Conversion of Benzaldehyde Imines into Isocyanides at a Low-Valent Molybdenum Center. Preparation and Reactivities of Isocyanide-**Dinitrogen Complexes** *trans***-[Mo(RNC)(N₂)(Ph₂PCH₂CH₂PPh₂)₂] (R = Aryl, Alkyl)1**

Hidetake Seino,^{2a} Chirima Arita,^{2a} Daigo Nonokawa,^{2a} Goh Nakamura,^{2a,b} Yuji Harada,^{2a,b} Yasushi Mizobe,^{*,2a,c} and Masanobu Hidai^{*,2b}

Institute of Industrial Science, The University of Tokyo, Roppongi, Minato-ku, Tokyo 106-8558, Japan, Department of Chemistry and Biotechnology, Graduate School of Engineering, The University of Tokyo, Hongo, Tokyo 113-8656, Japan, and Institute for Molecular Science, Myodaiji, Okazaki 444-8585, Japan

Received June 14, 1999

The molybdenum dinitrogen complex *trans*- $[Mo(N_2)_2(dppe)_2]$ (1; dppe $= Ph_2PCH_2CH_2PPh_2)$) reacts with benzaldehyde imines PhCH=NR ($R = Ph$, C_6H_4Me -*p*, C_6H_4OMe -*p*, C_6H_4F -*p*, C_6H_4 -NMe₂-p, Buⁿ, Prⁱ, CH₂Ph) in benzene at reflux to give a series of isocyanide-dinitrogen
complexes *trans*-[Mo(RNC)(N₂)(dppe)₂] (7) with the concurrent formation of benzene. The complexes *trans*-[Mo(RNC)(N2)(dppe)2] (**7**) with the concurrent formation of benzene. The structure of **7a** ($R = Ph$) has been determined by X-ray crystallography. Treatment of **7a** and **7f** ($R = Bu^n$) with CO (1 atm) or p -MeOC₆H₄CN results in the replacement of the coordinated N_2 by these ligands, affording *trans*-[Mo(RNC)(L)(dppe)₂] (L = CO (9), *p*-MeOC6H4CN (**10**)). The complex containing two different isocyanides *trans*-[Mo(PhNC)- (Bu^tNC)(dppe)₂] was obtained analogously from 7a and Bu^tNC. On the other hand, when benzene or toluene solutions of **7a** and **7e** ($R = C_6H_4NMe_2$ *-p*) were first heated under Ar up to 80–90 °C for a short period to dissociate the coordinated N₂ and then treated with H₂ at room temperature, dihydrogen complexes *trans*-[Mo(RNC)(*η*2-H2)(dppe)2] (**13**) were produced. The X-ray analyses have revealed the detailed structures for **9a** $(R = Ph)$, **10a** $(R = Ph)$, and **13e** ($R = C_6H_4NMe_2$ -*p*) along with the bis(isocyanide) complex *trans*-[Mo(PhNC)₂(dppe)₂].

Introduction

Since the isolation of the first Mo dinitrogen complex *trans*-[Mo(N₂)₂(dppe)₂] (**1**; dppe = Ph₂PCH₂CH₂PPh₂) in this laboratory,³ intensive efforts have been devoted to investigate reactivities of the N2 ligand in **1** and related Mo and W complexes such as $trans\text{-}[W(N_2)_2(\text{dppe})_2]$ (2) and *cis*- $[M(N_2)_2(PMe_2Ph)_4]$ (M = Mo, W), aiming at development of a new type of N_2 -fixing systems.⁴ Now it has become apparent that these Mo and W dinitrogen complexes are distinctive from other numerous N_2 complexes hitherto reported in the reactivity of the coordinated N_2 which gives rise to the formation of nitrogenous ligands and compounds under ambient conditions.4,5

One of the other intriguing reactions promoted by **1** is the decarbonylation of various oxygen-containing organic compounds including aldehydes, formamides, carboxylic acid esters, and alcohols,⁶ where 1 serves as a versatile precursor to generate a highly reactive, coordinatively unsaturated species by dissociating the N_2 ligands. Reactions of this type are exemplified by the formation of *trans*-[Mo(CO)(DMF)(dppe)₂] (3; DMF = *N*,*N*-dimethylformamide) upon treatment of **1** with an excess of DMF in benzene at reflux.⁷ Interestingly, the DMF ligand in **3** is so labile that **3** is readily converted to *trans*- $[Mo(CO)(N₂)(dppe)₂]$ (4) and $[Mo(CO)(dppe)₂]$ (**5**).6a From these carbonyl complexes as well as their W analogues derived similarly from **2**, a number of carbonyl complexes of the types $[M(CO)(L)(dppe)_2]$ (M = Mo, L = Lewis bases, nitrile, olefin, $6a, 8H_2, 9$ hydrosi-
langs:¹⁰ M = Mo, W: J = vinvlidene¹¹) and [M(CO)H. lanes;¹⁰ M = Mo, W: L = vinylidene¹¹) and [M(CO)H-

⁽¹⁾ Preparation and Properties of Molybdenum and Tungsten Dinitrogen Complexes. 64. Part 63: see ref 15. (2) (a) Institute of Industrial Science. (b) Department of Chemistry

and Biotechnology. (c) Institute for Molecular Science. (3) (a) Hidai, M.; Tominari, K.; Uchida, Y.; Misono, A. *J. Chem. Soc., Chem. Commun.* **1969**, 1392. (b) Hidai, M.; Tominari, K.; Uchida, Y. *J. Am. Chem. Soc.* **1972**, *94*, 110.

^{(4) (}a) Hidai, M.; Mizobe, Y. *Chem. Rev.* **1995**, *95*, 1115. (b) Hidai, M.; Ishii, Y. *Bull. Chem. Soc. Jpn.* **1996**, *69*, 819.

^{(5) (}a) Richards, R. L. *Coord. Chem. Rev.* **1996**, *154*, 83. (b) Bazhenova, T. A.; Shilov, A. E. *Coord. Chem. Rev.* **1995**, *144*, 69. (c) Gambarotta, S. *J. Organomet. Chem.* **1995**, *500*, 117. (d) Leigh, G. J. *Acc. Chem. Res.* **1992**, *25*, 177.

^{(6) (}a) Sato, M.; Tatsumi, T.; Kodama, T.; Hidai, M.; Uchida, T.; Uchida, Y. *J. Am. Chem. Soc.* **1978**, *100*, 4447. (b) Tatsumi, T.; Tominaga, H.; Hidai, M.; Uchida, Y. *J. Organomet. Chem.* **1981**, *215*, 67. (c) Tatsumi, T.; Tominaga, H.; Hidai, M.; Uchida, Y. *J. Organomet. Chem.* **1981**, *218*, 177.

⁽⁷⁾ Mizobe, Y.; Ishida, T.; Egawa, Y.; Ochi, K.; Tanase, T.; Hidai, M. *J. Coord. Chem.* **1991**, *23*, 57.

⁽⁸⁾ Tatsumi, T.; Tominaga, H.; Hidai, M.; Uchida, Y. *J. Organomet. Chem.* **1980**, *199*, 63.

