occur in pyridine- d_5 , which is a good donor to tungsten and presumably blocks coordination of the weaker nitrile donor. Also the siloxide-bridged dimeric compound [(Bu^tMe₂SiO)₃-W \equiv N]₂¹⁸ does not react with MeC \equiv ¹⁵N.

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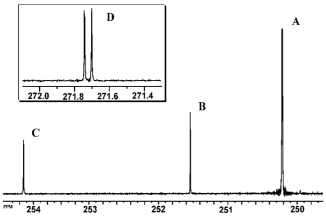


Figure 1. ¹⁵N NMR spectra of Cl(CH₂) $_4$ C=¹⁵N (**B**), p-Cl(C $_6$ H $_4$)C=¹⁵N (**C**), and o-F(C $_6$ H $_4$)C=¹⁵N (**D**), byproducts of a scrambling reaction with (1 BuO) $_3$ W=N and MeC=¹⁵N (**A**).

The chiral menthoxide complexes, both [(+)-men- $O]_3W\equiv N^{19}$ and [(-)-men- $O]_3W\equiv N^{19}$, were prepared and shown to catalyze exchange of N between MeC= ^{15}N and the bulky enantiomerically pure (S)-2-(6-methoxy-2-napthyl)propionitrile (see reaction 5). This raises the possibility of enantioselective N-atom exchange, but we have not investigated this further at this time.

Estimation of Activation Parameters. The atom exchange reaction proceeds at convenient rates under ambient conditions to allow its study by a variety of conventional techniques. However, the specifics of the system proved problematic. For example, the limited solubility and the high air sensitivity of (Bu^tO)₃W≡N made following the course of the reaction by react-IR unreliable. Similarly, mass spectrometry was complicated by relative intensities of RCN⁺ and RCNH⁺ ions due to experimental conditions, and ¹⁵N NMR required too long acquisition times. We finally resorted to the use of ¹³C NMR spectroscopy to follow the reaction between Me¹³C \equiv N and (Bu^tO)₃W \equiv ¹⁵N. The ¹³C NMR signal for Me¹³C \equiv N appears as a singlet at $\delta = 116.96$ ppm and upon ¹⁵N-atom exchange coupling to ¹⁵N, $I = \frac{1}{2}$, leads to a doublet at $\delta = 117.01$ and 116.77 ppm. The line-shape analysis program in XWIN-NMR makes it possible to extrapolate relative concentrations of Me¹³C≡N and Me¹³C≡¹⁵N with time and, hence, obtain data for kinetics. By keeping the concentration of (Bu^tO)₃W≡¹⁵N constant and varying the concentration of nitrile, it was determined that the reaction is first order in nitrile (see the Experimental Section). A variable-temperature study (20–50 °C) following the decay of Me¹³C≡N allowed for the production of the Eyring plot shown in Figure 3. Based on this, an estimation of the activation parameters can be made: $\Delta H^{\ddagger} = +13.4(7)$ kcal/mol and $\Delta S^{\ddagger} = -32(2)$ eu.

Although the alkyne metathesis reaction has been well explored in terms of its synthetic utility and reaction pathway, to our knowledge, the kinetics and activation parameters have only been determined for a related reaction involving the exchange of alkyne units and the metallacyclobutadiene complex, (DIPP)₃W(C₃R₃), where R = Et and Pr^n and DIPP = 2,6-diisopropyl phenoxide, with 3-hexyne- d_{10}^{20} (see reaction 6, where R = Et and Pr^n and $R' = CD_2CD_3$).

$$(DIPP)_3W(C_3R_3) + R'C \equiv CR' \rightarrow (DIPP)_3W(C_3RR'_2) + RC \equiv CR (6)$$

For R = Et, a variable-temperature study led to the determination of $\Delta H^{\ddagger} = +26.1(4)$ kcal/mol and $\Delta S^{\ddagger} = +15.2(15)$ eu, while for R = Prⁿ, $\Delta H^{\ddagger} = +25.4(5)$ kcal/mol and $\Delta S^{\ddagger} = +16.3(16)$ eu.²⁰ The reaction was also shown to be first order in tungsten and independent of 3-hexyne- d_{10} in the concentration range studied. Thus, the activation parameters reported by Schrock and co-workers pertain to a rate-limiting elimination of RC=CR from the metallacy-clobutadiene, whereas in our study, the parameters most likely reflect an associative 2 + 2 reaction. Qualitatively, we can see that this is reflected in the positive entropy of activation for reaction 6 and the negative ΔS^{\ddagger} value for the N-atom metathesis reaction.

