

# The Triple-Bond Metathesis of Aryldiazonium Salts: A Prospect for Dinitrogen Cleavage

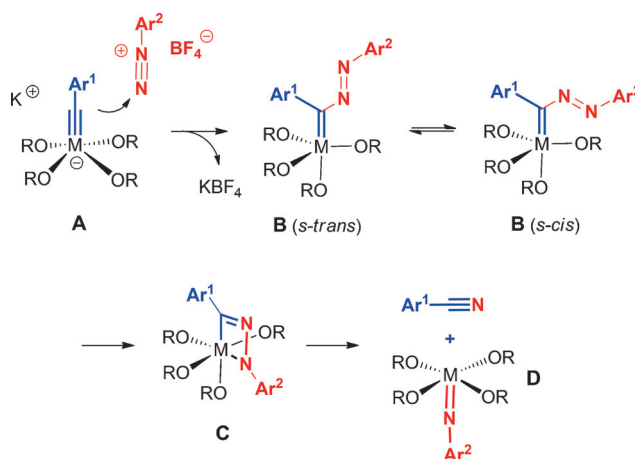
Aaron D. Lackner and Alois Fürstner\*

Dedicated to Professor R. R. Schrock on the occasion of his 70th birthday

**Abstract:** The  $\{N_2\}$  unit of aryldiazonium salts undergoes unusually facile triple-bond metathesis on treatment with molybdenum or tungsten alkylidyne ate complexes endowed with triphenylsilylanolate ligands. The reaction transforms the alkylidyne unit into a nitrile and the aryldiazonium entity into an imido ligand on the metal center, as unambiguously confirmed by X-ray structure analysis of two representative examples. A tungsten nitride ate complex is shown to react analogously. Since the bonding situation of an aryldiazonium salt is similar to that of metal complexes with end-on-bound dinitrogen, in which  $\{N_2\} \rightarrow M$   $\sigma$  donation is dominant and electron back donation minimal, the metathesis described herein is thought to be a conceptually novel strategy toward dinitrogen cleavage devoid of any redox steps and, therefore, orthogonal to the established methods.

The chemistry of aryldiazonium salts is largely dominated by two distinct reactivity modes:<sup>[1,2]</sup> loss of  $N_2$  by heterolytic or homolytic pathways or attack of a suitable nucleophile at the terminal N atom with formation of an azo derivative.<sup>[3]</sup> Metal catalysts or promoters enhance these inherent patterns by oxidative insertion into the weak C–N bond by single-electron transfer, or by end-on coordination to the  $Ar-N_2^+$  unit with formation of (transient) diazenido complexes.<sup>[4–6]</sup> We now report a fundamentally different transformation which is, to the best of our knowledge, without precedent in the literature.

Specifically, we conceived a metathetic cleavage of the  $+>N \equiv N$  bond of aryldiazonium salts that might occur preferentially over nitrogen extrusion, even though the latter process is extremely facile. This expectation arose from the properties of Schrock alkylidynes, which comprise transition-metal centers in their highest possible oxidation state.<sup>[7,8]</sup> Therefore, such complexes will neither serve as single-electron donors nor will they undergo formal oxidative insertion reactions. By virtue of the nucleophilicity of the alkylidyne C atom, however, they might engage the terminal N atom of the diazonium unit in a way that mimics the behavior of conventional nucleophiles (Scheme 1).<sup>[1]</sup> Rather than giving ordinary azo compounds, Fischer carbene complexes of type **B** are expected to form.<sup>[9]</sup> If the barrier for



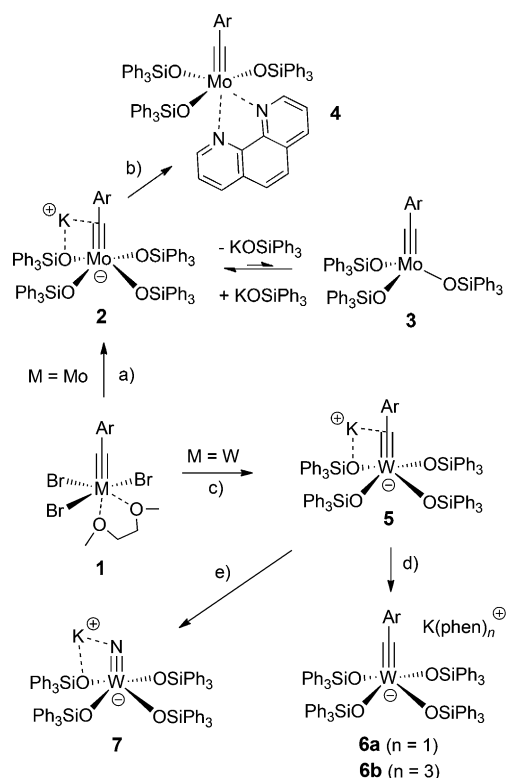
**Scheme 1.** Conceived metathetic cleavage of an aryldiazonium salt; the escorting ions and the derived salt (here  $KBF_4$ ) are arbitrarily chosen;  $M = Mo, W$ .

