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### Asymmetric total synthesis of (+)-xestoquinone and (+)-adociaquinones A and B†

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The asymmetric total synthesis of (+)-xestoquinone and (+)-adociaquinones A and B was achieved in 6–7 steps using an easily accessible meso-cyclohexadienone derivative. The [6,6]-bicyclic decalin B–C ring and the all-carbon quaternary stereocenter at C-6 were prepared via a desymmetric intramolecular Michael reaction with up to 97% ee. The naphthalene diol D–E ring was constructed through a sequence of Ti(Oi-Pr)4-promoted photoenolization/Diels–Alder, dehydration, and aromatization reactions. This asymmetric strategy provides a scalable route to prepare target molecules and their derivatives for further biological studies.

Various halenaquinone-type natural products with promising biological activity have been isolated from marine sponges of the genus *Xestospongia*<sup>1</sup> from the Pacific Ocean.  $(+)$ -Halenaquinone  $(1)$ ,<sup>2,3</sup> (+)-xestoquinone  $(2)$ , and  $(+)$ -adociaquinones A  $(3)$ and B  $(4)^{4,5}$  bearing a naphtha $[1,8{\text -}bc]$ furan core (Fig. 1) are the most typical representatives of this family. Naturally occurring  $(-)$ -xestosaprol N (5) and O  $(6)^{6,7}$  have the same structure as 3 and 4 except for a furan ring, while a naphtha[1,8-bc]furan core can also be found in fungus-isolated furanosteroids  $(-)$ -viridin (7) and (+)-nodulisporiviridin E  $(8)^{8,9}$  (Fig. 1). Halenaquinone  $(1)$ was first isolated from the tropical marine sponge Xestospongia  $exigua<sup>2</sup>$  and it shows antibiotic activity against *Staphylococcus* aureus and Bacillus subtilis. Xestoquinone (2) and adociaquinones A  $(3)$  and B  $(4)$  were firstly isolated, respectively, from the Okinawan marine sponge Xestospongia sp.<sup>4a</sup> and the Truk Lagoon sponge Adocia sp.,<sup>4b</sup> and they show cardiotonic,<sup>4a,c</sup> cytotoxic,<sup>4b,i</sup> antifungal,<sup>4*i*</sup> antimalarial,<sup>4*j*</sup> and antitumor<sup>4*l*</sup> activities. These compounds inhibit the activity of pp60v-src protein tyrosine kinase,<sup>4d</sup> topoisomerases I<sup>4e</sup> and II,<sup>4f</sup> myosin Ca<sup>2+</sup> ATPase,<sup>4c,g</sup> and phosphatases Cdc25B, MKP-1, and MKP-3.<sup>4h,k</sup> **EDGE ARTICLE**<br> **(a)** Check for updates<br> **Asymmetric total synthesis of (+)-xestoquinone a**<br> **Asymmetric total synthesis of (+)-xestoquinone a**<br> **(a)** Check for updates<br> **Consistence**<br> **Consistence**<br> **Consistence**<br> **Consi** 

Owing to their diverse bioactivities, the synthesis of this family of natural compounds has been extensively studied, with published pathways making use of Diels-Alder,<sup>3a,d,e,5a-c,e,g</sup> furan

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ring transfer,<sup>5b</sup> Heck,<sup>3b,c,5f,7,9b,d</sup> palladium-catalyzed polyene cyclization,<sup>5d</sup> Pd-catalyzed oxidative cyclization, $\frac{3}{3}$  and hydrogen atom transfer (HAT) radical cyclization<sup>9c</sup> reactions. In this study, we report the asymmetric total synthesis of (+)-xestoquinone (2),  $(-)$ -xestoquinone  $(2')$ , and  $(+)$ -adociaquinones A  $(3)$  and B  $(4)$ (Fig. 1).

