

HOKKAIDO UNIVERSITY

Title	Cobalt(I)-Catalyzed Reductive Cyclization of Enynes and Diynes Using Hydrogen Gas as a Reductant
Author(s)	Isoda, Kaho; Sato, Yoshihiro
Citation	Organic letters, 25(12), 2103-2107 https://doi.org/10.1021/acs.orglett.3c00524
Issue Date	2023-03-31
Doc URL	http://hdl.handle.net/2115/91879
Rights	This document is the Accepted Manuscript version of a Published Work that appeared in final form in Organic Letters, copyright © American Chemical Society after peer review and technical editing by the publisher. To access the final edited and published work see https://pubs.acs.org/articlesonrequest/AOR-RR5P3TD2XX2QE749UHM7.
Туре	article (author version)
File Information	OL_Isoda.pdf



Cobalt(I)-Catalyzed Reductive Cyclization of Enynes and Diynes Using Hydrogen Gas as a Reductant

Kaho Isoda and Yoshihiro Sato*

Faculty of Pharmaceutical Sciences, Hokkaido University, Sapporo 060-0812, Japan Supporting Information Placeholder



ABSTRACT: Reductive cyclization of enynes and divnes by using H_2 gas as a reductant was realized, and the corresponding cyclized products were obtained in good yields without olefin-isomerization and over-reduction of the products. By an experiment using D_2 instead of H_2 , it was confirmed that H_2 unambiguously operates as a reductant in this reaction. The protocol of the reaction is very economical and user-friendly by using air- and moisture-stable CoBr₂·6H₂O and 1 atm pressure of H_2 .

Reductive coupling between two different π components (e.g., alkenes, alkynes, and carbonyl groups) is one of the unique reactions catalyzed by transition metal catalysts.¹ Reductive cyclization between diynes or an alkyne and an alkene is a promising methodology for regio- and stereoselective construction of cyclic compounds. Most of the reactions proceed via a metallacycle intermediate formed through oxidative cycloaddition of C-C multiple bonds to a low valent metal complex (M^n) , and it is known that various transition metals such as $Pd_{1}^{2} Ni_{1}^{3,4}$ and $Co^{3c,5}$ can catalyze this type of reaction (Scheme 1). As a typical scenario of reductive cyclization of enynes shown in Scheme 1 (a), cleavage of the C-M bond in the metallacycle occurs by a hydrogen source (e.g., HX), giving a cyclized product along with formation of $M^{n+2}X_2$. To establish a catalytic cycle of this reaction, more than a stoichiometric amount of a reductant is necessary to reduce Mⁿ⁺²X₂ to Mⁿ. Thus, more than a stoichiometric amount of waste originating from the reductant is also produced after the reaction, and that is a serious drawback of this reaction.

Hydrogen gas (H_2) is one of the ideal reductants in various transformations including the above-mentioned reductive couplings because it is cheap and easily accessible and there is no production of waste after the reaction. Furthermore, with increasing utilization of H_2 in various fields, technology for production of H_2 without CO_2 emission is being developed to replace the traditional methods such as steam reforming.⁶ However, the utilization of H_2 as a reductant in transition metal-catalyzed C-C-bond forming reactions has been limited, probably due to competition with catalytic hydrogenation. In 2002, Krische and coworkers reported Rh-catalyzed reductive coupling between an enone and an aldehyde with H_2 (1 atm).⁷ After that, they showed that H_2 can be used as a reductant in various types of reductive coupling, including reductive cyclization of enynes,⁸ in a series of their excellent studies.⁹ However, reductive cyclizations of enynes or diynes using H₂ as a reductant have been limited, and there have been only a few reports so far.¹⁰ We herein report Co(I)-catalyzed reductive cyclization of enynes and diynes by using 1 atm of H₂ as a reductant (Scheme 1 (b)).¹¹

Scheme 1. Transition Metal-Catalyzed Reductive Cyclization of Enynes

(a) A typical scenario of reductive cyclization of enynes



Mⁿ: Ru, Rh, Pd, Ni, Co, etc.

