

Metal-Metal Quadruple Bonds Supported by 5-Ethynylthiophene-2carboxylato Ligands: Preparation, Molecular and Electronic Structures, Photoexcited State Dynamics, and Application as **Molecular Synthons**

Samantha E. Brown-Xu, Malcolm H. Chisholm,* Christopher B. Durr, and Thomas F. Spilker

Department of Chemistry and Biochemistry, The Ohio State University, Columbus, Ohio 43210, United States

S Supporting Information



ABSTRACT: From the reaction between M2(T'PB)4 and 2 equiv of 5-ethynylthiophene-2-carboxylic acid (H-ThCCH) in toluene, the complexes trans- $M_2(T^iPB)_2(ThCCH)_2$, where M = Mo (I) or W (II) and $T^iPB = 2,4,6$ -triisopropyl benzoate, have been isolated and characterized by ¹H NMR, IR, MALDI-TOF MS, UV-vis, steady-state emission, transient absorption, and time-resolved infrared (TRIR) spectroscopies and single-crystal X-ray crystallography for I. The molecular structure of I confirms the *trans*-substitution pattern and the extended conjugation of the ethynylthienyl ligands via interaction with the Mo₂ δ orbital. The HOMO of both I and II is the $M_{3}\delta$ orbital, and the intense color of the compounds (I is red and II is blue) is due to the $M_2\delta$ -to-ThCCH ¹MLCT transition. The S₁ states for I and II are ¹MLCT. The T₁ state is ³MLCT for II, but ³MoMo $\delta\delta^*$ for I. The TRIR spectra of the $\nu(C \equiv C)$ stretch in the MLCT states are consistent with the delocalization of the electron over both ThCCH ligands. Compound I is shown to be a synthem for the preparation of $trans-Mo_2(T^{i}PB)_2(ThCCPh)_2$ (III) and trans- $Mo_2(T'PB)_2(ThCCAuPPh_3)_2$ (IV). Both III and IV have been characterized spectroscopically and by single-crystal X-ray diffraction. The structure of III indicates the extended π -conjugation of the trans-ethynyl-thienyl units extends to the added phenyl rings.

1. INTRODUCTION

Conjugated organic polymers and even small oligomers are important in a range of rapidly emerging technologies including light-emitting diodes, field effect transistors, photovoltaic devices, and sensors.¹⁻⁶ The introduction of metal ions into these systems affords a variety of additional features, as is evident from fundamental studies by Hagihara 7 and the work of Wong in the introduction of platinum⁸⁻¹⁴ and other metals^{15,16} into conjugated ethynylthiophenes. The metal can enhance emission from triplet states by intersystem crossing and may allow hole transport by $d\pi - p\pi$ conjugation.¹⁷ We have for some time had an interest in the introduction of metal-metal (M_2) quadruply bonded units into organic π -systems and in the preparation of metalated polymers. The $M_2\delta$ orbital can interact directly with a π -conjugated organic molecule that is attached to the M_2 unit via a carboxylate or the related amidinate ligand.^{18–23} We have previously attempted to prepare polymers/oligomers containing M₂ units by the reaction between the homoleptic carboxylates $M_2(O_2CR)_4$ and conjugated dicarboxylic acids. For M = Mo and ter- and pentathienyl ($\alpha_{,}\alpha'$ -linked) dicarboxylic acids, oligomers were obtained suitable for spin coating and, when fabricated, showed electroluminescence.²⁴ However, this procedure gave only lowmolecular-weight materials having 6-8 M₂ units and containing molecular loops, triangles, and other species. Indeed, reactions involving the smallest of dicarboxylic acids, namely oxalic acid, gave almost exclusively triangles.²⁵ We are therefore seeking alternate strategies for the synthesis of oligomers and polymers by employing ethynyl and vinyl end groups that may be used in more well-developed carbon-carbon coupling reactions and eliminate the reliance on metathetic reactions at the M₂ center.

