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# ARTICLE TYPE

## **Using Singlet Oxygen to Synthesise the CDE-Ring System of the Pectenotoxins**

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**A non-classical route to the key CDE-ring fragment of the pectenotoxins has been developed which showcases a remarkable singlet oxygen-mediated cascade reaction sequence to install the complete DE ring system.**

- The pectenotoxin (PTX, Figure 1) family of natural products has attracted a great deal of attention from different groups of researchers in recent years, not only because of their biological activity which has been shown to possess certain new and exciting features, but also by reason of the structural challenges 15 to synthesis that they exhibit. Originally isolated<sup>1a</sup> from the host shellfish (a scallop named, *Patinopecten yessoensis*), the pectenotoxins are actually produced by dinoflagellates found
- worldwide in coastal waters. <sup>1</sup> However, the scarcity of samples available from natural sources has hindered the pursuit of further 20 studies into the PTXs' potentially useful cytotoxicity<sup>2</sup> (achieved via a novel mode of F-actin disruption<sup>2b</sup>). Thus, finding effective





Figure 1 The pectenotoxin family of natural products.

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In general, synthetic strategies towards polyoxygenated targets frequently rely heavily on cumbersome redox shuttling (manipulation and differentiation of the oxygen functionalities by <sup>30</sup> repeated back and forth alterations to their oxidation states often

using ungreen reagents) and on the use of an excessive number



**Scheme 1** Proposed synthetic plan.

<sup>35</sup> of unconstructive oxygen functionality protections and deprotections.<sup>3</sup> The primary focus of our research of late<sup>4</sup> is to develop new singlet oxygen-based methods for the synthesis of just such polyoxygenated motifs, which, by virtue of the ability of singlet oxygen to orchestrate cascade reaction sequences and its <sup>40</sup> inherent selectivity, help us to avoid many of these common pitfalls. Herein, we present one such effort in the form of a nonclassical route affording the key CDE-ring<sup>5</sup> fragment of the PTXs. When combined with our earlier succesful singlet oxygenmediated one-pot "super-cascade" which synthesised the ABC- $45$  ring section,<sup>6</sup> this serves to illustrate the full synthetic potential of this powerful and green oxidant.<sup>4a</sup>

PTXs 4 and 8 are the only members of the family which have been successfully synthesised<sup>7</sup> in the laboratory; however, many efforts have been reported that are directed towards synthesis of 50 specific fragments of these molecules.<sup>6,8</sup> Our current CDE-ring work joins the recent elegant efforts of Pihko,<sup>9</sup> Brimble,<sup>10</sup> and Micalizio<sup>11</sup> who have published their own investigations (following on from earlier work undertaken by the Roush  $group<sup>12</sup>)$  directed towards PTX fragments including these <sup>55</sup> particular rings. Furthermore, it should be noted that the groups of both Paquette<sup>13</sup> and Fujiwara<sup>14</sup> have completed PTX sections which include the necessary but, as yet, uncyclised backbone of the DE-bicycle section.

Our own synthetic blueprint (Scheme 1) was based on the idea <sup>60</sup> that a simple furanyl butanone unit could be united readily with an intact pre-synthesised C-ring fragment (using an adol condensation) and then methylated in such a way as to leave hydroxyl functionalities at the γ- and ε-positions of the alkyl side chain appended to the 2-position of the furan **B** (Scheme 1).

