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## Total Synthesis of (+)-Cavicularin: The Pyrone Diels-Alder Reaction in Enantioselective Cyclophane Synthesis

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We dedicate this paper to Prof. Kenneth W. Hedberg on the occasion of his 95th birthday.

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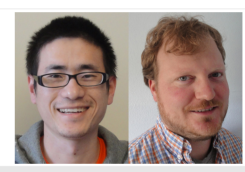
Abstract A pyrone Diels-Alder strategy was developed for the synthesis of the cyclophane natural product, cavicularin. The strategy uses a vinyl sulfone as an alkyne equivalent dienophile. An enantioselective variant delivered (+)-cavicularin.

**Key words** Diels-Alder, pyrone, cyclophane, chirality, total synthesis

Recently, we have been investigating natural products that display conformational chirality.1 In particular, we became interested in families of cyclophane natural products in which the presence of molecular chirality could not be deduced by inspection of the molecular structure. In the course of these studies we were attracted to the macrocyclic bis(bibenzyl) natural products.<sup>2</sup> These natural products, which have the general structure shown in Figure 1, contain two bibenzyl moieties that are joined by either biphenyl or diphenyl ether linkages. The linkages of the bibenzyl units can result in para-, meta-, or ortho-substituted phenyl rings (cf. isoplagiochin A and marchantin A).

Cavicularin possesses a strained molecular architecture containing a dihydrophenanthrene.3 Crystallographic studies indicated that the A-ring of cavicularin was distorted from planarity. The molecule was isolated as an optically active substance, which indicated that it was a chiral nonracemic molecule in Nature. The beautiful molecular structure has attracted the attention of several groups and, other than our work,4 there have been three syntheses of racemic cavicularin<sup>5</sup> and one synthesis of (-)-cavicularin by using a chiral auxiliary approach.6

We desired a strategy for preparing cavicularin that satisfied the following three criteria: (1) The strategy could assemble cyclophane architectures with considerable ring



Peng Zhao was born in Zhengzhou, P. R. of China. After receiving his B.S. in chemistry from Xiamen University (P. R. of China), he went to Zhengzhou University and obtained a M.S. in organic chemistry with Prof. Junbiao Chang. Currently, he is a Ph.D. candidate in the chemistry department of Oregon State University working with Prof. Beaudry on the synthesis of macrocyclic bis(bibenzyl) natural products. Christopher M. Beaudry joined the department of chemistry at Oregon State University in 2009 as an Assistant Professor. He received his B.S. degree from the University of Wisconsin in 2000. As an undergraduate, he conducted research with Prof. Steven D. Burke. He obtained his Ph.D. at the University of California, Berkeley under the direction of Prof. Dirk Trauner. He was an NIH postdoctoral fellow in the laboratory of Prof. Larry Overman at the University of California, Irvine.

strain; (2) It could be applicable to other macrocyclic bis(bibenzyl) natural products that contain different connectivity in the A-ring (i.e., meta- or para-substitution or diphenyl ether and biphenyl linkages); (3) It would deliver (+)-cavicularin in nonracemic form.

In 2006, Baran and Burns reported an elegant strategy for the formation of haouamine A, which contains a bent aromatic ring, by using a pyrone-alkyne Diels-Alder reaction (Scheme 1).7 The strategy benefits from the fact that the bent arene is prepared during the macrocyclization by way of a bicyclic intermediate that is a conformational

mimic of the final strained arene. This precedent suggested that such a process could be used to prepare a strained molecule such as cavicularin.

Figure 1 Selected macrocyclic bis(bibenzyl) natural products

Consideration of the pyrone–alkyne Diels–Alder reaction in the context of (+)-cavicularin, initially led to terminal alkyne 1. In the forward sense, bond formation should occur between the nucleophilic C6 of the pyrone and the terminal C5 of the alkyne to give intermediate 2. Retro-Diels–Alder reaction would deliver the desired connectivity in 3. However, this synthetic strategy was not without potential pitfalls. Simple hand-held models revealed that both regiochemical outcomes of the Diels–Alder cycloaddition could be possible;<sup>8</sup> the constraints of the molecular tether would not guarantee bond formation between C6 and C5 in 1, and the regiochemical alternative (Scheme 2, bottom) would deliver 4 and lead to a *meta*-substituted A-ring (5). Moreover, although the pyrone–alkyne Diels–Alder reac-

tion can be well-behaved, it commonly requires high temperatures (>200 °C) and often gives modest yields, potentially making the reaction unsuitable for use as a general strategy for other macrocyclic bis(bibenzyl) family members