^{(9) (}a) Kubas, G. J.; Ryan, R. R.; Wrobleski, D. A. *J. Am. Chem. Soc.* **1986**, *108*, 1339. (b) Kubas, G. J.; Burns, C. J.; Eckert, J.; Johnson, S. W.; Larson, A. C.; Vergamini, P. J.; Unkefer, C. J.; Khalsa, G. R. K.; Jackson, S. A.; Eisenstein, O. *J. Am. Chem. Soc.* **1993**, *115*, 569.

 $(X)(dppe)_2$] (M = Mo, W: X = alkynyl;¹¹ M = W, X = carbonate, carbamate, 12 hydride¹³) have been prepared.

The mechanism for the formation of **3** from **1** involves the cleavage of the aldehydic C-H bond in DMF at the coordinatively unsaturated Mo center to give a hydridocarbamoyl species followed by the elimination of dimethylamine (Scheme 1). This has been verified by the isolation of $[WH(\eta^2\text{-}\text{CONMe}_2)(\text{dppe})_2]$ (6) from the reaction of **2** with DMF under controlled conditions. As expected, when heated in benzene in the presence of DMF, 6 is converted to *trans*-[W(CO)(DMF)(dppe)₂] with concurrent formation of HNMe₂.¹³

Now the study has been extended to the reactions of **1** with benzaldehyde imines $PhCH=NR$ ($R = \text{aryl}$, alkyl) instead of aldehydic compounds, and we have found that unprecedented conversion of the imines into isocyanides takes place at the Mo center to give a series of isocyanide-dinitrogen complexes *trans*-[Mo(RNC)(N₂)(dppe)₂] (**7**). In this paper, details of the syntheses and characterization of **7** are described, together with the reactivities of **7** toward a series of neutral ligands including CO, nitriles, isocyanides, and H_2 . A part of this work has been reported recently as a communication.¹⁴

Results and Discussion

Preparation of Isocyanide-**Dinitrogen Complexes 7.** When reacted with a range of PhCH=NR in benzene at reflux for 2 h, the dinitrogen complex **1** afforded **7** in moderate yields (eq 1). Employment of the

imines with $R = C_6H_4X$ - p (X = H, Me, OMe, F, NMe₂), Buⁿ, Prⁱ, and CH₂Ph gave the well-defined complexes

7; however, use of the imines having the substituents C_6H_4X -*p* ($X = CF_3$, COOEt) and Bu^t for R did not afford the corresponding compounds. Due to the thermal instability, **7h** ($R = CH_2Ph$) once produced proved to decompose rapidly under the conditions employed. Thus, **7h** was obtained in satisfactory yield by refluxing the reaction mixture for only 5 min. The conversion of imines at the Mo center into the corresponding isocyanides appears to proceed via the scission of both the ^C-Ph and C-H bonds on the benzylidene carbon. This was confirmed by the concurrent formation of benzene with **7a** and **7b** in the reaction of **1** with PhCH=NPh or PhCH=NC₆H₄Me- p (0.71 and 0.73 mol of C₆H₆/mol of **1**, respectively).

Although the reactions of several aliphatic aldehyde imines R'CH=NR ($R =$ alkyl, aryl; $R' =$ alkyl) with **1** were attempted, complexes containing the RNC ligand were not produced. Interestingly, from the reaction mixtures of **1** with the imines that remained intact under these conditions, a Mo(0) complex with a novel tetradentate phosphine $[Mo{6}C_6H_4(PPhCH_2CH_2PPh_2)_2]$ (dppe)] has been isolated in low yield. Characterization and the improved method for preparation of this complex from **1** by thermolysis have already been reported separately.15 It is to be noted that the reactions of **1** with various isocyanides give bis(isocyanide) complexes *trans*- $[Mo(RNC)_2(dppe)_2]$ (R = alkyl, aryl) exclusively, and the formation of the monosubstituted complexes **7** was not observed.16

To our knowledge, little is known about the direct transformation of imines into isocyanides. In contrast, insertion of isocyanides into M-R bonds leading to *^η*1 or η^2 -iminoacyls is a well-precedented process.¹⁷ For certain metal iminoacyl species, formation of imines by succesive alkyl migration onto the iminoacyl carbon is also known.¹⁸ The novel conversion of benzylideneanilines into isocyanides reported here might proceed by following these processes in the reverse direction (Scheme 2). Thus, oxidative addition of the benzylidene ^C-H bond occurs initially at the coordinatively unsaturated Mo(0) center generated by thermolysis of **1**, giving a hydrido-iminoacyl intermediate $[MoH(C)]$ NR)Ph}(dppe)₂]. Deinsertion of RNC from the iminoacyl ligand and the following reductive elimination of ben-

31, 4481.

(13) Ishida, T.; Mizobe, Y.; Tanase, T.; Hidai, M. *J. Organomet. Chem.* **1991**, *409*, 355.

(14) Nakamura, G.; Harada, Y.; Arita, C.; Seino, H.; Mizobe, Y.; Hidai, M. *Organometallics* **1998**, *17*, 1010.

(15) Arita, C.; Seino, H.; Mizobe, Y.; Hidai, M. *Chem. Lett.* **1999**, 611.

(16) (a) Chatt, J.; Elson, C. M.; Pombeiro, A. J. L.; Richards, R. L.; Royston, G. H. D. *J. Chem. Soc., Dalton Trans.* **1978**, 165. (b) Pombeiro, A. J. L.; Pickett, C. J.; Richards, R. L.; Sangokoya, S. A. *J. Organomet. Chem.* **1980**, *202*, C15. (c) Pombeiro, A. J. L.; Richards, R. L. *Coord. Chem. Rev.* **1990**, *104*, 13.

(17) Durfee, L. D.; Rothwell, I. P. *Chem. Rev.* **1988**, *88*, 1059. (18) (a) Durfee, L. D.; Fanwick, P. E.; Rothwell, I. P.; Folting, K.; Huffman, J. C. *J. Am. Chem. Soc.* **1987**, *109*, 4720. (b) Koschmieder, S. U.; Hussain-Bates, B.; Hursthouse, M. B.; Wilkinson, G. *J. Chem. Soc., Dalton Trans.* **1991**, 2785. (c) Laı¨, R.; Desbois, O.; Zamkotsian, F.; Faure, R.; Feneau-Dupont, J.; Declercq, J.-P. *Organometallics* **1995**,
*14, 2145. (d) Cámpora, J.; Buchwald, S. L.; Gutiérrez-Puebla, E.;
Monge, A. <i>Organometallics* **1995**, *14, 2039.*

^{(10) (}a) Luo, X.-L.; Kubas, G. J.; Bryan, J. C.; Burns, C. J.; Unkefer, C. J. *J. Am. Chem. Soc.* **1994**, *116*, 10312. (b) Luo, X.-L.; Kubas, G. J.; Burns, C. J.; Bryan, J. C.; Unkefer, C. J. *J. Am. Chem. Soc.* **1995**, *117*, 1159.

