

Published on Web 07/22/2010

Practical New Silyloxy-Based Alkyne Metathesis Catalysts with Optimized Activity and Selectivity Profiles

Johannes Heppekausen, Robert Stade, Richard Goddard, and Alois Fürstner*

Max-Planck-Institut für Kohlenforschung, D-45470 Mülheim/Ruhr, Germany

Received June 2, 2010; E-mail: fuerstner@kofo.mpg.de

Abstract: Triphenylsilanolate ligands were found to impart excellent reactivity and outstanding functional group tolerance on molybdenum alkylidyne complexes, which catalyze alkyne metathesis reactions of all sorts. The active species either can be obtained in high yield by adaptation of the established synthesis routes leading to Schrock alkylidynes or can be generated in situ from the molybdenum nitride complex 11, which itself is readily accessible in large quantity from inexpensive sodium molybdate. Complexation of the active silanolate complexes 12 and 24 with 1,10-phenanthroline affords complexes 15 and 25, respectively, which are stable in air for extended periods of time. Although these phenathroline adducts are per se unreactive vis-à-vis alkynes, catalytic activity is conveniently restored upon exposure to MnCl₂. Therefore, the practitioner has the choice of different alkyne metathesis (pre)catalysts, which are easy to handle yet broadly applicable and exceedingly tolerant. A host of representative inter- as well as intramolecular alkyne metathesis reactions, including applications to a considerable number of bioactive and, in part, labile natural products, shows the remarkable scope of these new tools. Moreover, it was found that the addition of molecular sieves (5 Å \geq 4 Å \gg 3 Å) to the reaction mixture significantly improves the chemical yields while simultaneously increasing the reaction rates. This benefit is ascribed to effective binding of 2-butyne, which is released as the common byproduct in reactions of alkynes bearing a methyl end-cap. Thus, alkyne metatheses can now be performed at ambient temperature with neither the need to apply vacuum to drive the conversion nor recourse to tailor-made substrates. The structures of representative examples of this new generation of alkyne metathesis catalysts in the solid state were determined by X-ray analysis.

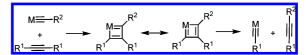
Introduction

A major reason for the overwhelming success of olefin metathesis is the ready availability of catalysts that combine high activity with a good to excellent tolerance toward functional groups other than alkenes. As some of these key catalysts are modular, well accessible, and easy to handle, this transformation was rapidly embraced by the synthetic community and has profoundly changed the way contemporary organic and polymer chemistry are practiced.

Compared to the omnipresence of alkene metathesis, the related metathesis of alkynes is much less commonly used.^{4,5} Initially described as early as 1968,⁶ the principle underlying

 Schrock, R. R.; Murdzek, J. S.; Bazan, G. C.; Robbins, J.; DiMare, M.; O'Regan, M. J. Am. Chem. Soc. 1990, 112, 3875.

Scheme 1. Basic Mechanism of Alkyne Metathesis Catalyzed by Metal Alkylidynes



this scrambling process was swiftly discerned⁷ and experimentally proven (Scheme 1).^{8,9} The active catalysts were shown to be Schrock alkylidyne complexes,¹⁰ which either can be generated in situ or, preferentially, are administered to the reaction mixtures as structurally well-defined, preformed species. Despite this excellent background knowledge, it was only after a considerable lag period¹¹ that the potential of alkyne metathesis

^{(2) (}a) Nguyen, S. T.; Grubbs, R. H.; Ziller, J. W. J. Am. Chem. Soc. 1993, 115, 9858. (b) Schwab, P.; Grubbs, R. H.; Ziller, J. W. J. Am. Chem. Soc. 1996, 118, 100.

 ^{(3) (}a) Trnka, T. M.; Grubbs, R. H. Acc. Chem. Res. 2001, 34, 18. (b) Fürstner, A. Angew. Chem., Int. Ed. 2000, 39, 3012. (c) Connon, S. J.; Blechert, S. Angew. Chem., Int. Ed. 2003, 42, 1900. (d) Nicolaou, K. C.; Bulger, P. G.; Sarlah, D. Angew. Chem., Int. Ed. 2005, 44, 4490. (e) Deiters, A.; Martin, S. F. Chem. Rev. 2004, 104, 2199. (f) Hoveyda, A. H.; Zhugralin, A. R. Nature 2007, 450, 243.

^{(4) (}a) Fürstner, A.; Davies, P. W. Chem. Commun. 2005, 2307. (b) Zhang, W.; Moore, J. S. Adv. Synth. Catal. 2007, 349, 93. (c) Schrock, R. R.; Czekelius, C. Adv. Synth. Catal. 2007, 349, 55. (d) Mortreux, A.; Coutelier, O. J. Mol. Catal. A: Chem. 2006, 254, 96.

^{(5) (}a) Fürstner, A. In *Handbook of Metathesis*; Grubbs, R. H., Ed.; Wiley-VCH: Weinheim, 2003; Vol. 2, p 432. (b) Davies, P. W. In *Handbook of Cyclization Reactions*; Ma, S., Ed.; Wiley-VCH: Weinheim, 2009; Vol. 1, p 599.

⁽⁶⁾ Pennella, F.; Banks, R. L.; Bailey, G. C. *Chem. Commun.* **1968**, 1548.

⁽⁷⁾ Katz, T. J.; McGinnis, J. J. Am. Chem. Soc. **1975**, 97, 1592.

⁽⁸⁾ Wengrovius, J. H.; Sancho, J.; Schrock, R. R. J. Am. Chem. Soc. 1981, 103, 3932.

⁽⁹⁾ Churchill, M. R.; Ziller, J. W.; Freudenberger, J. H.; Schrock, R. R. Organometallics 1984, 3, 1554.

^{(10) (}a) Schrock, R. R. Chem. Rev. 2002, 102, 145. (b) Schrock, R. R. Angew. Chem., Int. Ed. 2006, 45, 3748. (c) Schrock, R. R. J. Chem. Soc., Dalton Trans. 2001, 2541. (d) Schrock, R. R. Acc. Chem. Res. 1986, 19, 342.

was recognized and explored.^{12,13} The fact that nonterminal alkynes are required as the substrates may be one reason for the smaller impact compared with olefin metathesis, ^{14,15} yet a growing number of applications in the recent literature to natural product synthesis, ⁴ coordination chemistry, ¹⁶ and material science ^{17,18} show that this inherent drawback can be (partly) compensated for or even turned to advantage by the excellent selectivity of alkyne metathesis, as well as by the possibility of further elaborating the acetylenic products primarily formed in a diverse fashion by a host of different postmetathetic transformations. ^{19–24}

- (11) For early applications to fairly unfunctionalized substrates, see refs 26, 27 and the following: (a) Villemin, D.; Cadiot, P. *Tetrahedron Lett.* 1982, 23, 5139. (b) Kaneta, N.; Hirai, T.; Mori, M. *Chem. Lett.* 1995, 627. (c) Kaneta, N.; Hikichi, K.; Asaka, S.; Uemura, M.; Mori, M. *Chem. Lett.* 1995, 1055.
- (12) Fürstner, A.; Seidel, G. Angew. Chem., Int. Ed. 1998, 37, 1734.
- (13) Fürstner, A.; Guth, O.; Rumbo, A.; Seidel, G. J. Am. Chem. Soc. 1999, 121, 11108.
- (14) (a) Terminal alkynes are mainly polymerized in the presence of the standard catalysts, cf.: Bray, A.; Mortreux, A.; Petit, F.; Petit, M.; Szymanska-Buzar, T. J. Chem. Soc., Chem. Commun. 1993, 197. (b) Moreover, terminal alkynes are known to degrade Schrock alkylidynes via formation of deprotiometallacyclobutadiene intermediates, which could be unambiguously characterized, cf. ref 32a and the following: McCullough, L. G.; Listemann, M. L.; Schrock, R. R.; Churchill, M. R.; Ziller, J. W. J. Am. Chem. Soc. 1983, 105, 6729.
- (15) For a report claiming metathesis of terminal alkynes by a modified in situ catalyst, see: Coutelier, O.; Nowogrocki, G.; Paul, J.-F.; Mortreux, A. Adv. Synth. Catal. 2007, 349, 2259.
- (16) (a) Bauer, E. B.; Szafert, S.; Hampel, F.; Gladysz, J. A. Organometallics 2003, 22, 2184. (b) Bauer, E. B.; Hampel, F.; Gladysz, J. Adv. Synth. Catal. 2004, 346, 812. (c) Kotora, M.; Neèas, D.; Štipnièka, P. Collect. Czech. Chem. Commun. 2003, 68, 1897. (d) Bobula, T.; Hudlicky, J.; Novák, P.; Gyepes, R.; Cisaøova, I.; Štipnièka, P.; Kotora, M. Eur. J. Inorg. Chem. 2008, 3911. (e) Sato, M.; Watanabe, M. Chem. Commun. 2002, 1574.
- (17) (a) Weiss, K.; Michel, A.; Auth, E.-M.; Bunz, U. H. F.; Mangel, T.; Müllen, K. Angew. Chem., Int. Ed. Engl. 1997, 36, 506. (b) Kloppenburg, L.; Song, D.; Bunz, U. H. F. J. Am. Chem. Soc. 1998, 120, 7973. (c) Zhang, X.-P.; Bazan, G. C. Macromolecules 1994, 27, 4627. (d) Krouse, S. A.; Schrock, R. R. Macromolecules 1989, 22, 2569. (e) Bunz, U. H. F.; Kloppenburg, L. Angew. Chem., Int. Ed. 1999, 38, 478. (f) Bly, R. K.; Dyke, K. M.; Bunz, U. H. F. J. Organomet. Chem. 2005, 690, 825. (g) Brizius, G.; Kroth, S.; Bunz, U. H. F. Macromolecules 2002, 35, 5317. (h) Zhang, W.; Moore, J. S. Macromolecules 2004, 37, 3973. (i) Brizius, G.; Pschirer, N. G.; Steffen, W.; Stitzer, K.; zur Loye, H.-C.; Bunz, U. H. F. J. Am. Chem. Soc. 2000, 122, 12435. (j) Carnes, M.; Buccella, D.; Siegrist, T.; Steigerwald, M. L.; Nuckolls, C. J. Am. Chem. Soc. 2008, 130, 14078.
- (18) (a) Miljanić, O. Š.; Vollhardt, K. P. C.; Whitener, G. D. Synlett 2003, 29. (b) Hellbach, B.; Gleiter, R.; Rominger, F. Synthesis 2003, 2535.
 (c) Johnson, C. A.; Lu, Y.; Haley, M. M. Org. Lett. 2007, 9, 3725.
 (d) Ge, P.-H.; Fu, W.; Herrmann, W. A.; Herdtweck, E.; Campana, C.; Adams, R. D.; Bunz, U. H. F. Angew. Chem., Int. Ed. 2000, 39, 3607. (e) Pschirer, N. G.; Fu, W.; Adams, R. D.; Bunz, U. H. F. Chem. Commun. 2000, 87.
- (19) Ring-closing alkyne metathesis in combination with Lindlar-type semireduction provides a stereoselective entry into macrocyclic (Z)-alkenes that remain difficult to make by conventional RCM; cf. refs 12, 13. For recent advances toward highly Z-selective alkene metathesis catalysts, see: Jiang, A. J.; Zhao, Y.; Schrock, R. R.; Hoveyda, A. H. J. Am. Chem. Soc. 2009, 131, 16630.
- (20) Ring-closing alkyne metathesis in combination with *trans*-hydrosily-lation and protodesilylation provides a stereoselective entry into macrocyclic (E)-alkenes, see: (a) Fürstner, A.; Radkowski, K. Chem. Commun. 2002, 2182. (b) Lacombe, F.; Radkowski, K.; Seidel, G.; Fürstner, A. Tetrahedron 2004, 60, 7315.
- (21) For the formation of an aromatic heterocycle by an RCAM/alkyne activation sequence, see: Fürstner, A.; Castanet, A.-S.; Radkowski, K.; Lehmann, C. W. J. Org. Chem. 2003, 68, 1521.
- (22) For the formation of 1,3-dienes by an alkyne metathesis/enyne cross-metathesis sequence, see: (a) Fürstner, A.; Larionov, O.; Flügge, S. Angew. Chem., Int. Ed. 2007, 46, 5545. (b) Fürstner, A.; Flügge, S.; Larionov, O.; Takahashi, Y.; Kubota, T.; Kobayashi, J. Chem.—Eur. J. 2009, 15, 4011. (c) Groaz, E.; Banti, D.; North, M. Eur. J. Org. Chem. 2007, 3727.

