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Proof-of-Principle Direct Double Cyclisation of a Linear C₁₅-Precursor to a Dibrominated Bicyclic Medium-Ring Ether Relevant to Laurencia **Species**[†]

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Bicyclic dibrominated C₁₅ medium-ring ether hexahydrolaureoxanyne was produced directly from an acyclic model C₁₅-epoxide when treated with NBS with water as the solvent.

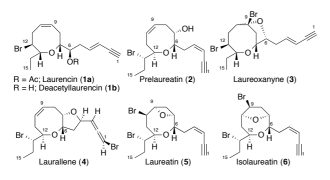
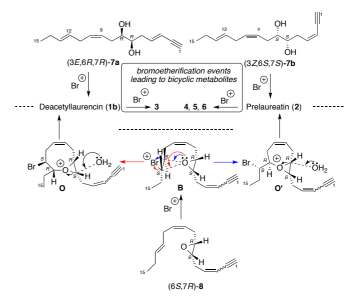


Figure 1. Representative monocyclic and bicyclic halogenated medium-ring ethers of formulae $C_{15}H_{21}BrO_2$ (1b, 2) and $C_{15}H_{20}Br_2O_2$ (3-6) from Laurencia species that are oxygenated at C-6 and C-7. Laurencin 1a is related as the acetate of 1b.

Since the original isolation of Laurencin (1a) in 1965,¹ marine red algae of Laurencia species have provided a wide variety of C15-acetogenic halogenated diastereo- and constitutional isomeric monocyclic (C15H21BrO2) and bicyclic (C15H20Br2O2) mediumring ethers that are oxygenated at both C-6 and C-7 (Figure 1).² Both the monocyclic and bicyclic metabolites have received considerable synthetic attention, with numerous necessarily different strategies used to forge the 7-, 8-, or 9-membered medium-ring, control the *cis* or *trans* α, α' -ether stereochemistry, install the requisite halogen(s), and – in the case of the bicyclic ethers – to fashion the second ring.³⁻⁵ Various recent studies have also been directed at the further understanding of their biogenesis,6 where the early pioneering work of Murai demonstrated enzymatic bromoetherifications of straight-chain co-isolated unsaturated C_{15} -diols – laurediols (3E,6R,7R)-7a and (3Z, 6S, 7S)-7b⁸ – to monocyclic medium-ring ethers deacetyl laurencin 1b and prelaureatin 2 respectively, albeit in very low yields (Scheme 1, top).⁹ We have recently advanced an alternative biogenesis for the monocyclic (C15H21BrO2) mediumring ethers from Laurencia species from (6S,7R)-epoxide 8 via an intramolecular bromonium ion assisted epoxide ring-opening (IBIAERO) reaction with water functioning as the external

[†] Electronic Supplementary Information (ESI) available: Experimental procedures, characterising data and ¹H and ¹³C NMR spectra for all compounds; a comparison of ¹H NMR data for (\pm) -[H₆]-3 with laureoxanyne.

nucleophile (Scheme 1, bottom, $8 \rightarrow B \rightarrow O/O' \rightarrow 1b/2$), and experimentally corroborated this with a model epoxide for the concurrent formation of 7-, 8- and 9-ring ethers corresponding to the halogenated medium-ring ethers of known metabolites from Laurencia species.^{10,11} The *bicyclic* metabolites are generally considered to originate by further bromoetherification of the residual unsaturation of the monocyclic compounds - the Zconfigured medium-ring alkene or the pendant envne – using the free alcohol of the original monocyclic compound located either at C-6 or C-7 as the nucleophile (Scheme 1, top).⁷ Several laboratory demonstrations of these later transformations have successful. either as enzymatic-mediated been bromoetherifications of naturally occurring monocycles,¹² or as part of the synthetic strategy in a total synthesis of the bicyclic natural products.¹³ Interestingly, although bromocyclisation events had been postulated for both monocycle and bicycle formation, prior to our 2012 report¹⁰ and Snyder's recent elegant work,6b,c a non-enzymatic bromonium-ion induced cyclisation process to directly form medium-ring ether cores relevant to Laurencia species had not been reported. Moreover, to the best of our knowledge, there has been no report of a C₁₅-dibrominated bicyclic medium-ring ether relevant to Laurencia species being formed directly from a linear unsaturated C₁₅-precursor by two successive bromination events in the same pot. Herein we report on a successful strategy to effect such a transformation.



