Site-Selective Hydrosilylation of Botryococcene - The Algal Biomass Hydrocarbon Oil

Dr. Hidehisa Kawashima,*[a, b, c] Mami Umezawa^[d] and Prof. Masashi Kijima*[a, b, c, e]

Abstract: An algal acyclic triterpene botryococcene ((*E*)-2,3,7,10,13,16,20,21-octamethyl-6,17-dimethylene-10-vinyldocosa-1,11,21-triene), new biomass for material sources, which has three-types of six carbon-carbon double bonds, *i.e.*, four vinylidenes at C2,6,17,21-positions, one C10-vinyl, and one C11-12 internal olefin, in the molecule underwent hydrosilylation of various organosilanes in the presence of Pt(0) 1,3-divinyl-1,1,3,3-tetramethyldisiloxane complex (Karstedt's catalyst). Various hydrosilylated botryococcene derivatives were site- and regioselectively obtained in high yields by an *anti*-Markovnikov type addition at the C10-vinyl moiety without any isomerizations of the unsaturated triterpenoid skeleton.

Introduction

Botryococcus braunii, a freshwater microalga, produces a variety of acyclic and monocyclic terpene hydrocarbons known as botryococcene. The production efficiency of botryococcene from *B. braunii* is high, and the oil contents are 17-90 w% in the dry alga body,^[1] which has accelerated researches on large-scale production of the algae biomass for the utilization of renewable energy and material sources.^[2] Among them, M. Watanabe et al. have reported that *B. braunii* (the strain Bot-22) dominantly produces a specific botryococcene (1) shown in Figure 1.^[3,4]

[a]	Dr. H. Kawashima, Prof. Dr. M. Kijima
	Division of Materials Science, Faculty of Pure and Applied Sciences
	University of Tsukuba
	1-1-1 Tennodai, Tsukuba, Ibaraki 305-8573, Japan
	E-mail: kawashima@ims.tsukuba.ac.jp (H. Kawashima),
	kijima@ims.tsukuba.ac.jp (M. Kijima)
[b]	Dr. H. Kawashima, Prof. Dr. M. Kijima
	Interdisciplinary Research Center for Catalytic Chemistry
	National Institute of Advanced Industrial Science and Technology
	1-1-1 Higashi, Tsukuba, Ibaraki, 305-8565, Japan.
[c]	Dr. H. Kawashima, Prof. Dr. M. Kijima
	Algae Biomass and Energy System R & D Center
	University of Tsukuba
	1-1-1 Tennodai, Tsukuba, Ibaraki 305-8573, Japan
[d]	M. Umezawa
	School of Science and Engineering, College of Engineering Science
	University of Tsukuba
	1-1-1 Tennodai, Tsukuba, Ibaraki 305-8573, Japan
[b]	Prof. Dr. M. Kijima
	Tsukuba Research Center for Interdisciplinary Materials Science
	University of Tsukuba
	1-1-1 Tennodai, Tsukuba, Ibaraki 305-8573, Japan
	Supporting information for this article is given via a link at the end of

Supporting information for this article is given via a link at the end of the document.

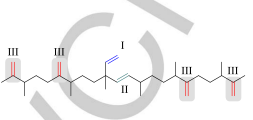


Figure 1. The chemical structure of botryococcene 1.

Botryococcene 1 has three types of six C-C double bonds of vinyl (I), internal trans-olefin (II), and vinylidene (III) in the molecule. Unlike most other biomass sources, 1 has no heteroatoms similar to main structures of petroleum components. From the perspective of synthetic organic chemistry, 1 should be a target for natural product synthesis because it has a peculiar chemical structure including a quaternary asymmetric carbon. Histrically, botryococcene analogues have been studied in organic chemistry for structural determination and elucidation of biosynthetic pathways,[5,6] and the first total synthesis of 1 was achieved by J. D. White et al. in 1988.^[7] However, except for the structure determination, little has been known about reactivity and chemical transformations of 1. It is necessary to develop methods for selective modifications of 1 to use as a source of chemical materials, because 1 merely has the three types of plural inactive C-C double bonds. It is thought that the reactivity of 1 in common organic reactions is usually low, *i.e.*, (i) the type-I vinyl moiety that positions in the middle of the terpenoid, whose reactivity should be suppressed by steirc suroundings, (ii) the type-II internal olefin between the tertially and quaternary carbons was specifically unreactive due to the buried environment, and (iii) the reactivity of the type-III four vinylidene moieties should be identical to that of similar terpenoids.^[8]