Alkyne metathesis was originally discovered using a heterogeneous catalyst composed of tungsten oxide on silica which was operative only at 200–450 °C.^{6,25} Shortly thereafter, Mortreux and co-workers showed that homogeneous mixtures comprising Mo(CO)₆ and resorcinol (or other phenols) in highboiling solvents are active at more manageable temperatures (ca. 130–160 °C).²⁶ Despite many rounds of optimization of the molybdenum source, the phenol additive, the solvent, and the reaction conditions, this simple and cheap system has not reached the level of activity and selectivity that better-defined (pre)catalysts can offer.^{27–29} It was the advent of well-defined d⁰-alkylidyne complexes of tungsten, ^{8,30,31} molybdenum, ^{32,33} and rhenium^{34,35} that set new standards in the field. Among them,

- (23) In addition to the established chemistry of alkynes, recent advances in the carbophilic activation of triple bonds with noble metal catalysts hold particular promise in this regard; see: (a) Fürstner, A.; Davies, P. W. Angew. Chem., Int. Ed. 2007, 46, 3410. (b) Gorin, D. J.; Toste, D. Nature 2007, 446, 395. (c) Jiménez-Núñez, E.; Echavarren, A. M. Chem. Commun. 2007, 333. (d) Hashmi, A. S. K. Chem. Rev. 2007, 107, 3180. (e) Fürstner, A. Chem. Soc. Rev. 2009, 38, 3208.
- (24) The established alkyne metathesis catalysts do not react with alkenes, whereas alkene metathesis catalysts can attack alkynes. For an early application exploiting this orthogonal selectivity profile, see: Fürstner, A.; Dierkes, T. Org. Lett. 2000, 2, 2463.
- (25) For further studies on heterogeneous catalysts for alkyne metathesis, see ref 53 and the following: (a) Mortreux, A.; Blanchard, M. Bull. Soc. Chim. Fr. 1972, 1641. (b) Moulijn, J. A.; Reitsma, H. J.; Boelhouwer, C. J. Catal. 1972, 25, 434. (c) Mortreux, A.; Petit, F.; Blanchard, M. J. Mol. Catal. 1980, 8, 97.
- (26) (a) Mortreux, A.; Blanchard, M. J. Chem. Soc., Chem. Commun. 1974, 786. (b) Mortreux, A.; Dy, N.; Blanchard, M. J. Mol. Catal. 1975/76, I, 101. (c) Mortreux, A.; Delgrange, J. C.; Blanchard, M.; Lubochinsky, B. J. Mol. Catal. 1977, 2, 73. (d) Mortreux, A.; Petit, F. Tetrahedron Lett. 1978, 4967.
- (27) (a) Du Plessis, J. A. K.; Vosloo, H. C. M. J. Mol. Catal. 1991, 65, 51. (b) Vosloo, H. C. M.; du Plessis, J. A. K. J. Mol. Catal. A: Chem. 1998, 133, 205. (c) Bencheick, A.; Petit, M.; Mortreux, A.; Petit, F. J. Mol. Catal. 1982, 15, 93. (d) Pschirer, N. G.; Bunz, U. H. F. Tetrahedron Lett. 1999, 40, 2481. (e) Brizius, G.; Bunz, U. H. F. Org. Lett. 2002, 4, 2829. (f) Sashuk, V.; Ignatowska, J.; Grela, K. J. Org. Chem. 2004, 69, 7748.
- (28) For a direct comparison of a Mortreux-type catalyst with more defined catalysts, see refs 13 and 46d. For an example in which an attempted alkyne metathesis with a Mortreux-type catalyst failed because of a competing reaction of the substrate with the phenol additive, see: Ma, J.; Kühn, B.; Hackl, T.; Butenschön, H. Chem.—Eur. J. 2010, 16, 1859.
- (29) In most cases the nature of the active catalyst in solution is unknown, even though it might be guessed on the basis of the extensive knowledge of the chemistry of metal alkylidynes. Therefore, the expressions "catalyst" and "precatalyst" are not rigorously distinguished in this publication.
- (30) (a) Schrock, R. R.; Clark, D. N.; Sancho, J.; Wengrovius, J. H.; Rocklage, S. M.; Pedersen, S. F. Organometallics 1982, I, 1645. (b) Pedersen, S. F.; Schrock, R. R.; Churchill, M. R.; Wasserman, H. J. J. Am. Chem. Soc. 1982, 104, 6808. (c) Freudenberger, J. H.; Schrock, R. R.; Churchill, M. R.; Rheingold, A. L.; Ziller, J. W. Organometallics 1984, 3, 1563. (d) Listemann, M. L.; Schrock, R. R. Organometallics 1985, 4, 74. (e) Tonzetich, Z. J.; Lam, Y. C.; Müller, P.; Schrock, R. R. Organometallics 2007, 26, 475.
- (31) (a) Beer, S.; Brandhorst, K.; Hrib, C. G.; Wu, X.; Haberlag, B.; Grunenberg, J.; Jones, P. G.; Tamm, M. Organometallics 2009, 28, 1534. (b) Beer, S.; Hrib, C. G.; Jones, P. G.; Brandhorst, K.; Grunenberg, J.; Tamm, M. Angew. Chem., Int. Ed. 2007, 46, 8890. (c) Beer, S.; Brandhorst, K.; Grunenberg, J.; Hrib, C. G.; Jones, P. G.; Tamm, M. Org. Lett. 2008, 10, 981.
- (32) (a) McCullough, L. G.; Schrock, R. R.; Dewan, J. C.; Murdzek, J. C. J. Am. Chem. Soc. 1985, 107, 5987. (b) McCullough, L. G.; Schrock, R. R. J. Am. Chem. Soc. 1984, 106, 4067.
- (33) (a) Tsai, Y.-C.; Diaconescu, P. L.; Cummins, C. C. *Organometallics* **2000**, *19*, 5260. (b) Blackwell, J. M.; Figueroa, J. S.; Stephens, F. H.; Cummins, C. C. *Organometallics* **2003**, *22*, 3351.
- (34) Weinstock, I. A.; Schrock, R. R.; Davis, W. M. J. Am. Chem. Soc. 1991, 113, 135.
- (35) Chabanas, M.; Baudouin, A.; Copéret, C.; Basset, J.-M. J. Am. Chem. Soc. 2001, 123, 2062.

the tungsten neopentylidyne complex ${\bf 1}$ found the broadest use and is now commercially available. 8,30

