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J. Am. Chem. Soc., **2005**, 127 (41), 14202-14203• DOI: 10.1021/ja0556023 • Publication Date (Web): 23 September 2005 Downloaded from http://pubs.acs.org on March 25, 2009



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Published on Web 09/23/2005

Unprecedented Coordination Modes and Demetalation Pathways for Unbridged Polyenyl Ligands. Ruthenium η^1, η^4 -Cycloheptadienyl Complexes from Allyl/Alkyne Cycloaddition

Christina M. Older, Robert McDonald, and Jeffrey M. Stryker*

Department of Chemistry, University of Alberta, Edmonton, Alberta, Canada T6G 2G2 Received August 16, 2005; E-mail: jeff.stryker@ualberta.ca

Metal-mediated cycloaddition reactions provide a range of cyclic polyenyl ligands coordinated to transition metals. The coordination mode adopted by such ligands determines to a considerable extent the strategies available for subsequent elaboration and demetalation of the organic moiety. For the most part, cyclic polyenyl ligands unconstrained by bridged structures adopt fully conjugated coordination modes, typically as a direct consequence of thermodynamic preferences. New coordination modes and, in particular, *less* conjugated coordination modes raise new possibilities for post-cyclization functionalization and demetalation pathways, both essential to the development of synthetically valuable transformations.

All unbridged cycloheptadienyl complexes of the transition metals exhibit fully conjugated η^5 -coordination.^{1–5} Transient formation of η^1, η^4 -coordinated intermediates has been proposed for cobalt- and iridium-mediated allyl/alkyne [3 + 2 + 2] cycloaddition reactions,^{4–6} but no evidence to support this conjecture has been obtained. Here, however, we report the selective formation of unprecedented η^1, η^4 -cycloheptadienyl complexes from ruthenium-mediated allyl/alkyne cycloaddition. The unique coordination mode promotes unique oxidative demetalation pathways, leading to the efficient construction of synthetically valuable seven-membered carbocycles.

To induce ruthenium to mediate seven-membered ring formation over cyclization to the kinetically favored six-membered ring, the ancillary η^6 -benzene ligand used in previous investigations⁷ was replaced with the sterically larger η^6 -hexamethylbenzene ligand. In the event, treatment of allyl complex **1a**⁸ with excess ethyne leads to the clean formation of two inseparable cycloaddition products, strongly favoring seven- over six-membered ring formation (Scheme 1).⁹ The major product was identified as the unexpected *symmetric* η^1, η^4 -cycloheptadienyl complex **2** by twodimensional NMR spectroscopy.¹⁰ The resonance for the unique σ -bonded methine appears characteristically shifted upfield in both ¹H NMR (δ -0.09) and ¹³C NMR (δ -38.1) spectra. Surprisingly, complex 2 is both indefinitely stable *and* fluxional at room temperature, as established by double irradiation experiments,⁹ rapidly equilibrating with the otherwise undetected cycloheptatriene hydride complex 2'. At 80 °C, however, slow isomerization occurs to give the more thermodynamically stable η^{5} -coordinated isomer **4**.^{9,11} The formation and subsequent isomerization of the η^{1} , η^{4} -complex 2 strongly supports prior mechanistic proposals rationalizing η^{5} -cycloheptadienyl ring formation in the cobalt and iridium series.^{4a,b}

A lower activation barrier for η^1, η^4 - to η^5 -isomerization is apparent in the reaction of *tert*-butylacetylene, which proceeds directly to η^5 -cycloheptadienyl complex **5**^{9,12} even under mild reaction conditions. No η^1, η^4 -coordinated isomer is detected in the crude reaction mixture. Among several minor products, the simple acyclic 1-*tert*-butyl- η^5 -pentadienyl complex **6**⁹ (~4%, not shown) can be isolated by chromatography.

Although some reactions of complex **1a** with alkynes are complicated,¹³ treatment with dimethylacetylene dicarboxylate (DMAD) provides η^1, η^4 -cycloheptadienyl complex **7** free from significant byproducts.^{9,14} The substituent array in complex **7** can arise only by extensive isomerization of the initial η^1, η^4 -cycloheptadienyl intermediate, presumably via iterative β -hydride elimination and reinsertion (Scheme 2). Notably, the system reacts *with perfect fidelity for migration to* η^1, η^4 -coordination rather than fully conjugated η^5 -hapticity. No further reaction is induced upon heating, strongly suggesting that the η^1, η^4 -coordination mode is thermodynamically favored.