Theoretical and Computational Considerations. Following the initial theoretical considerations of Bursten²¹ in 1983, considerable attention has been given to the relative stabilities and bonding in metallacyclobutadiene and metallatetrahedrane complexes and their role in the alkyne metathesis reaction. ^{22–24} This work has led up to the most recent 2006 publication²⁵ dealing with the total reaction pathway for alkyne metathesis by molybdenum and tungsten (RO)₃M≡CMe complexes as determined by density functional theory employing B3LYP and the effective core potentials of Hay and Walt with a double-ξ valence basis set including one f polarization function for Mo and W atoms

We have employed density functional theory using exchange and gradient-corrected correlation functional PW91 to investigate the reaction pathway leading to the N-atom metathesis reactions described above. Two kinds of basis set systems, BS-I (LanL2DZ basis sets augmented by an f polarization function for metal atoms and the 6-31G* basis sets for nonmetal atoms) and BS-II (SDD basis sets for metal atoms and cc-pVTZ basis sets for nonmetal atoms), were used to study the potential energy surfaces of the metathesis reactions between (MeO)₃M≡N (where M = W, Mo) and MeC≡N. The reaction pathways found by employing a larger basis set system, BS-II, are similar to those found by using

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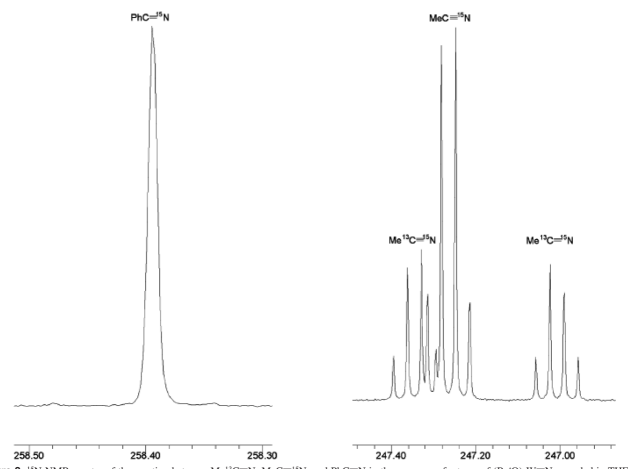


Figure 2. 15 N NMR spectra of the reaction between Me 13 C \equiv N, MeC \equiv 15 N, and PhC \equiv N in the presence of a trace of (Bu 10 O) $_3$ W \equiv N recorded in THF- d_8 at 298 K, 50.6 MHz. The PhC \equiv 15 N signal shows enhancement due to 15 N-atom exchange and appears as a singlet due to lack of coupling to 14 H or 13 C, whereas the MeC \equiv N 15 N signal shows coupling to 14 H, $^{3}J^{1}$ H $^{-15}N = 1.7$ Hz, and for Me 13 C \equiv 15 N coupling to 13 C, $^{1}J^{13}$ C $^{-15}N = 17$ Hz. The signal thus appears as a central 1:3:3:1 quartet flanked by 13 C satellites. The unsymmetrical nature of the 13 C satellites arises from 12 C 13 C isotopic chemical shift perturbation.

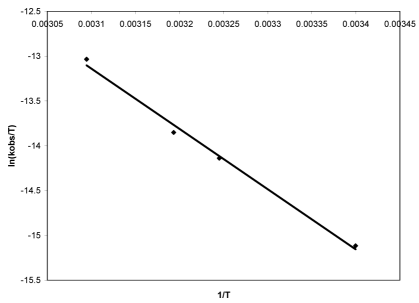


Figure 3. Eyring plot of $\ln(k_{\text{obs}}/T)$ vs 1/T where y = -6727x + 7.7142 and $R^2 = 0.9937$. The enthalpy parameters were calculated to be $\Delta H^{\ddagger} = +13.4(7)$ kcal/mol and $\Delta S^{\ddagger} = -32(2)$ eu for the ^{15}N isotope exchange reaction between (^{15}BuO)₃W¹⁵N and Me¹³CN.