Early on, this catalyst was shown to induce certain alkyne metathesis reactions, even at ambient temperature, within minutes. 1 turned out to be compatible with surprisingly many functional groups, although donor sites such as basic nitrogen, divalent sulfur, polyether fragments, or various heterocycles block its activity, presumably by coordination to the fairly Lewis acidic tungsten center. ^{12,13} Moreover, carbonyl groups may react with 1 and analogues in a Wittig-like manner. 36 Subsequent variations of the ligand sphere, as represented, for example, in complex 2, allowed the activity to be improved even further.³¹ The related molybdenum alkylidynes are generally considered to be less reactive but may exhibit an increased functional group tolerance. This fact is usually explained by the lower Lewis acidity of Mo(6+) compared to W(6+), though the chosen set of ancillary ligands must be taken into consideration.³² Catalytically active and highly tolerant molybdenum species can also be prepared in situ by reacting the now also commercial trisamido complex $\bf 6$ with CH_2Cl_2 or other *gem*-dihalides. Their remarkable selectivity profile is evident from many applications right through to the total synthesis of structurally complex and, in part, labile target molecules, including epothilone A and C,^{38,40} the latrunculins,⁴¹ prostaglandin E_2 1,15-lactone,⁴² cruentaren A,⁴³ amphidinolide V,^{22a,b} sophorolipid lactone,⁴⁴ and myxovirescin.⁴⁵⁻⁴⁷

- (36) Freudenberger, J. H.; Schrock, R. R. Organometallics 1986, 5, 398.
- (37) Fürstner, A.; Mathes, C.; Lehmann, C. W. J. Am. Chem. Soc. 1999, 121, 9453.
- (38) Fürstner, A.; Mathes, C.; Lehmann, C. W. Chem.—Eur. J. 2001, 7, 5299.
- (39) (a) Zhang, W.; Kraft, S.; Moore, J. S. J. Am. Chem. Soc. 2004, 126, 329. (b) Zhang, W.; Kraft, S.; Moore, J. S. Chem. Commun. 2003, 832.
- (40) Fürstner, A.; Mathes, C.; Grela, K. Chem. Commun. 2001, 1057.
- (41) (a) Fürstner, A.; De Souza, D.; Parra-Rapado, L.; Jensen, J. T. Angew. Chem., Int. Ed. 2003, 42, 5358. (b) Fürstner, A.; Turet, L. Angew. Chem., Int. Ed. 2005, 44, 3462. (c) Fürstner, A.; Kirk, D.; Fenster, M. D. B.; Aïssa, C.; De Souza, D.; Müller, O. Proc. Natl. Acad. Sci. U.S.A. 2005, 102, 8103. (d) Fürstner, A.; De Souza, D.; Turet, L.; Fenster, M. D. B.; Parra-Rapado, L.; Wirtz, C.; Mynott, R.; Lehmann, C. W. Chem.—Eur. J. 2007, 13, 115. (e) Fürstner, A.; Kirk, D.; Fenster, M. D. B.; Aïssa, C.; De Souza, D.; Nevado, C.; Tuttle, T.; Thiel, W.; Müller, O. Chem.—Eur. J. 2007, 13, 135.
- (42) (a) Fürstner, A.; Grela, K.; Mathes, C.; Lehmann, C. W. J. Am. Chem. Soc. 2000, 122, 11799. (b) Fürstner, A.; Grela, K. Angew. Chem., Int. Ed. 2000, 39, 1234.
- (43) (a) Fürstner, A.; Bindl, M.; Jean, L. Angew. Chem., Int. Ed. 2007, 46, 9275. (b) Bindl, M.; Jean, L.; Herrmann, J.; Müller, R.; Fürstner, A. Chem.—Eur. J. 2009, 15, 12310.
- (44) Fürstner, A.; Radkowski, K.; Grabowski, J.; Wirtz, C.; Mynott, R. J. Org. Chem. **2000**, 65, 8758.

These successes notwithstanding, all available catalysts—except for Mortreux's in situ recipe and its variants—suffer from high sensitivity toward oxygen, moisture, and, in the case of complex **6**, ⁴⁸ even molecular nitrogen. The necessary expert knowledge in handling such compounds may be another reason why alkyne metathesis has not yet become more popular. Convinced by the power of this transformation, however, our group is committed to change this situation by developing catalysts that retain the excellent selectivity of complexes such as 1-6 while being significantly more user-friendly. Ideally, the next generation of alkyne metathesis catalysts should be applicable to complex and polysubstituted targets while being cheap, easy to make, and air stable. In a recent Communication we described a first step toward this goal;⁴⁹ outlined below, we present a full account of our work in this area, which led to a set of new catalysts that meet these stringent criteria very well.

Results and Discussion

Molybenum Nitrido Complexes with Ancillary Silanolate Ligands. Inspiration for the development of novel user-friendly yet effective alkyne metathesis catalysts was provided by Johnson and co-workers, who reported that certain nitride complexes of molybdenum and tungsten endowed with fluorinated alkoxide ligands react with sacrificial alkynes to generate metal alkylidynes in situ (Scheme 2). ⁵⁰ However, the preparation of 7 as a prototype precursor complex requires the use of azides as well as the handling of air-sensitive intermediates.

In an attempt to find a more convenient entry point, we investigated if the same intermediate 8 could be generated by alcoholysis of the much more accessible precursor 11⁵¹ with hexafluoro-*tert*-butanol (14). Even though the mixture of 11 and

- (45) Fürstner, A.; Bonnekessel, M.; Blank, J. T.; Radkowski, K.; Seidel, G.; Lacombe, F.; Gabor, B.; Mynott, R. Chem.—Eur. J. 2007, 13, 8762.
- (46) For other applications to natural product chemistry from our laboratory, see: (a) Fürstner, A.; Seidel, G. J. Organomet. Chem. 2000, 606, 75.
 (b) Fürstner, A.; Rumbo, A. J. Org. Chem. 2000, 65, 2608. (c) Aguilera, B.; Wolf, L. B.; Nieczypor, P.; Rutjes, F. P. J. T.; Overkleeft, H. S.; van Hest, J. C. M.; Schoemaker, H. E.; Wang, B.; Mol, J. C.; Fürstner, A.; Overhand, M.; van der Marel, G. A.; van Boom, J. H. J. Org. Chem. 2001, 66, 3584. (d) Fürstner, A.; Stelzer, F.; Rumbo, A.; Krause, H. Chem.—Eur. J. 2002, 8, 1856. (e) Song, D.; Blond, G.; Fürstner, A. Tetrahedron 2003, 59, 6899. (f) Ghalit, N.; Poot, A. J.; Fürstner, A.; Rijkers, D. T. S.; Liskamp, R. M. Org. Lett. 2005, 7, 2961.
- (47) For applications by other groups, see: (a) Vintonyak, V. V.; Calà, M.; Lay, F.; Kunze, B.; Sasse, F.; Maier, M. E. Chem.—Eur. J. 2008, 14, 3709. (b) Vintonyak, V. V.; Maier, M. E. Angew. Chem., Int Ed. 2007, 46, 5209. (c) Vintonyak, V. V.; Maier, M. E. Org. Lett. 2007, 9, 655. (d) Doyle, D.; Murphy, P. V. Carbohydr. Res. 2008, 343, 2535. (e) Kraft, P.; Berthold, C. Synthesis 2008, 543. (f) Groothuys, S.; van den Broek, S. A. M. W.; Kuijpers, B. H. M.; IJsselstijn, M.; van Delft, F. L.; Rutjes, F. P. J. T. Synlett 2008, 111. (g) Schulz, S.; Yildizhan, S.; Stritzke, K.; Estrada, C.; Gilbert, L. E. Org. Bioorg. Chem. 2007, 5, 3434. (h) Ghalit, N.; Rijkers, D. T. S.; Liskamp, R. M. J. J. Mol. Catal. A: Chem. 2006, 254, 68. (i) Chan, J.; Jamison, T. F. J. Am. Chem. Soc. 2004, 126, 10682. (j) IJsselstijn, M.; Aguilera, B.; van der Marel, G. A.; van Boom, J. H.; van Delft, F. L.; Schoemaker, H. E.; Overkleeft, H. S.; Rutjes, F. P. J. T.; Overhand, M. Tetrahedron Lett. 2004, 45, 4379.
- (48) For a review on the activation of small molecules by complex 6, see: Cummins, C. C. Chem. Commun. 1998, 1777.
- (49) Bindl, M.; Stade, R.; Heilmann, E. K.; Picot, A.; Goddard, R.; Fürstner, A. J. Am. Chem. Soc. 2009, 131, 9468.
- (50) (a) Geyer, A. M.; Wiedner, E. S.; Gary, J. B.; Gdula, R. L.; Kuhlmann, N. C.; Johnson, M. J. A.; Dunietz, B. D.; Kampf, J. W. J. Am. Chem. Soc. 2008, 130, 8984. (b) Gdula, R. L.; Johnson, M. J. A. J. Am. Chem. Soc. 2006, 128, 9614.
- (51) Chiu, H.-T.; Chuang, S.-H.; Lee, G.-H.; Peng, S.-M. Adv. Mater. 1998, 10, 1475.

Scheme 2. Metal Nitride/Metal Alkylidyne Interconversion as an Alternative Concept for the in Situ Generation of Alkyne Metathesis Catalysts^a

Scheme 3. Preparation of a New Metathesis Precatalyst That Is Air-Stable for Limited Periods of Time^a

 a Conditions: (a) TMSCl, 1,2-dimethoxyethane (DME), reflux; (b) LiHMDS, hexane, 64% (over both steps); (c) Ph₃SiOH (3 equiv), toluene, 80 °C; (d) pyridine (5 equiv), 81% (over both steps). Cf. ref 49.