(b) This work: Co(I)-catalyzed reductive cyclization of enynes using H₂



Initially, the reaction of enyne **1a** with 10 mol% of the cationic Co(I) catalyst, prepared in situ from $CoBr_2$ (10 mol%) and various ligands in the presence of Zn (25 mol%) and NaBAr^F₄ (10 mol%) in a manner similar to the reported procedure, ^{5a} was carried out in dichloroethane (DCE) at 80 °C under 1 atm of H₂, and the results are

summarized in Table 1. The use of monodentate ligands, PPh₃ and PCy₃, was not effective for producing the desired compound **2a**, and most of **1a** was recovered in each reaction (Entries 1 and 2). Co(I) catalysts bearing bidentate ligands such as dppe, dppp, dppb, dcpe, and DPEphos promoted the cyclization, but a large amounts of isomerized product **3a** and/or alkyne-reduced starting material **4a** were obtained along with desired product **2a** (Entries 3-7). On the other hand, it was found that Xantphos-type ligands were effective for selective formation of the desired product **2a** without isomerized product **3a**. Thus, reactions of **1a** using Xantphos, N-Xantphos, and Xyl-Xantphos gave **2a** in 68%, 75%, and 83% yields, respectively, with a small amount of the reduced starting material **4a** (Entries 8-10).

Table 1. Ligand Screening in Reaction of 1a^a



					1a (%)
1 ^{b, c}	PPh ₃	-	-	trace	87
2^d	PCy ₃	-	-	-	quant.
3	dppe	23	17	5	5
4	dppp	21	44	trace	_
5	dppb	27	46	-	-
6	dcpe	48	33	-	-
7	DPEPhos	17	34	3	-
8	Xantphos	68	-	7	_
9	N- Xantphos	75	-	1	-
10	Xyl- Xantphos	83	-	6	-

^{*a*}Unless noted otherwise, the yields are isolated ones. ^{*b*}The yield was determined by ¹H-NMR analysis with 1,1,2,2-tetrachloroethane as an internal standard. ^c10 mol% of $CoBr_2(PPh_3)_2$ was used. ^{*d*}20 mol% of PCy₃ was used.



Next, we commenced further optimization of the reaction conditions in the reaction of **1a** using Xyl-Xantphos as a ligand (Table 2). When the reaction temperature was lowered from 80 °C to 60 °C, the yield was not affected, and the cyclized product **2a** was obtained in 82% yield (entry 1). On the other hand, in the reaction at 40 °C, the yield of **2a** was decreased to 52%, and over-reduced product **2a**' was also obtained in 15% yield, although the reason was not clear (Entry 2). For solvents, the reaction in toluene at 60 °C produced **2a** in 76% yield, while the use of THF and CH₃CN at the same temperature decreased the yield of **2a** to 36% and 4%, respectively (Entries 3-5). In these reactions, we used a cationic Co(I) catalyst formed in situ from CoBr₂ (10 mol%) and various ligands in the presence of Zn (25 mol%) and NaBAr^F₄ (10 mol%). When the amount of Zn was decreased from 2 to 10 mol%, the time required for complete consumption of the substrate **1a** was extended to 39 hours, but **2a** was obtained in good yield (Entry 6). On the other hand, the reaction in the absence of Zn or NaBAr^F₄ did not proceed at all, and the starting material **1a** was almost completely recovered (Entries 7 and 8), meaning that these reagents are necessary for formation of the active Co catalyst. It is noteworthy that the reaction under an atmosphere of N₂ instead of H₂ also did not proceed, indicating that H₂ plays a critical role for the reaction to proceed (Entry 9).

Table 2. Optimization for Reductive Cyclization of 1a^a



"Unless noted otherwise, the yields are isolated ones. **^b2a'** was obtained in 15% yield. 'Reaction time was 39 h. ^{*d*}The yield was determined by ¹H-NMR analysis with 1,1,2,2-tetrachloroethane as an internal standard.

