Rhenium-catalyzed [2 + 2] Cycloadditions of Norbornenes with Internal and Terminal Acetylenes

Yoichiro Kuninobu,* Peng Yu, and Kazuhiko Takai*

Division of Chemistry and Biochemistry, Graduate School of Natural Science and Technology,

Okayama University, Tsushima, Okayama 700-8530

(Received June 27, 2007; CL-070686; E-mail: kuninobu@cc.okayama-u.ac.jp; ktakai@cc.okayama-u.ac.jp)

Treatment of norbornenes with internal and terminal acetylenes in the presence of a catalytic amount of [ReBr(CO)₃(thf)]₂ gave cyclobutene derivatives in good to excellent yields.

[2+2] Cycloadditions of olefins with acetylenes is one of the most powerful tools to synthesize cyclobutene derivatives. Following Woodward–Hoffman rules, [2+2] cycloaddition reactions usually proceed under UV-irradiation conditions. However, using transition-metal complexes sometimes enables the reaction to be promoted without UV-irradiation. There have been many reports on metal-mediated or catalyzed [2+2] cycloadditions of norbornenes with acetylenes; the following metal complexes have been used: ruthenium, iron, cobalt, holdium, nickel, palladium, and copper. Recently, we have been investigating the catalytic abilities of rhenium complexes. In the process, we found that a rhenium complex also has the ability to catalyze [2+2] cycloadditions between norbornenes and acetylenes.

By treatment of norbornene (1a) with dimethyl but-2-ynedioate (2a) in the presence of a rhenium complex, [Re-Br(CO)₃(thf)]₂, as a catalyst, [2 + 2] cycloaddition reaction proceeded and cyclobutene derivative 3a was formed in 30% yield (Table 1, Entry 1). Another rhenium complex, ReBr(CO)₅, also provided 3a in the same yield. ¹¹ To the best of our knowledge, this is the first example of rhenium-catalyzed [2 + 2] cycloaddition between a norbornene derivative and an acetylene.

To improve the yield of **3a**, several additives were examined (Table 1). The yield of **3a** did not increase by addition of various Lewis acids and bases (Table 1, Entries 2–10). However, when *tert*-butyl isocyanide was added, the yield of **3a** was increased (Table 1, Entry 11). Benzyl isocyanide and 2,6-dimethylphenyl isocyanide also provided **3a** in moderate yields, respectively (Table 1, Entries 12 and 13). By using a bulky isocyanide, 2,6-diisopropylphenyl isocyanide, cyclobutene derivative **3a** was obtained in 71% yield (Table 1, Entry 14). ¹²

By increasing the amounts of the rhenium catalyst, [Re-Br(CO)₃(thf)]₂, and 2,6-diisopropylphenyl isocyanide, the yield of cyclobutene derivative **3a** was increased slightly (Table 2, Entry 1). Norbornadiene (**1b**) also gave the corresponding cyclobutene **3b**; however, the yield of **3b** was low (Table 2, Entry 2). In this reaction, a 1:2-adduct was not formed. Treatment of benzonorbornadiene (**1c**) with acetylene **2a** provided cyclobutene **3c** in 68% yield (Table 2, Entry 3). When the reaction was conducted at higher temperature (150 °C), the yield of **3c** increased to 92% yield (Table 2, Entry 4).

Next, we investigated the reactivities of several acetylenes (Table 3). Diethyl but-2-ynedioate (**2b**) also afforded **3d**; however, the yield of **3d** was low (Table 3, Entry 1). By using acetylenes having a phenyl or an alkyl group, **2c** and **2d**, also pro-

Table 1. Investigation of various additives^a

A	√ + MeO₂C——	[R CO ₂ Me —	ReBr(CO) ₃ (thf)] ₂ (2.5 mol %) additive (5.0 mol %)			
1a			toluene, 115 °C, 24 h 3a			
Entry	Additive	Yield/% ^b	Entry	Additive	Yield/%b	
1	none	30	8	benzonitrile	5	
2	Sc(OTf) ₃	30	9	N,N-dimethylimidazolidinone	10	
3	Y(OTf) ₃	28	10	tetrahydrothiophene	1	
4	In(OTf) ₃	36	11	t-BuNC	43	
5	PPh_3	5	12	PhCH ₂ NC	52	
6	N,N-dimethylaniline	5	13	$2,6\text{-Me}_2\text{C}_6\text{H}_3\text{NC}$	58	
7	pyridine	1	14	2,6- <i>i</i> -Pr ₂ C ₆ H ₃ NC	71	

^a2a (2.0 equiv.). ^{b1}H NMR yield.