*s-trans/s-cis* isomerization is low, such intermediates might evolve into a metallacycle of type **C**, which would ultimately furnish a nitrile and an arylimido-metal complex **D** upon formal [2+2] cycloreversion. We reasoned that this pathway could be promoted by not using neutral Schrock alkylidynes but rather the corresponding negatively charged ate complexes **A**: in this case, Coulombic attraction is expected to result in an increased affinity to a cationic aryldiazonium partner, which in turn will favor substrate uptake. Moreover, a salt by-product ( $KBF_4$  in Scheme 1) will be formed concomitantly, which should pay valuable thermodynamic dividends. From a conceptual viewpoint, it is noteworthy that the consequent metathesis event gains a “unimolecular” character, different from the canonic [2+2] mechanism.<sup>[10–12]</sup>

It was previously shown that molybdenum alkylidyne ate species such as **2** endowed with triarylsilanolates are highly competent catalysts for alkyne metathesis (Scheme 2).<sup>[12–15]</sup> However, we attribute the reactivity of **2** to its demonstrated equilibrium under the reaction conditions with the neutral alkylidyne **3**, which exhibits excellent catalytic performance even in highly demanding synthetic applications. Furthermore, the formation of ate complexes is fully reversible upon coordination of donor ligands such as 1,10-phenanthroline, which affords adduct **4** in high yield.<sup>[13,14]</sup> Since the proposed cleavage of a diazonium salt was thought to require an intact ate complex, we initially avoided the use of **2**, instead resorting to more stable variants. Formal replacement of molybdenum by an inherently more Lewis-

[\*] Dr. A. D. Lackner, Prof. A. Fürstner  
Max-Planck-Institut für Kohlenforschung  
45470 Mülheim/Ruhr (Germany)  
E-mail: fuerstner@kofo.mpg.de

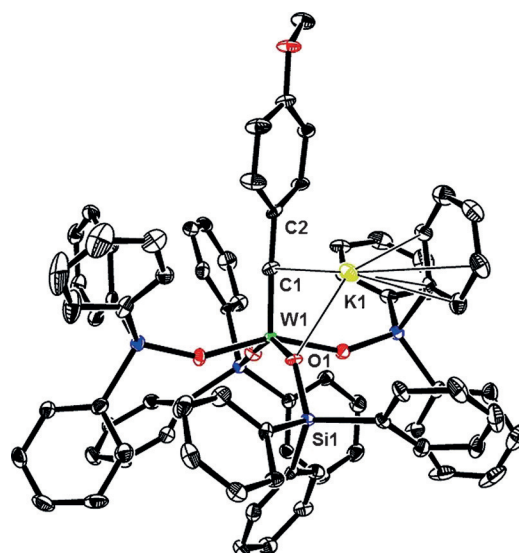
Supporting information for this article is available on the WWW under <http://dx.doi.org/10.1002/anie.201506546>.