BS-I. For both basis set systems, the same number of stationary points with similar structures on the reaction pathway was found for each reaction. The slightly higher activation free energies obtained by using BS-II than by using BS-I are mainly due to the lack of an f polarization function

for metal atoms in BS-II. This is evident by comparing the activation free energies for the reaction between (MeO)₃W≡N and MeC≡N obtained from BS-I, BS-II, BS-III (LanL2DZ for metal atoms and 6-31G* for nonmetal atoms), and BS-IV (SDD for metal atoms and 6-31G* for

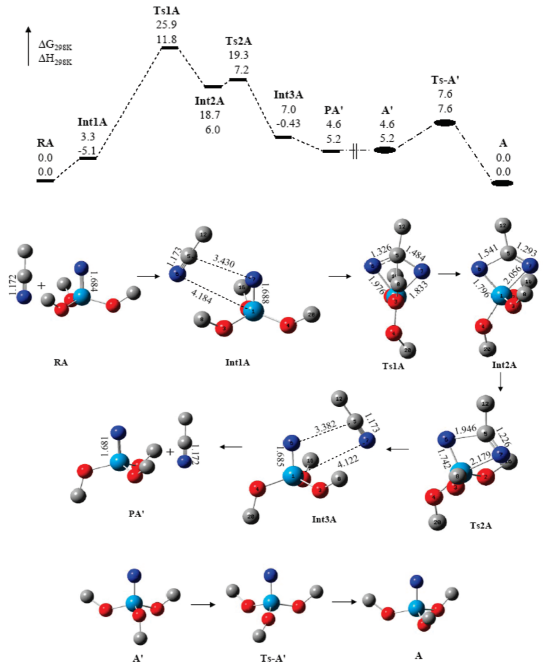


Figure 4. Free energy (at 298 K) profiles for the N-atom exchange reaction between (MeO)₃W≡N and MeC≡N and structures of transition states and intermediates with selected calculated structural parameters (Å).

nonmetal atoms). The activation free energy is defined by the difference of free energies between the high-lying transition structure (TS1A) and the starting materials. The calculated activation free energies from BS-I to BS-IV are 25.9, 31.7, 29.4, and 29.3 kcal/mol, respectively. The almost same values based on BS-III and BS-IV indicate that the different effective core potentials seem to have no effect on the activation free energy for this calculation. Employment of an the f function brings down the energy by about 3.5 kcal/mol, while employment of the triple basis sets cc-pVTZ for nonmetal atoms slightly increases the activation free energy but does not have a significant effect on the reaction pathway. The calculated activation free energy

by using BS-I is closest to the experimental value. On the basis of the above facts, we presented our results calculated by using BS-I.

The calculated free energy profiles for the N-atom exchange reaction between $(MeO)_3M\equiv N$ and $MeC\equiv N$ and structures of transition states and intermediates with selected calculated structural parameters (Å) for W and Mo using BS-I are shown in Figures 4 and 5, respectively.

The overall reaction between $(MeO)_3W\equiv N$ and $MeC\equiv N$ involves N-atom exchange and a conformation change. The N-atom exchange is endothermic $(\Delta G = 4.6 \text{ kcal/mol}, \text{ which}$ is principally caused by the presence of the different conformations of $(MeO)_3W\equiv N$ in RA and PA'). In this step,

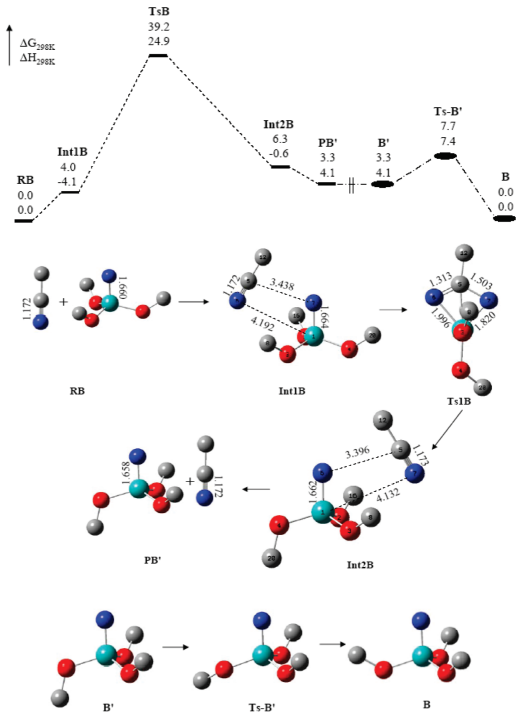


Figure 5. Free energy (at 298 K) profiles for the N-atom exchange reaction between $(MeO)_3Mo \equiv N$ and $MeC \equiv N$ and structures of transition states and intermediates with selected calculated structural parameters (\mathring{A}) .