TsN 2a'

In all of the experiments shown in Tables 1 and 2, commercially available "anhydrous CoBr2" was used as received. Since anhydrous CoBr₂ is known to be hygroscopic, we decided to use it after drying by heating at 130 °C in vacuo for 24 hours. As a result, in the reaction of 1a with dried CoBr₂ (10 mol%) and Xantphos (10 mol%), the yield of 2a significantly decreased to 20%, and the starting material 1a was recovered in 80% yield (Table 3, Entry 1). With the addition of 20 mol% water to the reaction under the same conditions, the yield of 2a was slightly improved to 36% (Entry 2). Surprisingly, the use of CoBr₂·6H₂O, which is the most stable hydrate complex of CoBr₂, increased the yield of **2a** to 90% (Entry 3). With the use of a Co(I) catalyst under the same conditions with Xyl-Xantphos, the yield of 2a reached 95% (Entry 4). Furthermore, in the reaction of 1 mmol scale of 1a, catalyst loading can be reduced to 2 mol%, and the product 2a was obtained in a high yield (Entry 5). As for the reason why CoBr₂·6H₂O was more effective than "anhydrous" or dried CoBr₂, we speculate that the presence of H₂O promoted the reduction of Co(II) species to Co(I) species by zinc. Similarly, Chirik recently reported that the presence of MeOH accelerated the reduction of a Co(II)-diphosphine complex to Co(I)-diphosphine complex by zinc as a reductant.12



"Unless noted otherwise, the yields are isolated ones. ^bThe yield was determined by ¹H-NMR analysis with 1,1,2,2-tetrachloroethane as an internal standard. 'Xantphos was used as a ligand. ^dXyl-Xantphos was used as a ligand. ^c1 mmol of **1a**, Xyl-Xantphos (2 mol%), Zn (5 mol%), and NaBAr^F₄ (2 mol%) were used in this reaction.

With the optimal conditions in hand, we next investigated the scope of substrates in this cyclization (Table 4). Various substituents, including electron-donating and electron-withdrawing groups, at the *p*-position of the aromatic ring at the terminus of the alkyne of enynes 1 were tolerated to give the corresponding cyclized products 2b-2f in 77%-95% yields. The reaction of 1g having a methyl group at the *m*-position of the aromatic ring took 24 hours to complete, but the desired product 2g was obtained in an excellent yield (97%). On the other hand, the reaction of 1h having a methyl group at the oposition did not proceed under the optimal conditions, probably due to the steric repulsion. In this case, however, the use of Xantphos instead of Xyl-Xantphos was effective and the product 2h was obtained in 91% yield. The existence of a heteroaromatic ring, thiophene, at the terminus of the alkyne somewhat retarded the reaction, but the cyclized product **2i** was obtained in a modest yield (36%). The reaction of 1j having a methyl group at the terminus of the alkyne gave 2j in 98% yield, indicating that the existence of an aromatic ring at the alkyne terminus is not essential. The presence of oxygen in the chain is also tolerated, but the yield of 2k was diminished to 46%. With respect to substrates having longer chains, the reaction of 1,7-enyne 11 gave the desired product 21 in 86% yield, while the reaction of 1,8-enyne 1m did not proceed, recovering 1m in 87% NMR yield. Next, the reaction of 1,6-diynes were checked under the same conditions. In the case of 1,6-diyne **1n** having a Ts amide moiety and phenyl substituents at each terminus of alkynes, the corresponding product 2n was obtained in 87% yield. Changing the Ts amide moiety in **1n** to oxygen did not affect the yield of the product, giving 20 in 70% yield. On the other hand, the malonate derivative 1p did not seem to be a good substrate in this reaction, giving the corresponding product 2p in 27% yield. Compared to diynes having phenyl substituents on the alkynes, 1,6-diynes 1q and 1r having methyl substituents on the alkynes are not suitable for this reaction, and the corresponding products 2q and 2r were obtained in 21% and 11% yields, respectively. Although it is a preliminary result, it is noteworthy that the reaction of allenyne 5 under the optimal conditions gave the product 6 in an excellent yield (99%)(Scheme 2).