Received: January 16, 2013

Scheme 1. Synthesis of $I-IV^a$



"Ar = 2,4,6-triisopropylphenyl. Reagents: (a) toluene, (b) excess phenyl iodide, THF: Et_3N , Pd(PPh₃)₂Cl₂, CuI; (c) 2 equiv of Ph₃PAuCl, Et₃N, THF:DCM.

We describe here the synthesis of thienylcarboxylates bearing terminal ethypyl groups bound to the M_2 center and show that these are viable synthons for the synthesis of extended π -systems and molecular triads. The two principal quadruply bonded complexes are also shown to exhibit very interesting photophysical properties.

2. RESULTS AND DISCUSSION

2.1. Preparation. The synthesis of the new compounds reported in this work is outlined in Scheme 1.

The compounds $trans-M_2(T^iPB)_2(ThCCH)_2$, where M = Mo (I) and W (II), $T^iPB = 2,4,6$ -triisopropyl benzoate, and ThCCH = 5-ethynylthiophene-2-carboxylate, are soluble in toluene, THF, and dichloromethane (DCM). They are airsensitive and intensely colored: I is red, and II is blue. They give molecular ions in the MALDI-TOF MS, and their ¹H NMR and other characterization data are given in the Supporting Information.

The compound *trans*-Mo₂(TⁱPB)₂(ThCCPh)₂ (III) was prepared by Sonogashira coupling involving phenyl iodide. Compound III is purple, in contrast to its precursor I. The gold-containing complex, *trans*-Mo₂(TⁱPB)₂(ThCCAuPPh₃)₂ (IV), is red like its precursor I. Further characterization data for III and IV are given in the Supporting Information.

2.2. Single-Crystal and Molecular Structures. The molecular structures of the dimolybdenum-containing complexes I, III, and IV are shown in Figures 1, 2, and 3, respectively.

Each structure has a molecular center of inversion, and the phenyl rings of the TⁱPB ligands are essentially out of conjugation with the carboxylate π -system. The dihedral planes involving the CO₂ and C₆ rings are 86.7° (I), 85.2° (III), and 81.9° (IV). In contrast, the thienyl ring and its attendant carboxylate are nearly planar, with dihedral angles of 3.8° (I), 7.7° (III), and 1.4° (IV). For compound III, the dihedral angle between the phenyl and thienyl rings is 9.3°. The central Mo₂(O₂R)₄ units are typical of Mo₂⁴⁺ quadruply bonded complexes, with Mo–Mo and Mo–O bond distances of ~2.10 Å.²⁶ All three compounds crystallized with donor solvent



Figure 1. Thermal ellipsoid plot of I shown at 50% probability. Hydrogens and solvent molecules have been omitted for clarity. Color code: gray, carbon; blue, nitrogen; scarlet, oxygen; yellow, sulfur; green, molybdenum.

molecules, THF (I and III) and DMF (IV), along the Mo–Mo axis.

Further crystallographic details can be found in the Supporting Information.

2.3. Electronic Structure Calculations. In order to aid in the interpretation of the spectral data, we performed electronic structure calculations on the model compounds I', II', and III', where formate is substituted for the TⁱPB ligands. This assists in reducing the computational time and resources and has proven useful for related compounds and interpretation of their spectral data. The calculations employed Density Functional Theory (DFT) and time-dependent DFT (TD-DFT). We also performed calculations on the anions $[I']^-$ and $[II']^-$ to assist in the determination of electron delocalization in the MLCT states.



Figure 2. Thermal ellipsoid plot of III shown at 50% probability. Hydrogens and solvent molecules have been omitted for clarity. Color code: gray, carbon; blue, nitrogen; scarlet, oxygen; yellow, sulfur; green, molybdenum.