two transition states, Ts1A and Ts2A, are located. The higher-lying transition state Ts1A connects minima Int1A and Int2A. The intermediate Int2A is transformed into another intermediate Int3A through transition state Ts2A. However, we shall not overemphasize the existence of intermediate Int2A and Ts2A because the barrier for transformation from Int2A to Int3A is only about 0.6 kcal/mol and the analogous intermediate and transition state were not found in the reaction pathway involving the molybdenum analogue. Dissociation of Int1A and Int3A, leading to the formation of separate tungsten nitride and acetonitrile, can

occur without barriers. Int1A has a free energy of 3.3 kcal/mol, which is essentially caused by the loss of entropy associated with the interaction between (MeO)₃W≡N and MeC≡N. The higher free energy of Int3A than that of PA' is also caused by such a loss of entropy. As can be seen in Figure 4, different conformations of (MeO)₃W≡N are present in the product of the N-atom exchange step, PA', and the reactant, RA. The conformer A' in PA' is calculated to be higher in free energy than conformer A in RA by 4.6 kcal/mol, which accounts for the difference of the free energies between RA and PA'. A' can be readily transformed into

the more stable conformer A by crossing an energy barrier of about 3 kcal/mol. A similar metathesis pathway was calculated for the reaction between (MeO)₃Mo≡N and MeC≡N except that neither the analogue of Int2A nor the analogue of TS2A was found in this pathway. The transition state Ts1B connects minima Int1B and Int2B and further dissociates to form (MeO)₃Mo≡N and MeC≡N. Similar conformation transfer from B' to B was also observed.

The transition state Ts1A was calculated to be higher in free energy than the starting materials by 25.9 kcal/mol (about 12 kcal/mol in enthalpy). This calculated free-energy barrier for the N-atom exchange reaction between (MeO)₃W≡N and MeC≡N is in good agreement with the experimental observations, while for the reaction involving (MeO)₃Mo≡N and MeC≡N, the transition state Ts1B was calculated to be higher in free energy than the starting materials by 39.2 kcal/mol, which is about 13 kcal/mol higher than that for the tungsten analogue. This result is also consistent with the experimental observation that N-atom exchange between (Bu¹O)₃Mo≡N and MeC≡N only occurs at elevated temperatures.

The calculated pathways leading to N-atom metathesis and alkyne metathesis bear the similarity that reactions proceed through 2 + 2 cycloaddition but that differences exist between them. In both types of metathesis reactions, the ringclosing step is the rate-determining step. However, no metal alkyne adduct can be located along the pathway for alkyne metathesis, and this was attributed to the fact that the d⁰ metal center cannot stabilize an alkyne complex by back-donation. Metal nitrile complexes (i.e., Int1A and Int1B) were found along the reaction coordinates for the N-atom exchange reactions, and the formation of metal nitrile complexes is probably due to the dipole—dipole interactions and weak van der Waals interactions between the metal nitride fragment and nitrile because the distance between the two fragments is fairly large and both fragments are almost unperturbed. Density functional theory may have a problem handling such interactions, and a higher level of theoretical considerations might be needed to study these weakly associated complexes.

Concluding Remarks

The facility of N-atom metathesis described herein together with the recent report by Johnson et al. ^{7,8} on the metathesis involving Mo \equiv N bonds and alkynes to form molybdenum alkylidynes and organic nitriles shows the kinetic lability of these strong M \equiv N bonds. If these reactions can be coupled to Cummins and co-workers ^{26–29} reductive cleavage of N \equiv N by Mo(NR₂)₃ compounds, then a relatively simple and inexpensive route to ¹⁵N-labeled organic compounds will be

achieved. Interestingly, the compound (${}^{1}BuO$) $_{3}CrN^{30}$ does not appear to entertain this type of chemistry, though in reactions with $M_{2}(O^{1}Bu)_{6}$, redox chemistry is observed with the formation of the blue volatile compound $Cr(O^{1}Bu)_{4}^{31}$ along with the compounds (${}^{1}BuO$) $_{3}MN$ as identified by their molecular ions in the mass spectrometer.

Experimental Section

General Procedures. All operations were carried out under an inert atmosphere of argon with dry and deoxygenated solvents using standard Schlenk-line and drybox procedures. All chemicals were purchased from commercial sources and were used as received unless duly noted. Aromatic and aliphatic hydrocarbon solvents were dried over either sodium or potassium, and THF and diethyl ether were distilled from sodium benzophenone. CH₃CN was dried over CaH₂ and distilled prior to use. All distilled solvents were stored over 4 Å molecular sieves. The nitriles employed in the N-atom exchange reactions were stored over 4 Å sieves and were subjected to three freeze—thaw—degas cycles prior to use. Deuterated solvents and ¹³C/¹⁵N-labeled isotopically enriched compounds were obtained from Cambridge Isotope Laboratories, Inc., and were similarly dried and degassed prior to use.

