First enantiospecific synthesis of marine sesquiterpene quinol akaol A

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The first enantiospecific synthesis of akaol A, a marine sesquiterpene quinol, has been achieved. Key steps of the synthetic sequence are the oxidative degradation of (-)-sclareol to 10 a dinorlabdane ketoester, mediated by the ozone – lead (IV) acetate system, the diastereoselective α -methylation of a ketoaldehyde, followed by an intramolecular aldol condensation and the further Diels-Alder cylcoaddition of a dienol ether.

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Merosesquiterpenes are natural products of mixed biosynthetic ¹⁵ origin (polyketide-terpenoid) containing a sesquiterpene unit joined to a phenolic or quinone moiety. The most important group of this family of compounds, due to the wide variety of structural types and to their important, potent biological activities, are those bearing a bicyclic terpene (drimane) moiety. Smenodiol (1)¹ and ²⁰ siphonodictyal C,² which exhibits CDK4/cyclin D1 complex inhibitory activity,³ are examples of drimenyl phenols. The sphingosine kinase inhibitor (-)-F-12509⁴ is a representative drimenyl quinone.



Figure 1 Some representative merosesquiterpenes

Another important group of compounds belonging to this family of metabolites is that of marine puupehenones, which possess a ³⁰ tetracyclic structure, including a pyran ring.⁵ During the last decade a new type of merosesquiterpene, presenting a tetracyclic structure, including a cyclopentane ring, has been isolated from marine sponges and terrestrial fungi. Examples of this include

pelorol (2),⁶ akaol A (3),² dasyscyphin B (4),⁷ C (5),⁷ D (6)⁸ and E ³⁵ (7).⁸ Though the bioactivities of this family of compounds have yet to be examined comprehensively, recent studies have revealed that pelorol (2) is an activator of the inositol 5-phosphatase SHIP,⁹ whereas dasyscyphin B (4) and C (5) present potent cytotoxic activities in several human cell lines,⁷ and dasyscyphin D (6) and E ⁴⁰ (7) exhibits antifungal properties.⁸

Many efforts have been made to synthesise drimenyl phenols and puupehenone-related compounds.10 In most cases, the carbon skeleton of these compounds has been elaborated through a twosynthon strategy, involving the reaction of an aryllthium derived from a suitably protected polyphenol with a bicyclic sesquiterpene (drimane derivative) electrophile. However, little work has been done concerning tetracyclic merosesquiterpenes bearing the cyclopentance C ring, such as compounds 2-7. Andersen et al. recently reported the first synthesis of pelorol (2) starting from (+)-⁵⁰ sclareolide.¹¹ These authors utilized the two-synthon strategy to construct the carbon skeleton of the target compound, by condensation of an aryllithium with a drimane hydroxy aldehyde. Andersen's group came up against some difficulties in creating the cyclopentane C ring, via intramolecular Friedel-Crafts alkylation, which required a sufficiently activated aromatic moiety.¹² As was to be expected, a tetracyclic precursor with the appropriate configuration on C-8 was achieved in good vield after careful selection of aromatic substrate and cyclization conditions.

Continuing our research into the synthesis of bioactive 60 merosesquiterpenes, we examined the synthesis of sesquiterpene quinols such as compounds 3-7, which have not been yet synthesized, probably because the B/C cis fused system is unattainable utilizing the strategies previously reported. The results reported by Andersen's group in their synthesis of pelorol (2), 65 corroborated by our preliminary studies, revealed that the twosynthon strategy followed by intramolecular Friedel-Crafts alkylation led to the tetracyclic intermediate bearing a C8ß methyl group as the major diastereomer. Considering the above arguments, we planned the synthetic strategy shown in Scheme 1 to achieve 70 akaol A (3). The cyclopentane C ring of the target compound will be obtained through the intramolecular aldol condensation of a ketoaldehyde. The aromatic D ring of precursor 8 will be elaborated after the Diels-Alder cycloaddition of silyl dienol ether 9 derived from the α,β -enone resulting from the intramolecular aldol condensation of ketoaldehyde 10. The C8 α methyl group of compound 3 will be introduced after the diastereoselective Cmethylation of enol derived from the corresponding suitably protected ketoaldehyde; this will be obtained from ketone 11, after oxidative hydroboration. Ketone 11 will be prepared from ⁸⁰ commercial sclareol (12).

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[†] Electronic Supplementary Information (ESI) available: Full experimental procedures, spectroscopic data and copies of ¹H and ¹³CNMR. See http://dx.doi.org/10.1039/b000000x/





First, the preparation of ketone 11 was investigated. This compound has been widely utilized as the key intermediate in the synthesis of many natural products and other compounds of interest.¹³ However, only a total synthesis of compound 11, 90 utilizing a long synthetic sequence which involves optical resolution, has been reported.^{13c} The most efficient methods for preparing ketone 11 use natural products as starting materials; the most common procedure entails the degradative oxidation of the side chain of a diterpene labdane, which possesses the exocyclic 95 carbon-carbon double bond of the target compound.13a,13d,13e Procedures starting from (-)-sclareol (12), a naturally abundant, inexpensive commercial diterpene, have also been reported; these utilize as an intermediate the corresponding 8-acetyloxy ketone prepared by selective acetylation of the 8-hydroxy group and 100 subsequent side chain oxidation, or otherwise the palladium catalyzed allylic rearrangement of sclareol diacetate followed by ozonolysis.13b Even though a moderate yield (60%) for the monoacetylation of (-)-sclareol (12) has been reported, we encountered serious difficulties in attaining this result.