Figure 3. Thermal ellipsoid plot of **IV** shown at 50% probability. Hydrogens and solvent molecules have been omitted for clarity. Color code: gray, carbon; blue, nitrogen; scarlet, oxygen; yellow, sulfur; maroon, phosphorus; orange, gold; green, molybdenum.

Figure 4 displays the frontier molecular orbital energy level diagram comparing I' and II' and isosurface contour plots for II', which are very similar to those for I'. For each molecule, the



Figure 4. Energy diagram of compounds \mathbf{I}' and $\mathbf{II}'.$ Orbitals for \mathbf{II}' were generated with Gaussview 5.0.

Article

highest occupied molecular orbital (HOMO) is $M_2\delta$ with a significant degree of mixing with the ligands. The lowest unoccupied molecular orbital (LUMO) in both cases is the symmetrical combination of the ethynylthienyl carboxylate π^* orbital that cannot mix with the $M_2\delta$. For molybdenum, the HOMO-1 and HOMO-2 are filled combinations of the ethynylthienyl carboxylate, which lie higher in energy relative to the $M_2\pi$ and $M_2\sigma$ orbitals. In the case of tungsten, the order of the ligand and metal π orbitals is reversed as the metal-based orbitals, δ and π for tungsten, lie roughly 0.5 eV higher relative to their molybdenum counterparts. For molybdenum, the LUMO+1 is the Mo₂ δ^* orbital, and the LUMO+2 is the ligandbased π^* orbital of the ethynylthienyl carboxylate that mixes with the M₂ δ . For tungsten, M₂ δ^* is the LUMO+2, and it is well separated from the ligand π^* combinations, the LUMO and LUMO+1. The HOMO-LUMO gap for tungsten is also 0.5 eV smaller than that of its molybdenum analogue.

A similar comparison of the frontier orbitals of I' and III' along with the isosurface contour plots of III' are shown in Figure 5. The calculations for both I' and III' were done in C_{2h}



Figure 5. Energy diagram for I' and III'. Orbitals for III' are shown.

symmetry, which emphasizes the maximum ligand $\pi - M_2\delta$ conjugation. For III' we see a greater mixing of the ligand filled π orbitals with the $M_2\delta$ as a result of the higher energy of the ligand filled π orbitals in III' relative to I'. This has the effect of raising the energy of the HOMO, while the LUMO in III' is lowered due to the extended π conjugation. The energy separation between the LUMO and LUMO+2 orbitals, representing the two ligand π^* systems, is also greater for III', 0.33 versus 0.26 eV, as a result of the greater $M_2\delta$ ligand π orbital mixing.

2.4. Electronic Absorption Spectra. The UV-vis-NIR spectra of compounds I and II in THF at room temperature are compared in Figure 6. At high energy, ~300 nm, we see the ethynylthienyl carboxylate ligand-based $\pi - \pi^*$ transition that is hardly influenced by the metal. In contrast, to lower energy we see the intense ¹MLCT (metal-to-ligand charge-transfer) bands arising from the M₂ δ to ethynylthienyl carboxylate π^* , HOMO \rightarrow LUMO, that track with the relative energy of the M₂ δ orbital. We also see evidence of a vibronic progression in the spectrum of II for which the (0,0) transition is the most intense. The vibronic features appear better resolved at low temperature in 2-MeTHF, and the spectrum at 125 K is shown



Figure 6. Electronic absorption spectra for I (red) and II (blue). MLCT λ_{max} = 499 (I) and 712 nm (II).

in the Supporting Information. This feature is common to many tungsten complexes of this type and has been discussed elsewhere.^{27–29} Also, for both compounds we see the $M_2\delta$ -to- $CO_2\pi^*$ transition associated with the TⁱPB ligand: 402 nm for M = W and 335 nm for M = Mo.

An analogous comparison of the absorption spectra of the Mo_2 -containing compounds, I, III, and IV, is given in Figure 7.



Figure 7. Electronic absorption spectra of I, III, and IV in THF at room temperature. MLCT λ_{max} = 499 (I), 517 (III), and 489 nm (IV).