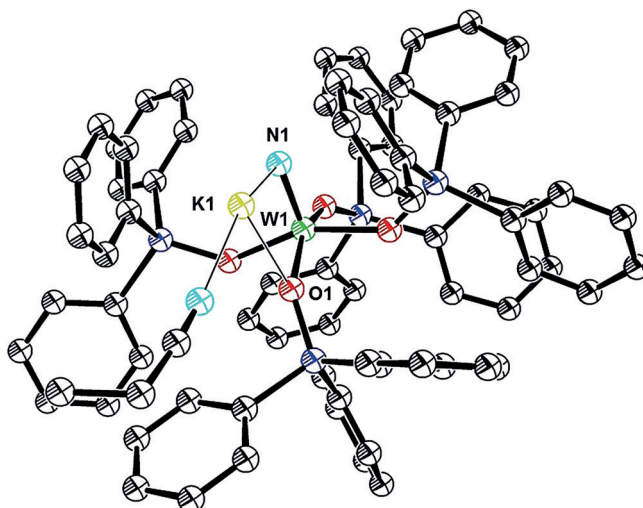
**Scheme 2.** Preparation of different ate complexes for the metathesis of aryldiazonium salts: a)  $\text{Ph}_3\text{SiOK}$ , toluene,  $0^\circ\text{C} \rightarrow \text{RT}$ , 68%, see Ref. [14]; b) phen, toluene, 84%, see Ref. [14]; c)  $\text{Ph}_3\text{SiOK}$ , toluene, 58%; d) phen (1 or 3 equiv), toluene, 90% ( $n=1$ ), 81% ( $n=3$ ); e) propionitrile (5 equiv), toluene,  $60^\circ\text{C}$ , 70%.  $\text{Ar} = 4\text{-MeOC}_6\text{H}_4$ ; phen = 1,10-phenanthroline.

acidic tungsten center proved effective. Thus, the addition of  $\text{Ph}_3\text{SiOK}$  to a solution of **1** ( $M = \text{W}$ ) in toluene furnished the analogous ate complex **5**, which shows no signs of dissociation of the fourth silanolate unit in solution. Complex **5** even persists in the presence of (excess) 1,10-phenanthroline, which fails to bind the tungsten center but rather chelates the escorting potassium cation in the resulting complexes **6a** ( $n=1$ ) and **6b** ( $n=3$ ). Stoichiometric nitrile metathesis<sup>[16,17]</sup> allowed **5** to be converted in good yield into the corresponding nitride ate complex **7**,<sup>[18]</sup> which crystallizes with an equivalent of propionitrile ligated to the potassium counterion (**7-EtCN**).

The structure of complex **5** in the solid state is depicted in Figure 1, and those of the derived phenanthroline adducts **6a** and **6b** are contained in the Supporting Information. As expected, **5** adopts a square-pyramidal coordination geometry, with the alkylidyne forming the apex and the silanolates the basal plane. The  $\text{W} \equiv \text{C}$  distance at  $1.769(7) \text{ \AA}$  is slightly longer than that of neutral alkylidyne complexes of this element<sup>[19]</sup> and the  $\text{W1-C1-Ar}$  unit is almost linear ( $177.1(6)^\circ$ ). The potassium counterion is held in place by weak contacts to an oxygen atom, the negatively polarized alkylidyne C atom, and the  $\pi$  cloud of one of the lateral phenyl rings. The structure of complex **7-EtCN** is largely similar (Figure 2); the key contacts to the escorting cation are now made by one of



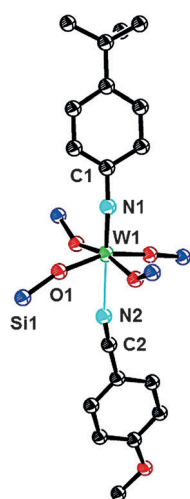
**Figure 1.** Structure of the tungsten alkylidyne ate complex **5**; cocrystallized  $\text{CH}_2\text{Cl}_2$  is not shown for clarity; color code: W = green, O = red, Si = blue, K = yellow.



**Figure 2.** Structure of the tungsten nitride ate complex **7-EtCN** in the solid state; color code: W = green, O = red, Si = blue, K = yellow, N = cyan.

the silanolate O atoms and the terminal N atom, indicative of the nucleophilicity of the nitrido group.