⁽¹¹⁾ Nakamura, G.; Harada, Y.; Mizobe, Y.; Hidai, M. *Bull. Chem. Soc. Jpn.* **1996**, *69*, 3305. (12) Ishida, T.; Hayashi, T.; Mizobe, Y.; Hidai, M. *Inorg. Chem.* **1992**,

Scheme 2

zene affords [Mo(CNR)(dppe)2] (**8**), which rapidly binds N2 to yield **7**. Many attempts to isolate or detect **8** were carried out, but failed.

Since the hydrido-carbamoyl intermediate **⁶** was able to be isolated only for W in the related decarbonylation reaction of DMF (Scheme 1), W complex **2** was allowed to react with benzaldehyde imines under the analogous conditions. However, neither the isocyanide-dinitrogen complexes nor the hydrido-iminoacyl complexes were obtained.

Since a variety of benzaldehyde imines are readily available, the reactions reported here are attractive as a route to isocyanides. However, the reaction proceeded only stoichiometrically, and the liberation of free isocyanides was not observed in the reactions with an excess amount of benzaldehyde imines.

Characterization of 7. The structure of **7a** has been determined in detail by single-crystal X-ray analysis. The ORTEP drawing is shown in Figure 1, while the important bond lengths and angles are listed in Table 1. Complex **7a** has a slightly distorted octahedral structure with mutually trans PhNC and N_2 ligands bound to the Mo center in an end-on fashion. In the PhNC ligand, the $Mo-C(1)-N(1)$ linkage is almost linear (176.7(3)°), while the $C(1)-N(1)-C(2)$ bond is slightly bent, with an angle of $167.4(4)^\circ$. The $C(1)-N(1)$ bond length at 1.179(4) Å is unexceptional for the isocyanide ligand of this coordination mode. With respect to the N_2 ligand, the Mo-N(2)-N(3) linkage is nearly linear (177.1(3)°), and the N(2)-N(3) distance at 1.102(3) Å is comparable to those in free N_2 (1.0976(2) Å)¹⁹ and **4** (1.09(2) Å).^{6a}

Spectroscopic and analytical data for **7a** are in good agrement with the solid-state structure. Thus, the IR spectrum shows two intense bands at 2049 and 1910 cm⁻¹ assignable to *ν*(NN) of the coordinated N₂ and *ν*-(NC) of the PhNC ligand, respectively, while the 31P- {1H} NMR spectrum exhibits one singlet at *δ* 68.9 due to the two dppe ligands constituting the basal plane of the octahedral structure. The other new complexes **7b**-**7h** show the analogous spectral features, indicating that the structures are essentially identical.²⁰

Now a series of the N2 complexes of the type *trans*- $[Mo(L)(N_2)(dppe)_2]$ are available, where the ligand L

Figure 1. ORTEP diagram of **7a**. Hydrogen atoms are omitted for clarity.

trans to the N_2 is CO,^{6a} N_2 ,³ nitriles,²¹ or isocyanides coordinated to the Mo(0) center in an end-on manner. Comparison of the IR data for these complexes provides the order of the $\nu(NN)$ values decreasing as $L = CO$ (2128 cm^{-1}) > PhNC (2049 cm^{-1}) > N₂ (2020_{sym}) 1970_{asym} cm⁻¹) > p -MeOC₆H₄CN (1946 cm⁻¹). Force constants calculated from these *ν*(NN) values are 18.7, 17.3, 16.4, and 15.6 mdyn/Å, respectively.22 This indicates that the π -acidity of L decreases according to this order, which results in the increase in the net *π*-donating nature of the ${Mo(L)(dppe)_2}$ moiety toward the antibonding orbitals of the N_2 ligand.

^{(19) (}a) Wilkinson, P. G.; Houk, N. B. *J. Chem. Phys.* **1956**, *24*, 528. (b) Gordon, A. J.; Ford, R. A. *The Chemist's Companion*; John Wiley & Sons: New York, 1972; p 107.

⁽²⁰⁾ The X-ray analyses have also been undertaken for **7c**, **7f**, and **7g**. Although the refinements were not completed to the satisfactory level, the preliminary results have clearly demonstrated that the structures of these three complexes are essentially analogous to that of **7a**. See Experimental Section.

⁽²¹⁾ Tatsumi, T.; Hidai, M.; Uchida, Y. *Inorg. Chem.* **1975**, *14*, 2530. (22) (a) Lazarowych, N. J.; Morris, R. H.; Ressner, J. M. *Inorg. Chem.* **1986**, *25*, 3926. (b) Morris, R. H.; Earl, K. A.; Luck, R. L.; Lazarowych, N. J.; Sella, A. *Inorg. Chem.* **1987**, *26*, 2674, and references therein.

The $\nu(NN)$ value (2049 cm⁻¹) is considerably lower than that of free N_2 (2331 cm⁻¹) but still higher than the parent N_2 complex **1** (2020 and 1970 cm⁻¹).³ Thus, the N_2 ligand in **7a** is so labile as to be converted readily into the related Mo(0) isocyanide complexes *trans*-[Mo- $(PhNC)(L)(dppe)_2$] (L = CO (**9a**), p -MeOC₆H₄CN (**10a**), and But NC (**11**)).

Preparation and Characterization of *trans***-[Mo- (RNC)(CO)(dppe)2] (9),** *trans***-[Mo(RNC)(***p***-MeOC6H4- CN)(dppe)2] (10), and** *trans***-[Mo(PhNC)(But NC)- (dppe)2] (11).** When **7a** suspended in THF was treated with CO at room temperature, an isocyanide-carbonyl complex **9a** was readily obtained in excellent yield. Although the reaction of **7f** with CO appears to proceed more slowly, **9f** ($R = Bu^n$) was able to be isolated in satisfactory yield by treatment of a benzene solution of **7f** with CO at 50 °C. Reactions of **7a** or **7f** with ca. 2 equiv of p -MeOC₆H₄CN in benzene at room temperature also resulted in the replacement of the N_2 ligand by the nitrile to give isocyanide-nitrile complexes **¹⁰** in high yields. Detailed structures of **9a** and **10a** have been determined by X-ray analyses as described below. On the other hand, the reaction of **7a** with an equimolar amount of Bu^tNC at room temperature gave 11, which was characterized by the spectroscopic and microanalytical data. This finding indicates that **7** can serve as the versatile precursor for preparing a wide range of Mo complexes containing two different isocyanides *trans*-[Mo(RNC)(R'NC)(dppe)₂]. The reactions forming **⁹**-**¹¹** are summarized in eq 2.

Table 2 lists the selected bond distances and angles in **9a** and **10a** determined by X-ray analyses. Since the attempt to prepare high-quality single crystals was unsuccessful for **11**, a single crystal of *trans*-[Mo- $(PhNC)_2$ (dppe)₂] $(12)^{16a}$ was prepared and subjected to the X-ray diffraction study, whose results are also shown in Table 2. As depicted in Figures 2-4, structures of **9a**, **10a**, and **12**, respectively, are well comparable to that of **7a**, viz., the CO, p -MeOC₆H₄CN, and PhNC ligands occupy the trans position of the PhNC ligand in the octahedral coordination sphere. The $Mo-C(8)$ O, $Mo-N(2)-C(8)$, and $Mo-C(1)-N$ linkages in each complexes are linear with angles of 177.9(4)°, 179.3(3)°, and 176.5(5)°, while the C(8)-O, N(2)-C(8), and C(1)-N bond distances in these moieties at 1.086(4), 1.153(4), and 1.171(6) Å, respectively, are unexceptional for these molecules of this coordination mode.