14 was devoid of any appreciable catalytic activity, we were pleased to observe that the use of triphenylsilanol resulted in a catalytically active mixture. 49,52,53 Complex **12** was readily identified as the active component (Scheme 3). Although 12 itself is hydrolytically labile and needs to be handled under an inert atmosphere, the corresponding pyridine adduct 13 turned out to be sufficiently stable to be weighed in air (Scheme 3).⁴⁹ At temperatures around 80 °C, however, solutions of this adduct in toluene exhibit appreciable catalytic activity, most likely by slow decomplexation of the pyridine. The fact that the presence of the N-heterocyclic ligand does not quench the catalytic activity of the molybdenum core may explain why this particular precatalyst, which has recently been made commercially available, was found compatible with many other polar groups (ester, ketone, amide, carbamate, sulfonate, nitro, ether, thioether, silyl ether, alkene, acetal, glycoside, thiazole, pyridine, thiophene, etc.); only epoxides, aldehydes, and acid chlorides were found to react stoichiometrically with the nitride function of 12 and hence consume the catalyst. 49,54,55

Although 13 combines favorable chemical attributes with a reasonable stability, it will eventually hydrolyze and hence must still be stored under an inert atmosphere. In an attempt to find an even more robust alternative, several ligands other than pyridine were tested. During this screening process, it was found

Scheme 4^e

^a Reactions and conditions: (a) Ph₃SiOH (3 equiv), toluene; (b) 1,10-phenanthroline, 82%; (c) MnCl₂, toluene, 80-100 °C.

that carefully dried 1,10-phenanthroline leads to the nicely crystalline complex **15**, which seems to be indefinitely stable on the benchtop without any precautions whatsoever (Scheme 4).⁵⁶ However, judging from the lack of catalytic activity in the metathesis of 1-phenyl-1-propyne as the model reaction, the bidentate ligand does not seem to come off the metal template to any noticeable extent at temperatures below 110 °C.

Gratifyingly, though, the catalytic activity can be restored upon treatment of 15 with metal salts that are able to form stable complexes with phenanthroline, including MnCl₂, FeCl₂, FeCl₃, CoCl₂, CuCl₂, ZnCl₂, MgCl₂, and NiCl₂ (see the Supporting Information). Among them, MnCl₂ is preferred for practical reasons, because this salt is cheap, benign, nontoxic, readily available, hardly Lewis acidic, and nonhygroscopic; hence, commercial samples can be used as such without further drying. The activation of complex 15 with MnCl₂ can be performed either prior to the addition of the substrate or in its presence. As expected, the combination of 15/MnCl₂ shows roughly the same activity and selectivity profile as the pyridine complex 13, since both systems are thought to release an identical active fragment, i.e., complex 12. Tables 1-3 provide a detailed survey of the performance of the 15/MnCl₂ system in a host of interand intramolecular alkyne metathesis reactions (see below).

In view of its impressive scope and unparalleled stability, complex 15 is considered a highly practical yet broadly applicable precatalyst for alkyne metathesis. Although its reactivity after treatment with MnCl₂ or related salts is clearly lower than that of preformed Schrock alkylidynes such as 1–5 or the powerful precursor complex 6, the handling of 15 is trivial and the functional group tolerance uncompromised, rivaling or even surpassing that of 1–6; note that storage on the benchtop in air is totally inconceivable for 1–6 or any other structurally defined alkyne metathesis catalyst known to date, and even the pyridine adduct 13 will eventually degrade in moist air. An optimized synthesis (see experimental details in the Supporting Information) allows multigram amounts of 15 to be prepared from inexpensive Na₂MoO₄.

The reason why even the only weakly Lewis acidic $MnCl_2$ is capable of pulling the phenanthroline ligand off the Mo(6+) template may be found in the structure of complex 15 in the solid state. As can be seen from Figure 1, the strongly distorted octahedral coordination geometry brings the bulky Ph_3SiO groups in a *mer* arrangement. The $N(1) \equiv Mo - N(3)$ angle of

⁽⁵²⁾ For alkyne metathesis catalysts modified by different silanols, see: (a) Cho, H. M.; Weissman, H.; Wilson, S. R.; Moore, J. S. J. Am. Chem. Soc. 2006, 128, 14742. (b) Villemin, D.; Héroux, M.; Blot, V. Tetrahedron Lett. 2001, 42, 3701.

⁽⁵³⁾ For alkyne metathesis catalysts grafted onto silica, see: (a) Coutelier, O.; Gauvin, R. M.; Nowogrocki, G.; Trébosc, J.; Delevoye, L.; Mortreux, A. Eur. J. Inorg. Chem. 2007, 5541. (b) Cho, H. M.; Weissman, H.; Moore, J. S. J. Org. Chem. 2008, 73, 4256. (c) Chabanas, M.; Baudouin, A.; Copéret, C.; Basset, J.-M. J. Am. Chem. Soc. 2001, 123, 2062. (d) Weissman, H.; Plunkett, K. N.; Moore, J. S. Angew. Chem., Int. Ed. 2006, 45, 585. (e) Gauvin, R. M.; Coutelier, O.; Berrier, E.; Mortreux, A.; Delevoye, L.; Paul, J.-F.; Mamède, A.-S.; Payen, E. Dalton Trans. 2007, 3127. (f) Merle, N.; Taoufik, M.; Nayer, M.; Baudouin, A.; Le Roux, E.; Gauvin, R. M.; Lefebvre, F.; Thivolle-Cazat, J.; Basset, J. M. J. Organomet. Chem. 2008, 693, 1733.

⁽⁵⁴⁾ For a recent application of complex 13 in alkaloid total synthesis, see: Smith, B. J.; Sulikowski, G. A. Angew. Chem., Int. Ed. 2010, 49, 1599.

⁽⁵⁵⁾ For the acylation chemistry of terminal metal nitrides, see the following for leading references: (a) Curley, J. J.; Sceats, E. L.; Cummins, C. C. J. Am. Chem. Soc. 2006, 128, 14036. (b) Clough, C. R.; Greco, J. B.; Figueroa, J. S.; Diaconescu, P. L.; Davis, W. M.; Cummins, C. C. J. Am. Chem. Soc. 2004, 126, 7742. (c) Figueroa, J. S.; Piro, N. A.; Clough, C. R.; Cummins, C. C. J. Am. Chem. Soc. 2006, 128, 940. (d) Sarkar, S.; Abboud, K. A.; Veige, A. S. J. Am. Chem. Soc. 2008, 130, 16128. and literature cited therein.

⁽⁵⁶⁾ A year-old batch shows unchanged appearance and catalytic activity.

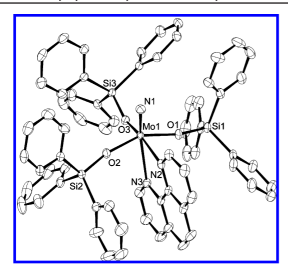


Figure 1. Structure of $Mo(\equiv N)(OSiPh_3)_3(phen)$ (15) in the solid state $(N1-Mo1-N2=91.2(1)^\circ)$. Disordered solute toluene is omitted for clarity; phen =1,10-phenanthroline.

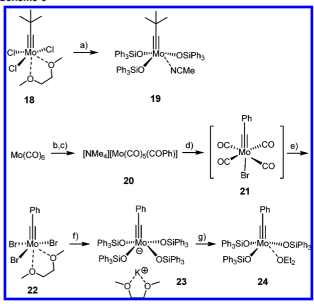
160.24(7)° deviates significantly from 180° and the Mo atom lies 0.33(1) Å above the plane defined by N(2), O(1), O(2) and O(3). Importantly, the Mo-N(3) distance of 2.512(2) Å is much larger than the distance from the metal to the phenanthroline nitrogen atom in the equatorial plane (Mo-N(2) 2.292(2) Å). The fact that the second N-atom of the chelate ligand obviously binds very weakly may explain why 12, even in the presence of excess pyridine, affords only the monoadduct 13 (Scheme 3). Two of the three Mo-O-Si angles are surprisingly obtuse (Mo-O(1)-Si(1) 165.26(8)°, Mo-O(2)-Si(2) 168.04(8)°, Mo-O(3)-Si(1) 145.78(8)°), which likely reduces the congestion in the periphery. The Mo \equiv N(1) bond (1.659(2) Å) is slightly longer than that observed in the pyridine adduct 13 (1.653(2) Å)49 and near the longer end of reported Mo≡N distances $(1.563-1.688 \text{ Å},^{57} \text{ with the exception of one outlier of } 1.786$ Å).⁵⁸

Molybdenum Alkylidyne Complexes with Ancillary Silanolate Ligands. As already mentioned, inspiration for the development of the novel silanolate complexes presented here was the work of Johnson et al., who showed that certain metal nitride species can convert into the corresponding alkylidyne complexes on reaction with a nonterminal alkyne (Scheme 2).⁵⁰ An analogous metathetic exchange process is believed to account for the catalytic activity exhibited by nitrides 12, 13, and 15. Yet, on NMR inspection of a mixture comprising $(Ph_3SiO)_3Mo \equiv N (12)^{59}$ and 5-decyne (2 equiv) in toluene d_8 , we were not able to detect appreciable amounts of valeronitrile (17), which must form in a quantity equal to that of the presumed alkylidyne complex 16 (Scheme 5). Only after prolonged heating of the mixture in a sealed tube (100 °C, 6 d) were small amounts of 17 discernible, in addition to massive polymerization of the mixture (see the Supporting Information).

Scheme 5^a

^a Reagents and conditions: (a) 5-decyne, toluene-d₈, 100 °C, cf. text.

Scheme 6ª



^a Reagents and conditions: (a) Ph₃SiOLi (3 equiv), Et₂O, −40 °C → rt, then MeCN, 85%; (b) PhLi, Et₂O, reflux; (c) NMe₄Br, H₂O, 52% (over both steps); (d) oxalyl bromide, CH₂Cl₂, −78 °C → −15 °C; (e) Br₂, 1,2-dimethoxyethane (dme, 5 equiv), CH₂Cl₂, −78 °C → rt, 88% (over both steps); (f) Ph₃SiOK (4 equiv), toluene; (g) Et₂O, 92%.