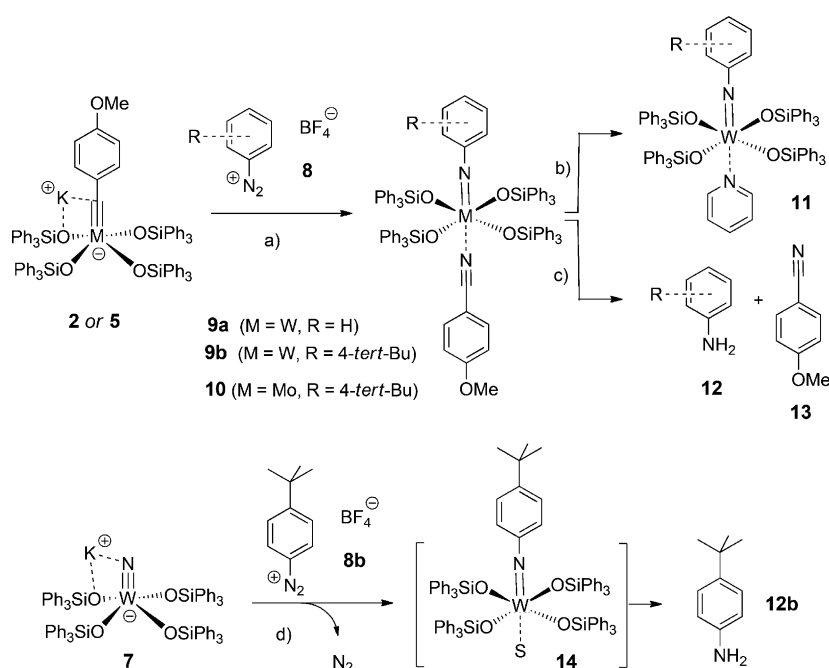
Next, the reactivity of the tungsten alkylidyne ate complex **5** toward aryldiazonium salts was examined. Addition of a bright pink solution of **5** in  $\text{CH}_2\text{Cl}_2$  to a solution of diazonium tetrafluoroborate **8b** ( $R = 4\text{-tert-butyl}$ ) in the same solvent at  $-78^\circ\text{C}$  leads to an instantaneous appearance of an intense purple color, which is tentatively ascribed to the formation of a carbene complex of type **B**, which comprises a diazenido substructure as an “azo-like” chromophore.<sup>[20]</sup> If the ate complex is instead added dropwise to the diazonium salt solution at  $0^\circ\text{C}$ , the purple color appears upon contact and fades within seconds. The mixture gradually turns yellow upon stirring; GC analysis after 30 min indicated 4-*tert*-



**Figure 3.** The structure of complex **9b** in the solid state; only the core region is depicted for clarity (for the entire structure, see the Supporting Information); color code: W = green, O = red, Si = blue, K = yellow, N = cyan.

butylaniline (**12b**) and 4-MeOC<sub>6</sub>H<sub>4</sub>CN (**13**) had formed as the only volatile organic products.<sup>[21]</sup> These compounds, as well as the recorded NMR spectra, are consistent with the formation of the tungsten imido complex **9b** by a purported metathesis reaction. Its presence and constitution was ultimately proven by single-crystal X-ray diffraction (Figure 3). **9b** features the expected N-aryl substituent derived from the diazonium salt and also ligates the substituted benzonitrile originating from the former alkylidyne unit; it is coordinated to the tungsten center *trans* to the imido group to complete a quasi-octahedral ligand sphere. The interaction with the Lewis-acidic metal site leads to a slight elongation of the nitrile bond (N2–C2 1.160(8) Å).<sup>[22]</sup> As expected, the W=N–Ar group is slightly tilted (167.4(4)°) and the W=N bond length of 1.749(5) Å falls into the typical range.<sup>[23]</sup> In any case, the presence of both predicted fragments in a single complex proves that the [N≡N] unit of the aryldiazonium salt must have been cleaved by a formal triple-bond metathesis event. The use of <sup>15</sup>N-labeled diazonium salts confirmed these conclusions (for details, see the Supporting Information).

A selection of other aryldiazonium salts were cleaved analogously, thus showing that this novel reactivity mode is general (Scheme 3). In some cases, the imido complexes were best isolated after exchange of the nitrile by pyridine; alternatively, the crude material was hydrolyzed and the amount of the released aniline **12** and 4-methoxybenzonitrile (**13**) determined. Importantly, the reaction is not confined to the tungsten complex **5**. Specifically, the nitrido ate complex **7** was found to react analogously with release of N<sub>2</sub> (not detected). Although no single crystals of the expected imido complex **14** have been obtained so far (it lacks the stabilizing nitrile present in **9b**), the high yield of the corresponding aniline **12b** after hydrolytic work up leaves no doubt that a metathetic cleavage of the substrate must have taken place.<sup>[24]</sup> Furthermore, the molybdenum ate complex **2** proved equally competent, even though it is capable of reverting to the neutral alkylidyne **3** (see above);<sup>[13,14]</sup> the resulting product complex **10** was unambiguously characterized by X-ray diffraction and found to be isostructural to **9b** (see the Supporting Information). A control experiment proved that it is the ate complex **2** which accounts for the observed N<sub>2</sub> cleavage: thus, treatment of **8b** with authentic **3**<sup>[14]</sup> was