Syntheses. The compounds $W_2(OR)_6$, where $R = OBu^{t,10}$ (+)-menthyl, ¹⁹ and (-)-menthyl ¹⁹ were prepared by alcoholysis reactions involving $W_2(NMe_2)$, ³² (Bu^tO)₃CrN, ³⁰ (Bu^tO)₃MoN, ⁹ and (Bu^tO)₃MoCPh, ³³ were prepared according to literature procedures.

Spectroscopic Methods. ¹H, ¹⁹F, ¹³C{¹H}, and ¹⁵N NMR spectra were recorded on Bruker Avance 400 and 500 MHz spectrometers. IR spectra were recorded on a Perkin-Elmer GX FT-IR spectrophotometer as KBr pellets. Mass spectra were recorded on a Kratos MS25 RFA double-focusing magnetic sector mass spectrometer as positive ions.

Computational Methods. All calculations were performed using density functional theory as implemented in the *Gaussian 03* suite of programs.³⁴ The PW91PW91³⁵ [Perdew and Wang's 1991 exchange and gradient-corrected correlation functional] density functional theory was used for all calculations. Two types of basis set systems, BS-I and BS-II were used for potential energy surface calculations. In BS-I, the LanL2DZ³⁶ basis sets augmented by an f polarization function [ζ (f) = 0.823, W; 1.043, Mo]³⁷ were used for transition-metal atoms and the 6-31G* basis sets³⁸ were used for all other nonmetal atoms in the model compounds. In BS-II,

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the SDD basis sets³⁹ were used for transition-metal atoms and the cc-pVTZ basis sets⁴⁰ were used for all other nonmetal atoms. Two other basis set systems, BS-III (LanL2DZ for metal atoms and 6-31G* for nonmetal atoms) and BS-IV (SDD for metal atoms and 6-31G* for nonmetal atoms), were also used for activation energy calculations to assess the influences of different effective core potential and the f polarization functions. All of the structures were fully optimized. Frequency calculations were also performed to confirm that all of the stationary points were minima or transition states (no imaginary frequency for the minimum and one imaginary frequency for the transition state). Intrinsic reaction coordinate⁴¹ calculations were carried out on transition states to confirm that these structures are indeed connecting two minima. The discussed energies are relative Gibbs free energies ($\Delta G_{298 \text{ K}}$). The relative enthalpies ($\Delta H_{298 \text{ K}}$) are also provided for reference. All of the relative energies were defined with respect to the starting materials.

Synthesis of $(O^tBu)_3W\equiv^{15}N$. $W_2({}^tBuO)_6$ (1.00 g, 1.24 mmol) was dissolved in hexane (250 mL) with rapid stirring. A hexane solution (10 mL) containing MeC¹⁵N (0.4 g, 9.5 mmol) was added to the dark-red solution with rapid stirring. Immediately after the addition of acetonitrile, the color changed to dark amber and a brown precipitate was formed. The brown precipitate [(${}^tBuO)_3W\equiv^{15}N$] was collected over an air-sensitive filter frit, washed with three 10 mL portions of hexane, and dried in vacuo (500 mg, 96% yield). tH NMR (500 MHz, THF- d_8): δ 1.49 (s, 9H, OC(CH_3)₃). tSN NMR (500 MHz, THF- d_8): δ 732 [(s), $J({}^{t83}W={}^{tSN})$ = 54 Hz]. IR (KBr plates, Nujol mull): 934 cm⁻¹ [ν ($W^{15}N$)]; the 1020 cm⁻¹ band attributed to ν (WN) for the unlabeled derivative was absent.

Synthesis of [(+)-**Men-O**]₃**W** \equiv **N.** [(+)-Men-O]₃**W** \equiv CEt (500 mg, 0.72 mmol) was dissolved in hexane (150 mL) with rapid stirring. Benzonitrile (1 mL) was added via a gastight syringe to the stirring hexane solution. The vessel was placed under vacuum, sealed, and stirred for 24 h. After stirring, the volume was reduced to ca. 2 mL of hexane. Acetonitrile (5 mL) was added, and a brown jellylike solid was formed. The solid was titurated (15 min) until a brown powder formed. The brown solid was filtered, washed with acetonitrile five times, and dried in vacuo (389 mg, 81% yield). 1 H NMR (500 MHz, $C_{6}D_{6}$): δ 4.77 (m, 1H), 2.91 (d, 1H), 2.54 (m, 1H), 1.82 (d, 2H), 1.58 (qt, 2H), 1.34 (d, 3H), 1.19 and 1.15 (d of

d, 8H), 1.0 (m, 1H). 13 C NMR (500 MHz, C_6D_6): δ 88.41, 51.59, 45.49, 35.12, 32.28, 25.87, 23.23, 23.13, 21.59, 16.45. IR (KBr plates, Nujol mull): 1023 cm $^{-1}$ [ν (WN)].