Here we see the similarity of compounds I and IV, which both appear red. This is not too surprising since the $[AuPPh_3]^+$ cation is commonly referred to as a "large proton".³⁰ In contrast, the ¹MLCT of compound III is notably red-shifted to lower energy compared to I as a result of the extended π system. The intraligand $\pi \rightarrow \pi^*$ transition in III (321 nm) also occurs at lower energy than in I (290 nm).

2.5. Solvent Dependence and Steady-State Emission. The absorption and steady-state emission spectra associated with the ¹MLCT states have been recorded in THF, toluene, and DCM at room temperature. These are shown for I and II in Figure 8, where it can be seen that while there is a significant solvent dependence in the absorption spectra, there is very little in the emission from the S₁ state between toluene and DCM, but in the case of THF we do see a significant shift, which is not surprising because THF can behave as a good ligand in axial coordination to the M_2^{5+} center in the MLCT state. This essentially stabilizes the MLCT state in THF. The emission of



Article

Figure 8. Solvent-dependent absorption and emission spectra of I (a) and II (b) in THF (red), DCM (black), and toluene (blue).

II in toluene was measured as a function of concentration and displays no shift in the spectrum, ruling out any self-absorption (SI Figure 1).

At room temperature, the absorption spectra correspond to a Boltzmann distribution of rotamers along the ethynylthienyl carboxylate, and both *trans*-ligands are relatively weakly coupled. However, upon photoexcitation, the promoted electron formally occupies the LUMO, which has a stronger preference for extended planarity. Thus, photoexcitation may well be expected to lead to an initially polar excited state that relaxes to one that is nonpolar. The steady-state emission spectra support this view that the S₁ state has little if any dipole moment. Alternately stated, S₁ is a delocalized excited state with the positive charge on the M_2 center and the electron delocalized over both ethynylthienyl carboxylate ligands. This view is further supported by the TRIR spectra, *vide infra*.

We have not determined the quantum yield for the emission, but it is akin to other dimetal tetracarboxylates for which the efficiency was estimated to be less than 0.1%.³¹

Compound I also shows phosphorescence from its triplet (T₁) state at ~1100 nm which is typical of Mo₂-containing carboxylates and corresponds to a ³MoMo $\delta\delta^*$ state (SI Figure 2).^{18,20–22,32} At 77 K, the emission intensity increases notably on cooling, and a vibronic progression corresponding to the Mo–Mo stretch (~350 cm⁻¹) is visible. Compound II has singlet (S₁) emission with $\lambda_{max} \approx 870$ nm, which is followed by tailing to ~1100 nm. There is a definite weak emission at 1260 nm which we suggest arises from weak phosphorescence from the ³MLCT state at room temperature (SI Figure 2). We have seen this type of behavior previously for tungsten, and it leads to ~3500 cm⁻¹ in terms of the energy separation of the S₁ and

T₁ MLCT states (SI Figure 2, inset).³³ It should be noted that the ³WW $\delta\delta^*$ state is notably higher in energy than the ³MoMo $\delta\delta^*$ state because of the relative energies of the M₂ δ orbitals. This is nicely seen in the absorption spectra of M₂(TⁱPB)₄ compounds, where the λ_{max} occurs at longer wavelength for M₂ = W₂, whereas the phosphorescence is in the inverse order.³²

2.6. Time-Resolved Spectroscopy. Compounds I and II have been examined by femtosecond time-resolved infrared (fs-TRIR) and nanosecond transient absorption (ns-TA) spectroscopies. From these we can estimate the S₁ lifetimes as 3.57 ps for I and 4.4 ps for II (SI Figure 3), while the T₁ lifetimes are 78.8 \pm 5.9 μ s for I and 10–30 ns for II (SI Figure 4). Most pertinent to the matter of charge location in the excited states are the TRIR spectra associated with the ethynyl group. In the ground state, this is a weak IR-active stretch seen at $\nu = 2099$ cm⁻¹ for both compounds. Upon photoexcitation, the vibration has enhanced intensity and is shifted to lower energy due to population of the π^* orbital. For complex II (Figure 9), the IR