**Scheme 3.** a) **8**, CH<sub>2</sub>Cl<sub>2</sub>, 0°C → RT, 84% (**9a**), 84% (**9b**), 81% (**10**); b) pyridine, 82% (**11c**, R = 4-MeO), 71% (**11d**, R = 4-F); c) H<sub>2</sub>O, 73% (**12e**, R = 2-Br), 60% (**12f**, R = 2,6-dibromo), 51% (**12g**, R = 2-phenyl); d) **8b**, CH<sub>2</sub>Cl<sub>2</sub>, –20°C → RT, then H<sub>2</sub>O, 72%.

unproductive, despite the remarkable performance of this particular neutral alkylidyne in regular alkyne metathesis.<sup>[25–27]</sup>

Although the metathesis of alkynes has gained considerable momentum in recent years,<sup>[12,28]</sup> not least because of the excellent application profile of **2** and relatives, the reactions typically require ambient temperature or higher.<sup>[29]</sup> In contrast, the metathesis of the aryldiazonium salts with **2**, **5**, or **7** proceeds even at ≤ –20°C. This ease is even more remarkable when compared to the difficulty of cleaving the C≡N bond, which limits the preparative significance of nitrile metathesis at this stage.<sup>[16]</sup> The ease can be attributed to a match between the properties of the diazonium substrates and the early transition metal alkylidyne ate complexes as reaction partners. Facile and likely irreversible nucleophilic attack leads to an intermediate which is uniquely poised for metathetic transformation to result in full N–N scission, with only a single bond rotation from *s-trans* to *s-cis* as the kinetic barrier. The robustness of this reactivity is reflected in the successful transformation of even fairly hindered diazonium derivatives (see Scheme 3).

This auspicious reactivity profile is, therefore, believed to provide opportunities beyond the aryldiazonium series. Such substrates can be thought of as consisting of a [N≡N] “ligand” acting as a σ donor to an aryl cation partner, with very little π character in the resulting C–N bond.<sup>[30,31]</sup> This bonding situation is analogous to that of certain (cationic) N<sub>2</sub>-ligated metal complexes in which σ donation of the end-on-bound dinitrogen to the metal center is the dominant interaction, whereas electron back donation from the metal into the π\* orbital of the N<sub>2</sub> ligand is minimal.<sup>[32–34]</sup> The net effect is a polarization of the {N≡N} unit, arguably increasing its



susceptibility to nucleophilic attack by an alkylidyne (or nitride).<sup>[35]</sup> The transfiguration of the newly discovered mode of triple bond metathesis into a (stoichiometric) nitrogen cleavage process is, therefore, a tempting prospect.<sup>[36]</sup> Under this proviso, the reaction would be orthogonal to the established means of dinitrogen activation in that it does not comprise any redox steps;<sup>[37]</sup> it might excel in kinetic terms if the favorable reactivity profile observed with aryldiazonium salts were even partly retained. Finally, such a process would obviate ammonia as the product of N<sub>2</sub> fixation and allow nitrogen to be “stitched” directly into an organic product.<sup>[38]</sup> Although attempts to reduce this tantalizing prospect to practice will almost certainly require a broad screening campaign in view of the vast number of dinitrogen complexes that might qualify as possible substrates, efforts along these lines seem worthwhile and are underway in our laboratory.

## Acknowledgements

Generous financial support by the MPG and the Alexander von Humboldt-Foundation (fellowship to A.D.L.) is gratefully acknowledged. We thank Dr. J. Heppkeausen for preliminary studies into the tungsten alkylidyne series, Dr. R. Goddard for solving the X-ray structures, and Prof. M. Alcarazo for helpful discussions.