Synthesis of [(−)-Men-O]₃W≡N. [(−)-Men-O]₃W≡CEt (750 mg, 1.09 mmol) was dissolved in hexane (150 mL) with rapid stirring. Benzonitrile (1 mL) was added via a gastight syringe to the stirring hexane solution. The vessel was placed under vacuum, sealed, and stirred for 24 h. After stirring, the volume was reduced to 2 mL of hexane. Acetonitrile (5 mL) was added, and a brown jellylike solid was formed. The solid was titurated (15 min) until a brown powder formed. The brown solid was filtered, washed with acetonitrile five times, and dried in vacuo (512 mg, 72% yield). 1 H NMR (500 MHz, C_6D_6): δ 4.77 (m, 1H), 2.91 (d, 1H), 2.54 (m, 1H), 1.82 (d, 2H), 1.58 (qt, 2H), 1.34 (d, 3H), 1.19 and 1.15 (d of d, 8H), 1.0 (m, 1H). 13 C NMR (500 MHz, C_6D_6): δ 88.41, 51.59, 45.49, 35.12, 32.28, 25.87, 23.23, 23.13, 21.59, 16.45. IR (KBr plates, Nujol mull): 1021 cm $^{-1}$ [ν (WN)].

Reaction of ('BuO)₃Mo \equiv N with W₂(O'Bu)₆. W₂(O'Bu)₆ (0.25 g) and hexanes (10 mL) were added to a Schlenk flask and allowed to stir. Separately ('BuO)₃Mo \equiv N (0.20 g) and hexanes (10 mL) were added to a second Schlenk flask and allowed to stir. The ('BuO)₃Mo \equiv N solution was added to the W₂(O'Bu)₆ solution via a cannula. The mixture was left to stir at room temperature overnight. A white precipitate formed, and the solution was filtered. The white solid was identified as (Bu^lO)₃WN by NMR and mass spectrometry. The filtrate was concentrated and yielded a red microcrystalline solid. By mass spectrometry, the ions Mo₂(OBu^l)₆⁺ and Mo-W(OBu^l)₆⁺ were identified.

Reaction of $({}^{t}BuO)_{3}Cr \equiv N$ with $Mo_{2}(O{}^{t}Bu)_{6}$. $({}^{t}BuO)_{3}Cr \equiv N$ (0.057 g, 0.2 mmol) was added with hexanes (10 mL) to a Schlenk flask and allowed to stir. Separately, Mo₂(O^tBu)₆ (0.126 g, 0.2 mmol) was added with hexanes (10 mL) to a Schlenk flask and allowed to stir. Both flasks were placed in a dry ice/acetone bath and were allowed to stir for 1 h until the solution had cooled to -78 °C. The (¹BuO)₃Cr≡N solution was slowly (15 min) added via a cannula to the Mo₂(O^tBu)₆ solution. The mixture was kept at −78 °C and was allowed to stir for 2 h. The solvent was removed in vacuo. As the flask slowly warmed, a bluish-green liquid formed around the stopper of the flask. The bluish-green volatile compound was identified as Cr(O^tBu)₄ by comparison of the IR spectrum with that of known Cr(O^tBu)₄. ¹H NMR of the reaction mixture (C₆D₆ solvent, C_6D_5H reference, ppm): δ 1.59 (bs), 1.57 (s), 1.55 (s), 1.48 (s, $({}^{t}BuO)_{3}Mo \equiv N$). The mass spectrum of the dried reaction product showed the $({}^{t}BuO)_{3}Mo \equiv N)^{+}$ ion.

Reaction of $({}^{t}BuO)_{3}Cr \equiv N$ with $W_{2}(O{}^{t}Bu)_{6}$. $({}^{t}BuO)_{3}Cr \equiv N$ (0.063 g, 0.22 mmol) was added with hexanes (10 mL) to a Schlenk flask and allowed to stir. Separately, W₂(O^tBu)₆ (0.177 g, 0.22 mmol) was added with hexanes (10 mL) to a Schlenk flask and allowed to stir. Both flasks were placed in a dry ice/acetone bath and were allowed to stir for 1 h until the solutions had cooled to -78 °C. The (¹BuO)₃Cr≡N solution was slowly (15 min) added via a cannula to the W₂(O^tBu)₆ solution. The mixture was kept at -78 °C and was allowed to stir for 2 h. The solvent was removed in vacuo. As the flask slowly warmed, a bluish-green liquid formed around the stopper of the flask. The bluish-green volatile compound was identified as Cr(OtBu)4 by comparison of the IR spectrum with that of known Cr(O^tBu)₄.³¹ ¹H NMR of the reaction mixture (C₇D₈ solvent, C_7D_7H reference, ppm): δ 3.42 (s), 2.1 (s), 1.59 (s, $(^{t}BuO)_{3}W\equiv N$), 1.48 (s), 1.45 (s). The mass spectrum of the dried reaction product has the correct peak range for (¹BuO)₃W≡N).

