Total Synthesis of the Dihydrooxepine-Spiroisoxazoline Natural Product Psammaplysin A

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ABSTRACT: We report a general synthetic entry to dihydrooxepine-spiroisoxazoline (DOSI) natural products that culminated in the first racemic total synthesis of psammaplysin A. For the synthesis of the unique spirocyclic fragment we employed a strategy that features two key transformations: (1) a diastereoselective Henry reaction/cyclization sequence to access the C7 hydroxylated arene oxide and (2) a selective Baeyer–Villiger ring expansion to install the fully substituted dihydrooxepine and avoid the risk of a previously observed oxepine-arene oxide rearrangement. The overall synthesis proceeds in 13 steps from an inexpensive starting material.

During preliminary studies in our laboratories, we found that aldehyde 6, obtained from the dehydration with Martin’s sulfurane and oxidation of diol 5 with Dess–Martin periodinane (DM), undergoes spontaneous arene oxide-oxepine rearrangement to oxepine 7 in 30% yield (Scheme 1B). We initially considered oxepine 7 as a valuable precursor for 1, but it proved to be unstable upon storage and underwent slow decomposition to unidentified aromatic byproducts. Unexpectedly, attempted conversion of oxepine 7 to ester 8b was even more problematic as arene oxide 9b was obtained instead. Two key observations were made when we investigated the hydroxylation of isoxazoline 10, the product of a preceding nitrile oxide [3 + 2]-cycloaddition reaction (see Supporting Information for details). In this case, hydroxylation to isoxazoline 11 was outcompeted by rapid ring opening of theaza enolate to give isoxazole 12 as the sole product in 65% yield. These key insights were crucial for our further synthetic planning and ultimately led us to the development of the revised retrosynthesis outlined in Scheme 2.

We began with the disconnection of the known linker side chain molokaine (13) from 1 and focused our further analysis on the resulting DOSI 14. We anticipated that the lack of a fully decorated oxepine should eliminate the risk of unwanted arene oxide formation. Sequential removal of the enol ether decoration of ester 14 revealed structurally simplified lactone 15. The lactone acetal contains the retron for a Baeyer–Villiger oxidation of C6 producing spirocyclic ketone 16a. To avoid potential ring opening as observed for isoxazoline 10, we considered simultaneous installation of...
the requisite C7 hydroxy group and the isoxazoline scaffold in one step employing a powerful diastereoselective Henry reaction/cyclization protocol. To this end, nitronate \(16a\) was traced back to commercially available 2-hydroxymethylenecyclohexanone (17).

We began our forward synthesis with the bromination of compound 17 (Scheme 3A), which was obtained on large scale from the formylation of cyclohexanone in 89% yield. The use of N-bromosuccinimide (NBS) afforded the crude \(\alpha\)-bromo ketoaldehyde in excellent yields; however, we found it to be air-sensitive. For this reason, we decided to telescope the following step and conduct the overall process as a one-pot reaction. According to the seminal report of Rosini, the intermediate bromide was treated with ethyl nitroacetate and triethylamine to give nitronate \(16a\) and its C7 diastereomer \(16b\) as a 3:1 mixture as determined from the crude \(1^H\) NMR (see Supporting Information for details). After chromatographic separation of the highly polar \(16b\), we obtained nitronate \(16a\) in 51% yield. The relative configuration of the desired diastereomer \(16a\) was validated by single crystal X-ray analysis. Performing the reaction in acetonitrile at \(-25^\circ C\) was crucial, as lower diastereoselectivities were observed when performing the reaction at higher temperatures or changing the solvent to methanol in which an equimolar mixture of nitronates \(16a\) and \(16b\) was obtained. We then proceeded with the deoxygenation of nitronate \(16a\) by employing triethyl phosphite in 1,4-dioxane at 100\(^\circ\)C to isolate the spirocyclic isoxazoline 18 in up to 99% yield.

Having installed the 4-hydroxyisoxazoline motif and with multigram quantities of ketone 18 in hand, our attention was directed toward investigation of the Baeyer–Villiger oxidation to enable installation of the DOSI motif. We were pleased to see that exposure of ketone 18 to \(m\)-chloroperoxybenzoic acid (\(m\)-CPBA) in the presence of disodium hydrogen phosphate afforded the corresponding lactone 19 in nearly quantitative yield. To our surprise, the protection of the hydroxy group in intermediate 19 with tert-butyldimethylsilyl trifluoromethanesulfonate (TBSOTf) and triethylamine failed to give the desired silyl ether. Instead, formation of carbonate 20 was observed. This product might be derived from the reaction of lactone 19 with ethyl cyanoformate, which could form via decomposition of lactone 19. As a part of the biosynthesis of ceratinamine A, a similar isoxazoline fragmentation was studied by Ganem. In parallel, we took advantage of the reaction of nitronate 16a with dimethylvinylsilyl chloride in the presence of imidazole. Under these conditions, clean silylation of the C7 hydroxy group followed by intramolecular 1,3-dipolar cycloaddition of the vinyl group with the nitronate took place to furnish the tricyclic acetal 21 in 95% yield. This motif serves a dual role as it (1) protects the hydroxy group and (2) enables one-step unmasking of the isoxazoline as described by Rosini.

The following Baeyer–Villiger oxidation proceeded with excellent regioselectivity and furnished the desired lactone 15 in 87% yield over two steps on a 6-g scale. When performing the ketene acetal phosphate formation at \(-78^\circ C\), varying yields between 36–67% of 24 were obtained and large amounts of lactone 15 were recovered. After further optimization, we found that slow addition of potassium hexamethyldisilazide (KHMDS) solution in THF at \(-95^\circ C\)
was crucial for complete consumption of the starting material and to give ketene acetal phosphate 24 in reproducible 86% yield.