<sup>65</sup> It was hoped that this newly acquired homochiral substrate would then succumb to an ambitious singlet oxygen-initiated reaction



**Scheme 2** Synthesis of C-ring fragment.

cascade sequence (in which these two pendant hydroxyl functionalities would participate) that would in a rapid and <sup>5</sup> extremely efficient manner construct the requisite DE-ring motif, and, thus, afford the completed CDE-ring section of the pectenotoxins (**A**, Scheme 1). We had previously investigated such a cascade for accessing simple 2,8 dioxabicyclo[3.2.1]octane model compounds and had recorded 10 successes both in organic solvents and in water.<sup>15</sup>

To explore the practicality of this proposal we begun by developing a C-ring synthesis in which we chose to implement fairly obvious disconnections in the forward synthetic sense. Thus, it was anticipated that a 5-*exo*-epoxide opening (Scheme 1)

- <sup>15</sup> of an appropriately substituted epoxide by a hydroxyl group would be the easiest means by which to construct the desired tetrahydrofuran ring, and, before this cyclisation step, the requisite vicinal diol unit could be generated using a Sharpless asymmetric dihydroxylation reaction.
- <sup>20</sup> To this end, the commercially available and cheap homoallylic alcohol **1** was oxidised using IBX and a solution of the resultant volatile aldehyde (not isolated) was treated directly with the stabilised ylide Ph3PCHCO2Bn to afford diene **2** (*trans*:*cis* isomers  $= 9:1$ ) in an overall yield for the two steps of 83%
- <sup>25</sup> (Scheme 2). Chemoselective epoxidation of diene **2** with *m*-CPBA in buffered (NaHCO<sup>3</sup> ) DCM, afforded epoxide **3** in 94% yield. AD-mix-β was then used to install the requisite vicinal diol unit at the site of the more electron deficient double bond  $(3 \rightarrow 4,$ 62% with 16% recovered starting material). The benzoate ester

<sup>30</sup> had been chosen for inclusion in the substrate (from the earlier



**Scheme 3** Synthesis of photooxidation substrate **10**.

Wittig reaction) to improve the difficult isolation of the diol product from this aqueous dihydroxylation reaction; indeed, the <sup>35</sup> analogous (to **4**) methyl ester had been synthesised, but could only be isolated in low yields, and, furthermore, its quantities were further depleted on purification and manipulation due to its polarity and poor solubility in organic solvents. Catalytic acid (PPTS, 10 mol%) was then used to effect the desired cyclisation <sup>40</sup> reaction to afford the now separable diastereoisomers **5a** and **5b** (**5a**:**5b**, 1:1) in 88% overall yield. The relative stereochemistry of the two cyclised compounds was deconvoluted using NOE studies of selected derivatives. More specifically, both diastereoisomers were advanced separately by, firstly, <sup>45</sup> undertaking a double protection of alcohol residues as TBS ethers (yield from **5a** 93%, and from **5b**, 46%), then by DIBALHmediated reduction of the ester to the primary alcohol (yielding either **6a** or **6b**, in yields of 82 and 89%, respectively). The reason for the low yield obtained for the double protection of **5b** <sup>50</sup> is not immediately clear, but was anyway of little consequence because **6b** was needed only for NOE studies and not for progression of the synthesis. With a key area within the <sup>1</sup>HNMR now simplified and extra helpful protons also now introduced, extensive NOEs studies were undertaken on both these <sup>55</sup> compounds (**6a** and **6b**) revealing that the relative stereochemistry was that which is shown in Scheme 2. In addition, the primary alcohol of the desired diastereoisomer **5a** was selectively protected as the TBS ether using standard conditions (yield 63%), the remaining secondary alcohol was <sup>60</sup> then converted into each of the two MPA esters separately and the resulting NMR spectra analysed. This confirmed that the expected absolute stereochemistry had been obtained from the Sharpless asymmetric dihydroxylation (for full details, see: Supporting Information).