Figure 9. fs-TRIR of $\nu(C \equiv C)$ of **II** in THF at room temperature, $\lambda_{ex} = 720$ nm.

band assignable to $\nu(C\equiv C)$ is initially shifted by 52 cm⁻¹ in the ¹MLCT state. This peak decays with time to leave an IR band assignable to the ³MLCT state of II shifted 68 cm⁻¹ to lower energy. The appearance of a single IR-active mode is consistent with a delocalized excited state and contrasts with the spectra of compounds of the form *trans*-PtL₂((C \equiv CAr)₂ that show two IR stretches in their T₁ states: one shifted to lower energy and the other to higher energy relative to the ground state.^{34,35} These have been ascribed to broken or $C_{2\nu}$ symmetry in the excited state. The further shift of the $\nu(C\equiv C)$ to lower energy in II with the change from S₁ to T₁ suggests that there is slightly greater $C\equiv C \pi^*$ character in the triplet state. The molybdenum complex I (SI Figure 5) shows similar behavior, with $\Delta\nu(C\equiv C) = -66$ cm⁻¹ in the S₁ state. Following decay to the T₁ state, however, the ethynyl stretch becomes IR silent, consistent with a ³MOMo $\delta\delta^*$ state.

Since the IR bands associated with ligands in MLCT states are very similar to those of the reduced form of the ligand, we have calculated the values of $\nu(C\equiv C)$ for $[I']^-$ and $[II']^-$. The calculations pertain to a delocalized anion with the negative charge distributed principally upon the two *trans*-ethynylthienyl carboxylates. A comparison of the experimental values for compounds I and II in their ground and photoexcited states is given in Table 1, along with those calculated for the ground states of I' and II' and their respective anions $[I']^-$ and $[II']^-$. The calculated shifts for the anions (~55 cm⁻¹) compare

Table 1. Experimental and Computational ν (CC) Frequencies (in cm⁻¹) and Corresponding Shifts for I and II

	I	II		I'	II'
S ₀	2099	2099	neutral	2130	2128
$S_{1}(T_{1})$	2033	2047 (2031)	anion	2073	2073
$\Delta \nu(CC)$	-66	-52 (-68)	$\Delta \nu(\text{CC})$	-57	-55

favorably with those observed in the $\rm S_1$ states, supporting the view of a delocalized excited state. 36

Time-resolved spectroscopic studies of III and IV will be reported later in conjunction with other compounds of similar nature. Evaluating the shift of $\nu(C\equiv C)$ in comparison to the parent complexes should provide a measure of the electronic interaction of their appended groups in the excited states.

3. CONCLUDING REMARKS

The spectroscopic data for compounds I and II indicate that the S₁ states and the T₁ state for II are fully delocalized. In terms of excited-state mixed valency,³⁷ this is Class III on the Robin and Day scheme.³⁸ There are, to our knowledge, no detailed TRIR studies of transition metal alkynyl complexes in their S₁ states, despite the fact that some fluoresce and have picosecond lifetimes. The T₁ states that have been examined have broken symmetry where the charge is centered upon one ligand. The delocalized ³MLCT states of II thus appear very different from all studies of *trans*-bisalkynyl Pt(II) complexes to date, which show lowest energy electronic transitions that are at least in part MLCT in character.^{39–47}

In terms of the synthetic strategy for the synthesis of higher order assemblies, the preparation of III and IV from I is very encouraging. Although the C–C coupling reactions employed are "standard" in organic synthesis, and have been successfully employed by others in the synthesis of later transition metalcontaining organic conjugated oligomers and polymers,^{48–54} the application to the kinetically labile M_2^{4+} center is notable, as many reactions may lead to undesired products due to metathetic reactions of the carboxylates. For example, the use of organomagnesium or organozinc reagents as in Kumada or Negishi coupling reactions is not possible due to carboxylate scrambling. Further work involving C–C coupling reactions of this type involving vinyl groups is currently underway.