"Isolated yields. ^bThe reaction time was 24 h. ^c10 mol% of Xantphos was used instead of Xyl-Xantphos. ^dThe reaction time was 48 h. ^eThe reduced starting material **4i** was also obtained in 27% yield. ^fThe residual starting material **1k** was observed as the mixture with unidentified byproducts. ^gThe starting material was almost consumed, and unidentified byproducts were also formed. ^hMost of the starting material remained but could not be separated from unidentified byproducts.

Scheme 2. Reaction of Allenyne^a



^aIsolated yields.

To gain an insight into the reaction mechanism, we conducted the reaction of **1a** under the optimal conditions using deuterium gas (D_2) instead of H_2 (Figure 1, upper part). As a result, the product **2a**-*d* was obtained in 87% yield, with deuterium being incorporated at each of the expected positions in **2a**-*d* with high D-contents (>95%). This result indicates that H_2 (or D_2) unambiguously acts as a reductant in the reaction. Also, it means that H_2O in CoBr₂·6H₂O and a catalytic amount of Zn (25 mol%) do not operate as reductants for the cyclization, and those would be necessary for preparation of an active Co(I) catalyst from a Co(II) complex.



Figure 1. Reaction using D2 and Proposed Mechanism

Based on the above-described result, the proposed mechanism of this reductive cyclization is shown in the lower part of Figure 1. A cationic Co(I) catalyst 7 was formed by reduction of CoBr₂.6H₂O by Zn in the presence of a ligand and NaBAr^F₄. Oxidative addition of substrate 1 to Co(I) catalyst 7 produced cobaltacycle intermediate I, which is the common intermediate in reductive cyclization catalyzed by a transition metal catalyst.¹⁻⁶ Then, σ -bond metathesis between I and H₂ occurred to give intermediate II,^{13,14} from which reductive elimination afforded product 2 along with regeneration of Co(I) catalyst 7. We speculate that a Xantphos-type ligand operates as a tridentate ligand in this reaction, which results in the prevention of side reactions such as isomerization and over-reduction by occupation of three coordination sites on the cobalt center.¹⁵

In conclusion, we succeeded in developing Co(I)-catalyzed reductive cyclization of enynes and diynes using H₂ as a reductant. The remarkable features of this reaction are as follows: (1) The reaction is economical and convenient owing to the use of the base metal Co and easily accessible H₂ and (2) the experimental protocol is userfriendly with easy handling owing to the use of an air-and moisturestable CoBr₂·6H₂O complex under 1 atm pressure of H₂. Further studies including extension to an asymmetric variant are in progress.

ASSOCIATED CONTENT

Data Availability Statement

The data underlying this study are available in the published article and its online Supporting Information.

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Experimental details and spectral data of substrate, cyclized product, and copies of NMR of the key compounds (PDF).

AUTHOR INFORMATION

Corresponding Author

Yoshihiro Sato – Faculty of Pharmaceutical Sciences, Hokkaido University, Sapporo 060-0812, Japan; E-mail: biyo@pharm.hokudai.ac.jp

Author

Kaho Isoda - Faculty of Pharmaceutical Sciences, Hokkaido University, Sapporo 060-0812, Japan

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENT

This work was financially supported in part by Grant-in-Aid for Scientific Research (B) (No. 20H03360) from the Japan Society for the Promotion of Science. This work was also partly supported by Hokkaido University, Global Facility Center (GFC), Pharma Science Open Unit (PSOU), funded by MEXT under "Support Program for Implementation of New Equipment Sharing System".