Reaction of (${}^{t}BuO$)₃W \equiv ¹⁵N with (${}^{t}BuO$)₃Mo \equiv N. (${}^{t}BuO$)₃W \equiv ¹⁵N (0.0209 g, 0.05 mmol) and (${}^{t}BuO$)₃Mo \equiv N (0.0165 g, 0.05 mmol) were added to a NMR tube and allowed to react in THF- d_8

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at room temperature for 1 week. The ¹H NMR spectrum of the reaction mixture shows no change from the starting materials. The ¹⁵N NMR spectrum of the reaction mixture (THF- d_8 solvent, THF- d_7 reference, ppm): δ 828.8 (s, ('BuO)₃Mo \equiv ¹⁵N), 731.8 ('BuO)₃-W \equiv ¹⁵N). Mass spectral analysis of the reaction products verifies the presence of ('BuO)₃Mo \equiv ¹⁵N.

Reaction of ('BuO)₃W≡CMe with ('BuO)₃Mo≡N. ('BuO)₃W≡CMe (0.0215 g, 0.05 mmol) was made in an NMR tube according to a previously reported method. ('BuO)₃Mo≡N (0.0165 g, 0.05 mmol) was added to the tube and allowed to react at room temperature for 1 week. ¹H NMR of the reaction mixture (C_6D_6 solvent, C_6D_5H reference, ppm): δ 3.57 (s, 3H, J_{HW} = 7.2 Hz, ('BuO)₃W≡CMe), 2.58 (s, 3H, ('BuO)₃Mo≡CMe), 1.48 (s, 27H, ('BuO)₃Mo≡N), 1.46 (s), 1.44 (s, 27H, ('BuO)₃W≡CMe), 1.44 (s), 1.42 (s). $^{13}C\{^{1}H\}$ NMR (C_6D_6 solvent, C_6D_5H reference, ppm): δ 333.1 (s), 279.6 (s, 1H, ('BuO)₃Mo≡CMe), 254.1 (s, 1H, J_{CW} = 306.5 Hz, ('BuO)₃W≡CMe).

Reaction of ('BuO)₃W≡CMe with ('BuO)₃Mo≡CPh. ('BuO)₃W≡CMe (0.0054 g, 0.0125 mmol) was made in an NMR tube according to a previously reported method. ('BuO)₃Mo≡CPh (0.0051 g, 0.0125 mmol) was added to the tube and allowed to react at room temperature for 1 week. ¹H NMR of the reaction mixture (C_6D_6 solvent, C_6D_5H reference, ppm): δ 7.49 (d, 2H, ('BuO)₃Mo≡CPh), 7.33 (d, 2H, ('BuO)₃W≡CPh), 7.23 (t, 2H, ('BuO)₃W≡CPh), 7.09 (t, 2H, ('BuO)₃Mo≡CPh), 6.87 (t, 1H, ('BuO)₃Mo≡CPh), 6.82 (t, 1H, ('BuO)₃W≡CPh), 3.57 (s, 3H, J_{HW} = 7.2 Hz, ('BuO)₃W≡CMe), 2.58 (s, 3H, ('BuO)₃Mo≡CMe), 1.49 (s, 27H, ('BuO)₃Mo≡CPh), 1.48 (s), 1.44 (s, 27H, ('BuO)₃W≡CMe), 1.42 (s, 27H, ('BuO)₃W≡CPh). $^{13}C\{^{1}H\}$ NMR of reaction mixture (C_6D_6 solvent, C_6D_5H reference, ppm): δ 333.1 (s), 307.8 (s), 279.8 (s, 1H, ('BuO)₃Mo≡CPh), 254.1 (s, 1H, J_{CW} = 306.5 Hz, ('BuO)₃W≡CMe).