It is envisaged that hydrosilylation of **1** with hydrosilanes can proceed by selecting an appropriate transition metal catalyst, because inactive alkenes and alkynes have been reported to react with various hydrosilanes when the certain catalyst was used such as Pt, Ru, Rh, Fe, Co and Ni.^[9-12] In particular, the Ptbased catalysts showing high reactivity have been extensively studied and used industrially. Furthermore, a site-selective reaction for **1** is expected to proceed, because terminal olefins should preferentially undergo hydrosilylation compared to branched olefins.^[13]

In this paper, it is reported that a series of botryococcene derivatives are succesfully synthesized in high yields by a site-selective hydrosilylation of **1** with various organosilanes using a Pt-based catalyst. The selective modification of **1** can indicate applicability of the algal substance to chemical materials.

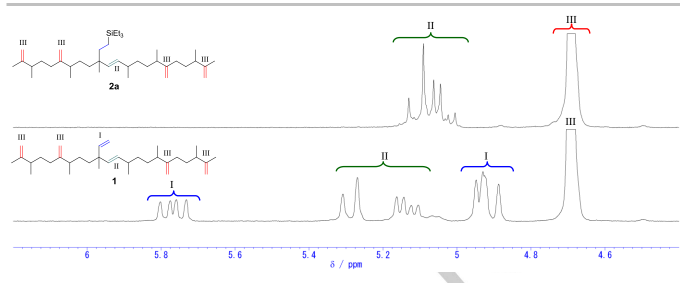


Figure 2. The ¹H NMR spectra of botryococcene 1 and the hydrosilylated product 2a.

Results and Discussion

First, to clarify the reactivity of 1 for hydrosilylation, 1 was reacted with Et₃SiH in the presence of various Pt-catalysts (Table 1). The hydrosilylation of 1 (0.2 mmol, 1 equiv.) with Et₃SiH (2 equiv.) was carried out in 2-propanol (1 mL) at 80 °C for 24 h in the presence of PtCl₂ (1 mol%) under an Ar atmosphere in a homogeneous system (entry 1). The products were analyzed by ¹H NMR. The isolated dominant product was a monohydrosilylated botryococcene derivative at the C10 vinyl moiety (2a) as shown in the reaction formula in Table 1, which was identified by characteristic signals (0.31-0.39 (multiplet (m), 2 H) ppm) for CH₂ adjacent to the inserted SiEt₃ moiety (see electronic supplementary information) in addition to the change of characteristic alkene signals of 1 and 2a in the area of 4.7-5.8 ppm. In the presence of $PtCl_2$, the conversion of 1 was 75% while 2a was dominantly produced in 8% yield. The yields and conversions were determined by ¹H NMR analyses using 1,1,2trichloroethane as an internal standard. The criteria ¹H NMR signals of 1,1,2-trichloroethane at 4.0 (doublet(d), 2 H) and 5.8 (triplet(t), 1 H) ppm did not overlap all the signals of 1 (0.9-2.2, 4.7-5.76 ppm) and 2a (0.3-0.55, 0.8-2.2, 4.7-5.2 ppm) (see electronic supplementary information).

Comparing the ¹H NMR spectra between **1** and the isolated **2a** in detail, it is revealed that the hydrosilylation of **1** with Et₃SiH proceeds with high site-selectivity. The ¹H NMR spectra of **1** and **2a** in the area of 4.4–6.2 ppm for the olefinic protons were shown in Figure 2. The ¹H NMR signals for the type-I vinyl moiety of **1** observed at 4.91 (d, 1 H), 4.94 (d, 1 H) and 5.77 (dd, 1 H) ppm completely dissapeared from the spectrum of **2a**. On the other hand, the signals for the type-II internal olefin of **2a** were observed in the region of 5.0–5.1 (d and dd, 2 H) ppm. The ¹H NMR signals of internal olefin were shifted from 5.1–5.3 ppm of **1** to 5.0–5.1 ppm of **2a**, because the protons of the internal olefin of **2a** were shielded by the C10-(triethylsilyl)ethyl moiety. It was suggested that the type-I vinyl is fully consumed by the hydrosilylation.