**Keywords:** alkylidynes · aryldiazonium salts · metathesis · molybdenum · tungsten

**How to cite:** *Angew. Chem. Int. Ed.* **2015**, *54*, 12814–12818  
*Angew. Chem.* **2015**, *127*, 13005–13009

- [1] a) *The Chemistry of Diazonium and Diazo Groups*, pt. 1–2 (Ed.: S. Patai), Wiley, Chichester, **1978**; b) K. H. Saunders, R. L. M. Allen, *Aromatic Diazo Compounds*, 3rd ed., Edward Arnold, London, **1985**.
- [2] F. Mo, G. Dong, Y. Zhang, J. Wang, *Org. Biomol. Chem.* **2013**, *11*, 1582–1593.
- [3] Depending on the very nature of the delivered N substituent, the resulting azo compound will either lose N<sub>2</sub> or can be indefinitely stable.
- [4] A. Roglans, A. Pla-Quintana, M. Moreno-Mañas, *Chem. Rev.* **2006**, *106*, 4622–4643.
- [5] C. Galli, *Chem. Rev.* **1988**, *88*, 765–792.
- [6] a) D. Sutton, *Chem. Rev.* **1993**, *93*, 995–1022; b) R. B. King, *J. Organomet. Chem.* **1995**, *500*, 187–195.
- [7] R. R. Schrock, *Angew. Chem. Int. Ed.* **2006**, *45*, 3748–3759; *Angew. Chem.* **2006**, *118*, 3832–3844.
- [8] a) R. R. Schrock, *Chem. Rev.* **2002**, *102*, 145–179; b) R. R. Schrock, *J. Chem. Soc. Dalton Trans.* **2001**, 2541–2550.
- [9] For the formation of some remotely related carbenes on reaction of tungsten alkylidyne ate complexes carrying calixarene ligands with electrophiles such as COCl<sub>2</sub>, I<sub>2</sub>, Ph<sub>3</sub>SnCl, or R<sub>2</sub>PCl, see S. Dovesi, E. Solari, R. Scopelliti, C. Floriani, *Angew. Chem. Int. Ed.* **1999**, *38*, 2388–2391; *Angew. Chem.* **1999**, *111*, 2545–2547.
- [10] T. J. Katz, J. McGinnis, *J. Am. Chem. Soc.* **1975**, *97*, 1592–1594.
- [11] a) J. H. Wengrovius, J. Sancho, R. R. Schrock, *J. Am. Chem. Soc.* **1981**, *103*, 3932–3934; b) S. F. Pedersen, R. R. Schrock, M. R. Churchill, H. J. Wasserman, *J. Am. Chem. Soc.* **1982**, *104*, 6808–6809.
- [12] A. Fürstner, *Angew. Chem. Int. Ed.* **2013**, *52*, 2794–2819; *Angew. Chem.* **2013**, *125*, 2860–2887.
- [13] J. Heppkeausen, R. Stade, R. Goddard, A. Fürstner, *J. Am. Chem. Soc.* **2010**, *132*, 11045–11057.
- [14] J. Heppkeausen, R. Stade, A. Kondoh, G. Seidel, R. Goddard, A. Fürstner, *Chem. Eur. J.* **2012**, *18*, 10281–10299.
- [15] a) V. Hickmann, A. Kondoh, B. Gabor, M. Alcarazo, A. Fürstner, *J. Am. Chem. Soc.* **2011**, *133*, 13471–13480; b) K. Micoine, A. Fürstner, *J. Am. Chem. Soc.* **2010**, *132*, 14064–14066; c) W. Chaladaj, M. Corbet, A. Fürstner, *Angew. Chem. Int. Ed.* **2012**, *51*, 6929–6933; *Angew. Chem.* **2012**, *124*, 7035–7039; d) S. Benson, M.-P. Collin, A. Arlt, B. Gabor, R. Goddard, A. Fürstner, *Angew. Chem. Int. Ed.* **2011**, *50*, 8739–8744; *Angew. Chem.* **2011**, *123*, 8898–8903; e) L. Brewitz, J. Llaveria, A. Yada, A. Fürstner, *Chem. Eur. J.* **2013**, *19*, 4532–4537; f) S. M. Rummelt, J. Preindl, H. Sommer, A. Fürstner, *Angew. Chem. Int. Ed.* **2015**, *54*, 6241–6245; *Angew. Chem.* **2015**, *127*, 6339–6343.