With the above concerns in mind, we considered using vinyl sulfone **6** as an alkyne-equivalent dienophile (Scheme 3). The vinyl sulfone dienophile is more electronically polarized than a terminal alkyne, and would likely result in a high regiochemical preference for **7** in the Diels-Alder event. Phenylsulfinic acid could be eliminated under the conditions of the Diels-Alder reaction to give **2**, followed by loss of CO<sub>2</sub> to give the cavicularin architecture **3**.

Vinyl sulfone regioisomer **8** could also be prepared, which would give the opposite regiochemical outcome in the Diels–Alder reaction (**9**), allowing for control of the connectivity in the macrocyclization. Intermediate **9** would un-

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dergo elimination to give *meta*-substituted **5**. This regiocontrolled strategy may also be applicable to other macrocyclic bis(bibenzyl) natural products with both *meta*- and *para*-substituted A-rings. Moreover, because the vinyl sulfone was more electrophilic than an alkyne, we anticipated that the reaction would occur at lower temperatures (and potentially with higher yields) relative to the alkyne substrate.

Enantioselective Diels–Alder reactions of pyrone dienes have been explored. These reactions form oxabicyclo[2.2.2]octanones with high to excellent enantioselectivities, and generally fall into three types: (1) Normal-electron-demand Diels–Alder reactions of 3-hydroxy-2-pyrones and electron-deficient alkenes promoted by cinchona alkaloid or amino-indanol-derived catalysts (Scheme 4, eq. 1); <sup>10,11</sup> (2) Inverse-electron-demand Diels–Alder reactions of 3-acyl-2-pyrones and electron-rich alkenes (eq. 2); <sup>12</sup> (3) Normal-electron-

$$\begin{array}{c} \text{OO} \\ \text{OO} \\ \text{HO} \\ \text{SO}_2\text{Ph} \\ \end{array} \begin{array}{c} \text{Chiral catalyst} \\ \text{PhO}_2\text{S} \\ \text{PhO}_2\text{S} \\ \end{array} \begin{array}{c} \text{OH} \\ \text{-PhS(O)OH} \\ \text{-CO}_2 \\ \end{array}$$

**Scheme 4** Enantioselective pyrone Diels–Alder reactions

demand Diels–Alder reactions of 2*H*-pyran-2,5-diones with electron-deficient alkenes catalyzed by a cinchona-based thiourea (eq. 3).<sup>13</sup>

Our retrosynthetic strategy uses a vinyl sulfone, which precluded the inverse-electron-demand Diels-Alder reaction. The substitution pattern of cavicularin more closely matched the substitution present in the Diels-Alder reactions exemplified by eq. 1, and our overall plan was to pursue the enantioselective (+)-cavicularin synthesis by using an application of this reaction in which the initial cycloaddition would give a nonracemic oxabicyclo[2.2.2]octanone, which would undergo sequential eliminations of phenylsulfinic acid and CO<sub>2</sub> to deliver the chiral cyclophane structure in nonracemic form (eq. 4).

The first objective was to develop an efficient synthesis of the achiral Diels–Alder substrate. The synthesis began with dibromostyrene **10**.<sup>14</sup> It was envisioned that **10** could serve directly in regioselective Suzuki reactions without the need for protecting groups to differentiate the bromine atoms.

In the event, we were delighted to observe complete regioselectivity in the Suzuki reaction of dibromide **10** with **11** (Scheme 5). We believe the regioselectivity can be attributed to the alkene: bromides with proximal vinyl groups undergo oxidative addition at a reduced rate. Related reactions of dibrominated heteroaromatic systems have been reported. However, such Suzuki reactions of dibromobenzenes are quite rare; Indeed, we believe this is the first example of using an alkene to control the site selectivity in a dibromobenzene Suzuki reaction.

CO<sub>2</sub>t-Bu

(3)

Although Suzuki product 12 could be isolated, it was more convenient to perform the subsequent Suzuki reaction with 13 in situ. When TLC indicated consumption of 10, boronic ester 13 was added and the reaction proceeded to completion. A single regioisomer of terphenyl 14 was isolated. Two additional transformations gave 15 and three transformations produced 6. By using a related synthetic strategy, the second vinyl sulfone isomer 8 and alkyne 1 were prepared.