SiEt₃ cat Et₃Si-2a entry catalyst yield of conv. solvent 2a /%[b] of 1 /%^[b] 1 PtCl₂ 2-propanol 8 75 H₂PtCl₆/6H₂O 2 2-propanol 21 <1 3 Pt(NH₃)₄(NO₃)₂ 2-propanol 7 23 trans-Pt(NH₃)₂Cl₂ 2-propanol 43 >99 5 $[(C_6H_5)_3P]_2Pt(H_2C=CH_2)$ 2-propanol 53 89 6 Karstedt's catalyst 2-propanol 63 >99 7 Karstedt's catalyst toluene 89 92 THF 8 Karstedt's catalyst 91 >99 9 Karstedt's catalyst CICH₂CH₂CI 91 >99 10 Karstedt's catalyst none 91 >99

Table 1. Hydrosilylation of botryococcene 1 with Et₃SiH. [a]

[a] Conditions: [1] = 0.2 mol dm⁻³, [Et₃SiH] = 0.4 mol dm⁻³ (2 eq), [catalyst] = 0.002 mol dm⁻³ (1 mol%) , in solvent (1 mL) at 80 °C for 24 h under Ar. [b] The yields and conversions were determined by ¹H NMR.

The signals of the type-III four vinylidene moieties of **1** and **2a** were observed at 4.69 (singlet(s), 8 H) and did not change after the hydrosilylation. In addition, it has been known that the Pt catalyzed hydrosilylation of several branched olefins such as 2-hexene and 3-hexene frequently gives unordinary adducts through isomerization of the olefins to more reactive terminal ones such as 1-hexene.^[14] In the case of hydrosilylation of **1**, the

reaction site-selectively proceeded only at the vinyl moiety (type-I) without detection of isomerized products, although **1** has a branched structure involving the type-II and type-III carboncarbon double bonds in the molecule. The reason is because the internal *trans*-olefin and four vinylidene moieties that are inactive to the hydrosilylation do not isomerize into reactive terminal vinyl moieties. In summary, the appearance of CH₂ signals at 0.31–0.39 ppm after addition of Et₃SiH at the vinyl moiety (I), the change of the olefinic protons' signals between **1** and **2a**, and little detection of byproducts signals can elucidate that hydrosilylation of **1** proceeded with high site- and regioselectivity at the type-I vinyl moiety.

When H₂PtCl₆/6H₂O (Speier's catalyst), an effective hydrosilylation catalyst,^[9] was used instead of PtCl₂ (entry 2), the reaction was retarded, resulting in the lower conversion of 1 and little production of 2a under the same conditions. Similarly, Pt(NH₃)₄(NO₃)₂ was also not effective, showing low conversion of 1 (entry 3). The use of a divalent amine coordinated Pt catalyst, trans-Pt(NH₃)₂Cl₂,^[15] enhanced the reactivity of 1 in large in the conversion of 1 (>99%) and production of 2a (43% vield) (entry 4). More effective to increase vield of 2a was using Pt catalysts with π -ligands, such as $[(C_6H_5)_3P]_2Pt(H_2C=CH_2)^{[16]}$ and Karstedt's catalvst (Pt(0) 1.3-divinvl-1.1.3.3tetramethyldisiloxane complex)^[10,17,18] (entries 5 and 6). However, a variety of unidentified minor products were detected in all cases, which might be due to competitive formations against 2a and presence of impurities in 1.

The difference in reactivity of each catalyst observed in Table 1 is considered from the viewpoint of activity in each catalytic process based on the proposed mechanism.^[19,20] As shown in Figure 3, the Pt catalyst with ligands (Ln) (A) undergoes oxidative addition of hydrosilane (H-SiR₃), giving a reactive species (B) as usual, which must coordinate 1 rapidly to give species (C) for progress of the catalysis cycle. However, the formation step of C by coordinating 1 with B is a rate determining step due to large steric hindrance of 1 in the reactive type-I vinyl site vicinal to the quaternary C in 1. Consequently, conversion of 1 should fall to a low level when the ligand exchange velocity is slow, because the confined **B** should be destined for inactivation such as degradation to metal.^[19] Meanwhile, in order for hydrosilylation to proceed efficiently, at least the L in the species B must promptly be exchanged with botryococcene 1.