The striking inefficiency of the crucial nitride → alkylidyne exchange compared with the excellent catalytic performance of 15 in our preparative experiments (Tables 1-3) implies that the small amounts of alkylidynes, such as 16, formed in the mixture must be superbly active. We sought to clarify this aspect by preparing the hitherto unknown molybdenum alkylidynes of the general structure (Ph₃SiO)₃Mo≡CR and derivatives thereof. This goal was readily attained by adaptation of the established entry routes to related d⁰-alkylidynes (Scheme 6). Specifically, the desired neopentylidyne complex 19 · MeCN was obtained by ligand exchange of the corresponding trichloride 18, which, in turn, was prepared according to the route previously reported by the Schrock group. 32 Much more efficient overall was the synthesis of the corresponding benzylidyne analogue 24 · Et₂O, which was formed analogously from cheap Mo(CO)₆ via the known building block 22.^{60,61} As described in the Supporting Information, 24 • Et₂O can be obtained in only three operations, as the intermediates 21 and 23 need not be isolated.

^{(57) (}a) Chisholm, M. H.; Davidson, E. R.; Pink, M.; Quinlan, K. B. *Inorg. Chem.* **2002**, *41*, 3437. (b) Ritleng, V.; Yandulov, D. V.; Weare, W. W.; Schrock, R. R.; Hock, A. S.; Davis, W. M. *J. Am. Chem. Soc.* **2004**, *126*, 6150.

⁽⁵⁸⁾ Dilworth, J. R.; Dahlstrom, P. L.; Hyde, J. R.; Zubieta, J. Inorg. Chem. Acta 1983, 71, 21.

⁽⁵⁹⁾ This experiment was deliberately performed with 12 generated in situ from 11 and Ph₃SiOH, as such a solution is free of any N-donor ligand that might retard the metathetic nitride/alkylidyne interconversion by complexation to the metal center.

^{(60) (}a) Fischer, E. O.; Maasböl, A. Chem. Ber. 1967, 100, 2445. (b) Fischer, E. O.; Maasböl, A. Ger. Offen. DE 1214233, 1966; Chem. Abstr. 1966, 65, 12474.

^{(61) (}c) Mayr, A.; McDermott, G. A. J. Am. Chem. Soc. 1986, 108, 548.
(d) McDermott, G. A.; Dorries, A. M.; Mayr, A. Organometallic 1987, 6, 925.

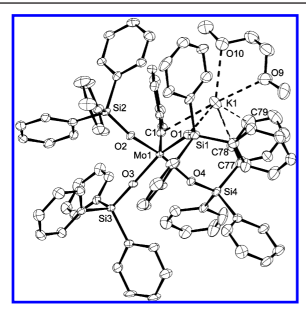


Figure 2. Structure of the ate-complex $[(Ph_3SiO)_4Mo \equiv CPh]^ [K^+ \cdot (dme)]$ (23) in the solid state. Only one of the two independent molecules in the unit cell is depicted for clarity; dme =1,2-dimethoxyethane.

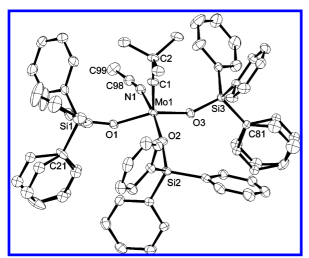


Figure 3. Structure of the complex Mo(≡CtBu)(OSiPh₃)₃(MeCN) (19·MeCN) in the solid state (C1−Mo1−N1 94.4(1)°). Two of the phenyl groups (at C21 and C81) are disordered over two positions.

Interestingly, we noticed that the primary product formed during the ligand-exchange process of 22^{62} and Ph_3SiOK was the ate-complex 23, in which four rather than three silanolates are covalently bound to the molybdenum center (Figure 2). This may be taken as an indication for the poor donor capacity of the silanolates, which allows the molybdenum to retain an appreciable Lewis acidity, which is essential for high activity in alkyne metathesis. Even though the Ph_3SiO unit seems very bulky, the complex accommodates four of them in a square pyramidal environment about the metal center, with the alkylidyne forming the apex and the silanolates the basal plane. The bond angles of the individual Si-O-Mo units differ quite significantly from each other and can be almost linear $(Mo(1)-O(3)-Si(3)\ 177.8(3)^\circ; Mo(1)-O(2)-Si(2)\ 151.8(2)^\circ;$

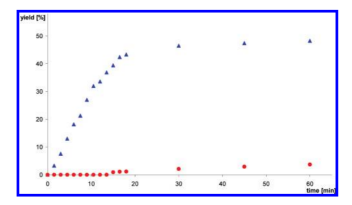


Figure 4. Metathetic conversion of 1-phenyl-1-propyne to tolane catalyzed by 1 mol % of either Mo(≡CPh)(OSiPh₃)₃(Et₂O) ($24 \cdot$ Et₂O) (blue triangles) or (tBuO)₃W≡CtBu (1) (red circles) in toluene at ambient temperature and ambient pressure. The yields were determined by GC using biphenyl as an internal standard.

Mo(1)–O(4)–Si(4) 149.0(3)°). This suggests that the Si–O–Mo groups serve as hinges which, through facile bending, allow for optimal use of the available space in the periphery. The potassium cation escorts the molybdate core by binding to the fourth silanolate oxygen, which is more pyramidalized than the other oxygen atoms in order to make its lone pairs accessible for binding to the cation $(Mo(1)-O(1)-Si(1)\ 134.5°)$. The $Mo \equiv C$ distances of the two independent molecules of 23 at 1.747(6) and 1.756(5) Å are slightly longer than the equivalent distance of the corresponding neutral analogue $19 \cdot MeCN (1.741(2) \text{ Å}; see below)$.

Workup of the crude mixtures by evaporation of the solvent and trituration of the residues with MeCN or Et₂O delivered the neutral alkylidyne complexes 19 and 24, respectively, in the form of the corresponding adducts. Their NMR data were in excellent agreement with the proposed structures. The constitution of 19 · MeCN was unambiguously confirmed by crystal structure analysis (Figure 3), which revealed a distorted square pyramidal coordination geometry about the Mo center. The Mo≡C(1) bond length of 1.741(2) Å is slightly shorter than in the ate-complex 23. Interestingly, the acetonitrile ligand does not bind in a strictly linear fashion, as evident from the Mo-N(1)-C(98) angle of only $166.31(15)^{\circ}$; moreover, the C≡N triple bond is extended to 1.145(3) Å as a result of the coordination to the high-valent metal center (reference bond length for RC≡N, 1.136 Å).⁶⁴ However, no spontaneous metathesis takes place between the ligated nitrile and the alkylidyne unit as a consequence of the still largely end-on rather than side-on coordination mode of the C≡N triple bond.

As expected for prototype Schrock alkylidynes, the acetonitrile and ether adducts of **19** and **24** are both air- and moisture-sensitive. Even though they do not provide any advantage over other alkylidynes in terms of handling, they turned out to be exquisitely active alkyne metathesis catalysts which are operative at low loadings and distinguished by an outstanding selectivity profile. Whereas the preparative details are outlined in a later section of this paper, the plot for the conversion of 1-phenyl-1-propyne to tolane catalyzed by $24 \cdot \text{Et}_2\text{O}$ against the rate recorded for the classical tungsten neopentylidyne $(tBuO)_3W \equiv CCMe_3$ (1), which defines the standard in the field, clearly demonstrates the extraordinary performance of the novel catalyst (Figure 4). Specifically,

⁽⁶²⁾ Complex 22 was prepared according to ref 60. A detailed procedure is described in the Supporting Information.

⁽⁶³⁾ This ate-complex crystallizes from the reaction mixture even if only 3 equiv of Ph₃SiOK is added to complex **22**.

⁽⁶⁴⁾ Allen, F. H.; Kennard, O.; Watson, D. G.; Brammer, L.; Orpen, A. G.; Taylor, R. J. Chem. Soc., Perkin Trans. 2 1987, S1.

Scheme 7^a

 $^{\it a}$ Reagents and conditions: (a) 1,10-phenanthroline, toluene/Et₂O, 81%; (b) MnCl₂, toluene, 80 °C.

the equilibrium is reached after 25 min at ambient temperature; at this point, complex 1 gave less than 5% conversion. Although molybdenum alkylidynes are generally considered less active than their tungsten counterparts, 31,32,65 these data show that the triphenylsilanolate ligands impart a truly remarkable activity onto the operative molybdenum alkylidyne unit, rendering 24. Et₂O one of the most active metathesis catalysts known to date. This, in turn, corroborates the view that even small amounts of (Ph₃SiO)₃Mo≡CR generated in situ from $(Ph_3SiO)_3Mo \equiv N \cdot L$ (L = pyridine, phenanthroline) and an appropriate alkyne can account for the excellent results obtained with such robust precatalyst systems. Actually, we currently believe that nitride complexes of the type (Ph₃SiO)₃Mo≡N·L act as a reservoir, from which tiny amounts of active alkylidyne species are constantly released during the reaction. This may explain why solutions of 13 and 15 retain catalytic activity at 80 °C over the course of days, 49 whereas all known metal alkylidynes degrade when kept under such forcing conditions for prolonged periods of time.

