Enantio- and diastereocontrolled synthesis of (+)-juvabione employing organocatalytic desymmetrisation and photoinduced fragmentation[†]

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(+)-Juvabione, a natural sesquiterpene exhibiting insect juvenile hormone activity, has been synthesized from σ -symmetric 4-(2-formylethyl)cyclohexanone by employing organocatalytic asymmetric aldolisation and Norrish I-type fragmentation as the key steps.

The development of a catalytic enantioselective transformation that enables facile access to useful chiral platforms is a major goal in current synthetic organic chemistry.¹ We have recently reported the highly enantioselective construction of both enantiomeric forms of endo-8-hydroxybicyclo[3.3.1]nonan-2-one (2) from the σ -symmetric keto-aldehyde 1 via intramolecular asymmetric aldolisation employing a chiral amino acid or its tetrabutylammonium salt as a catalyst,² and demonstrated the synthetic utility of the aldol product 2 as a chiral cyclohexanoid block based on a novel aldolisation/retroaldolisation interconversion.³ As an alternative use of the building block 2, we envisage the applicability of a photochemical process known as the Norrish I reaction⁴ to the production of 3 from 2, which brings about a formal intramolecular asymmetric redox transformation of 1. We confirmed that 2 smoothly isomerises to 3 on irradiation with light (300 nm) in degassed MeOH at ambient temperature for 90 min in 60% yield (Scheme 1).



Scheme 1 Organocatalytic aldolisation and photoinduced fragmentation.

To demonstrate the synthetic utility of this methodology, we performed an enantio- and diastereocontrolled synthesis of (+)-juvabione (4), a natural sesquiterpene isolated together with (+)-epijuvabione (5) and exhibiting insect juvenile hormone activity;⁵ the presence of two contiguous stereogenic centers on a ring and a side chain and the fact that the (+)-(4R, 1'R) isomer exhibits the highest biological activity make this compound a fascinating target in enantio- and diastereocontrolled synthesis (Fig. 1).⁶



Fig. 1

A highly enantiomerically enriched (+)-2 was obtained *via* catalytic asymmetric aldolisation using **6** as a catalyst, and the scaffold for the installation of the C1 methyl group was generated *via* a two-step sequence involving MOM protection and IBX-mediated oxidation to give (+)-7 (Scheme 2).^{2,3}



Scheme 2 Reagents and conditions: a) 6, MeCN, rt, 23 h; b) MOMCl, i-Pr₂NEt, CH₂Cl₂, rt, 18 h; c) IBX, toluene–DMSO, 55–75 °C, 11 h.

To set the stage for the Norrish I reaction, the enone (+)-7 (> 99% ee) was treated with Me₂CuLi to give the 1,4-adduct 8 in 98% yield exclusively. The bicyclic ketone 8 was then transformed to the TES ether 10^7 via a deprotection/reprotection sequence in 76% yield over 2 steps. Having obtained the substrate for the key step, we attempted its transformation to a cyclohexenol derivative with a correct stereochemistry to (+)-juvabione (4) under photochemical conditions. It was found that the selective cleavage occurred smoothly to give the desired aldehyde 11 containing a cyclohexenol moiety in a good reproducible yield (66%) together with a trace amount of mechanistically predictable 3-(4-TESoxycyclohexyl) butanoic acid methyl ester via ketene generated in situ as a byproduct. Note that structurally related compounds such as MOM ether 8 and free alcohol 9 furnished the corresponding aldehydes such as 12 and 13 in 69% and 60% yields, respectively, under the same conditions, indicating the potential use of this process in the synthesis of chiral cyclohexanoids. Also, note that the cleavage occurred selectively at the carbon-carbon bond connected to the bridgehead (Scheme 3).⁸

The isobutyl side chain of juvabione was introduced under Imamoto's conditions⁹ using the Grignard reagent–CeCl₃ system to give the secondary alcohol **14** in 81% yield as a diastereomeric mixture (1 : 1).¹⁰ Upon sequential reactions involving BOM

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Scheme 3 Reagents and conditions: a) MeLi, CuI, THF, -40 °C, 1.5 h; b) LiBF₄, 1,4-dioxane, H₂O, 50–70 °C, 7 h; c) TESCl, imidazole, DMF, rt, 12 h; d) *hv* (300 nm), MeOH, rt, 1.5 h.



Scheme 4 Reagents and conditions: a) *i*-BuMgBr, CeCl₃, THF, 0 $^{\circ}$ C, 2 h; b) BOMCl, *i*-Pr₂NEt, TBAI, THF, rt, 47 h; c) TBAF, THF, rt, overnight; d) MnO₂, CH₂Cl₂, rt, 12 h; e) (methoxymethyl)triphenylphosphonium chloride, *n*-BuLi, THF, $-30 \,^{\circ}$ C, 2 h; f) 10% aq. HCl, THF, rt, 2 days; g) Dess–Martin periodinane, CH₂Cl₂, rt, 1 h or cat. 1-Me-AZADO (19), BAIB, CH₂Cl₂, rt, 7.5 h; h) NaCN, MnO₂, AcOH, MeOH, rt, 24 h.

protection, TBAF-mediated removal of the TES group, and MnO₂ oxidation, **14** furnished the enone **15** in 92% yield. The crucial C-1 homologation of the enone **15** was attained *via* the Wittig reaction using (methoxymethyl)triphenylphosphonium chloride and *n*-BuLi in THF at -30 °C to give the methyl dienol ether **16**, which was immediately treated with aqueous 10% HCl at ambient temperature for 2 days to give the corresponding hydroxy-α,β-unsaturated aldehyde **17** in 64% yield. While the oxidation of resultant secondary alcohol **17** was carried out with Dess–Martin periodinane¹¹ to give the penultimate ketone **18**^{6α,6b} in 85% yield, it was found that using 5 mol% 1-methyl-2-azaadamantane *N*-oxyl [1-Me-AZADO (**19**)],¹² a stable nitroxyl-radical-type oxidation catalyst that has recently been developed by

our laboratory, with bis(acetoxy)iodobenzene; this improved the yield of oxidation up to 94%. Finally, according to Trost's synthesis⁶⁶ **18** was subjected to Corey's conditions¹³ using NaCN, MnO₂, and AcOH in MeOH at ambient temperature to give (+)-juvabione (4), $[\alpha]_D^{28} = +69.1$ (*c* 1.00, benzene) [lit.:^{6g} [$\alpha]_D^{25} = +66.9$ (*c* 2.57, benzene)], in 78% yield and complete the synthesis (Scheme 4).

In conclusion, we have described the efficient synthesis of (+)juvabione (4) with excellent stereocontrol from the σ -symmetric keto-aldehyde 1 based on "asymmetric aldolisation/Norrish I cleavage" methodology, in which the temporarily generated chiral aldol motif in 2 plays essential roles in stereochemical control. The present strategy is complementary to the aldolisation/retroaldolisation interconversion that we established³ and offers a versatile use of 2 as a chiral cyclohexanoid block.

Notes and references

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