The construction of the fused tetracyclic B–C–D–E skeleton and the all carbon quaternary stereocenter at C-6 is a major challenge towards the total synthesis of xestoquinone (2) and adociaquinones A (3) and B (4). Based on our retrosynthetic analysis (Scheme 1), the all-carbon quaternary carbon center at C-6 of *cis*-decalin 12 could first be prepared stereoselectively from the achiral aldehyde 13 via an organocatalytic



Fig. 1 Structure of halenaquinone-type natural products and viridintype furanosteroids.

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Scheme 1 Retrosynthetic analysis of (+)-xestoquinone and (+)-adociaquinones A and B.

desymmetric intramolecular Michael reaction.<sup>10,11</sup> The tetracyclic framework 10 could then be formed via a Ti(Oi-Pr)<sub>4</sub>promoted photoenolization/Diels–Alder (PEDA) reaction<sup>12-16</sup> of

11 and enone 12. Acid-mediated cyclization of 10 followed by oxidation state adjustment could be subsequently applied to form the furan ring A of xestoquinone (2). Finally, based on the biosynthetic pathway of (+)-xestoquinone (2) <sup>4</sup>b,5<sup>c</sup> and our previous studies, $7$  the heterocyclic ring F of adociaquinones A (3) and B (4) could be prepared from 2  $via$  a late-stage cyclization with hypotaurine (9).

The catalytic enantioselective desymmetrization of meso compounds has been used as a powerful strategy to generate enantioenriched molecules bearing all-carbon quaternary stereocenters.<sup>10,11</sup> For instance, two types of asymmetric intramolecular Michael reactions were developed using a cysteinederived chiral amine as an organocatalyst by Hayashi and coworkers,<sup>11a,b</sup> while a desymmetrizing secondary amine-catalyzed asymmetric intramolecular Michael addition was later reported by Gaunt and co-workers to produce enantioenriched decalin structures.<sup>11c</sup> Prompted by these pioneering studies and following the suggested retrosynthetic pathway (Scheme 1), we first screened conditions for organocatalytic desymmetric intramolecular Michael addition of meso-cyclohexadienone 13 (Table 1) in order to form the desired quaternary stereocenter at C-6. Compound 13 was easily prepared on a gram scale via a four-step process (see details in the ESI†).

We initially investigated the desymmetric intramolecular Michael addition of 13 using  $(S)$ -Hayashi-Jørgensen catalysts,<sup>17</sup>

Table 1 Attempts of organocatalytic desymmetric intramolecular Michael addition<sup>4</sup>





 $^a$  All reactions were performed using 13 (5.8 mg, 0.03 mmol, 1.0 equiv., and 0.1 M) and a catalyst at room temperature in analytical-grade solvents, unless otherwise noted.  $^b$  The yields and diastereoisomeric ratios (d.r.) were determined from the crude  $^1$ H NMR spectrum of  $\bf{14}$  using CH $_2$ Br $_2$  as an internal standard, unless otherwise noted. The enantiomeric excess (e.e.) values were determined by chiral high-performance liquid chromatography (Chiralpak IG-H).  $^d$  Compound 13: 9.6 mg, 0.05 mmol, and 0.1 M.  $^e$  Isolated combined yield of 14a + 14b.  $^f$  Compound 13: 192 mg, 1.0 mmol, and 0.1 M. <sup>g</sup> The d.r. values decreased to 1:1 after purification by silica gel column chromatography. <sup>h</sup> Compound 13: 1.31 g, 6.82 mmol, and 0.1 M. *i* Isolated combined yield of  $12a + 12b$ . *j* The d.r. values were determined from the crude <sup>1</sup>H NMR spectrum of 12 obtained from the one-pot process.