<sup>65</sup> With our C-ring stereochemistry now fully analysed, the synthesis was advanced by oxidation of the primary alcohol of **6a** using IBX (98%, Scheme 3). The resultant aldehyde **7** was then used as the electrophile in an aldol reaction with the kinetic enolate of furanyl butanone **8**. This aldol reaction proceeded in <sup>70</sup> 70% yield and afforded the desired isomer as the major product  $(dr 10:1)<sup>16</sup>$  The conditions recently developed by Pihko and coworkers<sup>9a</sup> were then employed to effect the last remaining transformation prior to the key cascade reaction sequence,



**Scheme 4** Key photooxidation cascade reaction.

namely, a stereoselective methylation reaction. The reaction yielded photooxidation substrate **10**, as the major <sup>5</sup> diastereoisomer, in a yield of 65% (the minor diastereoisomer accounted for a further 17%). 16

The stage was now set for implementation of the crucial cascade reaction sequence (Scheme 4). Thus, photooxidation substrate **10** was subjected to a set of singlet oxygen reaction <sup>10</sup> conditions that have been honed in our laboratory. These include bubbling oxygen gently through the reaction solution containing small quantities  $(10^{-4}$  M) of a photosensitiser, in this case, methylene blue, whilst irradiating the cooled solution (ice bath) with a Variac Eimac Cermax 300W lamp. Following the

- <sup>15</sup> completion of the initial oxidation steps, as monitored by tlc, an excess of dimethyl sulfide was introduced into the reaction vessel to effect the reduction of the newly formed peroxy residue. Finally, and, once again when tlc monitoring indicated the reduction was complete, trace acid (*p*-TsOH) was added to
- <sup>20</sup> complete the cascade reaction sequence (Scheme 4). Gratifyingly, each of these stages proceeded exceptionally smoothly and the final product, the completed pectenotoxin CDE-ring unit **11**, was cleanly formed in an extraordinary yield of 82%, as a single stereoisomer. Extensive NOE experiments, as shown in Scheme
- <sup>25</sup> 4, confirmed the stereochemistry of the final cyclised compound **11**. As expected, a set of NOEs similar to those observed for **6a** were shown for the C-ring of **11**. More importantly, the existence of NOEs between  $H_6$  and both  $H_9$  and  $H_{10}$  proves the stereochemistry of the DE-bicyclic ketal in the final compound
- <sup>30</sup> **11**. In addition, this remarkable cascade reaction sequence could also be adapted so that it could be conducted in water (in this case, using rose bengal as sensitiser), albeit with a slightly reduced yield of 74%, adding to its already excellent green credentials.
- <sup>35</sup> The mechanistic rationale for this highly complex cascade reaction sequence is given in Scheme 5. Thus, initial [4+2] cycloaddition of singlet oxygen to the furan fleetingly yields ozonide **C**, which is rapidly opened by the pendant internal nucleophile (a hydroxyl group) to afford the [5,5]-spirocyclic
- $\mu_0$  hydroperoxide  $\mathbf{D}^{(4,6,15,17)}$  All the characteristic peaks  $\mathbf{D}^{4,6,15,17}$ corresponding to intermediate **D** were observed in crude <sup>1</sup>H NMR



spectra of the photooxidation mixture, prior to the addition of <sup>45</sup> dimethyl sulfide. Reduction of the hydroperoxy residue of **D** to hemiketal **E** is followed by acid catalysed rearrangement of the spirocycle and *cis* – *trans* isomerisation of the double bond, to give the final product **11** via intermediate **F**.

To summarise, an ambitious singlet oxygen-mediated cascade <sup>50</sup> reaction sequence has been successfully implemented in a "real" system example, giving us rapid access to the complex and complete CDE-ring fragment of the pectenotoxins. The cascade reaction sequence itself showcases a remarkable increase in molecular complexity through an easily implemented laboratory <sup>55</sup> protocol and uses a green and highly selective oxidant, thereby

making it a powerful solution to the instransigent problem of how to significantly improve our efficiency when we seek to synthesise complex polyoxygenated polycyclic molecules.

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- † Electronic Supplementary Information (ESI) available: [Experimental 70 procedures, full spectroscopic data and copies of <sup>1</sup>H and <sup>13</sup>C NMR spectra for all new compunds, as wellas copies of the NOE experiments]. See DOI: 10.1039/b0000000x/
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