REFERENCES

(1) For selected reviews, see: (a) Broene, R. D. Reductive Coupling of Unactivated Alkenes and Alkynes. *Top. Curr. Chem.* **2007**, *279*, 209-248. (b) Holmes, M.; Schwartz, L. A.; Krische, M. J. Intermolecular Meta-Catalyzed Reductive Coupling of Dienes, Allenes, and Enynes with Carbonyl Compounds and Imines. *Chem. Rev.* **2018**, *118*, 6026-6052. (c) Agrawal, T.; Sieber, J. D. Recent Developments in C-C Bond Formation Using Catalytic Reductive Coupling Strategies. *Synthesis* **2020**, *52*, 2623-2638.

(2) (a) Trost, B. M.; Rise, F. A Reductive Cyclization of 1,6- and 1,7-Enynes. J. Am. Chem. Soc. **1987**, 109, 3161-3163. (b) Trost, B. M.; Lee, D. C. A Catalytic Reductive Cyclization of 1,6-Diynes. J. Am. Chem. Soc. **1988**, 110, 7255-7258. (c) Oh, C. H.; Rhim, C. Y.; Hwang, S. K. Chemoselectivities in Palladium Catalyzed Enyne Cyclizations. Chem. Lett. **1997**, 26, 777-778. (d) Oh, C. H.; Jung, H. H. Reductive elimination of alkylpalladium formate intermediates formed in enyne cyclizations. Tetrahedron Lett. **1999**, 40, 1535-1538. (e) Oh, C. H.; Jung, H. H.; Kim, J. S.; Cho, S. W. Highly Regio- and Stereoselective Cycloreductions of 1,6- and 1-7-Enynes Activated with Carbonyl Functionality. Angew. Chem. Int. Ed. **2000**, 39, 752-755.

(3) For reviews of Ni-catalyzed reductive cyclizations between alkynes and alkenes, see: (a) Montgomery, J. Nickel-Catalyzed Reductive Cyclizations and Couplings. *Angew. Chem. Int. Ed.* **2004**, 43, 3890-3908. (b) Tasker, S. Z.; Standley, E. A.; Jamison, T. F. Recent advances in homogeneous nickel catalysis. *Nature* **2014**, 509, 299-309. For reviews of Ni- and Co-catalyzed reductive cyclizations between alkynes and alkenes, see: (c) Jeganmohan, M.; Cheng, C.-H. Cobalt- and Nickel-Catalyzed Regio- and Stereoselective Reductive Coupling of Alkynes, Allenes, and Alkenes with Alkenes. *Chem. Eur. J.* **2008**, *14*, 10876-10886.

(4) Chen, M.; Weng, Y.; Guo, M.; Zhang, H.; Lei, A. Nickel-Catalyzed Reductive Cyclization of Unactivated 1,6-Enynes in the Presence of Organozinc Reagents. *Angew. Chem. Int. Ed.* **2008**, *47*, 2279-2282.

(5) (a) Chang, H.-T.; Jayanth T. T.; Wang, C.-C.; Cheng, C.-H. Cobalt-Catalyzed Reductive Coupling of Activated Alkenes with Alkynes. J. Am. Chem. Soc. 2007, 129, 12032-12041. For reviews, see: (b) Gandeepan, P.; Cheng, C.-H. Cobalt Catalysis Involving π -Components in Organic Synthesis. Acc. Chem. Res. 2015, 48, 1194-1206. (c) Biswas, S.; Parsutkar, M. M.; Jing, S. M.; Pagar, V. V.; Herbort, J. H.; RajanBabu, T. V. A New Paradigm in Enantioselective Cobalt Catalysis: Cationic Cobalt(I) Catalysts for Heterodimerization Reactions of Olefins. *Acc. Chem. Res.* **2021**, 54, 4545-4564. For recent examples of Co-catalyzed reductive cyclization of enynes, see: (d) Xi, T.; Lu, Z. Cobalt-Catalyzed Hydrosilylation/Cyclization of 1,6-Enynes. J. Org. Chem. 2016, 81, 8858-8866. (e) Yang, Z.; Hou, S.; Cheng, Y.; Sun, L.; Yang, C.-H. Co-Catalyzed Reductive Cyclization of Acrylate-Containing 1,6-Enynes. *J. Org. Chem.* **2022**, *87*, 13339-13345.