Scheme 2 PEDA reaction of 11 and enone 12

and found that the absolute configuration of the obtained cisdecalin was opposite to the required stereochemistry of the natural products (see Table S1 in the ESI†). In order to achieve

the desired absolute configuration of the angular methyl group at C-6,  $(R)$ -cat.I was used for further screening. In the presence of this catalyst, the intramolecular Michael addition afforded 14a (96% e.e.) and 14b (75% e.e.) in a ratio of 10.3 : 1 and 52% combined yield (entry 1, Table 1). We assumed that the enantioselectivity of the reaction was controlled by the more sterically hindered aromatic group of  $(R)$ -cat.I, which protected the upper enamine face and allowed an *endo*-like attack by the siface of cyclohexadienone, as shown in the transition state TS-A (Table 1). In order to increase the yield of this reaction and improve the enantioselectivity of 14b, we further screened solvents and additives. Increasing the catalyst loading from 0.5 to 1.0 equivalents and screening various reaction solvents did not improve the enantiomeric excess of 14b (entries 2–7, Table 1). Therefore, based on previous studies, $11d,e$  we added 5.0 equivalents of acetic acid (AcOH) to a solution of compound 13 and  $(R)$ -cat.I in toluene, which improved the enantiomeric excess of 14b to 95% with a 60% combined yield (entry 8, Table 1). And, the stability of  $(R)$ -cat.I has also been verified in the presence of AcOH (see Table S2 in the ESI†). Further adjustment of the (R)-cat.I and AcOH amount and ratio (entries 9–12, Table 1) indicated that 0.2 equivalents each of  $(R)$ -cat.I and AcOH were the best conditions to achieve high enantioselectivity for both 14a and 14b, and it also increased the reaction yield (entry 12, Table 1). The enantioselectivity was not affected when the optimized reaction was performed on a gram scale: 14a (97% e.e.) and 14b (91% e.e.) were obtained in 80% isolated yield (entry 12, Table 1). We also found that the gram-scale experiments needed a longer reaction time which led a slight decrease of the diastereoselectivity. The purification of the cyclized products by silica gel flash column chromatography indicated that the major product 14a was epimerized and slowly converted to the minor product 14b (entry 11, Table 1). Both 14a Edge Article<br>
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Scheme 3 Total synthesis of (+)-xestoquinone and (+)-adociaquinones A and B.

and 14b are useful in the syntheses because the stereogenic center at C-2 will be converted to  $sp<sup>2</sup>$  hybridized carbon in the following transformations. Therefore, the aldehyde group of analogues 14a and 14b was directly protected with 1,3-propanediol to give the respective enones 12a and 12b for use in the subsequent PEDA reaction.

Afterward, we selected the major cyclized cis-decalins 12a and  $12a'$  (obtained by using  $(S)$ -cat.I in desymmetric intramolecular Michael addition, see Table S1 in the ESI†) as the dienophiles to prepare the tetracyclic naphthalene framework 10 through a sequence of  $Ti(Oi-Pr)_4$ -promoted PEDA, dehydration, and aromatization reactions (Scheme 2). When using 3,6 dimethoxy-2-methylbenzaldehyde (11) as the precursor of diene, no reaction occurred between 12a/12a' and 11 under UV irradiation at 366 nm in the absence of  $Ti(Oi-Pr)_{4}$  (Scheme 2A). In contrast, the 1,2-dihydronaphthalene compounds 16a and 16a' were successfully synthesized when 3.0 equivalents of  $Ti(Oi-Pr)<sub>4</sub>$  were used. Based on our previous studies,<sup>13a,e</sup> the desired hydroanthracenol 15a was probably generated through the chelated intermediate TS-B and the cycloaddition occurred through an endo direction (Scheme 2B).<sup>18</sup> The newly formed  $\beta$ hydroxyl ketone groups in 15a and 15a' could then be dehydrated with excess Ti $\left( \mathrm{Oi\text{-}Pr} \right)_{4}$  to form enones **16a** and **16a** $^{\prime}$ . These results confirmed the pivotal role of  $Ti(Oi-Pr)_4$  in this PEDA reaction: it stabilized the photoenolized hydroxy-o-quinodimethanes and controlled the diastereoselectivity of the reaction. Open Access Article. Published on 19 February 2021. Downloaded on 12/27/2024 11:39:42 AM. This article is licensed under a [Creative Commons Attribution-NonCommercial 3.0 Unported Licence.](http://creativecommons.org/licenses/by-nc/3.0/) **[View Article Online](https://doi.org/10.1039/d0sc07089k)**