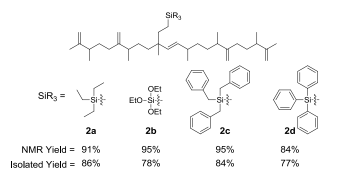
2 reductive elimination side-reaction H-[Pt]-SiR₃ C B deactivation

Figure 3. The supposed catalytic hydrosilylation cycle of 1.

In the case of PtCl₂ (entry 1), the coordination of the sterically hindered large molecule 1 to B proceeds smoothly, as it has no intended ligands. However, the octahedral complexes fully coordinated with valid ligands would not smoothly proceed the ligand exchange with 1, resulting in the considerable low conversions (entries 2 and 3). On the other hand, there is room for coordination of 1 for the square-planar complex, trans-Pt(NH₃)₂Cl₂, attaining the high conversion (entry 4). The catalysts having π -ligands are also proper, because the π ligands must be easily exchangeable with 1 (entry 5 and 6). In particular, the Karstedt's catalyst has a merit to stabilize B by the bidentate non-rigid π -ligand. After the ligand exchange of **B** with 1, reductive elimination of the species C should give 2, and this process is key to determine the reaction selectivity as well as production yield of 2. Activation and stabilization on C that is coordinated with 1 at the site of type-I vinyl must be important factors to give 2 in high yields with few byproducts, since several side reactions such as dehvdrogenative sillvlation and olefin hydrogenation, have been recognized.^[10] The Karstedt's catalyst worked most effectively in this step too (entry 6).

Next, solvent effect was examined to improve the production yield of **2a**. 2-Propanol which has been generally used in hydrosilylation^[9] was changed to another solvent under the same conditions of entry 6 using the Karstedt's catalyst. When toluene was used as the solvent, the yield increased but the conversion somewhat decreased (entry 7). The efficient production of **2a** in 91% yields and completion of conversions were attained in THF (entry 8), in 1,2-dichloroethane (entry 9), and under solvent-free conditions (entry 10). These results suggest that less donor environment than that in the isopropanol solvent is preferable to hold activity of **C** for progress of the selective hydrosilylation.

Lastly, the various organosilanes were applied to react with 1 under the best conditions of entry 9 in 1,2-dichloroethane, and the structures of hydrosilylated products 2 and their high production yields determined by ¹H NMR were shown in Figure 4. Triethoxysilane efficiently reacted with 1, and 2b was isolated in 78% yield. (PhCH₂)₃SiH and Ph₃SiH that have aromatic rings also site-selectively reacted well with 1, giving 2c and 2d in 84% and 77% isolated yields, respectively. Interestingly, 1 did not react with *i*Pr₃SiH at all. This suggests that the reactive vinyl site of 1 was difficult to access to the more sterically hindered *i*Pr₃Si linked B compared with the other R₃Si linked B.





Conclusions

It was found that botryococcene 1, an unsaturated hydrocarbon (C₃₄H₅₈) produced from Botryococcus braunii (Bot-22), could quantitatively be modified to a monohydrosilylated derivative 2. This is the first report that 1 was selectively converted to a botryococcene derivative as an isolable and dominant product. Difficulty against selective modifications of 1 is that 1 has three types of plural carbon-carbon double bonds, i.e., a vinyl (I), an internal trans-olefin (II), and four vinylidene moieties (III), in the molecule. Since hydrosilylation has usually been applied to terminal olefins to produce anti-Markovnikov addition alkanes in good yields, hydrosilylation of 1 with an organosilane was investigated in the presence of a Pt catalyst, for the first time. Unexpectedly, 1 showed low reactivity under general conditions for the hydrosilylation reactions. After optimization, the hydrosilvlation of 1 proceeded in high vields with high site- and regioselectivity at the reactive vinvl moiety (I) without isomerizations in the low-donor environment using the Karstedt's catalyst. It is demonstrated that various organosilanes could be applied to this reaction, which suggests that the terpenoid moiety of 1 can be embedded in materials via a silvlene linkage to add functionality such as bioactivity, lubricity, and cross-link activity. Moreover, these results provide the potential for biomass utilization of algal productions in chemical industry.

Supporting Information Summary

Experimental Section associated with this article can be found in electrical supporting information.