PhO<sub>2</sub>S

B(pin)

Grubbs II

EtOAc, 60 °C

83%

CH<sub>2</sub>Cl<sub>2</sub> 2. H<sub>2</sub>, Pd/C

11

D

MeO

В

 $\Box$ 

15

В

12 (not isolated)

PhO<sub>2</sub>S

PhO<sub>2</sub>S

С

D

Br

D

ÓMe

MeO

В

Pd(Ph<sub>3</sub>P)<sub>4</sub>, KBr

K<sub>3</sub>PO<sub>4</sub>, H<sub>2</sub>O dioxane, 55 °C

В

Scheme 5 Suzuki reactions for cavicularin

PhO<sub>2</sub>S

**Scheme 6** Synthesis of (±)-cavicularin

We began our Diels–Alder reaction investigations by using pyrone substrate **6** (Scheme 6). We found that the cycloaddition of **6** occurred with complete regiochemistry and in good chemical yield to give **3**. This reaction could be performed by using microwave irradiation (240 °C, 8 h) or by using a standard heating mantle at 170 °C for 6 days. Presumably, the cycloaddition delivers bicyclic intermediate **7**, which undergoes sequential elimination of phenylsulfinic acid and  $CO_2$  as outlined in Scheme 3. The order of the elimination is inconsequential, and no intermediates were observed or isolated from the reaction. Deprotection of **3** gave cavicularin.

then add:

С

63% (one flask)

3 steps

13

B(pin)

MeO

When isomeric vinyl sulfone **8** was subjected to microwave irradiation, Diels–Alder cascade product **5** was isolated in good yield. Again, the regioselectivity was high and only a single isomer of **5** was obtained. This indicates that the regiochemistry of the Diels–Alder cycloaddition can be controlled by the electronics of the substrate, and it is not strictly a result of constraints of the intramolecular tether.

Alkyne **1** also underwent the Diels–Alder cascade. The reaction occurred at 250 °C under microwave irradiation (no reaction occurred at temperatures of 200 °C using a heating mantle); the reaction requires more forcing conditions because the alkyne is less activated than vinyl sulfone **6**. The reaction was unselective, leading to a mixture of both regioisomeric Diels–Alder products **5** and **3**.

With access to synthetic samples of cavicularin, our attention turned to performing an enantioselective synthesis. The enantiomers of cavicularin cleanly separated on chiral HPLC (Diacel OD-H) and we resolved approximately 5 mg of each enantiomer. Preliminary experiments indicated that cavicularin is stable as a single enantiomer up to 150 °C,

Hydroxypyrone substrate  $16^{14}$  was then subjected to the cinchona-based alkaloid quinidine in EtOAc by following the report of Deng (Scheme 7). We were delighted to observe reaction at the milder temperature of  $100\,^{\circ}$ C. However, the reaction produced the undesired regioisomer 17. Addition of the hydroxyl group rendered the C3 position of the pyrone nucleophilic, and bond formation occurred at C5 of the vinyl sulfone to give 18. Elimination of phenylsulfinic acid and  $CO_2$  gave 17. As above, no intermediates were observed in this reaction. Interestingly, the reaction was modestly enantioselective, but no effort was made to improve this aspect of the reaction.

Isomeric vinyl sulfone **20** was prepared.<sup>14</sup> Mild heating of **20** in the presence of cinchona-based thiourea **21** gave **22** with good enantioselectivity (er = 89:11). The resulting phenol was a sensitive intermediate that was prone to decomposition, so this intermediate was immediately converted into the corresponding triflate **23**. The yield of the overall process was good (45% from **20**). Removal of the triflate and methyl groups gave (+)-cavicularin.

In conclusion, a vinyl sulfone–pyrone Diels–Alder strategy was developed for the synthesis of cavicularin. The substitution of the vinyl sulfone allows for control of the regiochemistry in the reaction. A hydroxypyrone substrate also participated in an enantioselective Diels–Alder reaction using a chiral cinchona-based thiourea. The regiochemistry of the reaction could again be controlled by the choice of the vinyl sulfone substitution pattern.

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Scheme 7 Synthesis of (+)-cavicularin

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