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Epoxidation of bromoallenes connects red algae metabolites by an intersecting bromoallene oxide – Favorskii manifold†

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DMDO epoxidation of bromoallenes gives directly α , β -unsaturated carboxylic acids under the reaction conditions. Calculated (@B97XD/ 6-311G(d,p)/SCRF = acetone) potential energy surfaces and ²H- and ¹³C-labeling experiments are consistent with bromoallene oxide intermediates which spontaneously rearrange via a bromocyclopropanone in an intersecting bromoallene oxide – Favorskii manifold.

The remarkably wide structural diversity and complexity of halogenated C₁₅ acetogenin metabolites isolated from marine red algae of *Laurencia* species¹ continue to stimulate innovative efforts in their target synthesis, 2 in the discovery of new synthetic transformations³ and in advancing biosynthetic hypotheses.⁴ A recent re-isolation 5 of obtusallene IV $\left(1\right)^6$ from *Laurencia marilzae* provided also 12-epoxyobtusallene IV (2) and unnamed α , β unsaturated carboxylate ester (3) with an identical macrocycle to epoxybromoallene 2 (Fig. 1). It seems reasonable to connect E-alkene 1 and trans-epoxide 2 biogenetically via enzymatic epoxidation, $⁷$ and on the basis of their co-isolation, we propose</sup> **EDOXIDATION**
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Fig. 1 Metabolites 1–3 from Laurencia marilzae and proposed biogenesis via epoxidation events.

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† Electronic supplementary information (ESI) available: Notes §, ¶, f, ††, **, ‡‡, §§, ¶¶, ff, †††, ***, ‡‡‡; general experimental; experimental details and characterising data for compounds leading to bromoallenes 4 (including ESI Scheme S1 for the synthesis of bromoallenes 4), $(1^{-2}H)-4$ and $(1^{-13}C-4)$ and epoxidation thereof leading to E - and Z-5, $(E$ -2⁻²H)- and $(Z$ -2⁻²H)-5, and $(E$ -2⁻¹³C)and (Z-2-¹³C)-5; Copies of ¹H and ¹³C spectra for all compounds showing ²H and ¹³C isotopic shifts and coupling constants where appropriate; ESI references. See DOI: 10.1039/c3cc46720a

that bromoallene 2 and α , β -unsaturated carboxylate 3 may also be connected biogenetically by epoxidation.

While the epoxidation of allenes^{8,9} and vinyl bromides¹⁰ has been studied, the epoxidation of bromoallenes has not been reported.¹¹ Herein, we report the hitherto unknown direct conversion of bromoallenes to α , β -unsaturated carboxylic acids via an initial epoxidation event and the presumed intermediacy of a bromoallene oxide. We also show by computational modeling and 2 H- and 13 C-labeling studies that the latter's spontaneous reorganization to an α , β -unsaturated carboxylic acid under the reaction conditions is consistent with a bromocyclopropanone intermediate in an intersecting allene oxide – Favorskii manifold.

Bromoallene 4^{12} was selected as a suitable substrate for investigating epoxidation and was synthesized by a standard sequence from heptanal (ESI[†]).¹³ Much to our delight, epoxidation of bromoallene 4 using dimethyl dioxirane (DMDO), generated either in $situ^{14}$ or as a solution (ESI[†])¹⁵ (Scheme 1), gave a mixture of Z and E - α , β -unsaturated carboxylic acids 5 directly in low but reproducible yields (note §, ESI†). The low yields can be attributed to decomposition of DMDO^{16a} under the reaction conditions to methyl radicals, $16b$ and subsequent radical attack on either of the products or starting materials (note ¶, ESI†).

Mechanistically, we invoke the following pathway for the formation of α , β -unsaturated carboxylic acids from DMDO mediated epoxidation of bromoallenes (Fig. 2). Initial epoxidation of the bromoallene would give bromoallene oxides of the type A and/or B (note \overline{f} , ESI[†]). Spontaneous epoxide opening^{8c} via bromo oxyallyl cations C and D (note $\dagger\dagger$, ESI \dagger) respectively converge on the same bromocyclopropanone E. This intermediate now intersects with the Favorskii rearrangement manifold of α, α - and α, α' -dibromoketones where the resulting bromocyclopropanones E are known to collapse after attack by water giving hydrate \bf{F} to α , β -unsaturated carboxylic acids 5 (note **, ESI \dagger).^{17,18} Evidently, there is sufficient water in the dioxirane solution to function as a nucleophile here (note ‡‡, ESI†).

Scheme 1 Epoxidation of bromoallene 4 using DMDO solution.

Fig. 2 Mechanistic rationale for conversion of bromoallenes into α . B-unsaturated carboxylic acids, with the carbon atoms of the functional groups numbered 1–3 showing an interchange of carbon atoms 1 and 2 (see also interactive Fig. 2 in HTML version of this article).

Interestingly, regardless of the initial site of epoxidation, this mechanism predicts that carbon atoms 1 and 2 in bromoallene 4 interchange positions in the α , β -unsaturated carboxylic acid products 5.