For the PhNC ligands in *trans*-[Mo(PhNC)(L)(dppe)₂], important bonding parameters associated with **9a**, **10a**, and **12** are summarized in Table 3, together with those of **7a** for comparison. With respect to the bond lengths, Table 3 clearly shows that the $C(1)-N(1)$ bonds are elongated, following the order **9a** < **¹²** < **7a** < **10a**, and

Table 2. Selected Bond Lengths (Å) and Angles (deg) in 9a, 10a, and 12

accordingly the $Mo-C(1)$ and $N(1)-C(2)$ bond distances decrease as $9a > 12 > 7a > 10a$. This finding is interpreted in terms of the net *π*-donating ability of the ${Mo(L)(dppe)_2}$ moieties toward the PhNC ligand, increasing in the order L = CO < PhNC < N_2 < p -MeOC₆H₄CN, which is in good agreement with the results obtained for a series of dinitrogen complexes $trans$ -[Mo(N₂)(L)(dppe)₂] (vide supra). In contrast to relatively small differences in the bond angles of the $Mo-C(1)-N(1)$ arrays $(179.2(4)-168.9(3)°)$, those of the $C(1)-N(1)-C(2)$ linkages change significantly, from 177.6(5)° in **9a** to 149.8(4)° in **10a**. In general, bent ^C-N-C linkages are found in the aliphatic isocyanides bound to the zerovalent metal center having no *π*-accepting ligands, where the electron density remains localized on the N atom in the NC group and the ^C-N-C linkage tends to be bent due to the pairing effects of electrons,²³ e.g., the average $C-N-C$ angle of 130° for the two equatorial isocyanide ligands in the trigonal bipyramidal [Ru(CNBu^t)₄(PPh₃)] and that of

Figure 2. ORTEP diagram of **9a**. Hydrogen atoms are omitted for clarity.

Figure 3. ORTEP diagram of **10a**. Hydrogen atoms are omitted for clarity.

135° for the two bent isocyanide ligands in [Fe(CN-Bu^t)₅].²⁴ On the other hand, significant C—N—C bending
bardly occurs for the aryl isocyanide ligands because of hardly occurs for the aryl isocyanide ligands because of their ability to delocalize the charge from the isocyanide

Figure 4. ORTEP diagram of **12**. Hydrogen atoms are omitted for clarity.

Table 3. Selected Bonding Parameters in *trans***[Mo(PhNC)(L)(dppe)**₂]

	9а	12	7а	10a					
L	CO	PhNC	N,	p -MeOC ₆ H ₄ CN					
(a) Bond Length (Å)									
$Mo-C(1)$	2.072(4)	2.031(6)	1.992(3)	1.964(3)					
$C(1) - N(1)$		$1.099(4)$ $1.171(6)$ $1.179(4)$		1.217(4)					
$N(1) - C(2)$	1.407(5)	1.399(7)	1.388(4)	1.386(4)					
		(b) Bond Angle (deg)							
$Mo-C(1)-N(1)$	179.2(4)	176.5(5)	176.7(3)	168.9(3)					
$C(1)-N(1)-C(2)$	177.6(5)		$172.7(7)$ $167.4(4)$	149.8(4)					

N atom into the aromatic rings; cis -[W(CNC₆H₄NC)₂- $(dppe)_2$] provides the rare example containing the bent aryl isocyanide ligands with C-N-C angles of 136.8- (11) ° and $141.4(8)$ °.²³ This tendency is consistent with the finding that the C-N-C angle in *trans*-[Mo(PhNC)₂-(dppe)2] (172.9(8)°) is larger than that in *trans*-[Mo- $(MeNC)_2$ (dppe)₂] (156(1)°).²⁵ In this respect, the substantially bent C-N-C angle of 149.8(4)° observed in the PhNC complex **10a** is noteworthy.

All of the 31P{1H} NMR spectra for **⁹**-**¹¹** show one singlet assignable to the dppe ligands, indicating that all have the trans structure, demonstrated by the X-ray analyses for **9a** and **10a**. In the IR spectra appear the intense bands characteristic of the C-O and C-^N multiple bonds in the carbonyl, nitrile, and isocyanide ligands. Splitting of the *ν*(NC) band in **9a** (2017 and 1991 cm^{-1}) might be explained by the packing effect as observed previously for 4 and its W analogue.^{6a,13} By using the *ν*(NC) frequencies observed for a series of trans-[Mo(PhNC)(L)(dppe)₂], force constants of the isocyanide N-C multiple bonds may be calculated to be 15.3 (av), 14.4, 13.9, and 12.6 mdyn/Å for $L = CO$, (23) Hu, C.; Hodgeman, W. C.; Bennett, D. W. *Inorg. Chem.* **1996**,

³⁵, 1621.

⁽²⁴⁾ Basset, J.-M.; Berry, D. E.; Barker, G. K.; Green, M.; Howard, J. A. K.; Stone, F. G. A. *J. Chem. Soc., Dalton Trans.* **1979**, 1003.

⁽²⁵⁾ Chatt, J.; Pombeiro, A. J. L.; Richards, R. L.; Royston, G. H. D.; Muir, K. W.; Walker, R. *J. Chem. Soc., Chem. Commun.* **1975**, 708.

PhNC, N_2 , and p -MeOC₆H₄CN, respectively, and this order is similar to that observed for the force constants of the N-N bonds in *trans*- $[M_0(N_2)(L)(dppe)_2]$ (vide supra).

Preparation and Characterization of *trans***-[Mo-** $(RNC)(\eta^2-H_2)(dppe)_2$ (13). Activation of the H-H bond on transition metal centers is of much interest,²⁶ and the binding of H_2 to the 16e species $[M(CO)(R_2 PCH_2CH_2PR_2$] (M = Mo, R = aryl, alkyl; M = W, R = aryl) and $[Mn(CO)(R_2PCH_2CH_2PR_2)_2]^+$ (R = alkyl, aryl) has been investigated extensively. From these studies, several *η*²-H₂ complexes have already been isolated and characterized in a well-defined manner, which include *trans*-[Mo(CO)(η^2 -H₂)(R₂PCH₂CH₂PR₂)₂] (R = Ph,⁹) $PhCH_2$, *m*-CH₃C₆H₄CH₂²⁷) and *trans*-[Mn(CO)(η ²-H₂)(R₂- $PCH_2CH_2PR_2)_2$ ⁺ (R = Et, Ph).²⁸ In contrast, it has been revealed that the reactions of Mo complexes containing more electron-donating diphosphines under H_2 afford dihydride complexes $[Mo(CO)H_2(R_2PCH_2CH_2PR_2)_2]$ (R = Et, Buⁱ).^{9b,29} The analogous electronic control has been
achieved, by replacing the Mo atom with the more achieved by replacing the Mo atom with the more electron-rich W, which results in the formation of dihydride complex $[W(CO)H_2(Ph_2PCH_2CH_2PPh_2)_2]$.¹³