Acknowledgements

We gratefully acknowledge the Algae Biomass and Energy System R&D Center and especially Prof. M. Watanabe in University of Tsukuba, Japan for the generous supply of botryococcene **1**. We thank Chemical Analysis Division, Research Facility Center for Science and Technology, University of Tsukuba, for facilities of the NMR analysis. This paper is based on results obtained from a project (P16010) commissioned by the New Energy and Industrial Technology Development Organization (NEDO).

Conflict of Interest

The authors declare no conflict of interest.

Keywords: Algae biomass hydrocarbon oil • Biomass • Botryococcene • Hydrosilylation • Siteselective reaction

- [1] T. G. Tornabene, *Experientia* **1982**, *38*, 49-53.
- [2] For industrial applications of algae research, we (Algae Biomass and Energy System R&D Center in University of Tsukuba) are promoting to develop an algae-based renewable energy and material source production system.
- [3] a) Y. Tanabe, S. Kato, H. Matsuura, M. M. Watanabe, *Procedia Environmental Sciences* 15 2012, 22-26; b) A. Ishimatsu, H. Matsuura, T. Sano, K. Kaya, M. M. Watanabe, *Procedia Environmental Sciences* 15 2012, 56-65.
- [4] R. E. Cox, A. L. Burlingame, D. M. Wilson, G. Eglinton, J. R. Maxwell, J. Chem. Soc., Chem. Commun. 1973, 284-285.
- [5] A. P. Pulis, V. K. Aggarwal, J. Am. Chem. Soc. 2012, 134, 7570-7574.
- [6] J. D. White, G. R. Nagabushana, G. O. Spessard, J. Chem. Soc. Perkin Trans. 1 1993, 7, 759-767.
- J. D. White, G. R. Nagabushana, G. O. Spessard, J. Am. Chem. Soc. 1988, 110, 1624-1626.
- [8] a) H. Kawashima, M. Sakai, Y. Kaneko, Y. Kobayashi, *Tetrahedon* 2015, *71*, 2387-2392; b) H. Kawashima, Y. Kaneko, M. Sakai, Y. Kobayashi, *Chem. Eur. J.* 2014, *20*, 272-278.
- [9] J. L. Speier, J. A. Webster, G. H. Bernes, J. Am. Chem. Soc. 1957, 79, 974-979.
- [10] Y. Nakajima, S. Shimada, RSC Adv. 2015, 5, 20603-20616.
- [11] P. B. Kettler, Org. Proc. Res. Dev. 2003, 7, 342-354.
- [12] X. Du, Z. Huang, ACS Catal. 2017, 7, 1227-1243.
- [13] F. R. Hartley, Chem. Rev. 1969, 69, 799-844.
- [14] H. M. Bank, J. C. Saam, J. L. Speier, *ibid.* 1964, 29, 792-794.
- [15] A. Albinati, W. R. Caseri, P. S. Pregosin, *Organometallics* **1987**, *6*, 788-793.
- [16] B. Marciniec, J. Guliński, W. Urbaniak, T. Nowicka, J. Mirecki, Appl. Organometal. Chem. 1990, 4, 27-34.
- [17] T. K. Meister, K. Riener, P. Gigler, J. Stohrer, W. A. Herrmann, F. E. Kühn, ACS Catal. 2016, 6, 1274-1284.
- [18] T. Galeandro-Diamant, M.-L. Zanota, R. Sayah, L. Veyre, C. Nikitine, C. Bellefon, S. Marrot, V. Meille, C. Thieuleux, *Chem. Commun.* 2015, *51*, 16194-16196.
- [19] A. J. Chalk, J. F. Harrod, J. Am. Chem. Soc. 1965, 87, 16-21.
- [20] S. Sakaki, N. Mizoe, M. Sugimoto, Organometallics 1998, 17, 2510-2523.

Entry for the Table of Contents (Please choose one layout)

Layout 2:

FULL PAPER

Dr. Hidehisa Kawashima*, Mami Site- & regio-selective hydrosilylation SiR₃ Umezaw and Prof. Masashi Kijima* Page No. – Page No. -SiRa н Microalgal product, botryococcene Site-Selective Hydrosilylation of **Botryococcene - The Algal Biomass** Hydrocarbon oil An unsaturated hydrocarbon biomass (C34H58) produced from Botryococcus braunii (a freshwater microalga), could quantitatively be modified to a monohydrosilylated derivatives with high site- and regio selectivity.