- [16] a) A. M. Geyer, E. S. Wiedner, J. B. Gary, R. L. Gdula, N. C. Kuhlmann, M. J. A. Johnson, B. D. Dunietz, J. W. Kampf, *J. Am. Chem. Soc.* **2008**, *130*, 8984–8999; b) R. L. Gdula, M. J. A. Johnson, *J. Am. Chem. Soc.* **2006**, *128*, 9614–9615.
- [17] In line with the metathetic course of the reaction, 1-(4-methoxyphenyl)but-1-yne and 4,4'-dimethoxytolane were detected by GC-MS.
- [18] For analogous Mo-nitride complexes, see Refs. [13,14] and the following: M. Bindl, R. Stade, E. K. Heilmann, A. Picot, R. Goddard, A. Fürstner, *J. Am. Chem. Soc.* **2009**, *131*, 9468–9470.
- [19] For example, the W≡C bond in [(tBuO)<sub>3</sub>W≡CPh] is 1.758(5) Å, see F. A. Cotton, W. Schwotzer, E. S. Shamshoum, *Organometallics* **1984**, *3*, 1770–1771.
- [20] We used terminally <sup>15</sup>N-labeled (2-phenyl)phenyldiazonium tetrafluoroborate in preliminary attempts to characterize a pertinent intermediate by spectroscopic means, as this substrate reacts somewhat more slowly. A carbon signal at δ<sub>C</sub> = 252.2 ppm with a distinctive J(<sup>13</sup>C,<sup>15</sup>N) coupling constant of 6 Hz was detected, which we tentatively ascribe to a Fischer-carbene intermediate of type **B**.
- [21] Products from ligand decomposition are also observed.
- [22] The typical bond length of an Ar-CN group is 1.138 Å, see F. H. Allen, O. Kennard, D. G. Watson, L. Brammer, A. G. Orpen, R. Taylor, *J. Chem. Soc. Perkin Trans. 2* **1987**, S1–S19.
- [23] These metrics are comparable to the W=N bond length (1.772(22) Å) and the W=N-Ar bond angle (169.5(19)°) in [(iPrO)<sub>4</sub>W=NPh], see J. T. Barry, M. H. Chisholm, K. Folting, J. C. Huffman, W. E. Streib, *Polyhedron* **1997**, *16*, 2113–2133.
- [24] Inspection of the crude material by GC-MS and IR suggest that small amounts of (tert-butyl)phenylazide are also present; studies into this interesting side reaction are underway.
- [25] a) P. Persich, J. Llaveria, R. Lhermet, T. de Haro, R. Stade, A. Kondoh, A. Fürstner, *Chem. Eur. J.* **2013**, *19*, 13047–13058; b) R. Lhermet, A. Fürstner, *Chem. Eur. J.* **2014**, *20*, 13188–13193.
- [26] a) C. M. Neuhaus, M. Liniger, M. Stieger, K.-H. Altmann, *Angew. Chem. Int. Ed.* **2013**, *52*, 5866–5870; *Angew. Chem.* **2013**, *125*, 5978–5983; b) K. J. Ralston, H. C. Ramstadius, R. C. Brewster, H. S. Niblock, A. H. Hulme, *Angew. Chem. Int. Ed.* **2015**, *54*, 7086–7090; *Angew. Chem.* **2015**, *127*, 7192–7196.
- [27] a) J. Willwacher, A. Fürstner, *Angew. Chem. Int. Ed.* **2014**, *53*, 4217–4221; *Angew. Chem.* **2014**, *126*, 4301–4305; b) K. Gebauer, A. Fürstner, *Angew. Chem. Int. Ed.* **2014**, *53*, 6393–6396; *Angew. Chem.* **2014**, *126*, 6511–6514; c) G. Valot, C. S. Regens, D. P. O'Malley, E. Godineau, H. Takikawa, A. Fürstner, *Angew. Chem. Int. Ed.* **2013**, *52*, 9534–9538; *Angew. Chem.* **2013**, *125*, 9713–9717; d) K. Lehr, R. Mariz, L. Leseurre, B. Gabor, A. Fürstner, *Angew. Chem. Int. Ed.* **2011**, *50*, 11373–11377; *Angew. Chem.* **2011**, *123*, 11575–11579; e) L. Hoffmeister, P. Persich, A.