Now the reactions of **7** containing a readily replaceable N_2 ligand with H_2 have been investigated to clarify the interaction of H_2 with the ${Mo(RNC)(dppe)_2}$ chromophore. This will demonstrate for the first time the effect of the ligand occupying the site trans to the H_2 upon the bonding of the incorporated H_2 . In contrast to the reactions with CO, nitriles, and isocyanides, initial attempts to prepare $[Mo(RNC)(H₂)(dppe)₂]$ through the direct substitution of the N_2 ligand in **7** by H_2 (1 atm) at room temperature or 50 °C were unsuccessful, since most of **7** was recovered even after the prolonged reaction time. However, when a toluene solution of **7a** was heated at 90 °C for 30 min or a benzene solution of **7e** was refluxed for 30 min under Ar to dissociate the N_2 ligand and then the resultant black solutions presumably containing the five-coordinate complexes [Mo- $(RNC)(dppe)_2$ (8) were treated with H_2 gas (1 atm), the desired H₂ complexes *trans*-[Mo(RNC)($η$ ²-H₂)(Ph₂PCH₂- $CH_2PPh_2)_2$] (13a, R = Ph; 13e, R = p -Me₂NC₆H₄) could be isolated in moderate yields (Scheme 3).

Since the N_2 ligands in the isocyanide complexes 7 are bound to the Mo center more firmly than the N_2 in **4** containing the more electron-withdrawing CO ligand, removal of the N_2 to give the reactive five-coordinate species **8** probably requires more forcing conditions than those for generating **5** from **4**. This well correlates to the previous findings that **5** is readily obtained by bubbling Ar gas through a benzene solution of **4** at 50 °C for several minutes or degassing it in vacuo, whereas the W analogue [W(CO)(dppe)2] (**14**) is isolable after refluxing a benzene solution of *trans*-[W(CO)(N₂)(dppe)₂] (**15**) under Ar, owing to the presence of the more

13e: $R = p$ -Me₂NC₆H₄ 13f: $Buⁿ$

Table 4. T_1 Values for the H_2 Signal in *trans***[Mo(CNR)(H₂)(dppe)₂] (13)**^{*a*}

	13a		13e		13f	
temp(K)		T_1 (ms)		T_1 (ms)		T_1 (ms)
290	-4.8	h	-5.4		-5.9	20
260		21		15		15
230		17		20		17

 a 400 MHz; toluene-*d*₈ solutions. Inversion recovery method. *b* Data are unreliable due to much broadening of the signal.

electron-rich W center. The IR data of **4**, **15**, and **7** support this observation; the *ν*(NN) bands appear at 2110 and 2080 cm-¹ for **4** and at 2070 and 2030 cm-¹ for **15**, indicating the presence of the stronger backdonation to the N2 ligand in **15**. The fact that the *ν*- (NN) values observed for **7** are almost the same or lower than those of **15** is consistent with the requirement of the drastic conditions to generate **8**.

Despite the repeated trials, characterization of **8** could not be completed possibly due to its extreme sensitivity to air and moisture. Thus, although dark brown crystals obtained from the black solution produced after thermolysis of **7e** showed the satisfactory analytical data to be formulated as $[Mo(p-Me_2NC_6H_4NC)(dppe)_2]$ (**8e**), reliable spectral data were unable to be collected. As observed for **5** and **14**, solutions possibly containing **8** were treated with N_2 to give 7 reversibly.

Taking into account the weaker *π*-acidity of the RNC ligands than the CO ligand, **13** had been expected to have dihydride ligands rather than a H_2 ligand. However, the 1H NMR spectra for **13a** and **13e** both exhibited broad signals at -4.8 and -5.2 ppm, which are characteristic of the η^2 -H₂ protons. As summarized in Table 4, the T_1 values in the range $15-21$ ms at 230 and 260 K are comparable to 20 ms at 200 K of the fully characterized dihydrogen complex *trans*-[Mo(CO)(*η*2- H_2)(dppe)₂] (16),⁹ indicating unambiguously that these can be identified as the η^2 -H₂ complexes.²⁶ Even for 13f $(R = Bu^n)$, possibly having the more *π*-donating {Mo- $(BuⁿNC)(dppe)₂$ } moiety, the η^2 -H₂ nature has been demonstrated by the NMR data at 230, 260, and 290 K as listed in Table 4, although isolation of the pure

^{(26) (}a) Heinekey, D. M.; Oldham, W. J., Jr. *Chem. Rev.* **1993**, *93*, 913. (b) Crabtree, R. H. *Angew. Chem., Int. Ed. Engl.* **1993**, *32*, 789. (c) Jessop, P. G.; Morris, J. H. *Coord. Chem. Rev.* **1992**, *121*, 155. (d) Kubas, G. J. *Acc. Chem. Res.* **1994**, *27*, 183. (27) Luo, X.-L.; Kubas, G. J.; Burns, C. J.; Eckert, J. *Inorg. Chem.*

¹⁹⁹⁴, *33*, 5219.

^{(28) (}a) King, W. A.; Scott, B. L.; Eckert, J.; Kubas, G. J. *Inorg. Chem.* **1999**, *38*, 1069. (b) King, W. A.; Luo, X.-L.; Scott, B. L.; Kubas, G. J.; Zilm, K. W. *J. Am. Chem. Soc.* **1996**, *118*, 6782.

⁽²⁹⁾ Kubas, G. J.; Ryan, R. R.; Unkefer, C. J. *J. Am. Chem. Soc.* **1987**, *109*, 8113.

Figure 5. ORTEP diagram of **13e**. Hydrogen atoms are omitted for clarity.

compound showing satisfactory microanalysis data was unsuccessful.

The X-ray analysis was carried out for **13e**, whose results are shown in Figure 5 and Table 5. Although the *η*2-H2 moiety could not be located, **13e** has a common octahedral structure and the η^2 -H₂ ligand is inferred to be present at the site trans to the isocyanide. The structure clarified here is well comparable to that of fully characterized **16**. ⁹ By contrast, seven-coordinate dihydride complexes $[W(CO)H_2(dppe)_2]^{13}$ and $[Mo(CO)$ - $H_2(Et_2PCH_2CH_2PEt_2)_2]^{29}$ tend to have a pentagonal bipyramidal geometry. Morris et al. claimed previously that treatment of the octahedral d^6 N₂ complexes exhibiting a $\nu(NN)$ band in the region 2060-2150 cm⁻¹ with H_2 results in the formation of stable η^2 - H_2 complexes.30,31 It is noteworthy that **13a** and **13e** have apparently the well-characterized η^2 -H₂ ligand despite

the appearance of the *ν*(NN) band at 2048 and 2018 cm-¹ for **7a** and **7e**, respectively.