The palladium-catalyzed reduction of the ketene acetal phosphate to enol ether 25 also required some optimization. The use of Pd(PPh$_3$)$_4$ and Et$_3$Al gave enol ether 25 as a complex mixture together with an unstable byproduct resulting from a competing cross-coupling reaction with Et$_3$Al. We then switched to LiBH$_4$ as the reductant, however, this led to competing reduction of the ester moiety. The chemoselectivity issue could be avoided by replacing LiBH$_4$ with formic acid/triethylamine buffer. This allowed for clean conversion of ketene acetal phosphate 24 to tetrahydrooxepine-spiroisoxazoline 25 for the first time. Further screening revealed PhMe$_2$SiH as the ideal reagent to give enol ether 25 in 66% yield.

With a scalable route to enol ether 25 in hand, we continued with the investigation of its sequential functionalization (Scheme 3B). We exposed enol ether 25 to a panel of oxidation conditions, most of which employed tert-butyl hydroperoxide (TBHP) in combination with a transition metal catalyst (CuBr$_2$, CuI, RuCl$_3$, Mn(OAc)$_3$, Mn(dpm)$_3$, Pd(OH)$_2$/C; see Supporting Information for details). The screening revealed that the desired vinylogous lactone 26 was formed under most conditions accompanied only by its regioisomeric $\alpha,\beta$-unsaturated lactone 27. The use of Pd(OH)$_2$/C as the catalyst afforded vinylogous lactone 26 and lactone 27 as a 2.5:1 mixture of regioisomers; however, the reaction suffered from low conversion. The best performing conditions in terms of conversion involved Mn(dpm)$_3$ and dropwise addition of TBHP overnight to give 26 in 32% isolated yield. Direct conversion of the undesired regioisomer 27 (19%) to ketene acetal phosphate 24 was unsuccessful in our hands but allowed for the isolation of lactone 15 in 67% yield.

Subsequent $\alpha$-bromination of vinylogous lactone 26 with CuBr$_2$ and 2,6-di-$t$-tert-butylypyridine (DTBP) in a mixture of chloroform and ethyl acetate at 100 °C proceeded cleanly to deliver the C4-brominated product in 78% yield. The second bromination was accomplished upon exposure to tetra-$n$-butylammonium tribromide (TBATB) and DTBP at 60 °C in

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*See the Supporting Information for detailed procedures and characterization data.*

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**Scheme 3. Synthesis of Psammaplysin A**

A) **Synthesis of the Spiroisoxazoline Motif**

B) **Completion of the Synthesis**
chloroform to afford vinylogous lactone 28 in 72% yield. We found that the overall yield of vinylogous lactone 28 could be improved by telescoping the three oxidations without chromatographic purification of the intermediates. In this way, we obtained vinylogous lactone 28 in 27% yield over three steps from 25. Finally, sequential treatment of vinylogous lactone 28 with KHMDS and methyl trifluoromethanesulfonate (MeOTf) gave the fully substituted DOSI 14 in 53% yield.

For the late-stage attachment of the protected moloka’i-amine unit 29, which was prepared from tyramine in four steps (see Supporting Information), we first attempted conversion of ethyl ester 14 into the corresponding carboxylic acid. Saponification under aqueous conditions (LiOH, THF, H2O, 23 °C) followed by isolation of the acid turned out to be impossible in our hands as decomposition, presumably via decarboxylation, was observed even at low temperatures. For this reason, we decided to treat ester 14 with potassium trimethylsilanolate (TMSOK) at −15 °C under nonaqueous conditions followed by direct addition of ammonium salt N'-methylimidazole (NMI), and chloro-N,N,N',N'-tetramethylformamidinium hexafluorophosphate (TCFH). This one-pot procedure bypassed the problematic isolation of the acid and allowed us to obtain the desired amide 30 in 79% yield from ester 14. The following deprotections of the C7 oxygen and C20 nitrogen proceeded smoothly to furnish psammaplysin A (1) in 60% yield. The analytical data (1H NMR, 13C NMR, HRMS) for 1 and its bisacetylated derivative fully matched those reported for the natural compound. Overall, psammaplysin A (1) was synthesized in 13 steps from commercially available starting materials.

In summary, we have accomplished the first synthesis of psammaplysin A (1) nearly 40 years after its first isolation. The key to success of the developed strategy was the use of a Henry addition/O-alkylation sequence instead of a nitrile oxide [3 + 2]-cycloaddition. This granted access to the fully substituted C7 hydroxylated heterocyclic scaffold in one step. For the construction of the 7-membered ring-system a high-yielding Baeyer–Villiger oxidation turned out to be the method of choice. This avoided the risk of a previously encountered oxepine-arene oxide rearrangement and paved the way for the selective installation of the fully intact DOSI motif. The late-stage attachment of the side chain ensures the high flexibility and modularity of the synthesis. This approach should allow for the preparation of several structurally related members as well as analogs with deep-seated structural modifications and facilitate future bioactivity studies.

■ ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/DOI/10.1021/jacs.2c10010. Experimental details, spectroscopic data, and X-ray data (PDF)

Accession Codes

CCDC 2207064–2207066 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

■ CONFLICT OF INTEREST

The authors declare no competing financial interest.

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Notes

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