ASSOCIATED CONTENT

S Supporting Information

Materials and methods, synthesis and characterization of I–IV, NIR emission spectra, kinetic traces of I and II from the fs-TRIR and of I from the ns-TA, TRIR spectrum of I, and crystallographic information (CIF). This material is available free of charge via the Internet at http://pubs.acs.org. CCDC records 919547–919549 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

AUTHOR INFORMATION

Corresponding Author

chisholm@chemistry.ohio-state.edu

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors acknowledge Dr. Judith Gallucci for helpful discussion and Prof. Claudia Turro for use of instrumentation. This work was supported by NSF grant CHE-0957191 and the OSU Institute for Materials Research. Computational resources were provided by the Ohio Supercomputing Center. S.E.B.-X. acknowledges support from the National Defense Science and Engineering Graduate (NDSEG) Fellowship. During this work we discovered that compound I has been prepared and structurally characterized independently by Thomas P. Robinson in the group of Prof. Paul R. Raithby in the University of Bath, UK.

REFERENCES

(1) Mishra, A.; Bäuerle, P. Angew. Chem., Int. Ed. 2012, 51, 2020–2067.

(2) Osaka, I.; Shimawaki, M.; Mori, H.; Doi, I.; Miyazaki, E.; Koganezawa, T.; Takimiya, K. J. Am. Chem. Soc. **2012**, 134, 3498–3507.

(3) Silvestri, F.; Marrocchi, A.; Seri, M.; Kim, C.; Marks, T. J.; Facchetti, A.; Taticchi, A. J. Am. Chem. Soc. 2010, 132, 6108–6123.

(4) McCulloch, I.; Ashraf, R. S.; Biniek, L.; Bronstein, H.; Combe, C.; Donaghey, J. E.; James, D. I.; Nielsen, C. B.; Schroeder, B. C.; Zhang, W. Acc. Chem. Res. **2012**, 45, 714–722.

(5) Bronstein, H.; Chen, Z.; Ashraf, R. S.; Zhang, W.; Du, J.; Durrant, J. R.; Shakya Tuladhar, P.; Song, K.; Watkins, S. E.; Geerts, Y.; Wienk, M. M.; Janssen, R. A. J.; Anthopoulos, T.; Sirringhaus, H.; Heeney, M.; McCulloch, I. J. Am. Chem. Soc. **2011**, *133*, 3272–3275.

(6) Tsao, H. N.; Cho, D.; Andreasen, J. W.; Rouhanipour, A.; Breiby, D. W.; Pisula, W.; Müllen, K. *Adv. Mater.* **2009**, *21*, 209–212.

(7) Sonogashira, K.; Takahashi, S.; Hagihara, N. Macromolecules 1977, 10, 879-880.

(8) Wong, W.-Y.; Wang, X.-Z.; He, Z.; Chan, K.-K.; Djurišić, A. B.; Cheung, K.-Y.; Yip, C.-T.; Ng, A. M.-C.; Xi, Y. Y.; Mak, C. S. K.; Chan, W.-K. J. Am. Chem. Soc. **200**7, 129, 14372–14380.

(9) Zhan, H.; Wong, W.-Y.; Ng, A.; Djurišić, A. B.; Chan, W.-K. J. Organomet. Chem. **2011**, 696, 4112–4120.

(10) Wong, W.-Y.; Ho, C.-L. Acc. Chem. Res. 2010, 43, 1246-1256.

(11) Wong, W.-Y. Macromol. Chem. Phys. 2008, 209, 14-24.

(12) Ho, C.-L.; Wong, W.-Y. Coord. Chem. Rev. 2011, 255, 2469–2502.