It is interesting that the $C(1)-N(1)$ bond in the isocyanide ligand in **13e** is fairly long at 1.213(4) Å and the $C(1)-N(1)-C(2)$ linkage is significantly bent with an angle of 148.4(4)°, which reflect the quite weak π -accepting nature of the η^2 -H₂ ligand in this complex. Considerably low *ν*(NC) frequencies observed for **13** (e.g., **13a**, 1875 and 1817; **13e**, 1891 and 1813 cm-1) also arise from this electronic feature of the η^2 -H₂ ligand.

Experimental Section

General Considerations. All manipulations were carried out under an atmosphere of nitrogen except for those stated otherwise. IR and NMR spectra were recorded on JASCO FT/ IR-420 and JEOL EX-270 or LA-400 spectrometers. The signals arising from the aromatic protons are omitted in the following 1H NMR data. Elemental analyses were done with a Perkin-Elmer 2400 series II CHN analyzer. Amounts of solvating molecules in the crystals were determined by the X-ray crystallography, NMR measurement, and/or GLC analysis. Complexes **1**³² and **12**16a were prepared according to the literature methods.

Preparation of *trans***-[Mo(RNC)(N2)(dppe)2] (7) from** Benzaldehyde Imines. Preparation of $7a$ $(R = Ph)$. Complex $1(166 \text{ mg}, 0.175 \text{ mmol})$ and PhCH=NPh $(33 \text{ mg}, 0.18)$ mol) were dissolved in benzene (2 mL), and the solution was heated at reflux for 2 h. After cooling, ether was layered on the resulting dark red solution. Dark red crystals of $7a·2C_6H_6$ deposited, which were filtered off and dried in vacuo (129 mg, 62%). Anal. Calcd for $C_{71}H_{65}N_3M_0P_4$: C, 72.26; H, 5.55; N, 3.56. Found: C, 71.99; H, 5.70; N, 3.23. IR (KBr): *ν*(NN), 2049; *ν*- (NC), 1910 cm⁻¹. ¹H NMR (C₆D₆): δ 2.35 (br, 8H, PCH₂). ³¹P- 1H NMR (C₆D₆): δ 68.9 (s).

Preparation of 7b ($R = C_6H_4Me$ **-***p***).** This compound was obtained from $1(166 \text{ mg}, 0.175 \text{ mmol})$ and $PhCH=NC_6H_4Me$ -p (68 mg, 0.35 mmol) by essentially the same procedure as that for preparing **7a** except that the red crystals of $7\mathbf{b} \cdot C_6H_6$ were isolated by addition of hexane to the product solution. Yield: 52%. Anal. Calcd for C₆₆H₆₁N₃MoP₄: C, 71.03; H, 5.51; N, 3.77. Found: C, 71.02; H, 5.73; N, 3.23. IR (KBr): *ν*(NN), 2053; *ν*- (NC), 1914 cm⁻¹. ¹H NMR (C₆D₆): *δ* 2.11 (s, 3H, Me), 2.40 (br, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): δ 69.1 (s).

The following compounds **7c**-**7g** were obtained similarly, except that **1** (0.175 mmol) was reacted with a larger excess of the imine (1.75 mmol).

 $7c \cdot 3/2C_6H_6$ ($\mathbb{R} = C_6H_4OMe$ -*p*): Red crystals, yield 46%. Anal. Calcd for C₆₉H₆₄N₃MoOP₄: C, 70.77; H, 5.51; N, 3.59. Found: C, 70.12; H, 5.74; N, 3.11. IR (KBr): *ν*(NN), 2054; *ν*- (NC), 1908 cm⁻¹. ¹H NMR (C₆D₆): δ 3.31 (s, 3H, OMe), 2.41 (br, 8H, PCH2). 31P{1H} NMR (C6D6): *δ* 69.0 (s). Repeated purification did not improve the analysis data.

7d·1/2C₆H₆ ($\mathbf{R} = \mathbf{C}_6\mathbf{H}_4\mathbf{F}$ - p): Orange crystals, yield 35%. Anal. Calcd for C₆₂H₅₅N₃FMoP₄: C, 68.89; H, 5.13; N, 3.89. Found: C, 68.23; H, 5.43; N, 3.58. IR (KBr): *ν*(NN), 2054; *ν*- (NC), 1907 cm⁻¹. ¹H NMR (C₆D₆): δ 2.3–2.5 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): δ 68.7 (s). Repeated purification did not result in the better carbon analysis.

7e ($\mathbf{R} = \mathbf{C}_6\mathbf{H}_4\mathbf{N}\mathbf{M}\mathbf{e}_2\mathbf{p}$ **):** Orange crystals, yield 38%. Anal. Calcd for C61H58N4MoP4: C, 68.67; H, 5.48; N, 5.25. Found: C, 68.70; H, 5.68; N, 5.22. IR (KBr): *ν*(NN), 2018; *ν*(NC), 1929 cm-1. 1H NMR (C6D6): *δ* 2.51 (s, 6H, NMe), 2.41 (br, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): δ 69.1 (s).

7f ($R = Bu^n$ **):** Orange crystals, yield 56%. Anal. Calcd for C57H57N3MoP4: C, 68.19; H, 5.72; N, 4.19. Found: C, 68.63; H, 5.60; N, 4.02. IR (KBr): *ν*(NN) and *ν*(NC), 1964 cm-1. 1H

⁽³⁰⁾ Morris, R. H.; Earl, K. A.; Luck, R. L.; Lazarowych, N. J.; Sella, A. *Inorg. Chem.* **1987**, *26*, 2674.

⁽³¹⁾ In the IR spectrum of **7f**, a very strong and broad band centered at 1963 cm-¹ appeared. The NN and NC stretching bands are probably overlapping, and it was impossible to assign the correct *ν*(NN) frequency.

⁽³²⁾ Hussain, W.; Leigh, G. J.; Mohd.-Ali, H.; Pickett, C. J.; Rankin, D. A. *J. Chem. Soc., Dalton Trans.* **1984**, 1703.

NMR (C₆D₆): *δ* 0.83 (t, *J* = 7.1 Hz, 3H, CH₂*Me*), 1.0-1.2 (m, 4H, C H_2 C H_2 Me), 2.90 (t, $J = 6.5$ Hz, 2H, NCH₂), 2.2-2.5 (br m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): δ 70.0 (s).

7g ($\mathbf{R} = \mathbf{Pr}^i$ **): Orange-red crystals, yield 60%. Anal. Calcd
C_{CC}H_{EC}N₀M₀P_C C 67.95; H 5.60; N 4.24. Found: C 67.80;** for C56H55N3MoP4: C, 67.95; H, 5.60; N, 4.24. Found: C, 67.80; H, 5.61; N, 3.96. IR (KBr): *ν*(NN), 2035; *ν*(NC), 1929 cm-1. 1H NMR (C_6D_6): *δ* 0.67 (d, $J = 6.4$ Hz, 6H, Me), 3.37 (sep, $J =$ 6.4 Hz, 1H, NCH), 2.3-2.5 (m, 8H, PCH₂). ³¹P{¹H} NMR (C6D6): *δ* 70.0 (s).