(6) For a recent review of this matter, see: Qureshi, F.; Yusuf, M.; Khan, M. A.; Ibrahim, H.; Ekeoma, B. C.; Kamyab, H.; Rahman, M. M.; Nadda, A. K.; Chelliapan, S. A State-of the-Art Review on the Latest trends in Hydrogen production, storage, and transportation techniques. *Fuel* **2023**, *340*, 127574.

(7) Jang, H.-Y.; Huddleston, R. R.; Krische, M. J. Reductive Generation of Enolates from Enones Using Elemental Hydrogen: Catalytic C-C Bond Formation under Hydrogen Conditions. *J. Am. Chem. Soc.* **2002**, *124*, 15156-15157.

(8) (a) Jang, H.-Y.; Krische, M. J. Rhodium-Catalyzed Reductive Cyclization of 1,6-Diynes and 1,6-Enynes Mediated by Hydrogen: Catalytic C-C Bond Formation via Capture of Hydrogenation Intermediates. *J. Am. Chem. Soc.* **2004**, *126*, 7875-7880. (b) Jang, H.-Y.; Hughes, F. W.; Gong, H.; Zhang, J.; Brodbelt, J. S.; Krische, M. J. Enantioselective Reductive Cyclization of 1,6-Enynes via Rhodium-Catalyzed Asymmetric Hydrogenation: C-C Bond Formation Precedes Hydrogen Activation. *J. Am. Chem. Soc.* **2005**, *127*, 6174-6175.

(9) For selected reviews, see: (a) Jang, H.-Y.; Krische, M. J. Catalytic C-C Bond Formation via Capture of Hydrogenation Intermediates. *Acc. Chem. Res.* **2004**, *37*, 653-661. (b) Nguyen, K. D.; Park, B. Y.; Luong, T.; Sato, H.; Garza, V. J. Krische, M. J. Metal-Catalyzed Reductive Coupling of Olefin-Derived Nucleophiles: Reinventing Carbonyl Addition. *Science* **2016**, *354*, No. aah5133. (c) Santana, C. G.; Krische, M. J. From Hydrogenation to Transfer Hydrogenation to Hydrogen Auto-Transfer in Enantioselective Metal-Catalyzed Carbonyl Reductive Coupling: Past, Present, and Future. *ACS Catal.* **2021**, *11*, 5572-5585.

(10) (a) For Pt-catalyzed reductive cyclization of diynes and enynes using H₂ as reductant: Jung, I. G.; Seo, J.; Lee, S. I.; Choi, S. Y.; Chung, Y. K. Reductive Cyclizations of Diynes and Enynes Catalyzed by Allyl Platinum N-Heterocyclic Carbene Complexes. *Organometallics* 2006, *25*, 4240-4242.
(b) For Fe-catalyzed reductive cyclization of diynes and enynes using H₂ as reductant: Sylvester, K. T.; Chirik, P. J. Iron-Catalyzed, Hydrogen-Mediated Reductive Cyclization of 1,6-Enynes and Diynes: Evidence for Bis(imino)pyridine Ligand Participation. *J. Am. Chem. Soc.* 2009, *131*, 8772.
(c) Pt-catalyzed reductive cyclization of allenynes using H₂ as reductant was reported: H.-T. Kim, H.-S. Yoon, W.-Y. Jang, Y. K. Kang, H.-Y. Jang, *European J. Org. Chem.* 2011, 2011, 3748.

(11) It has been reported that reductive cyclization occurred in competition with reductive Pauson-Khand reaction of enynes by using a stoichiometric amount of $Co_4(CO)_{12}$ under an atmosphere of H_{2} , see: Krafft, M. E.; Boñaga, L. V. R.; Wright, J. A.; Hirosawa, C. Cobalt Carbonyl-Mediated Carbocyclizations of Enynes: Generation of Bicyclooctanones or Monocyclic Alkenes. J. Org. Chem. **2002**, 67. 1233-1246.