Subsequent aromatization of compounds 16a and 16a' with 2,3-dichloro-5,6-dicyanobenzoquinone (DDQ) at 80 °C afforded compounds  $10a$  and  $10a'$  bearing a fused tetracyclic B–C–D–E skeleton. The stereochemistry and absolute configuration of 10a were confirmed by X-ray diffraction analysis of single crystals (Scheme 3). The synthesis of (+)-xestoquinone (2) and (+)-adociaquinones A (3) and B (4) was completed by forming the furan A ring. Compound 10 was oxidized using bubbling oxygen gas in the presence of t-BuOK to give the unstable diosphenol 17a, which was used without purification in the next step. The subsequent acid-promoted deprotection of the acetal group led to the formation of an aldehyde group, which reacted in situ with enol to furnish the pentacyclic compound 18 bearing the furan A ring. The stereochemistry and absolute configuration of 18 were confirmed by X-ray diffraction analysis of single crystals (Scheme 3). Further oxidation of 18 with ceric ammonium nitrate afforded (+)-xestoquinone (2) in 82% yield. Following the same reaction process,  $(-)$ -xestoquinone  $(2')$  was also synthesized from 10a' in order to determine in the future whether xestoquinone enantiomers differ in biological activity. Further heating of a solution of (+)-xestoquinone (2) with hypotaurine (9) at 50 °C afforded a mixture of  $(+)$ -adociaquinones A (3) (21% yield) and B (4) (63% yield). We also tried to optimize the selectivity of this condensation by tuning the reaction temperature and pH of reaction mixtures (see Table S3 in the ESI†). The  $^1\mathrm{H}$  and  $^{13}\mathrm{C}$  NMR spectra, high-resolution mass spectrum, and optical rotation of synthetic (+)-xestoquinone (2),  $(+)$ -adociaquinones A (3) and B (4) were consistent with those data reported by Nakamura,<sup>4a,g</sup> Laurent,<sup>4j</sup> Schmitz,<sup>4b</sup> Harada<sup>5a,c</sup> and Keay.<sup>5d</sup>

#### Conclusions

In summary, we developed a concise approach for the asymmetric total synthesis of (+)-xestoquinone (2) in 6 steps and of (+)-adociaquinones A (3) and B (4) in 7 steps from a known compound 13. Organocatalytic desymmetric intramolecular Michael addition was used to construct the cis-decalin skeleton bearing the all-carbon quaternary carbon center at the C-6 position. The B–C–D–E tetracyclic framework was then prepared through a  $Ti(Oi-Pr)_4$ -mediated PEDA reaction, and further modifications led to the desired naphtha $[1,8-bc]$ furan core. The application of this strategy to the synthesis of structurally related natural products is currently under investigation.

#### Conflicts of interest

There are no conflicts to declare.

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- 18 Based on our previous studies (ref. 13a and  $e$ ), we concluded that  $Ti(Oi-Pr)_4$  plays a key role in the PEDA reaction, which chelated with the photo-generated Z-dienol and ortho methoxy group of 11, forming a relatively stable complex. This complex may exist as a dimeric titanium complex, given the observed relationships between the Ti dosage and reaction yield (ref. 13a). The ortho methoxy may serve as a key neighbouring group that helps to stabilize the short-lived photoenolized hydroxy-o-quinodimethane diene, which then interacts with dienophile 12a to give a chelated intermediate TS-B. Then the activated enone reacts with the diene component from the endo direction. After dissociation, cycloaddition product 15a was generated stereospecifically.