This mechanism can be subjected to scrutiny via density functional level $(\omega B97XD/6-311G(d,p)/SCRF = \text{acetone})^{19}$ exploration of the potential energy surface $(R = H, Me,$ presented as an interactive version of Fig. 2 (ref. 20) via a digital data repository²¹). Oxygen transfer from dimethyldioxirane to form both A and B (TS1) have thermally accessible free energy activation barriers AG_{298}^{\dagger} (R = H, 26.8 for **A**, 27.3 for **B**; R = Me, 26.8 for **A**, 24.6 kcal mol⁻¹ for **B**), followed by a second, lower energy dyotropic rearrangement (TS2) to give E. An intrinsic reaction coordinate (IRC) reveals that TS2 $(R = H, Me)$ represents the concerted transformation of **A** or **B** to E, with C/D acting as "hidden intermediates" in the process.²² Such hidden intermediates can be potentially transformed to real ones by tuning the substituents, and in this instance changing R from H or Me to OMe is predicted to accomplish this by stabilization of C/D (see interactive Fig. 2). TS2 itself $(R = Me)$ has some early character of C/D; the C–Br bond is calculated to initially contract in length due to a significant stabilising resonance contribution of Br lone pairs, from 1.924/1.896 Å (A and B respectively) via 1.840/1.885 (TS2), 1.856/1.868 (C/D acting as hidden intermediates) to 1.921/1.922 Å (E).23 Calculations having demonstrated the thermal accessibility of the epoxidation-bromocyclopropanone sequence, ²H- and ¹³C-labeling experiments were necessary to verify the overall reorganization (4 to A/B to E to F to 5, Fig. 2) of the carbon framework.²⁴ Communication
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Deuterated bromoallene $(1^{-2}H)$ -4 was prepared by addition of ethynylmagnesium bromide to heptanal, in situ deprotonation of the propargylic alkoxide with n -butyllithium and quenching with

Scheme 2 Synthesis of deuterated bromoallene $(1 - 2H) - 4$.

MeOH- d_4 to give labeled propargylic alcohol $(1^{-2}H)$ -6 (Scheme 2). Subsequent alcohol trisylation²⁵ gave (1 - ^{2}H)-7, and S_{N2} ['] displacement of the trisylate with bromide under the action of LiCuBr₂ (ref. 26) provided bromoallene (1-2 H)-4 with 70% deuterium incorporation at the 1-position.†

 $13C$ -labeled bromoallene $(1^{-13}C)$ -4 was similarly targeted, commencing with silyl enol ether 8 formation²⁷ from octanal (Scheme 3). Oxidation using $mCPBA$ gave interrupted Rubottom²⁸ adduct 9, which could be acetylated to give acetate 10. Desilylation using buffered TBAF²⁹ revealed protected α -hydroxyaldehyde 11, which we planned to use in a Wittig reaction with a suitably ¹³C-labeled phosphorous ylid. To the best of our knowledge, there is only a single report³⁰ using methyltriphenylphosphonium iodide to generate the Stork–Wittig reagent³¹ using an in situ deprotonation– iodination–deprotonation procedure which we adapted using 13 C-labeled salt 12 – available from relatively inexpensive 99% atom ¹³C-labeled methyl iodide – to give vinyl iodides Z -(1⁻¹³C)-13, $E(1^{-13}C)$ -13 and diiodide $(1^{-13}C)$ -14.³² Acetate deprotection as a mixture gave the corresponding alcohols $Z(1^{-13}C)$ -15, $E(1^{-13}C)$ -15 and (1-¹³C)-16 all with 99% ¹³C at the alkene terminus.[†]

Dehydrohalogenation of Z - and E-iodides $(1^{-13}C)$ -15 in the presence of inseparable diiodide $(1^{-13}C)$ -16 with LDA gave propargylic alcohol $(1^{-13}C)$ -7 in good overall yield, with the unprecedented observation that LDA converts vinyl 1,1-diiodides into terminal alkynes also (note §§, ESI†). Interestingly, 4% of the alkyne product was found to be the 2^{-13} C isotopomer (ESI[†]), implicating a 1,1-elimination reaction pathway for diiodide 16

Fig. 3 ¹H NMR spectra of (a) (*E*-2-¹³C)-**5** and (b) (*Z*-2-¹³C)-**5** displaying the expected 1_{C} -values for the α -vinyl protons.

and competitive alkyl group migration from a vinylidene intermediate (note $\P\P$, ESI[†]). Alcohol (1-¹³C)-7 was then converted to the desired bromoallene (1-¹³C)-7 (as 4% of its 2-¹³C-isotopomer, ESI⁺) as previously described (cf., Scheme 2).

With $(1^{-2}H)$ -4 and $(1^{-13}C)$ -4 in hand, epoxidation with DMDO was conducted. For deuterated $(1 - 2H) - 4$, after the reaction was conducted in the usual manner (cf., Scheme 1), $E-(2^{-2}H)$ -5 and $Z-(2^{-2}H)$ -5 were isolated each showing 65% deuteration at the α -position only (note ‡, ff, ESI†). Evidently, this result is consistent with the proposed mechanism (cf., Fig. 2) (note †††, ESI†). More compellingly, epoxidation of bromoallene $(1^{-13}C)$ -4 gave $(E-2^{-13}C)$ -5³³ and $(Z-2^{-13}C)$ -5 (28% isolated yield) where carbon atoms 1 and 2 from the bromoallene have entirely interchanged positions, giving also 4% of each of the $(E-1^{-13}C)$ -5 and $(Z-1^{-13}C)$ -5 isotopomers (ESI[†]). The expected $\rm ^1J_{CH}$ coupling constants experienced by the α -vinyl protons of the major isotopomers are clearly apparent in their ¹H NMR spectra (Fig. 3). Chenchon m

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In conclusion we have established that the hitherto unknown direct conversion of bromoallenes to α , β -unsaturated carboxylic acids using DMDO is consistent with an initial epoxidation event (note ***, ESI†) followed by a spontaneous reorganization via a bromocyclopropanone, a mechanism supported by calculations, in an intersecting bromoallene oxide – Favorskii manifold. These experiments support the proposed biogenesis of α , β -unsaturated carboxylate 3 from bromoallene 2 by epoxidation (note ‡‡‡, ESI†).

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