The electron-deficient alkyne also allows for cycloaddition reactions incorporating two different alkynes. Unique among alkynes,⁸ treatment of complex **1a** with 1 equiv of DMAD leads to exclusive formation of the *acyclic* adduct, [(C₆Me₆)Ru(1,2-*syn*-dicarbomethoxypentadienyl)]OTf **8** (90%, not shown).⁹ In wet benzene, however, the reaction stops at the η^1, η^2 -vinyl olefin intermediate, providing cationic aquo complex **9** (Scheme 1).⁹ This



10.1021/ja0556023 CCC: \$30.25 © 2005 American Chemical Society





complex converts slowly and quantitatively at room temperature to η^5 -pentadienyl complex 8. Similarly, solvolysis of allyl chloride complex 1b15 in trifluoroethanol containing DMAD gives the analogous but neutral vinyl olefin complex 10 in near quantitative vield.9

Complexes 9 and 10 transform selectively to η^1, η^4 -cycloadducts on treatment with a second alkyne. Optimal yields and selectivity are obtained from neutral complex 10 under conditions of assisted ionization, providing η^1 , η^4 -cycloheptadienyl complexes **11** and **13** from the reactions of ethyne and 2-butyne, respectively (Scheme 1).9,16 Structurally distinct minor products are isolated from each reaction, including the unusual *acyclic* η^3 , η^2 -heptadienyl complex 12, which presumably arises from allylic C-H bond activation following migratory insertion of ethyne. Structural assignments for complexes 12, 13, and 14 have been confirmed by X-ray crystallography.9

The unique coordination of the η^1, η^4 -cycloheptadienyl ring suggested the use of oxidative bond heterolysis as a novel demetalation strategy, simultaneously effecting both scission of the metal-carbon σ -bond and minimization of metal-diene backbonding.¹⁷ In the event, exhaustive iodinolysis indeed mediates the decomplexation of cycloadducts 7 and 11, returning the unusual tricyclic lactones 17 in moderate to good yields (Scheme 3).9,18 The Ru(III) coproduct was identified crystallographically as triiodide salt 18,9 accounting for the unexpected stoichiometry in iodine. Direct chromatography of the crude reaction mixture without thiosulfate workup unmasks the latent seven-membered ring, providing cycloheptatriene 19 exclusively.^{9,19} Iodinolysis of complex 13 leads directly to cycloheptatriene anhydride 20 in as yet variable yields,9 a bicyclic ring system of considerable synthetic interest.20

The use of stoichiometric (but recyclable) ruthenium obviously limits the potential use of this cycloaddition reaction in organic synthesis. Nonetheless, the strong preference for η^1, η^4 -coordination of the seven-membered ring is unique in transition metal chemistry, offering mechanistic insight and raising new opportunities for developing synthetically valuable metal-mediated cycloaddition and demetalation reactions.

Acknowledgment. Financial support from NSERC and the University of Alberta is gratefully acknowledged. R.M. is affiliated with the Department of Chemistry X-ray Crystallography Laboratory.

Supporting Information Available: Experimental procedures and complete characterization data for all new compounds; details of the X-ray crystallography for complexes 12-14 and 18. This material is available free of charge via the Internet at http://pubs.acs.org.

