

Unified Synthesis of C1–C19 Building Blocks of Halichondrins via Selective Activation/Coupling of Polyhalogenated Nucleophiles in (Ni)/Cr-Mediated Reactions

Jingwei Li, Wuming Yan, and Yoshito Kishi*

Department of Chemistry and Chemical Biology, Harvard University, Cambridge, Massachusetts 02138, United States

S Supporting Information

ABSTRACT: A unified synthesis of the C1–C19 building blocks 8–10 of halichondrins A-C was developed from the common synthetic intermediates 26a,b. Acetylenic ketones 26a,b were in turn synthesized via selective activation/coupling of polyhalogenated nucleophiles 23a,b with aldehyde 11 in a (Ni)/Cr-mediated coupling reaction. Compared with Ni/Cr-mediated couplings of vinyl iodides and aldehydes, this (Ni)/Cr-mediated coupling exhibited two unique features. First, the coupling was found to proceed with a trace amount or no added Nicatalyst. Second, TES-Cl, a dissociating agent to regenerate the Crcatalyst, was found to give a better yield than $Zr(Cp)_2Cl_2$. An adjustment of the oxidation state was required to transform acetylenic ketones 26a,b into C1-C19 building blocks 8 and 9 of halichondrins A and B, respectively. In the halichondrin B series, a hydroxyl-directed $(Me)_4NBH(OAc)_3$ reduction of *E*- and *Z*- β -alkoxy-enones **30** was found



cleanly to achieve the required transformation, whereas a DMDO oxidation of E-vinylogous ester 27 allowed to introduce the C13 hydroxyl group with a high stereoselectivity in the halichondrin A series. In the halichondrin C series, $Hf(OTf)_4$ was used to convert the double oxy-Michael product 28 into C1-C19 building block 10.

INTRODUCTION

Halichondrins are polyether macrolides, originally isolated from the marine sponge Halichondria okadai by Uemura, Hirata, and co-workers (Scheme 1).¹ Several additional members, including





halistatin 1, were isolated from various marine sponges.² This class of natural products displays interesting structure diversities at two sites, one being the oxidation state at C10, C12, and C13 of the C8-C14 polycycle and the other being the length of the carbon backbone. Thus, the halichondrin class of natural products is subgrouped into halichondrins A-C series or the norhalichondrin/halichondrin/homohalichondrin series. Due to their intriguing structural architecture and extraordinary in vitro and in vivo antitumor activity, the halichondrin class of marine natural products has received much attention from the scientific communities.^{3,4} We have

been engaged with the synthetic studies in this field since the late 1980s, aiming at total syntheses, coupled with development of a new synthetic strategy and discovery of new synthetic methods, with particular focus on the Cr-mediated coupling reactions.5-8

We recently reported a total synthesis of halichondrin A (1), a phantom member in this class of natural products.^{5c} Scheme 2 schematically illustrates the high convergence incorporated in the synthesis. There are two appealing aspects recognized in the synthesis. First, because of its high degree of convergence, one can expect a high overall efficiency in synthesis. Interestingly, the key two couplings have been achieved efficiently with Ni/ Cr-mediated coupling reactions. Second, with a replacement of 8 with 9 and 10, this convergent strategy can be extended to a synthesis of halichondrins B and C, respectively. Thus, we are interested in establishing a unified synthesis of C1-C19 building blocks 8, 9, and 10. In this paper, we report a solution to achieve this goal, with the use of a selective activation/ coupling of polyhalogenated nucleophile 23a,b in the (Ni)/Crmediated coupling reaction as the key C-C bond-forming step.

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Scheme 2. Summary of the Synthesis of the Right Half of Halichondrin A and Requisite C1–C19 Building Blocks (BBs) of Halichondrins A–C



RESULTS AND DISCUSSION

Synthetic Plan. We noticed the possibility that all of C1–C19 building blocks 8-10 could be synthesized from acetylenic ketone 15a,b, by adjusting its oxidation state (Scheme 3). Indeed, C1–C19 building block 8 of halichondrin A was synthesized with this strategy.^{Sc}

In the halichondrin A synthesis, we synthesized acetylenic ketone 15a via two (Ni)/Cr-mediated couplings. In light of the successful selective activation/coupling of polyhalogenated nucleophile 17 in a Ni/Cr-mediated coupling, we recognized a possibility of synthesizing 15a,b in one step, cf., $11 + 16a,b \rightarrow b$ 15a,b. In the halichondrin B series, a selective activation/ coupling of the *trans-\beta*-bromoenone was realized with use of a trace amount of a polyether-type Ni-catalyst.9 For the present case, because activation of halo-acetylenes is known to require only a trace amount of Ni-catalyst or even no added Ni-catalyst, a selective activation of the halo-acetylene over the vinyl iodide or bromide and saturated chloride present in 16a,b should not present an issue.^{10,11} Thus, the remaining question was the coupling efficiency of the nucleophile generated from 16a,b with an aldehyde, and we first addressed this issue with model compounds listed in Scheme 4.