In an attempt to transfer the stabilizing effect of 1,10phenanthroline from the metal nitride to the metal alkylidyne series, the crude reaction mixture formed upon ligand exchange of Br₃Mo≡CPh•dme (22) with Ph₃SiOK was added to a solution of carefully dried 1,10-phenanthroline in toluene. The desired adduct 25 could be isolated from this mixture in good yield (Scheme 7). Its structure, as determined by X-ray crystallography (Figure 5), closely resembles the structure of the isolobal nitride complex 15. Once again, a strongly distorted coordination geometry about the metal center is observed, characterized by a significant deviation of the $C(1) \equiv Mo \cdots N(1)$ axis (168.34(11)°) from linearity. The Mo···N bond lengths are again very uneven (2.244(2) versus 2.408(2) Å), with the distance to the apical nitrogen being much longer. The alkylidyne bond length Mo \equiv C(1) (1.761(3) Å) exceeds those in the corresponding square pyramidal nitrile adduct 19·MeCN (1.7410(17) Å) and the ate-complex **23** (1.747(6)/1.756(5) Å).

Importantly, the 1,10-phenanthroline ligand stabilizes this particular Schrock alkylidyne to the extent that complex 25 is stable in air for hours; 66,67 however, complex 25 per se does not react with 1-phenyl-1-propyne to any appreciable extent under conditions in which the corresponding ether adduct 24·Et₂O leads to very fast conversions. Gratifyingly, though, addition of commercial MnCl₂ (1 equiv relative to 25) and heating of the reaction mixture for 30 min to about 80 °C restores an outstanding performance; the resulting catalyst solution containing complex 24 is then active even at ambient temperature. The ligand swap from molybdenum to manganese can easily be monitored by precipitation of MnCl₂•phen and

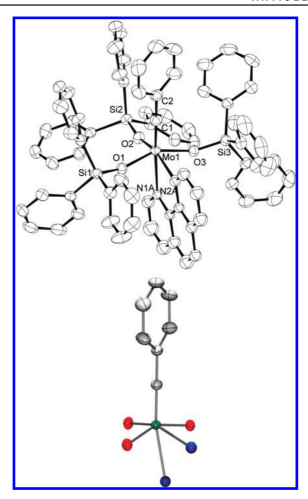


Figure 5. Top: Structure of Mo(≡CPh)(OSiPh₃)₃(phen) (25) in the solid state. The 1,10-phenanthroline ligand is disordered over two positions in the ligand plane (one conformation shown). Bottom: Core of the complex showing the distorted coordination geometry about the Mo center (green); O, red; N, blue.

was confirmed by mass spectrometry. Within experimental error, the resulting solutions are largely equipotent to those containing authentic $24 \cdot \text{Et}_2\text{O}$ (see Tables 1–3 for details) and are able to metathesize most alkyne substrates within unprecedentedly short periods of time. Likewise, the functional group tolerance is extraordinary (see below). Therefore, we conclude that complex 25 is an excellent compromise between the desirable activity, selectivity, and practicality aspects associated with alkyne metathesis.

The user may hence choose among (i) the air-sensitive Mo(≡CPh)(OSiPh₃)₃(Et₂O) (24 • Et₂O), which is superbly active and selective; (ii) the phenanthroline adduct 25 thereof, which, after activation with MnCl₂, is similarly potent but can be handled in air, even though it still needs to be stored under argon; and (iii) the totally air-stable nitride complex 15, which is exceptionally user-friendly. The preexisting alkylidyne unit present in 24 and 25 ensures that these complexes are active under notably mild conditions even at very low loadings (see

⁽⁶⁵⁾ For a computational study, see: Zhu, J.; Jia, G.; Lin, Z. Organometallics 2006, 25, 1812

⁽⁶⁶⁾ Crystalline samples of 25 seem partly intact after several days in air.

^{(67) (}a) Addition of hydrotris(3,5-dimethyl-1-pyrazolyl)borate to (dme)Cl₃W≡CPh leads to an air-stable but catalytically inactive Schrock alkylidyne; see: Blosch, L. L.; Abboud, K.; Boncella, J. M. J. Am. Chem. Soc. 1991, 113, 7066. (b) An analogous complex derived from (dme)Br₃W≡CPh was described as moderately air-stable but sensitive to moisture; see: Jeffery, J. C.; McCleverty, J. A.; Mortimer, M. D.; Ward, M. D. Polyhedron 1994, 13, 353.

Scheme 8ª

$$W(CO)_{6} \xrightarrow{a,b} [NMe_{4}][W(CO)_{5}(COPh)] \xrightarrow{c,d} Br^{\text{II}} COP^{\text{II}} CO$$

^a Reagents and conditions: (a) PhLi, Et₂O, reflux; (b) NMe₄Br, H₂O, 55% (over both steps); (c) oxalyl bromide, CH₂Cl₂, −78 °C → −15 °C; (d) Br₂, 1,2-dimethoxyethane (dme, 5 equiv), CH₂Cl₂, −78 °C → rt, 85% (over both steps); (e) Ph₃SiOK (4 equiv), toluene; (f) 1,10-phenanthroline, toluene, 79%.

below), whereas nitride 15 first needs to be converted into an operative alkylidyne in situ, a process that mandates higher loadings of the precatalyst and heating of the mixture to about 80 °C. Whatever precatalyst one might prefer, however, all systems are available in quantity from cheap and commercially available starting materials, lead to excellent preparative results, and compare favorably with all established alkyne metathesis catalysts (see below). Complex 25 is much more stable and practical than any other catalytically active Schrock alkylidyne known to date. Therefore, we believe that these systems, collectively, represent a significant step forward in our quest for truly user-friendly yet generally applicable alkyne metathesis catalysts.

Tris(triphenylsilyloxy)tungsten Alkylidyne Complexes. To study the generality of the concept, the corresponding tungsten alkylidynes endowed with triphenylsilanolate ancillary ligands were targeted. To this end, the known benzylidyne tribromide complex $27^{60,61}$ was subjected to ligand exchange with Ph₃SiOK in toluene, which furnished the corresponding ate-complex 28 (Scheme 8). In contrast to its molybdenum analogue 23, however, treatment with 1,10-phenanthroline in toluene did not afford a neutral adduct but merely replaced the dme ligated to the potassium counterion to give the surprisingly robust atecomplex 29. X-ray diffraction proved that the phenanthroline ligand binds to the potassium escort rather than the tungsten center (Figure 6). Overall, the structure of 29 in the solid state resembles the molybdenum counterpart 23, featuring again three obviously flexible metal-O-Si moieties, which manage the bulk in the periphery by appropriate bending. The $W \equiv C(1)$ bond length (1.758(2) Å) is identical within the error limits to the corresponding bond length in the molybdenum species 23 (1.747(6)/1.756(5) Å).

The corresponding neutral tungsten alkylidyne species could not, so far, be generated upon trituration of **28** or **29** with Et_2O nor on reaction with TMSCl. Not unexpectedly, the catalytic activity of these stable ate complexes themselves is marginal. Even though 1-phenyl-1-propyne slowly gets consumed, only small amounts of tolane ($\leq 25\%$) were detected. As polymeri-

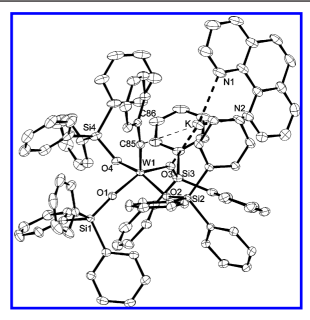


Figure 6. Structure of the ate-complex $[(Ph_3SiO)_4W \equiv CPh]^- [K^+ \cdot (phen)]$ (29) in the solid state.

Scheme 9. Formation of 2-Butyne as a Generic Byproduct in the Reaction of Alkynes End-Capped with a Methyl Group, As Exemplified by the Conversion of 1-Phenyl-1-propyne to Tolane

zation seems to prevail, this particular complex is not relevant in the present context; its reaction behavior as well as further attempts to secure neutral tungsten alkylidynes of the general type $[(Ph_3SiO)_3W \equiv CR] \cdot L$ are subject of a separate investigation

Beneficial Effect of 5 Å Molecular Sieves on Alkyne Metathesis. A priori, the metathetic scrambling of a pair of alkynes leads to an equilibrium, which needs to be shifted to one side in order to make the reaction preparatively useful. If one of the products is an alkyne of low molecular weight (2-butyne, 3-hexyne, etc.), this goal is usually accomplished by driving this compound out of the mixture.

Therefore, alkynes with capping methyl substituents are the most common substrates, and the reactions are usually per-

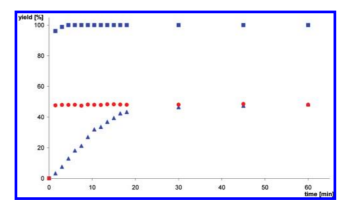


Figure 7. Metathesis of 1-phenyl-1-propyne catalyzed by $Mo(≡CPh)(OSiPh_3)_3(Et_2O)$ (24·Et_2O, 1 mol %) in toluene in the absence (blue triangles) or in the presence (blue squares) of powdered MS 5 Å at ambient temperature. Comparison with the plot obtained upon pretreatment of the catalyst with MS 5 Å, which was removed prior to the addition of the substrate (red circles). The yields were measured by GC against biphenyl as internal standard.

^{(68) (}a) Zhang, W.; Moore, J. S. J. Am. Chem. Soc. 2004, 126, 12796. (b) Zhang, W.; Moore, J. S. J. Am. Chem. Soc. 2005, 127, 11863. (c) Zhang, W.; Brombosz, S. M.; Mendoza, J. L.; Moore, J. S. J. Org. Chem. 2005, 70, 10198.

formed at elevated temperatures and/or by applying gentle vacuum (Scheme 9). However, the solubility of 2-butyne in hydrocarbon solvents is non-negligible, and the use of reduced pressure may lead to undesirable changes in the concentration of the mixture, as the solvent will eventually also distill off. Moreover, 2-butyne may be consumed by certain alkyne metathesis catalysts in a competitive polymerization process. To avoid such problems, Moore and co-workers have developed a "precipitation-driven" setup for alkyne metathesis; hough highly productive, this approach requires tailor-made starting materials and is less favorable from the viewpoint of atom economy.