(13) Wong, W.-Y.; Wang, X.-Z.; He, Z.; Djurišić, A. B.; Yip, C.-T.; Cheung, K.-Y.; Wang, H.; Mak, C. S. K.; Chan, W.-K. Nat. Mater. 2007, 6, 521–527.

(14) Wong, W.-Y.; Ho, C.-L. Coord. Chem. Rev. 2006, 250, 2627–2690.

(15) Li, P.; Ahrens, B.; Choi, K.-H.; Khan, M. S.; Raithby, P. R.; Wilson, P. J.; Wong, W.-Y. *CrystEngComm* **2002**, *4*, 405–412.

(16) Wong, W.-Y.; Choi, K.-H.; Lu, G.-L.; Lin, Z. Organometallics 2002, 21, 4475-4481.

(17) Holliday, B. J.; Swager, T. M. Chem. Commun. 2005, 23-36.

(18) Burdzinski, G. T.; Chisholm, M. H.; Chou, P.-T.; Chou, Y.-H.; Feil, F.; Gallucci, J. C.; Ghosh, Y.; Gustafson, T. L.; Ho, M.-L.; Liu, Y.;

Ramnauth, R.; Turro, C. Proc. Natl. Acad. Sci. U.S.A. 2008, 105, 15247-15252.

(19) Chisholm, M. H.; D'Acchioli, J. S.; Hadad, C. M. J. Cluster Sci. 2006, 18, 27–49.

(20) Alberding, B. G.; Chisholm, M. H.; Ghosh, Y.; Gustafson, T. L.; Liu, Y.; Turro, C. Inorg. Chem. **2009**, 48, 8536–8543.

(21) Brown-Xu, S. E.; Chisholm, M. H.; Gallucci, J. C.; Ghosh, Y.; Gustafson, T. L.; Reed, C. R. *Dalton Trans.* **2012**, *41*, 2257–2263.

(22) Brown-Xu, S. E.; Chisholm, M. H.; Durr, C. B.; Gustafson, T. L.;

- Naseri, V.; Spilker, T. F. J. Am. Chem. Soc. 2012, 134, 20820–20826. (23) Chisholm, M. H.; Patmore, N. J. Acc. Chem. Res. 2007, 40, 19– 27.
- (24) Chisholm, M. H.; Epstein, A. J.; Gallucci, J. C.; Feil, F.; Pirkle, W. Angew. Chem., Int. Ed. 2005, 44, 6537–6540.

(26) Cotton, F. A.; Murillo, C. A.; Walton, R. A. *Multiple Bonds Between Metal Atoms*, 3rd ed.; Springer Science and Business Media, Inc.: New York, 2005.

(27) Bursten, B. E.; Chisholm, M. H.; Clark, R. J. H.; Firth, S.; Hadad, C. M.; MacIntosh, A. M.; Wilson, P. J.; Woodward, P. M.; Zaleski, J. M. J. Am. Chem. Soc. **2002**, 124, 3050–3063.

(28) Alberding, B. G.; Chisholm, M. H.; Lear, B. J.; Naseri, V.; Reed, C. R. Dalton Trans. 2011, 40, 10658–10663.

(29) Chisholm, M. H.; Lear, B. J. Chem. Soc. Rev. 2011, 40, 5254–5265.

(30) Raubenheimer, H. G.; Schmidbaur, H. Organometallics **2012**, 31, 2507–2522.

(31) Chisholm, M. H.; Gustafson, T. L.; Turro, C. Acc. Chem. Res. 2013, 46, 529–538.

(32) Alberding, B. G.; Chisholm, M. H.; Chou, Y.-H.; Gallucci, J. C.; Ghosh, Y.; Gustafson, T. L.; Patmore, N. J.; Reed, C. R.; Turro, C. *Inorg. Chem.* **2009**, *48*, 4394–4399.