Scheme 3. Proposed One-Step Synthesis of Acetylenic Ketones 15a,b via Selective Activation/Coupling of Halo-Acetylenic Ketones 16a,b



Scheme 4. Substrates, Sulfonamide, and Ni-Catalyst Used for the Model Study a



^{*a*}The coupling was tested with the molar ratio of 18/19:20 = 1.7:1; the coupling product was isolated by silica gel chromatography and yields are based on aldehyde 20.

Model Study on Coupling Efficiency. In order to assess the coupling efficiency, we chose six halo-acetylenes 18a-c and 19a-c and aldehyde 20 (Scheme 4). In this study, we first compared the coupling efficiency of halo-acetylenic ketones 18a-c over halo-acetylenic ketone ketals 19a-c¹² under the coupling conditions: 10 mol % Cr-catalyst, prepared from (S)sulfonamide 21, Ni-catalyst 22 (0.05 mol %), or no added Nicatalyst, Zr(Cp)₂Cl₂ (1.5 equiv), Mn (2 equiv), and LiCl (2 equiv) in MeCN ([C] 0.4M) at room temperature.^{13,14} This experiment demonstrated: (1) halo-acetylenic ketones 18a-c gave the desired product in only modest yields, with the order of coupling efficiency being 18b (49%) > 18c (20%) > 18a (11%); (2) halo-acetylenic ketone ketals **19a,b** gave the desired product in good yields, with the order of coupling efficiency being $19b (93\%) > 19a (82\%) \gg 19c (8\%); (3)$ no significant difference was detected between 0.05 mol % and no added Nicatalyst. In addition, a brief study on solvents and concentration revealed: (1) the solvent choice being EtCN > MeCN > DME

> THF, but not DMF, and (2) the optimum concentration being a range of 0.4-0.8 M.

Coupling in the Real Series. With the results gained in the model study, we began the study on the proposed coupling. The requisite nucleophiles **23a,b** were readily prepared from the previously reported, optically pure aldehydes **14a,b** (Scheme 5).^{15,16} With respect to the electrophile, several possible C8,C9-protecting groups were screened, thereby showing bis-TBS aldehyde **11** as the best option.¹⁷

Scheme 5. Synthesis of Nucleophiles 23a,b



For the simplicity of presentation, we will discuss first the coupling experiments in the nucleophile 23a series, although the studies were carried out on both 23a,b simultaneously.

In the proposed coupling, one chiral center is introduced at the C11 position. Based on the previous work, we were aware that a nucleophilic addition to an aldehyde such as **11** predominantly, often exclusively, gives the desired C11 β alcohol.^{7e,11a} With this background, we subjected **11** and **23a** to the coupling reaction under the condition used in the model study, to furnish the desired product **26a** in 55% yield, with a 10:1 stereoselectivity. The structure of **26a** was established via correlation with the authentic sample obtained in the previous route.^{5c} As anticipated, we did not detect a product derived through activation of the vinyl bromide or saturated chloride present in **23a**.

In order to improve the observed stereoselectivity, we adopted the toolbox approach and screened a representative set of sulfonamides (Table 1).^{7e,18} This screening showed that (1)

Table 1. Sufonamides Tested for a Ligand Search with a Toolbox Approach



as previously observed, the stereochemistry outcome was dictated by the substrate structure rather than the chirality present in the Cr-catalyst and (2) for this coupling, sulfonamides in the (R)-series gave a better stereoselectivity than the corresponding sulfonamides in the (S)-series. Among the tested ligands, we chose sulfonamide (R)-21 and Ni-catalyst 22 for the following study.

Naturally, we were anxious to improve the coupling efficiency. Particularly, we wondered how we could enhance the efficiency of Cr-catalytic cycle. In a catalytic Cr-mediated carbonyl addition, it is required to dissociate a Cr/product

alcohol complex and regenerate a Cr-catalyst. TMS-Cl or TES-Cl and $Zr(Cp)_2Cl_2$ are known to be effective dissociating agents.¹⁹ Generally, the overall coupling rate with $Zr(Cp)_2Cl_2$ is significantly faster than that with TMS-Cl or TES-Cl, at least for the Ni/Cr-mediated coupling of vinyl iodides with aldehydes. In addition, it is noteworthy that, when the TMS-Cl procedure is applied for a readily enolizable aldehyde, a significant amount of aldehyde is recovered, due to a silyl enol ether formation *in situ*.