Scheme 1. Irie-Murai biogenesis of monocyclic medium-ring ethers from laurediols 7a and 7b (top); alternative biogenesis of deacetyllaurencin 1b and prelaureatin 2 via IBIAERO reaction with water functioning as the nm Accepted Manuscrip

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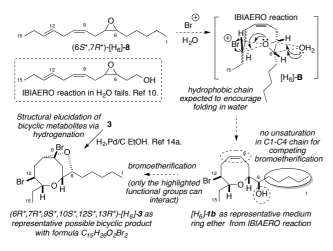
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external nucleophile (bottom). The other six possible monocyclic ethers of formulae $C_{15}H_{21}BrO_2$ are not shown.

To investigate the proof-of-principle demonstration of a direct double cyclisation of a C₁₅ unsaturated linear precursor to a bicyclic medium-ring ether relevant to Laurencia species we targeted hexahydroepoxide $(6S^*, 7R^*)$ -[H₆]-8, with the aim that this would undergo an initial IBIAERO reaction via [H₆]-B where water functions as both the solvent and the nucleophile (Scheme 2). The use of water in this manner thus guarantees a free hydroxyl group for any subsequent bromoetherification reaction (e.g., $[H_6]$ -1b \rightarrow [H₆]-3, Scheme 2) with a second equivalent of an electrophilic bromine source. While we had previously demonstrated successful IBIAERO reactions in water with NBS as the electrophilic bromine source,¹¹ the attempted IBIAERO reaction of a model epoxide as a truncated C₁₂ alcohol (inset, Scheme 2) under the same conditions had failed.^{10,§} We considered that hexahydroepoxide [H₆]-8 offered distinct benefits compared to this earlier model and also to epoxide 8 for the proposed experiment: (i) the hydrophilic hexahydro chain may encourage folding of the substrate in water thus inherently facilitating the IBIAREO reaction; (ii) post-IBIAERO reaction, the only region of unsaturation will be located in the medium ring and - compared with the hypothetical use of the putative biosynthetic precursor itself, epoxide 8 - there can be no complicating bromoetherifications to form bromoallene adducts cyclisation onto any C₁-C₄ enyne moiety; (iii) by hexahydrobicyclic compounds of formulae C15H26O2Br2 are known in the literature as a consequence of the structural elucidation of the naturally occurring compounds via hydrogenation,14 providing data for identification of bicyclic products.

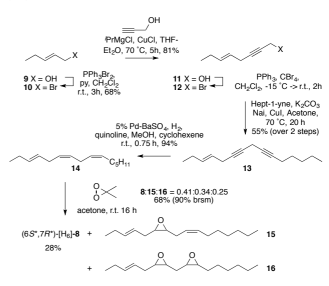


Scheme 2. Proposed proof-of-principle direct cyclisation of $(6S^*, 7R^*)$ - $[H_6]$ -**8** to bicyclic medium ring ethers via IBIAERO reaction and subsequent bromoetherification of the remaining unsaturation.

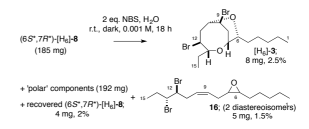
Accordingly, epoxide $(6S^*, 7R^*)$ - $[H_6]$ -8 was synthesised from bromide 12, itself prepared from (*E*)-2-penten-1-ol (9) via a known sequence^{15,10} with minor modifications. Subsequent copper-mediated coupling¹⁶ with hept-1-yne gave novel enediyne 13 (Scheme 3).[‡] Chemoselective and stereoselective hydrogenation¹⁷ afforded (*E*,*Z*,*Z*)-doubly skipped triene 14. Epoxidation of triene 14 with DMDO¹⁸ was found to be entirely selective for the *Z*-olefins,¹⁹ giving a mixture of mono epoxides (6*S**,7*R**)-[H₆]-8 and 15 which could be separated by chromatography. ¶.¥,††