Scheme 2 Synthesis of ketone 11, via formate 14

Scheme 2 shows a more efficient means of preparing ketone 11 ¹¹⁰ from (-)-sclareol (12). Treatment of hydroxy aldehyde 13, obtained in high yield after the ozonolysis of diterpene 12, with lead (IV) acetate in dichloromethane at room temperature for 30 min gave ketoester 14¹⁴ in 93% yield. Compound 12 was directly converted into formate 14 in almost quantitative yield, after treatment with the ¹¹⁵ ozone – lead (IV) acetate system, previously reported by our group;¹⁵ the scope and limitations of this reaction, which can be utilized in a multigram scale, are currently under study. Ketoester 14 underwent the regioselective elimination of formic acid to give the exocyclic ketone 11, in high yield,¹⁶ by heating with collidine.

Scheme 3 shows the transformation of ketone 11 into aldehyde 18, bearing the characteristic $C8\alpha$ methyl group of the target compound. After protecting the ketone group as ethylene ketal, the oxidative hydroboration of the exocyclic carbon-carbon double 125 bond proceeded with complete diastereoselectivity, affording alcohol 16, which was easily converted into aldehyde 17.17 However, the α -methylation of the latter involved some difficulties, because of the strong tendency of this aldehyde to undergo Oalkylation, probably due to steric factors. This forced us to essay 130 the methylation of aldehyde 17 under different reaction conditions. The use of a strong base, such as NaH, in a polar aprotic solvent, such as DMF, considerably favoured the O-methylation (18/19, 4:6). The highest proportion of C-methylation product was attained utilizing t-BuOK in benzene. As was to be expected, the addition of 135 small quantities of HMPA to the reaction medium increased the proportion of O-alkylation product.



Scheme 3 Synthesis of aldehyde 18

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After the above-described procedures, we addressed the construction of the cyclopentane C and aromatic D ring of akaol A. Scheme 4 shows the synthesis of intermediate **8**, bearing the ¹⁴⁵ tetracyclic skeleton of the final sesquiterpene quinol. The ketal



Scheme 4 Synthesis of the sesquiterpene quinol precursor 8

¹⁵⁰ aldehyde **18**, after treatment with 1M HCl in THF under reflux for 3 h underwent simultaneous ketone deprotection and intramolecular aldol condensation, affording the tricyclic α,β-

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enone 20. Treatment of this with TfOTMS and Pri_2NEt gave silyl dienol ether 9, which after refluxing with methyl propiolate in

¹⁵⁵ xylene and further oxidation with DDQ in dioxane under reflux afforded the tetracyclic compound **21** together with the phenol **8**. The silyl ether **21** was transformed into hydroxy ester **8** after treatment with 1M HCl in MeOH.

Finally, functionalization of the aromatic D ring was tackled. ¹⁶⁰ Scheme 5 shows the transformation of compound **8** into akaol A (**3**). First, the elaboration of the catechol unit was attempted; the treatment of compound **8** with Fremy's salt or benzeneseleninic anhydride gave the unaltered starting material. This result can be attributed to the deactivation of the aromatic ring by the ester ¹⁶⁵ group. In order to avoid this inconvenience, the latter was reduced to the hydroxymethyl group. The phenol **25**, resulting from the deprotection of benzyl ether **24**, was then easily converted into *o*quinone **26**, which was finally transformed into akaol A (**3**) by reaction with Raney nickel. The optical rotation of synthetic akaol

¹⁷⁰ A (**3**) ($[\alpha]_D^{25}$: -13.7; c 8.0, MeOH) was similar to that reported for the natural product ($[\alpha]_D^{25}$: -12; c 0.15, MeOH); the spectroscopic properties were identical to those previously described.²



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Scheme 5 Synthesis of akaol A (3)

In summary, the first synthesis of (-)-akaol A (**3**) has been achieved starting from (-)-sclareol (**12**). The synthetic sequence includes a ¹⁸⁰ new oxidative degradation of the latter, induced by the ozone – lead (IV) acetate system, which affords ketoester **14** in high yield. The suitable configuration on C-8 was attained after the diastereoselective α-methylation of aldehyde **17**. The cyclopentane C ring of the target compound was obtained after intramolecular ¹⁸⁵ aldol condensation and the aromatic D ring was constructed through a Diels-Alder cycloaddition involving silyl dienol ether **9**. The authors thank the Spanish Ministry of Science and Innovation (Project CTQ2009-09932) and the Regional Government of Andalusia (Project P07-FQM-03101 and assistance to the FQM-¹⁹⁰ 348 group) for financial support. A. F. thanks the Spanish Ministry

of Science and Innovation for the predoctoral grant provided.

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