7h·C₆H₆ ($\mathbb{R} = \mathbb{C}H_2\mathbb{P}$ h). A mixture of 1 (167 mg, 0.176 mmol) and PhCH=NCH₂Ph (166 μ L, 0.882 mmol) in benzene (3 mL) was refluxed for 5 min, and the resultant dark green solution was filtered. Addition of hexane (10 mL) to the filtrate gave orange-red crystals of $7h$ ⁻C₆H₆ (111 mg, 56%). Anal. Calcd for $C_{66}H_{61}N_3MoP_4$: C, 71.03; H, 5.51; N, 3.77. Found: C, 70.75; H, 5.47; N, 3.68. IR (KBr): *ν*(NN), 1972; *ν*(NC), 1933 cm-1. 1H NMR (C₆D₆): *δ* 4.13 (s, 2H, NCH₂), 2.3–2.5 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): *δ* 69.9 (s).

Quantitative Analysis of the Benzene Produced in the Reaction Forming 7. A mixture of **1** (104 mg, 0.110 mmol) and PhCH=NPh (22.7 mg, 0.125 mmol) in toluene (1 mL) was refluxed under N_2 for 1 h. After cooling to room temperature, the resultant solution was subjected to the GLC analysis using a Shimadzu GC-14A Gas Chromatograph equipped with a 25 $m \times 0.25$ mm fused silica capillary column, which showed the formation of 0.71 mol of benzene per Mo atom. Formation of benzene (0.73 mol/Mo atom) from the reaction of **1** with $PhCH=NC_6H_4Me-p$ in *p*-xylene was confirmed by the analogous method.

Preparation of *trans***-[Mo(PhNC)(CO)(dppe)₂]** (9a). Through a suspension of $7a \cdot 2C_6H_6$ (114 mg, 0.0966 mmol) in THF (5 mL) was passed CO gas for a few minutes, and the mixture was stirred under CO for 25 h at room temperature. The resultant orange solid was filtered off, washed with THF and ether, and then dried in vacuo (57 mg, 58%). Addition of hexane (10 mL) to the combined THF filtrate afforded the second crop of **9a** as orange prisms (18 mg, 18%). Anal. Calcd for C₆₀H₅₃NMoOP₄: C, 70.38; H, 5.22; N, 1.37. Found: C, 69.88; H, 5.19; N, 1.53. IR (KBr): *ν*(NC), 2017 and 1991; *ν*(CO), 1812 cm-1. 1H NMR (C6D6): *δ* 2.35 (br, 8H, PCH2). 31P{1H} NMR (C6D6): *δ* 72.1 (s).

Preparation of *trans***-[Mo(BuⁿNC)(CO)(dppe)₂] (9f).** A benzene solution (15 mL) of **7f** (307 mg, 0.306 mmol) was stirred under CO at 50 °C for 3 h, and the resulting orange mixture was filtered. Hexane was added to the concentrated filtrate, affording **9f** as orange crystals (190 mg, 62% yield). Anal. Calcd for C₅₈H₅₇NMoOP₄: C, 69.39; H, 5.72; N, 1.40. Found: C, 68.75; H, 5.69; N, 1.66. IR (KBr): *ν*(NC), 2075; *ν*- (CO), 1788 cm⁻¹. ¹H NMR (C₆D₆): δ 0.71 (t, *J* = 6.4 Hz, 3H, Me), 0.8-0.9 (m, 4H, MeC*H*2C*H*2), 2.40 (m, 2H, NC*H*2), 2.2- 2.3 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): δ 73.4 (s).

Preparation of *trans***-[Mo(PhNC)(***p***-MeOC6H4CN)- (dppe)₂] (10a).** A mixture of $7a.2C_6H_6$ (315 mg, 0.308 mmol) and p -MeOC₆H₄CN (78 mg, 0.59 mmol) in benzene (17 mL) was stirred at room temperature for 24 h. Addition of hexane to the resultant solution gave dark brown prisms of **10a**'1/ $2C_6H_6$ (290 mg, 84% yield). Anal. Calcd for $C_{70}H_{63}N_2MoOP_4$: C, 71.98; H, 5.44; N, 2.40. Found: C, 71.93; H, 5.27; N, 2.24. IR (KBr): *ν*(CN_{nitrile}), 2171; *ν*(NC_{isocyanide}), 1821 and 1715 cm⁻¹. ¹H NMR (C₆D₆): *δ* 3.13 (s, 3H, OMe), 2.3–2.7 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): *δ* 70.2 (s).

Preparation of *trans***-[Mo(BunNC)(***p***-MeOC6H4CN)- (dppe)2] (10f).** This complex was obtained similarly from **7f** and p -MeOC₆H₄CN as dark brown crystals in 78% yield. Anal. Calcd for C65H64N2MoOP4: C, 70.39; H, 5.82; N, 2.53. Found: C, 70.27; H, 5.85; N, 2.58. IR (KBr): $ν$ (CN_{nitrile}), 2154; *ν*- $(NC_{\text{isocvanide}})$, 1734 cm⁻¹. ¹H NMR (C_6D_6) : δ 0.93 (t, *J* = 7.4 Hz, 3H, CH₂*Me*), 1.26 and 1.40 (m, 2H each, NCH₂C*H*₂*CH*₂), 3.22 (t, $J = 6.6$ Hz, 2H, NCH₂), 3.10 (s, 3H, OMe), 2.35-2.55 (m, 8H, PCH2). 31P{1H} NMR (C6D6): *δ* 71.8 (s).

Preparation of *trans***-[Mo(PhNC)(But NC)(dppe)2] (11).**

A mixture of $7a·2C_6H_6$ (99 mg, 0.097 mmol) and Bu^tNC (11
 μ I 0.098 mmol) in benzene (5 mJ) was stirred at room *µ*L, 0.098 mmol) in benzene (5 mL) was stirred at room temperature for 24 h. Crystallization from benzene/hexane afforded $11 \cdot C_6H_6$ (57 mg, 51% yield) as dark brown crystals. Anal. Calcd for C₇₀H₆₈N₂MoP₄: C, 72.66; H, 5.92; N, 2.42. Found: C, 72.50; H, 5.87; N, 2.60. IR (KBr): *ν*(NC), 2010 (But - NC) and 1880 (PhNC) cm-1. 1H NMR (C6D6): *δ* 0.64 (s, 9H, Bu^t), 2.2–2.5 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): *δ* 73.3 (s).

Preparation of $[Mo(p-Me_2NC_6H_4NC)(dppe)_2]$ (8e). A benzene solution (5 mL) of **7e** (108 mg, 0.101 mmol) was refluxed under Ar for 40 min to give a yellow-black solution. After cooling to room temperature, hexane (10 mL) was added and the mixture was filtered. Addition of hexane (12 mL) to the concentrated filtrate (ca. 5 mL) afforded **8e** as dark brown crystals (40 mg, 38% yield), which were separated manually from undesired byproducts deposited as orange crystals. Anal. Calcd for $C_{61}H_{58}N_2MoP_4$: C, 70.52; H, 5.63; N, 2.70. Found: C, 70.38; H, 5.85; N, 2.89.