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With epoxide $(6S^*, 7R^*)$ -[H₆]-8 in hand, it was treated with two equivalents of NBS - a water stable reagent - under high dilution conditions in water (Scheme 4).^{‡‡} Here, various dibromination adducts, bromohydrin regioisomers, and dibromotetrahydrofurans are expected to be formed by competing processes.¹⁰ In the event, as expected, a complex mixture was obtained that was subjected to extensive chromatography, where 'non-polar' components could be separated away from 'polar' components.88 Much to our delight, by further chromatography of the non-polar components, hexahydrolaureoxanyne $[(\pm)-[H_6]-3]^{12a}$ was isolated as a bicyclic medium-ring ether with ¹H NMR data identical to that previously reported,^{‡,¶¶} along with dibromoepoxides 16. Thus the desired proof-of-principle has been achieved. This also constitutes the first synthetic route to the laureoxanyne bicyclic medium-ring ether scaffold, and the isolated yield of (\pm) -[H₆]-3 (2.5%) from (6S*,7R*)-[H₆]-8 compares well with the reported enzymatic conversion of deacetyl laurencin 1b (obtained from natural laurencin 1a) into 3 (3%).^{12a}



Scheme 3. Synthesis of (6*S**,7*R**)-[H₆]-8.



Scheme 4. Proof-of-principle direct double cyclisation of $(6S^*, 7R^*)$ -[H₆]-**8** into (±)-[H₆]-**3** via IBIAERO reaction and subsequent bromoetherification of the remaining unsaturation (*c.f.*, Scheme 2).

In conclusion, we have demonstrated the proof-of-principle direct cyclisation of a linear unsaturated C₁₅-precursor into a C₁₅-dibrominated bicyclic medium-ring ether relevant to *Laurencia* species – where hexahydrolaureoxanyne (\pm)-[H₆]-**3** has an identical bicyclic medium ring ether framework to laureoxanyne **3** – by two successive bromination events in the same pot. These studies are also consistent with epoxide (6*S*,7*R*)-**8** acting as the biogenetic precursor¹⁰ for bromocyclisation to bicyclic medium-ring ethers of *Laurencia* species via IBIAERO reactions followed by subsequent bromoetherification events.

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Notes and references

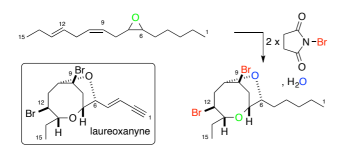
- Control to the terminal termin
- $\$ We speculate that the truncated C_{12} epoxide suffers from an intramolecular hydrogen bond from the alcohol functional group reducing its nucleophilicity.
- ¶ 25% of a bis-epoxide was also observed.
- ¥ Attempted epoxidation of 14 with *m*CPBA was unselective for the *Z*-olefins.
- ^{††} ¹H-¹³C and ¹H-¹H NMR correlation spectroscopy were used to distinguish between epoxides ($6S^*$, $7R^*$)-[H₆]-8 and 15.[‡]
- **‡** In an experiment with 1 equivalent of NBS in water, (\pm) -[H₆]-**3** was isolated in 1.8% yield after extensive chromatography.
- §§ The 'polar' components were expected to contain regioisomeric bromohydrins and dibromohydrins by reference to our earlier work (ref. 10) and were not further characterised.
- **¶** The medium-ring bicyclic structure of $[H_6]$ -**3** is also supported by a characteristic NOESY cross-peak between H_7 and H_9 as previously reported (as an nOe) for **3** (ref. 12a).[‡]
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GRAPHICAL ABSTRACT



Bicyclic medium ring ethers of relevance to *Laurencia* species have been obtained by direct double brominative cyclisation of an acyclic precursor.