- Fürstner, *Chem. Eur. J.* **2014**, *20*, 4396–4402; f) L. Hoffmeister, T. Fukuda, G. Pototschnig, A. Fürstner, *Chem. Eur. J.* **2015**, *21*, 4529–4533; g) F. Ungeheuer, A. Fürstner, *Chem. Eur. J.* **2015**, *21*, 11387–11392.
- [28] a) R. R. Schrock, *Chem. Commun.* **2013**, *49*, 5529–5531; b) W. Zhang, J. S. Moore, *Adv. Synth. Catal.* **2007**, *349*, 93–120; c) H. Yang, Y. Jin, Y. Du, W. Zhang, *J. Mater. Chem. A* **2014**, *2*, 5986–5993; d) A. Fürstner, P. W. Davies, *Chem. Commun.* **2005**, 2307–2320; e) C. Deraedt, M. d'Halluin, D. Astruc, *Eur. J. Inorg. Chem.* **2013**, 4881–4908; f) S. Shahane, C. Bruneau, C. Fischmeister, *ChemCatChem* **2013**, *5*, 3436–3459.
- [29] Ref. [13] provides a single example of an alkyne metathesis carried out at  $-10^{\circ}\text{C}$ .
- [30] a) R. Glaser, C. J. Horan, H. Zollinger, *Angew. Chem. Int. Ed. Engl.* **1997**, *36*, 2210–2213; *Angew. Chem.* **1997**, *109*, 2324–2328; b) R. Glaser, C. J. Horan, *J. Org. Chem.* **1995**, *60*, 7518–7528.
- [31] Analysis of the frontier orbitals of  $\text{PhN}_2^+$  indicates significant  $\sigma$  donation from  $\{\text{N}_2\}$  into the formal  $\{\text{Ph}\}^+$  fragment, thereby giving rise to a low-lying bonding orbital ( $E = -0.493\text{ eV}$ ); all other occupied frontier orbitals show antibonding interactions between these units, thus implying little back donation from the aromatic  $\pi$  cloud to the  $\text{N}_2$  group. The LUMO has the largest lobes at the C–N bond, which explains why nucleophiles or electron donors able to interact with this orbital will readily extrude  $\text{N}_2$ .
- [32] J. Chatt, J. R. Dilworth, R. L. Richards, *Chem. Rev.* **1978**, *78*, 589–625.
- [33] F. Studt, F. Tuczek, *J. Comput. Chem.* **2006**, *27*, 1278–1291.
- [34] The fact that aryldiazonium ions undergo facile  $\text{N}_\alpha/\text{N}_\beta$  rearrangement through in situ release and trapping of  $\text{N}_2$  by the concomitantly formed  $\text{Ar}^+$  corroborates the analogy to end-on-bound  $\{\text{N}_2\}$  complexes, see a) E. S. Lewis, R. E. Holliday, *J. Am. Chem. Soc.* **1969**, *91*, 426–430; b) I. Szele, H. Zollinger, *J. Am. Chem. Soc.* **1978**, *100*, 2811–2815.
- [35] For pioneering studies showing successful attack of an organometallic reagent onto end-on-coordinated nitrogen complexes, see a) D. Sellmann, W. Weiss, *Angew. Chem. Int. Ed. Engl.* **1977**, *16*, 880–881; *Angew. Chem.* **1977**, *89*, 918–919; b) D. Sellmann, W. Weiss, *Angew. Chem. Int. Ed. Engl.* **1978**, *17*, 269–270; *Angew. Chem.* **1978**, *90*, 295–296; c) D. Sellmann, W. Weiss, *J. Organomet. Chem.* **1978**, *160*, 183–196.
- [36] Even a catalytic scenario is a priori not excluded, although certainly difficult to realize. Arylimido ligands afford dichloro complexes on protonation, which open a way back to an alkylidyne; for leading references on replacement of imido ligands, see a) R. R. Schrock, I. A. Weinstock, A. D. Horton, A. H. Liu, M. H. Schofield, *J. Am. Chem. Soc.* **1988**, *110*, 2686–2687; b) R. R. Schrock, R. T. DePue, J. Feldman, K. B. Yap, D. C. Yang, W. M. Davis, L. Park, M. DiMare, M. Schofield, J. Anhaus, E. Walborsky, E. Evitt, C. Krüger, P. Betz, *Organometallics* **1990**, *9*, 2262–2275.
- [37] a) H.-P. Jia, E. A. Quadrelli, *Chem. Soc. Rev.* **2014**, *43*, 547–564; b) K. C. MacLeod, P. L. Holland, *Nat. Chem.* **2013**, *5*, 559–565; c) G. Ertl, *Angew. Chem. Int. Ed.* **2008**, *47*, 3524–3535; *Angew. Chem.* **2008**, *120*, 3578–3590; d) R. R. Schrock, *Acc. Chem. Res.* **2005**, *38*, 955–962; e) M. P. Shaver, M. D. Fryzuk, *Adv. Synth. Catal.* **2003**, *345*, 1061–1076; f) M. Hidai, *Coord. Chem. Rev.* **1999**, *185–186*, 99–108.
- [38] For a discussion, see M. Mori, *J. Organomet. Chem.* **2004**, *689*, 4210–4227.

Received: July 15, 2015

Published online: September 2, 2015