Since catalysts of increased activity, such as $24 \cdot \text{Et}_2\text{O}$, allow the reaction to be performed at ambient temperature, the problem of possible 2-butyne (bp = 27 °C) accumulation in solution becomes more pressing. This is evident from Figure 7, which shows that the metathesis of 1-phenyl-1-propyne effected by $24 \cdot \text{Et}_2\text{O}$ (1 mol %) in toluene is fast but levels off at about 50% conversion after 20 min when carried out at ambient temperature and atmospheric pressure (blue triangles).

We reasoned that 2-butyne might also be removed from the mixture with the aid of an appropriate molecular sieve (MS). Since activated MS 5 Å is capable of adsorbing C4 hydrocarbons, the reaction was repeated in the presence of 2 mg of MS 5 Å per μ mol of released 2-butyne. The corresponding plot (blue squares) in Figure 7 shows the dramatic effect of this additive on the reaction rate as well as the conversion, which is essentially complete within 5 min under otherwise identical conditions. This outcome cannot be explained by simple removal of moisture from the mixture, as the equilibrium is not affected by adventitious water. This notion is corroborated by the fact that MS 3 Å has no significant effect on the reaction, even though it is known to bind water effectively; MS 4 Å is slightly less efficient than MS 5 Å, which is the preferred additive (see Supporting Information).

The data in Figure 7 also show that added MS 5 Å not only improves the conversion but also significantly accelerates the reaction rate. This effect became particularly evident when a solution of 24 · Et₂O in toluene was stirred in the presence of powdered MS 5 Å for 30 min and the molecular sieves were then removed from the mixture prior to the addition of the substrate. In this case, the equilibrium is reached in <5 min (Figure 7, red circles), which is much faster than the reaction without this "preactivation". It is believed that the MS 5 Å helps liberate the required free coordination site at molybdenum by absorbing the only weakly bound diethyl ether ligand. ^{72,73}

In any case, this simple measure of adding powdered MS 5 \mathring{A} to the reaction mixture turned out to be general and was applied to all preparative experiments compiled in Tables 1–3,

unless stated otherwise.⁷⁵ Consequently, we recommend the use of MS 5 Å in future alkyne metathesis reactions independent of the chosen catalyst and hope that this convenient setup will make the preparation of tailor-made substrates largely unnecessary.

Survey of the Preparative Scope of the New Alkyne Metathesis Catalysts. With a set of new (pre)catalysts and an optimized procedure for alkyne metathesis at hand, we were committed to explore the scope of this methodology in detail. To this end, a representative set of alkyne homometathesis, alkyne cross-metathesis (ACM),⁷⁴ and ring-closing alkyne metathesis (RCAM) reactions was investigated, as well as a prototype cyclooligomerization. Even though not all substrates were tested with all available catalysts, care was taken to accumulate a sufficiently large data set for direct comparison. Furthermore, particular attention was paid to investigating the compatibility of the novel catalysts with polar functional groups, since this aspect is key for future applications in advanced organic synthesis and material science.

The results compiled in Tables 1–3 deserve further comment. With very few exceptions, complexes **15**, **24** • Et₂O, and **25** behaved similarly well, giving good to excellent results with a diverse set of functionalized substrates. Although it may not be surprising that the isolated yields tended to be highest when the preformed alkylidyne **24** • Et₂O was employed, the very stable molybdenum nitride **15** also performed remarkably well. The price to be paid for the use of this fully air stable precatalyst, however, is a larger loading (generally 10 mol %), a higher reaction temperature (usually 80 °C), ⁷⁵ and longer reaction times

As expected, the data compiled in the tables also show that the alkylidyne complex 24 · Et₂O and the much more robust phenanthroline adduct 25 thereof, after activation with MnCl₂, by and large lead to the same preparative results, although slightly higher loadings were applied in the latter case. However, it is emphasized that the catalyst loading has not been optimized for each entry reported here. In view of the spectacular reactivity of **24** (see Figure 7), we are optimistic that loadings well below the 1-2 mol % generally used during this investigation can be reached by proper adjustment of the reaction parameters, in particular since the half-life of 24 · Et₂O in toluene-d₈ was found to be ~30 h at ambient temperature (see Supporting Information). In fact, a single experiment, in which 1-phenyl-1-propyne was exposed to only 0.1 mol % of 24 • Et₂O and the usual loading of MS 5 Å, reached 95% conversion (GC) after 2 h reaction time. Likewise, the same model substrate was quantitatively converted to tolane within ≤20 min by 1 mol % of complex **24** • Et₂O, even at -10 °C. To the best of our knowledge, this is the lowest temperature at which an alkyne metathesis has so far been successfully performed, but it does not seem to mark the lower limit for our new catalyst.

The compatibility of the new catalysts with various functional groups is outstanding. Esters, ethers, various silyl ethers,

⁽⁶⁹⁾ Alkyne metathesis reactions using Mortreux-type catalyst mixtures have previously been run in the presence of MS 4 Å in order to remove traces of moisture. However, it was reported later that carefully dried solvents make the addition of MS 4 Å unnecessary. No effect of this additive on the conversion was noticed; see: (a) Huc, V.; Weihofen, R.; Martin-Jimenez, I.; Oulié, P.; Lepetit, C.; Lavigne, G.; Chauvin, R. New J. Chem. 2003, 27, 1412. (b) Maraval, V.; Lepetit, C.; Caminade, A.-M.; Majoral, J.-P.; Chauvin, R. Tetrahedron Lett. 2006, 47, 2155.

⁽⁷⁰⁾ MS 5 Å is slightly more effective than MS 4 Å, whereas the use of MS 3 Å has little effect, if any. Since MS 3 Å, however, is an efficient trap for water, this comparison shows that the observed rate acceleration and the improved yields are not caused by ensuring a rigorously dry medium.

⁽⁷¹⁾ As expected, powdered MS 5 Å is more effective than the use of pellets, and about 2 mg/µmol butyne turned out to be optimal.

⁽⁷²⁾ Control experiments, in which the metathesis of 1-phenyl-1-propyne was carried out in Et₂O, THF, or CH₂Cl₂ as the reaction medium, did not go to complete conversion, even in the presence of molecular sieves. Reactions in THF were much slower than those in the other solvents investigated.

⁽⁷³⁾ Notably, the metathesis of 1-phenyl-1-propyne induced by the molybdate complex 23 (1 mol%) in toluene in the presence of MS 5 Å proceeded with the same rate as that induced by 24·Et₂O/MS 5 Å.

⁽⁷⁴⁾ Fürstner, A.; Mathes, C. *Org. Lett.* **2001**, *3*, 221.

⁽⁷⁵⁾ As the reactions with complex 15 have to be performed at higher temperature, addition of molecular sieves is usually not necessary and sometimes even disadvantageous due to possible side reactions at this temperature.

Table 1. Intermolecular Alkyne Metathesis Reactions in the Presence of 5 Å Molecular Sieves

Entry	Substrate	Product		15 ^a	24 ∙Et₂O ^b	25 ^c
1	D.	D.	R = H	99%	99%	99%
2			R = OMe	96%	97%	97%
3		R	R = SMe	87%	98% ^d	96% ^d
4			R = COOMe	72% ^e	95%	97%
5	F ₃ C-\	F ₃ C-CF ₃		94%	93%	95%
6) H) H		NR	NR	NR
7	<u> </u>	_		< 40% ^{e,f}	84%	84%
8	N=	$\stackrel{\sim}{\longrightarrow} = \stackrel{\sim}{\longrightarrow}$		76% ^e	90% ^d	88% ^d
9	<u> </u>	[<u>\$</u> = -{\$]		86%	88%	87%
10	TsO-\	TsO—————OTs		95%	92%	92%
11	MeOOC	MeOOC		85%	89%	91%
12	H H O				92%	88%
13	N OH 12	N 12 12	[⊋] N	81%	87%	89%

 a **15** (10 mol %), MnCl₂ (10 mol %), MS 5 Å, toluene, 80 °C, 30 min, then addition of the substrate and reaction at 80 °C unless stated otherwise. b **24** ·Et₂O, (2 mol %), toluene, ambient temperature, MS 5 Å. c **25** (5 mol %), MnCl₂ (5 mol %), toluene, 80 °C, 30 min; then addition of the substrate and MS 5 Å, and reaction at ambient temperature. d At 50 °C. e At 100 °C. f Cf. text; NR = no reaction.

thioethers, sulfonates, amides, carbamates, ketones, acetals, epoxides (see below), nitro groups, and trifluoromethyl groups are generally well tolerated, even if they are oriented toward the reacting alkyne (Table 2, entry 12). Likewise, a nitrile was also found at least kinetically stable in the presence of all three novel molybdenum silanolate complexes, despite being a possible substrate for metathesis (Table 1, entry 13). Moreover, compounds containing various types of aromatic heterocycles (pyridine, thiophene, thiazole, carbazole) known to interfere with the activity of the classical tungsten alkylidyne $1^{13,38}$ were metathesized without problems. Chiral centers next to enolizable carbonyl groups were not racemized. An elimination-prone primary tosylate (Table 1, entry 10) as well as an acid- and base-sensitive aldol substructure (Table 2, entries 7) also remained intact. Although the orthogonal character of alkene and alkyne metathesis has previously been recognized,²⁴ the rigorous distinction of the catalysts between the π -systems of

alkynes and olefins is noteworthy: olefins are inert, independent of whether they are mono-, di-, or trisubstuted, terminal, internal, or conjugated to a carbonyl group. Likewise, only the acetylene motif of a 1,3-enyne will undergo productive metathesis (Table 2, entries 11 and 12, and Table 3, entry 2), and even highly base- and acid-sensitive skipped 1,4-enynes posed no problems whatsoever (Table 2, entries 9 and 10).