For the present case, it was found that the coupling rate with TES-Cl was slower than that with $Zr(Cp)_2Cl_2$, yet the coupling yield with TES-Cl was noticeably better than that with $Zr(Cp)_2Cl_2$, i.e., 85% with TES-Cl vs 70% with $Zr(Cp)_2Cl_2$.²⁰ Although its mechanistic reason was not clear, the TES-Cl condition made it possible to achieve the proposed coupling with the synthetically useful efficiency.

As noted before, Cr-mediated coupling of a halo-acetylene with an aldehyde is known to proceed with only a trace amount of Ni-catalyst or even no added Ni-catalyst.¹¹ For this reason, we studied the coupling of $11 + 23a \rightarrow 26a$ "with" and "without" added Ni-catalyst, thereby showing the coupling efficiency to be comparable (Scheme 6). There is no definite



^{*a*}Coupling conditions: Cr-catalyst prepared from (*R*)-21: 20 mol %; Ni-catalyst (0.05 mol %) or no added Ni-catalyst; TES-Cl (2.5 equiv); LiCl (4 equiv); Mn (4 equiv); EtCN ([*C*] 0.4 M); RT. The coupling was done with the molar ratio of 18/19:20 = 1.7:1; the coupling product was isolated by silica gel chromatography and yields are based on aldehyde 11.

experimental evidence to conclude whether this coupling involves activation of bromoacetylene with Ni-catalyst, followed by Cr-mediated coupling, or activation/coupling with only a Cr-catalyst. In this connection, it is worthwhile mentioning that the homodimer of bromoacetylene was isolated in ca. 0.3% yield (based on 23a) in the coupling without added Ni-catalyst. Formation of the dimer with no added Ni-catalyst may support that Ni-catalyst, or some unknown metal, activated haloacetylene 23a prior to the C–C bond formation.²¹ However, reflecting this ambiguity, we refer to the coupling as (Ni)/Crmediated reaction in this paper.

As mentioned, we carried out the coupling studies with both 23a,b simultaneously and obtained the virtually identical results in the both series, although a small reduction in yield was noticed in the 23b series.

Synthesis of C1–C19 Building Blocks of Halichondrins A–C from the Common Synthetic Intermediate 26. Synthesis of C1–C19 Building Block of Halichondrin A. In the halichondrin A synthesis, we already established the transformation of 26a into C1–C19 building block 8 (Scheme 7).





The key reactions in this transformation included: (1) a selective TBS-deprotection to form *E*-enone **27** and (2) a highly stereoselective DMDO-oxidation to introduce the C13 hydroxyl group. It should be noted that C1–C19 building block **8** bears the C19 vinyl bromide, because the corresponding vinyl iodide was not compatible with the DMDO oxidation. 5c,22

Synthesis of C1–C19 Building Block of Halichondrin C. In the halichondrin C synthesis, we reported a synthetic route to construct the polycyclic ring system from an acetylenic ketone.^{Sb} Although the transformation was carried out on the macrolactone framework, we felt confident in extending this synthetic method to the present acyclic system; indeed, we had no unexpected difficulty in the transformation of **26b** to **10** in 60% overall yield (Scheme 7). The key reactions in this transformation included: (1) double oxy-Michael addition of C8,C9-hydroxyl groups to the acetylenic ketone to form ketal **28** and (2) Hf(OTf)₄-induced conversion of the double oxy-Michael product **28** to polycycle **10** in allyl alcohol. The structure **10** was fully supported by the spectroscopic data (HR-MS, ¹H and ¹³C NMR).

Synthesis of C1–C19 Building Block of Halichondrin B. In order to synthesize C1–C19 building block 9 in the halichondrin B series from the common synthetic intermediate, we obviously needed an acetylene-to-olefin reduction and tested first the reactivity of **26b** and its C11-OTBS derivative against CuH, HN=NH, and CrCl₂ (Scheme 8), thereby indicating that the C11-OTBS substrate exhibited a very poor reactivity. Based on this observation, we used **26b** for a search of a satisfactory reducing reagent/condition. Among reagents tested, (BDP)CuH, a Stryker CuH modified by Lipshutz, gave a most promising result (Scheme 8).²³ Yet, we had two issues to address. First, this reduction gave a mixture of *E*- and *Z*-enones. As discussed in the preceding companion paper (DOI: 10.1021/jacs.Sb03498),⁹ *Z*-enone was found to form readily Scheme 8. Synthesis of C1–C19 Building Blocks (BBs) in the Halichondrin B Series



the furan.⁹ Thus, although it was a minor product, *Z*-enone was wasted. Second, this reduction gave the desired *E*-enone **30** as the major product, but the isolated yield varied from 55% up to 80%. Apparently, the problem was over-reduction. Despite extensive efforts, we were unable to identify a condition to achieve the reduction with a high reproducibility and high yield.