Table 2. Reactions between norbornene 1 and dimethyl but-2-ynedioate (2a)^a

Entry	Norbornene	Temp/°C	Yield/% ^b
1	1a	115	3a 79 (86)
2	1b	115	3b 37 (41)
3	1c	115	3c 68 (70)
4	1c	150	3c 92 (95)

^a2a (2.0 equiv.) ^bIsolated yield. The yield determined by ¹H NMR is reported in parentheses.

duced the corresponding cyclobutene derivatives **3e** and **3f** in 57 and 77% yields, respectively (Table 3, Entries 2 and 3). It is usually difficult to obtain cyclobutene derivatives from terminal acetylenes. ^{13,14} By using a rhenium catalyst, cyclobutene derivatives **3g–3j** were also obtained from terminal acetylenes **2e–2h** (Table 3, Entries 4–7). In these reactions, trimerization products of acetylenes were not detected.

The proposed reaction mechanism is as follows (Scheme 1):

Table 3. Reactions between norbornadienes 1 and acetylenes 2^a

Entry	Norbornene	Acetylene	Temp/°C	Yield/% ^b
1	1c	$EtO_2C -\!$	150	3d 40 (45)
2 ^c	1c	MeO ₂ C———Ph 2c	180	3e 57 (60)
3	1c	MeO_2C ——— Me 2d	150	3f 77 (82)
4	1c	MeO ₂ C	150	3g 76 (77)
5	1c	EtO ₂ C——H 2f	150	3h 72 (73)
6	1c	О = Н 2д	150	3i 22 (41)
7	1b	Ph───H 2h	115	3j 17 (20)

^a**2a** (2.0 equiv.) ^bIsolated yield. The yield determined by ¹H NMR is reported in parentheses. ^cBenzonorbornadiene (2.0 equiv.), acetylene (1.0 equiv.), ReBr(CO)₅ was used as a catalyst.

Scheme 1. Proposed mechanism of the formation of cyclobutene derivatives.

(1) coordination of a norbornene and an acetylene to a rhenium center; (2) formation of a rhenacyclopentene intermediate; ^{9f,15,16} (3) reductive elimination.

In summary, we have succeeded in the [2+2] cycloaddition of norbornenes with both internal and terminal acetylenes using a rhenium complex, $[ReBr(CO)_3(thf)]_2$, as a catalyst and an isocyanide, 2,6-diisopropylphenyl isocyanide, as an additive. Recently, we have reported rhenium-catalyzed insertion of acetylenes into a carbon–carbon single bond of non-strained cyclic compounds under mild conditions. In the first step of the ring-enlargement, we have postulated the formation of a rhenacyclopentene intermediate by the reaction of the rhenium catalyst, a β -keto ester and a terminal acetylene. To the best of our knowledge, this is the first example of rhenium-catalyzed [2+2] cycloaddition, and this result supports the mechanism for the ring-enlargement.

References and Notes

- a) M. Frank-Neumann, M. Miesch, L. Gross, *Tetrahedron Lett.* 1990, 31, 5027. b) D. Neville, W. S. Murphy, *Tetrahedron Lett.* 1996, 37, 5221.
- 2 a) T. Mitsudo, K. Kokuryo, Y. Takegami, J. Chem. Soc., Chem.