(33) Chisholm, M. H.; Chou, P.-T.; Chou, Y.-H.; Ghosh, Y.; Gustafson, T. L.; Ho, M.-L. Inorg. Chem. 2008, 47, 3415–3425.

(34) Emmert, L. A.; Choi, W.; Marshall, J. A.; Yang, J.; Meyer, L. A.; Brozik, J. A. J. Phys. Chem. A 2003, 107, 11340–11346.

(35) Batista, E. R.; Martin, R. L. J. Phys. Chem. A 2005, 109, 9856-9859.

(36) Alberding, B. G.; Chisholm, M. H.; Gallucci, J. C.; Ghosh, Y.; Gustafson, T. L. Proc. Natl. Acad. Sci. U.S.A. 2011, 108, 8152–8156.

(37) Plummer, E. A.; Zink, J. I. Inorg. Chem. 2006, 45, 6556-6558.
(38) Robin, M. B.; Day, P. In Advances in Inorganic Chemistry and Radiochemistry; Emeléus, H. J., Sharpe, A. G., Eds.; Academic Press: San Diego, 1968; Vol. 10, pp 247-422.

(39) Wilson, J. S.; Chawdhury, N.; Al-Mandhary, M. R. A.; Younus, M.; Khan, M. S.; Raithby, P. R.; Köhler, A.; Friend, R. H. J. Am. Chem. Soc. 2001, 123, 9412–9417.

(40) Silverman, E. E.; Cardolaccia, T.; Zhao, X.; Kim, K.-Y.; Haskins-Glusac, K.; Schanze, K. S. *Coord. Chem. Rev.* 2005, 249, 1491–1500.
(41) Prusakova, V.; McCusker, C. E.; Castellano, F. N. *Inorg. Chem.* 2012, 51, 8589–8598.

(42) Beljonne, D.; Wittmann, H. F.; Köhler, A.; Graham, S.; Younus, M.; Lewis, J.; Raithby, P. R.; Khan, M. S.; Friend, R. H.; Brédas, J. L. *J. Chem. Phys.* **1996**, *105*, 3868–3877.

(43) Muro, M. L.; Diring, S.; Wang, X.; Ziessel, R.; Castellano, F. N. Inorg. Chem. 2009, 48, 11533-11542.

(44) Whittle, C. E.; Weinstein, J. A.; George, M. W.; Schanze, K. S. Inorg. Chem. 2001, 40, 4053–4062.

(45) Archer, S.; Weinstein, J. A. Coord. Chem. Rev. 2012, 256, 2530–2561.

(46) Glik, E. A.; Kinayyigit, S.; Ronayne, K. L.; Towrie, M.; Sazanovich, I. V.; Weinstein, J. A.; Castellano, F. N. *Inorg. Chem.* **2008**, 47, 6974–6983.

(47) Ramakrishna, G.; Goodson, T.; Rogers-Haley, J. E.; Cooper, T.

M.; McLean, D. G.; Urbas, A. J. Phys. Chem. C 2009, 113, 1060–1066.

(48) Chen, W.-Z.; Fanwick, P. E.; Ren, T. Organometallics 2007, 26, 4115–4117.

(49) Chen, W.-Z.; Ren, T. Inorg. Chem. 2006, 45, 9175-9177.

(50) Xu, G.-L.; Ren, T. Organometallics 2005, 24, 2564-2566.

(51) Zhang, L.; Huang, Z.; Chen, W.-Z.; Negishi, E.; Fanwick, P. E.; Updegraff, J. B.; Protasiewicz, J. D.; Ren, T. *Organometallics* **2007**, *26*, 6526–6528.

(52) Iii, S. W. T.; Yagi, S.; Swager, T. M. J. Mater. Chem. 2005, 15, 2829–2835.

(53) Ren, T. Organometallics 2005, 24, 4854-4870.

(54) Ren, T. Chem. Rev. 2008, 108, 4185-4207.