References

- (1) See, for example: (a) Salzer, A.; Werner, H. J. Organomet. Chem. 1975, 87, 101-108. (b) Bennett, M. A.; Matheson, T. W. J. Organomet. Chem. 1978, 153, C25–27. (c) Salzer, A.; Bigler, P. Inorg. Chim. Acta 1981, 48, 199–203. (d) Williams, G. M.; Fisher, R. A.; Heyn, R. H. Organo-metallics 1986, 5, 818–819. (e) Chen, W.; Sheridan, J. B.; Cote, M. L.; Lalancette, R. A. Organometallics 1996, 15, 2700–2706 and references therein.
- (2) Nonconjugated pentahapticity (invariably as η^2, η^3 -coordination) occurs more commonly in larger rings with greater coordination strain and/or conformational barriers: (a) Lewis, J.; Parkins, A. W. J. Chem. Soc. (A)
 1969, 953–957. (b) Bennett, M. A.; Matheson, T. W.; Robertson, G. B.;
 Smith, A. K.; Tucker, P. A. Inorg. Chem. 1981, 20, 2353–2365. (c) Itoh,
 K.; Nagashima, H.; Ohshima, T.; Oshima, N.; Nishiyama, H. J. Organomet. Chem. 1984, 272, 179-188 and references therein
- (3) An early claim^{2a} that protonation of $(C_5H_5)Co(\eta^4$ -cycloheptatriene) provides $(C_5H_3)Co(\eta^2,\eta^3$ -cycloheptadienyl)⁺ is not supported by spectroscopic data and remains unsubstantiated. The isolated product of this reaction was subsequently identified as the expected η^5 -cycloheptadienyl complex.
- (a) Schwiebert, K. E.; Stryker, J. M. J. Am. Chem. Soc. 1995, 117, 8275-(4)8276. (b) Etkin, N.; Dzwiniel, T. L.; Schwiebert, K. E.; Stryker, J. M. J. Am. Chem. Soc. **1998**, 120, 9702–9703. (c) Dzwiniel, T. L.; Etkin, N.; Stryker, J. M. J. Am. Chem. Soc. **1999**, 121, 10640–10641. (d) From cyclopentenyl ring expansion: Dzwiniel, T. L.; Stryker, J. M. J. Am. Chem. Soc. 2004, 126, 9184-9185.
- (5) Mn/ η^3 -allyl/ α , ω -diyne: (a) Tang, J.; Shinokubo, H.; Oshima, K. Orga-Initian and Antonio Sciences (1998) 17, 290–292. (b) Nishikawa, T.; Kakiya, H.; Shinokubo, H.; Oshima, K. J. Am. Chem. Soc. 2001, 123, 4629–4630.
- Ostinia, K. J. Am. Chem. Soc. 2001, 125, 4029 4050.
 Other recent [3 + 2 + 2] cycloaddition reactions: (a) Baluenga, J.; Barrio, P.; López, L. A.; Tomás, M.; García-Granda, S.; Alvarez-Rúa, C. Angew. Chem., Int. Ed. 2003, 42, 3008–3011. (b) Saito, S.; Masudo, M.; Komegawa, S. J. Am. Chem. Soc. 2004, 126, 10540–10541. (6)
- (a) Lutsenko, Z. L.; Aleksandrov, G. G.; Petrovskii, P. V.; Shubina, E. S.; Andrianov, V. G.; Struchkov, Y. T.; Rubezhov, A. Z. J. Organomet. Chem. 1985, 281, 349–364. (b) Lutsenko, Z. L.; Petrovskii, P.; Bezrukova, A.; Rubezhov, A. Z. Bull. Acad. Sci. USSR 1988, 735–738.
- Older, C. M.; Stryker, J. M. Organometallics 1998, 17, 5596-5598
- Complete experimental details are provided as Supporting Information.
- (10) The minor product **3** is spectroscopically very similar to Rubezhov's $[(C_6H_6)Ru(2,3,4,5-tetramethyl-1-methanocyclohexadiene)]^+PF_6^- complex.^{7b} (11) Complex$ **4** $is spectroscopically identical to the known PF_6^- salt.^{2c}$
- (12) The substitution pattern for η^5 -cycloheptadienyl complex 5 is identical to that obtained from (C₅Me₅)Co(allyl)OTf and 3,3-dimethyl-1-butyne.^{4b}
- (13) Phenylacetylene undergoes oligomerization rather than cyclization. Disubstituted alkynes undergo exclusive [3 + 2] cycloaddition with concomitant demethylation of the hexamethylbenzene ligand.⁸ Reactions involving 2-butyne (≥ 2 equiv) diverge substantially and will be discussed elsewhere
- (14) Neither the cobalt nor the iridium system reacts with DMAD. No tractable product was obtained from the reaction of DMAD with (C6H6)Ru(allyl)+
- (15) Bennett, M. A.; Huang, T.-N.; Turney, T. W. J. Chem. Soc., Chem. Commun. 1979, 312–313.
- (16)That η^1, η^4 -complexes 7, 11, and 13 face no significant kinetic barrier to conjugation is strongly suggested by the isolation of η^5 -cycloheptadienyl complex 15 in modest yield (42%) as the major among many products obtained from the reaction of complex 10 with phenylacetylene:



- (17) This demetalation strategy fails in the cobalt η^5 -cycloheptadienyl series. This extensive transformation presumably involves intramolecular iodolactonization/iodide elimination, with concomitant iodide-mediated ester demethylation. Related, completely organic, transformations are known: Garratt, D. G.; Ryan, M. D.; Beaulieu, P. L. J. Org. Chem. 1980, 45, 839-845.
- (19) This process nominally involves lactone elimination followed, presumably,
- (20) See, for example: (a) Tanino, K.; Shimizu, T.; Miyama, M.; Kuwajima, L. J. Am. Chem. Soc. 2000, 122, 6116–6117. (b) Wockenfull, B.; Wolff, C.; Tochtermann, W. Tetrahedron 1997, 53, 13703–13708. (c) Bowden, S. L.; Harris, S. A.; Heller, H. G.; Hewlins, M. J. E. J. Chem. Soc., Perkin Trans. 1 1992, 725-728. (d) Rainer, H. Ugi, I. Angew. Chem., Int. Ed. Engl. 1985, 24, 594-596. JA0556023