- Commun. 1976, 722. b) T. Mitsudo, T. Kondo, Synlett 2001, 3, 309. c) P. Alvarez, J. Gimeno, E. Lastra, S. Garcia-Granda, J. F. Van der Maelen, M. Bassetti, Organometallics 2001, 20, 3762. d) A. Tenaglia, L. Giordano, Synlett 2003, 2333. e) K. Villeneuve, W. Tam, Angew. Chem., Int. Ed. 2004, 43, 610. f) P. Liu, R. W. Jordan, S. P. Kibbee, J. D. Goddard, W. Tam, J. Org. Chem. 2006, 71, 3793.
- A. Greco, A. Carbonaro, G. Dall'Asta, J. Org. Chem. 1970, 35, 271.
- 4 K. C. Chao, D. K. Rayabarapu, C.-C. Wang, C.-H. Cheng, J. Org. Chem. 2001, 66, 8804.
- 5 T. Shibata, K. Takami, A. Kawachi, Org. Lett. 2006, 8, 1343.
- 6 M. Lautens, L. G. Edwards, W. Tam, A. J. Lough, J. Am. Chem. Soc. 1995, 117, 10276.
- 7 B. M. Trost, M. Yanai, K. Hoogsteen, J. Am. Chem. Soc. 1993, 115, 5294.
- Y. Takenaka, H. Ito, M. Hasegawa, K. Iguchi, *Tetrahedron* 2006, 62, 3380.
- a) Y. Kuninobu, A. Kawata, K. Takai, J. Am. Chem. Soc. 2005, 127, 13498.
 b) Y. Kuninobu, A. Kawata, K. Takai, Org. Lett. 2005, 7, 4823.
 c) Y. Kuninobu, Y. Tokunaga, A. Kawata, K. Takai, J. Am. Chem. Soc. 2006, 128, 202.
 d) Y. Kuninobu, Y. Nishina, M. Shouho, K. Takai, Angew. Chem., Int. Ed. 2006, 45, 2766.
 e) Y. Kuninobu, Y. Nishina, K. Takai, Org. Lett. 2006, 8, 2891.
 f) Y. Kuninobu, A. Kawata, K. Takai, J. Am. Chem. Soc. 2006, 128, 11368.
 g) Y. Kuninobu, Y. Nishina, C. Nakagawa, K. Takai, J. Am. Chem. Soc. 2006, 128, 12376.
 h) Y. Kuninobu, Y. Inoue, K. Takai, Chem. Lett. 2006, 35, 1376.
- Rhenium-catalyzed reactions have also been reported by other groups, see: a) F. E. Kühn, A. Scherbaum, W. A. Herrmann, J. Organomet. Chem. 2004, 689, 4149. b) M. R. Luzung, F. D. Toste, J. Am. Chem. Soc. 2003, 125, 15760. c) K. A. Nolin, R. W. Ahn, F. D. Toste, J. Am. Chem. Soc. 2005, 127, 12462. d) H. Kusama, H. Yamabe, Y. Onizawa, T. Hoshino, N. Iwasawa, Angew. Chem., Int. Ed. 2005, 44, 468. e) L. L. Ouh, T. E. Müller, Y. K. Yan, J. Organomet. Chem. 2005, 690, 3774.
- 11 This reaction did not proceed using $Re_2(CO)_{10}$, $ReBr(CO)_3$ -(2,2'-bipyridyl), $Re(C_5Me_5)(CO)_3$, $ReCl_3$, $ReCl_3(PMe_2Ph)_3$, $ReCl_3O(PPh_3)_2$, and Re_2O_7 .
- 12 The role of isocyanides is not clear. However, addition of the isocyanide inhibited the polymerization of acetylenes.
- 13 Ruthenium complexes, RuCl₂(PPh₃)₃ and RuH₂(CO)(PPh₃)₃, which are sometimes employed as catalysts to promote [2 + 2] cycloadditions of norbornenes with acetylenes, did not give cyclobutene derivatives 3g and 3h.
- 14 For ruthenium-catalyzed [2+2] cycloadditions using terminal acetylenes, see: T. Mitsudo, H. Naruse, T. Kondo, Y. Ozaki, Y. Watanabe, Angew. Chem., Int. Ed. Engl. 1994, 33, 580. However, the yields of cyclobutenes are low because of cyclic trimerization of acetylenes.
- 15 There are some reports on transition-metal-catalyzed synthesis of [2+2] cycloadducts by the reactions between bicyclic alkenes and acetylenes via metalacyclopentene intermediates, see: a) T. Mitsudo, K. Kokuryo, T. Shinsugi, Y. Nakagawa, Y. Watanabe, Y. Takegami, J. Org. Chem. 1979, 44, 4492. b) D.-J. Huang, D. K. Rayabarapu, L.-P. Li, T. Sambaiah, C.-H. Cheng, Chem. Eur. J. 2000, 6, 3706. c) K. C. Chao, D. K. Rayabarapu, C.-C. Wang, C.-H. Cheng, J. Org. Chem. 2001, 66, 8804.
- 16 In this step, the orientation of the acetylene is not clear.
- 17 Supporting Information is available electronically on the CSJ-Journal Web site, http://www.csj.jp/journals/chem-lett/ index.html.