Reaction of (${}^{4}BuO$)₃W≡C ${}^{4}Pr$ with (${}^{4}BuO$)₃Mo≡N. (${}^{4}BuO$)₃W≡C ${}^{4}Pr$ (0.0229 g, 0.05 mmol) and (${}^{4}BuO$)₃Mo≡N (0.0165 g, 0.05 mmol) were added to a NMR tube and allowed to react at room temperature for 1 week. ${}^{4}H$ NMR of reaction mixture (C_6D_6 solvent, C_6D_5H reference, ppm): δ 4.05 (sept, 1H, J_{HW} = 6.85 Hz, (${}^{4}BuO$)₃W≡C ${}^{4}Pr$), 3.11 (sept, 1H, (${}^{4}BuO$)₃Mo≡C ${}^{4}Pr$), 1.48 (s, 27H, (${}^{4}BuO$)₃Mo≡N), 1.47 (s, 27H, (${}^{4}BuO$)₃Mo≡C ${}^{4}Pr$), 1.46 (s, 27H, (${}^{4}BuO$)₃W≡C ${}^{4}Pr$), 1.45 (s, 27H, (${}^{4}BuO$)₃W≡N), 1.24 (d, 6H, (${}^{4}BuO$)₃W≡C ${}^{4}Pr$), 1.16 (d, 6, (${}^{4}BuO$)₃Mo≡C ${}^{4}Pr$).

Preparation of Samples for the Determination of Kinetics. All samples were prepared in an NMR tube equipped with a J. Young adapter. For all runs, a stock solution of Me¹³CN in THF- d_8 was prepared by weighing 48.5 mg in a 5 mL volumetric flask to produce a 0.231 M solution. For determination of the enthalpy parameters, 890 μ L of THF- d_8 was added to 11.3 mg of (15 BuO) $_3$ W¹⁵N via a gastight syringe and transferred to the NMR tube. Addition of 110 μ L of the stock solution with a gastight syringe followed. Preparation occurred immediately before NMR

spectra were collected. In this instance, the catalyst and nitrile concentrations remained constant throughout the temperature range $(50, 40, 35, \text{ and } 21 \,^{\circ}\text{C})$. In each case, the sample was heated in the probe, and once the overnight run finished, the sample was removed and placed in its respective temperature oil bath for the time. In the case of the determination of the reaction order in nitrile, the amount of catalyst remained the same, while the amount of stock solution was changed according to the desired ratio (cat:nitrile = 1:1, 1:0.5, 1:2).

The nitrile cleavage reaction involving Me¹³CN was monitored by 13 C{ 1 H} NMR spectroscopy. The parameters used were as follows: number of scans (NS = 15), number of data points (TD = 65 536), receiver gain (RG = 2896), and relaxation time (D1 = 75 s). The deconvolution of 13 C NMR spectra was done using XWIN-NMR version 3.5 PL6. The T_1 relaxation times for acetonitrile (15.2 s) 13 C NMR signals were measured at 298 K. The Ernst equation (1) (where $t_1 = T_1$ relaxation time, pw 90 = 90° pulse width, at = acquistion time, and d1 = delay time between pulses). NMR probe temperatures were calibrated with a sample of ethylene glycol, and van Greet's equation was used to determine the optimal pulse widths for the delay time used in NMR experiments.

Calculation of the Kinetics and Activation Parameters. The absolute concentration of the C¹Bu resonance (s, 1.49 ppm) versus the Me¹³C \equiv N methyl resonance (d, 1.95 J= 10.7 Hz). The cleavage and atom exchange was measured by following the disappearance of a singlet at 115.9 ppm (Me¹³C \equiv N) and the appearance of a doublet at 115.95 and 115.82 ppm (Me¹³C \equiv 15N). The Lorentzian deconvolution of ¹³C NMR spectra was performed using XWIN-NMR 3.5 PL6. All of the cleavage and atom exchange reactions were of the type A + B \Leftrightarrow C + D, where the rate law was determined (eq 2) and where $\Delta = [A] - [A]_{eq}$.

When $\ln[(\Delta/(\Delta(1 - 1/K) + [A]_e + [B]_e + (1/K)([C]_e + [D]_e))]$ is plotted versus time, the slope is inserted into eq 2 to solve for k_{obs} . The enthalpy parameters were calculated by plotting $\ln(k_{\text{obs}}/T)$ versus 1/T, where the slope $= -\Delta H^{\ddagger}/R$ and the intercept $= \Delta S^{\ddagger}/R + 23.8$.

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Supporting Information Available: Free energy (at 298 K) profiles for a N-atom exchange reaction between $(MeO)_3M\equiv N$ (where M=W, Mo) and $MeC\equiv N$ and structures of transition states and intermediates with selected calculated structural parameters calculated based on BS-II (Figures S1 and S2). This material is available free of charge via the Internet at http://pubs.acs.org.

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