Preparation of *trans***-[Mo(PhNC)** $(\eta^2$ **-H₂)(dppe)₂] (13a).** A toluene solution (4.5 mL) of $7a \cdot 2C_6H_6$ (99 mg, 0.097 mmol) was heated at 90 °C under Ar for 30 min. After cooling, the resultant yellowish black solution was degassed and then kept under H_2 (1 atm). The solution gradually turned to orangered, from which red crystals of 13a·C₆H₅CH₃ precipitated (46 mg, 43% yield). Addition of hexane to the filtrate gave the second crop (26 mg, 25% yield). Anal. Calcd for C₆₆H₆₃NMoP₄: C, 72.72; H, 5.83; N, 1.28. Found: C, 72.32; H, 5.74; N, 1.58. IR (KBr): $\nu(NC)$, 1875 and 1817 cm⁻¹. ¹H NMR (C₆D₆): δ -4.8 (br, 2H, H₂), 2.15 (s, 3H, C₆H₅C*H*₃), 2.2-2.4 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): *δ* 74.5 (s).

Preparation of *trans***-[Mo(** p **-Me₂NC₆H₄NC)(** η **²-H**₂) **(dppe)2] (13e).** A benzene solution (3 mL) of **7e** (84 mg, 0.078 mmol) was refluxed under Ar for 30 min. Analogous treatment of the resultant solution with H2 afforded **13e**, which was obtained as orange-red crystals by recrystallization from benzene-hexane (60 mg, 74% yield). Anal. Calcd for $C_{61}H_{60}N_2$ -MoP4: C, 70.38; H, 5.81; N, 2.69. Found: C, 70.01; H, 5.86; N, 2.67. IR (KBr): *ν*(NC), 1891 and 1813 cm⁻¹. ¹H NMR (C₆D₆): *δ* -5.2 (br, 2H, H₂), 2.55 (s, 6H, NCH₃), 2.3-2.6 (m, 8H, PCH₂). ³¹P{¹H} NMR (C₆D₆): *δ* 75.1 (s).

Preparation of *trans***-[Mo(BuⁿNC)(** η **²·H₂)(dppe)₂] (13f).** This complex was obtained from **7f** by the same procedure as that for **13e**. However, the compound is always contaminated by a small amount of byproduct(s), and satisfactory analysis data were not available. Anal. Calcd for $C_{57}H_{59}NMoP_4$: C, 70.01; H, 6.08; N, 1.43. Found: C, 69.27; H, 5.90; N, 1.61. IR (KBr): *ν*(NC), 1800−1900 cm⁻¹ (vbr). ¹H NMR (C₆D₆): δ −5.9 (br, 2H, H₂), 0.86 (t, $J = 7.4$ Hz, 3H, CH₃), 1.0-1.2 (m, 2H each, NCH₂C*H*₂C*H*₂), 2.94 (t, *J* = 6.6 Hz, 2H, NCH₂), 2.2-2.3 (m, 8H, PCH2). 31P{1H} NMR (C6D6): *δ* 76.3 (s).

X-ray Diffraction Studies. Single crystals were sealed in glass capillaries under N2 for **7a**, **9a**, **10a**, and **12** or under Ar for **13e**, which were transferred to a Rigaku AFC7R diffractometer equipped with a graphite-monochromatized Mo $K\alpha$ source. Diffraction studies were carried out at room temperature. Orientation matrixes and unit cell parameters were determined by least-squares treatment of 25 reflections with 35° < ²*^θ* < 40°.The intensities of three check reflections were monitored every 150 reflections during data collection, which revealed no significant decay for all crystals. Intensity data were corrected for Lorentz and polarization effects and for absorption (ψ scans). Details of crystal and data collection parameters are summarized in Table 6.

Structure solution and refinements were carried out by using the teXsan program package.³³ The positions of the nonhydrogen atoms were determined by Patterson methods and

⁽³³⁾ *teXsan: Crystal Structure Analysis Package*; Molecular Structure Corp.: The Woodlands, TX, 1985 and 1992.

subsequent Fourier syntheses (DIRDIF PATTY),³⁴ which were refined anisotropically by full-matrix least-squares techniques. The hydrogen atoms in **13e** were found in the Fourier map except for the η^2 -H₂ hydrogens and refined isotropically, while unobserved η^2 -H₂ hydrogens were not included for refinements. For the other complexes, all hydrogen atoms were placed at the calculated positions and included in the final stages of refinements with fixed parameters.

X-ray diffraction studies were undertaken also for the following compounds. Although the analyses could not be completed as described below, the preliminary results clarified the atom connecting scheme in these compounds.

7c·**3/2C₆H₆:** $a = 13.278(3)$ Å, $b = 22.298(3)$ Å, $c = 11.497(2)$ Å, α = 92.34(1)°, β = 113.43(1)°, and $γ = 104.19(1)°$ with *Z* = 2 in space group \overline{PI} (No. 2). $R(R_w) = 0.067$ (0.053) for 3594 data with $I > 3.0 \sigma(I)$. Carbon atoms in the dppe ligands were treated only isotropically.

7f: $a = 10.905(3)$ Å, $b = 12.370(5)$ Å, $c = 18.996(5)$ Å, $\alpha =$ 98.69(3)°, β = 101.42(2)°, and γ = 94.78(3)° with *Z* = 2 in space group *P*¹ (No. 2). *R* (R_w) = 0.080 (0.082) for 3747 data with *I* > 3.0 *^σ*(*I*). An uncharacterizable large peak still remained in the final Fourier map.

7g: $a = 12.821(4)$ Å, $b = 17.594(3)$ Å, $c = 11.471(3)$ Å, $\alpha =$ 100.56(2)°, $\beta = 103.47(3)$ °, and $\gamma = 84.54(2)$ ° with $Z = 2$ in space group \overline{PI} (No. 2). $R(R_w) = 0.078$ (0.070) for 4825 data with $I > 3.0 \sigma(I)$. The least-squres refinements resulted in the unusually large temperature factors for the Pri carbons probably due to the disorder of this group.

Acknowledgment. This work was supported by the Ministry of Education, Science, Sports, and Culture of Japan (Grant Nos. 09102004 and 09238103).

Supporting Information Available: Tables of atomic coordinates and equivalent isotropic thermal parameters, anisotropic thermal parameters, and extensive bond lengths and angles, together with figures with full atom-numbering scheme, for **7a**, **9a**, **10a**, **12**, and **13e**. This material is available free of charge via the Internet at http://pubs.acs.org.

OM9904605

⁽³⁴⁾ PATTY: Beurskens, P. T.; Admiraal, G.; Beurskens, G.; Bosman, W. P.; Garcia-Granda, S.; Gould, R. O.; Smits, J. M. M.; Smykalla, C. *The DIRDIF program system*; Technical Report of the Crystallography Laboratory: University of Nijmegen, The Netherlands, 1992.