This excellent profile is particularly evident from a host of applications to compounds bearing more than one polar group (Table 2). We have largely relied upon substrates previously used in this laboratory for the total synthesis of bioactive natural products, including homoepilachnene (entry 6)¹³ and epothilone C (entry 7).^{38,40} The product shown in entry 10 leads to *ent*-amphidinolide V;^{22a,b} this example is particularly noteworthy, as it contains reactive vinylepoxide, hydroxyepoxide, allylic ether, and skipped enyne motifs packed in a dense array, in addition to an ester and a silyl ether group. Entry 12 shows

Table 2. Intramolecular Alkyne Metathesis Reactions

Entry	Substrate	Product	15 ^a	24 ·Et ₂ O ^b	25 ^c
1			70%	97%	94%
2	02N	O ₂ N		85%	
3			91%	73%	78%
4			85%	92%	90%
5	0=		67%	72%	
6	NFmoc	NFmoc		90%	
7	(R = TBS)	S OR OR		91%	
8	TBSO" OTHP MeO TBDPSO" TBDPSO"	TBSO Meo O TBDPSO		82% ^d	
9		0,11,10		90% ^e	
10	TBSQ Q, OTBS	H, OH OTBS		81%	
11	TESO	TESO		84% ^e	
12	Meo OMOM	O HN OMOM		79%	
13	H ₂₈ C ₁₄ —N	H ₂₉ C ₁₄ C ₁₄ H ₂₉	83%	82%	81%

 a **15** (10 mol %), MnCl₂ (10 mol %), toluene, 80 °C, 30 min, then addition of the substrate and reaction at 80 °C unless stated otherwise; no MS added, cf. ref 75. b **24** •Et₂O (2 mol %), toluene, ambient temperature, MS 5 Å. c **25** (5 mol %), MnCl₂ (5 mol %), toluene, 80 °C, 30 min, then addition of the substrate and MS 5 Å and reaction at ambient temperature. d At 80 °C. e With 4–5 mol % of catalyst.

Table 3. Alkyne Cross-Metathesis Reactions

Entry	Substr	ates	Product	15 ^a	24 ·Et ₂ O ^b	25°
1	N=>-=-	5-decyne	N=	76%	65%	72%
2	EtO	tolane	Eto Ph	50%	65%	62%
3	OTBDPS	tolane	OTBDPS Ph	d	62%	61%

^a **15** (10 mol %), MnCl₂ (10 mol %), MS 5 Å, toluene, 80 °C, 30 min, then addition of the substrate and reaction at 100 °C unless stated otherwise. ^b **24** · Et₂O (2 mol %), toluene, ambient temperature, MS 5 Å. ^c **25** (5 mol %), MnCl₂ (5 mol %), toluene, 80 °C, 30 min, then addition of the substrate and MS 5 Å and reaction at ambient temperature. ^d Only low conversion.

another highly adorned case, in which a ketone, ester, amide, two different acetals, a chelation-prone ether, seven chiral centers, and a 1,3-enyne entity are present; this particular product was instrumental for our synthesis of the antibiotic myxovirescin.45 Equally instructive is entry 8, which shows the key intermediate en route to the highly cytotoxic F-ATPase inhibitor cruentaren A.43 Whereas attempted ring closure of this densely substituted divne substrate with the aid of the tungsten alkylidyne 1 resulted only in the cleavage of the -OTHP group due to the Lewis acidity of this complex, 24 · Et₂O afforded the 12membered cycloalkyne in excellent yield. These applications are complemented by two additional examples from an ongoing synthesis project in this laboratory, which will be reported in separate publications in due time. ⁷⁶ They are included in Table 2 to further illustrate the exceptional power of the new alkylidyne complex 24 · Et₂O in forming even particularly fragile (entry 9) as well as fairly strained compounds (entry 11). It is of note that the product displayed in entry 9 could not be made at all with the tungsten alkylidyne 1, whereas complex 6 gave variable results and required high loadings (20–40 mol %), most likely due to competing side reactions with the very reactive oxirane ring of this particular compound. Apparently, the Mo center in 24 · Et₂O is not sufficiently Lewis acidic to cause any damage, and the silanolates do not engage in epoxide opening due to their low nucleophilicity. From the preparative viewpoint, it is also important to note that the RCAM reactions displayed in Table 2 were performed on scales ranging from a few milligrams to several grams of product (see the Supporting Information).

Another relevant case is the cyclooligomerization of the bispropynylated carbazole derivative depicted in Table 2, entry 13. The resulting macrocycle is of interest in material science and was previously best prepared by "precipitation-driven" alkyne metathesis in trichlorobenzene that required, however, a specially designed starting material and a highly sensitive complex of type 5 (81%, ⁶⁸ 61%, ⁷⁷). We were pleased to see that the new procedure, benefitting from the ability of MS 5 Å to drive the conversion of a propynylated substrate to completion at ambient temperature, was at least equally productive, independent of which of the newly prepared catalysts was chosen (81–83%).

Only a few limitations of the new molybdenum complexes have so far been encountered. As noticed previously for the pyridine adduct 13,⁴⁹ epoxides do react with the catalysts and

nitride species and hence destroy the activity of complexes 13 and 15. Importantly, however, these reactive groups seem to be tolerated by the preformed alkylidyne 24. Et₂O (Table 2, entries 9 and 10). Whether this compatibility is due to the fact that 24 · Et₂O reacts at ambient temperature, whereas 13 and 15 need heating, remains to be studied. Likewise, ketones posed no problem with catalysts **24** · Et₂O and **25** (Table 1, entries 7 and 12; Table 2, entry 12) but seem to interfere with the nitride complex 15. Although the product of intermolecular metathesis of the acetophenone derivative shown in Table 1, entry 7, was detected (<40%), full conversion could not be reached in this particular case. Aldehydes, in contrast, seem to mark the limit of the current methodology. Attempts to metathesize a propynylated benzaldehyde derivative were unsuccessful because the catalysts quickly got destroyed (Table 1, entry 6). Ongoing work in this laboratory studies the stoichiometric reaction behavior of these complexes in more detail.

Conclusions

Although alkyne metathesis may never reach the breadth of alkene metathesis because of a smaller substrate base, the potential of this transformation is nevertheless significant, as witnessed by a rapidly increasing number of applications to sophisticated targets. To fully explore its scope, however, it is mandatory to make the required catalysts more readily available and user-friendly. To this end, we present a new generation of alkyne metathesis (pre)catalysts that are optimized for activity on the one hand and practicality on the other. Specifically, the readily available Schrock alkylidyne complex 24. Et₂O constitutes one of the most active catalysts known to date, but retains an outstanding tolerance for functional groups. The only weakly donating triphenylsilanolate ligands impart a well-balanced level of Lewis acidity onto the d⁰-molybdenum center, which is required for high catalytic activity yet is not high enough to endanger polar substituents. At the same time, the sheer size of the Ph₃Si residues prevents more than one alkyne from binding to the metal center and hence disfavors competing polymerization pathways while likely facilitating the cycloreversion of the metallacyclobutadiene intermediates. In contrast to the previously used alkoxides or fluorinated alkoxides, the bulk of the Ph₃SiO- unit seems more "flexible", as bending of the Mo-O-Si angle is facile. "Stretching" of this hinge provides the necessary space about the molybdenum and, as a consequence, prevents substrate-binding from becoming a limiting factor.

Although **24**•Et₂O itself is air- and moisture-sensitive, the corresponding phenanthroline adduct **25** is stable in air for hours.

⁽⁷⁶⁾ Hickmann, V.; Alcarazo, M.; Fürstner, A. J. Am. Chem. Soc. 2010, in press (doi: 10.1021/ja104796a).

⁽⁷⁷⁾ Zhang, W.; Cho, H. M.; Moore, J. S. Org. Synth. 2007, 84, 177.

While solutions of this complex in toluene are inactive per se, reaction of 25 with MnCl₂ releases 24 into the solution by a ligand swap and thereby restores superb performance. The combination $25/\text{MnCl}_2$ is therefore considered a practical entry point into all kinds of alkyne metathesis reactions.

Even more facile is the manipulation of the corresponding phenanthroline nitrido complex 15, which seems indefinitely stable on the benchtop. 15 is accessible in multigram quantities from inexpensive starting materials. It is again by phenanthroline transfer to $MnCl_2$ that an active template is released from this precatalyst, which needs, however, yet to undergo an exchange of nitride for alkylidyne to become active. Although this process requires heating to about 80 °C, the combination $15/MnCl_2$ performed very well in a representative number of inter- and intramolecular alkyne metathesis reactions.

Finally, we report that the addition of 5 Å molecular sieves exerts a pronounced effect on the reaction rate as well as on the conversion in metathesis reactions of alkynes bearing a methyl end-cap. This beneficial influence is ascribed to the removal of 2-butyne from the mixture by absorption into the

pores. This operationally simple measure drives the conversion, allows the reactions to be conducted at ambient temperature and atmospheric pressure, and makes the use of designer substrates unnecessary.

Acknowledgment. Generous financial support by the MPG and the Fonds der Chemischen Industrie is gratefully acknowledged. We thank Dr. Martin Bindl, Dr. Eike Heilmann, and Dr. Daniel A. Clark for preliminary studies; Dr. A. Jantsch for preparing the substrate shown in Table 1, entry 12; V. Hickmann and Dr. K. Micoine for performing the reactions displayed in Table 2, entries 9 and 11; and the NMR, X-ray, and chromatography departments of our Institute for excellent support.

Supporting Information Available: Experimental section including spectroscopic data of all compounds, further information about the structure of the complexes in the solid state, and additional screening data. This material is available free of charge via the Internet at http://pubs.acs.org.

JA104800W