(12) Friedfeld, M. R.; Zhong, H.; Ruck, R. T.; Shevlin, M.; Chirik, P. J. Cobalt-Catalyzed Asymmetric Hydrogenation of Enamides Enabled by Single-Electron Reduction. *Science* **2018**, *360*, 888-893.

(13) We have recently found that $C(sp^2)$ -metal bond is more reactive than $C(sp^3)$ -metal bond in rhodacyclopentene intermediate, which is a congener of intermediate I, toward σ -bond metathesis and insertion of C=X bond. Thus, we speculated that σ -bond metathesis between I and H₂ occurred at the $C(sp^2)$ -Co bond in I. At present, however, other possibilities (e.g., σ -bond metathesis at the $C(sp^3)$ -Co bond in I, successive insertion of alkyne/alkene to Co-H species generated from Co(I) catalyst and H₂) cannot be excluded. For related references with respect to the reactivity of rhodacyclopentene intermediate, see ref. 14.

(14) For α -bond metathesis of rhodacyclopentene with C-H bond, see: (a) Oonishi, Y.; Kitano, Y.; Sato, Y. Csp³-H Bond Activation Triggered by Formation of Metallacycles: Rh(I)-Catalyzed Cyclopropanation/Cyclization of Allenynes. Angew. Chem. Int. Ed. 2012, 51, 7305-7308. (b) Huang, G. Mechanism of Rhodium-Catalyzed Cyclopropanation/Cyclization of Allenynes. Org. Lett. 2015, 17, 1994-1997. (c) Oonishi, Y.; Sakamoto, S.; Agata, S.; Sato, Y. Rh(I)-Catalyzed Enantioselective Cyclization of Enynes through Site-Selective C(sp³)-H Bond Activation. Synthesis 2021, 53, 2976-2983. For α -bond metathesis of rhodacyclopentene with O-H bond, see: (d) Oonishi, Y.; Masusaki, S.; Sakamoto, S.; Sato, Y. Rhodium(I)-Catalyzed Enantioselective Cyclization of Enynes by Intramolecular Cleavage of the Rh-C Bond by a Tethered Hydroxy Group. Angew. Chem. Int. Ed. 2019, 58, 8736-8739. For C=O bond insertion, see (e) Oonishi, Y.; Yokoe, T.; Hosotani, A.; Sato, Y. Rhodkum(I)-Catalyzed Cyclization of Allenynes with a Carbonyl Group through Unusual Insertion of a C=O Bond into a Rhodacycle Intermediate. Angew. Chem. Int. Ed. 2014, 53, 1135-1139. For C=N bond insertion, see (f) Oonishi, Y.; Hato, Y.; Sato, Y. Rhodium(I)-Catalyzed [2+2+2] Cycloaddition between Allene, Alkyne, and Imine via a Strained Azarhodacycle Intermediate. Adv. Synth. Catal. 2015, 357, 3033-3039. (g) Oonishi, Y.; Hato, Y.; Sato, Y. Rhodium(I)-Catalyzed Diastereoselective Cycloisomerization of Envnes with Tethered (S)-2-Methyl-2-propanesulfinyl Imine. Adv. Synth. Catal. 2016, 358, 2273-2279. (h) Hato, Y.; Oonishi, Y.; Yamamoto, Y.; Nakajima, K.; Sato, Y. Stereoselective Construction of Spiro-Fused Frameworks by Sequential Reaction of Enynes, Imines, and Diazoalkenes with Rh(I) and Rh(III) Catalysts. J. Org. Chem. 2016, 81, 7847-7854.

(15) It is known that Xantphos-type ligands can operate as tridentate ligands, see: (a) Adams, G. M.; Weller, A. S. POP-Type Ligands: Variable Coordination and Hemilabile Behavior. Coord. Chem. Rev. 2018, 355, 150-172.
(b) van Leeuwen, P. W. N. M.; Kamer, P. C. J. Featuring Xantphos. Catal. Sci. Technol. 2018, 8, 26-113.