Under this circumstance, we decided to focus on reduction of *E*-enone *E*-**30**. Once again, we attempted several known methods, including various CuH reagents, H_2 /Crabtree catalyst, and Na₂S₂O₄, but with only limited success.²⁴ Ultimately, we found that (Me)₄NBH(OAc)₃ reduced the vinylogous ester cleanly to give **31** in 80% yield as a 5:1 mixture of 12 α :12 β diastereomers.

(Me)₄NBH(OAc)₃ is well recognized as an excellent hydride donor in a so-called hydroxyl-directing setting.²⁵ However, a literature search revealed that no example was reported for reduction of a vinylogous ester with (Me)₄NBH(OAc)₃. Nonetheless, we would assume that, like the case of β -hydroxyl ketones,²⁶ the reduction was facilitated via a ligand exchange of (Me)₄NBH(OAc)₃ with the C11-hydroxyl group, followed by an intramolecular hydride delivery in a conjugated fashion, to yield the 12 α -alcohol as the major product. Consistent with this suggestion, we found that the substrate with the C11-OH masked with a TBS was inert to the reduction.

We anticipated that the intramolecular hydride delivery should yield predominantly 12α -stereoisomer, which is the undesired stereoisomer. However, we were aware that it should not be an issue, because the C12-configuration is prone to isomerization via a retro oxy-Michael/oxy-Michael process; indeed, the ratio of 12α : 12β stereoisomers varied from one experiment to other, likely due to a different degree of isomerization during workup.

As expected from the above consideration, we observed that $(Me)_4NBH(OAc)_3$ reduction of the corresponding Z-enone Z-30 gave 31 as a mixture of 12α : 12β stereoisomers. Thus, for the preparative purpose, it was not necessary to separate E- and Z-enones **30**.

On TBAF treatment, **31** furnished **32** as a ~1:1 mixture of $12\alpha:12\beta$ diastereomers. With an ion-exchange resin-based device,²⁷ this mixture was transformed cleanly to C1–C19 building block **9** of halichondrin B without isolation/separation/equilibration of intermediates. On comparison of spectroscopic data (¹H and ¹³C NMR, MS, TLC), **9** thus obtained was found to be superimposable on the authentic sample.⁹

CONCLUSION

A unified synthesis of the C1–C19 building blocks 8–10 of halichondrins A–C was developed from the common synthetic intermediates 26a,b. Acetylenic ketones 26a,b were in turn synthesized via selective activation/coupling of polyhalogenated nucleophiles 23a,b with aldehyde 11 in a (Ni)/Cr-mediated coupling reaction. Compared with Ni/Cr-mediated couplings of vinyl iodides and aldehydes, this (Ni)/Cr-mediated coupling exhibited two unique features. First, the coupling was found to proceed with a trace amount or no added Ni-catalyst. Second, TES-Cl, a dissociating agent to regenerate the Cr-catalyst, was found to give a better yield than Zr(Cp)₂Cl₂.

An adjustment of the oxidation state was required to transform acetylenic ketones **26a,b** into C1–C19 building blocks **8** and **9** of halichondrins A and B, respectively. In the halichondrin B series, a hydroxyl-directed $(Me)_4NBH(OAc)_3$ reduction of *E*- and *Z*- β -alkoxy-enones **30** was found cleanly to achieve the required transformation, whereas a DMDO oxidation of *E*-vinylogous ester **27** allowed to introduce the C13 hydroxyl group with a high stereoselectivity in the halichondrin A series. In the halichondrin C series, Hf(OTf)₄ was used to convert double oxy-Michael product **28** into C1–C19 building block **10**.

Lastly, we note that an application of the synthetic method developed in the halichondrin A synthesis^{5c} should allow us to transform C1–C19 building blocks 8-10 into halichondrins A–C, respectively.

ASSOCIATED CONTENT

S Supporting Information

Experimental procedures, characterization data, and copies of spectra data. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/ jacs.Sb03499.

■ AUTHOR INFORMATION

Corresponding Author

*kishi@chemistry.harvard.edu

Notes

The authors declare no competing financial interest.

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(13) The standard coupling conditions used in the toolbox approach; see ref 7e.

(14) Three Ni-complexes were screened for this coupling, thereby showing that the order of effectiveness was $22 \sim i > ii$. Based on this observation, 22 was chosen for this study.

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(17) C8,C9-Protecting groups tested included: cyclohexylidene ketal, anisylidene acetal, methylene acetal, bis-MOM ether, diacetate, dibenzoate, carbonate, and others.

(18) We originally used sulfonamide iii for ligand search in the toolbox approach (see ref 7e). In order to expand the space of ligand search, we have recently added sulfonamides iv and v to this approach; see the structure highlighted by a green box. Sulfonamides (S)- and (R)-24 and 25 were sulfonamides used in that study.



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