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Biocatalytic approaches for a more sustainable synthesis of sandalwood fragrances†

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The synthesis of campholenic-based fragrances requires the preservation of specific structural elements to capture the desired sandalwood scent. The most critical step of their preparation is the reduction of α,β -unsaturated carbonyl precursors while preserving the campholenic unsaturation. Classical reductions, especially hydrogenations, often lack complete chemoselectivity, leading to the formation of overreduced byproducts. In addition, the stereochemistry plays a key role in the olfactory perception of these chiral fragrances. However, none of the current industrial syntheses are stereoselective, resulting in wasteful production of non-contributory isomers. Herein, we explore the untapped potential of biocatalytic reductions using ene-reductases (ERs) and alcohol dehydrogenases (ADHs) to enhance the sustainability of four commercial sandalwood fragrances (Brahmanol®, Firsantol®, Sandalore®, and Ebanol®), focusing on the stereoselective synthesis of their most odorant isomers. A comparison of green metrics, including E-factors and EcoScale, between bio- and chemo-based reductions is presented.

Introduction

The hydro-distillation of *Santalum album* wood yields one of the most renowned essential oils in fine perfumery. This essence, composed of a complex mixture of natural products, is highly appreciated by consumers for its unique sweet, balsamic, and exquisite woody scent notes. However, over the last century, the sandalwood oil demand increased to the point where its availability became increasingly difficult, reaching a critical level by the late 70s. Such high demand led to unsustainable cultivation practices and even to illegal harvesting in nature reserves. As a result of its intensive exploitation, in 1998 *Santalum album* was added to the red list (IUCN) of endangered species; this is a compelling example of how the label "natural" does not necessarily mean sustainable!

Prompted by the rapid depletion of this raw material, many organic chemists were engaged in the challenging synthesis of its main components, *i.e.* the α - and the β -santalol sesquiterpenes (Scheme 1A), however, so far, none of these laboratory-scale syntheses appears to be exploitable on an industrial

Department of Chemistry, Materials and Chemical Engineering "Giulio Natta", Politecnico di Milano, p.zza L. da Vinci 32, Milano, 20133, Italy. E-mail: francesco.gatti@polimi.it level.³ In this regard, very recently, Ventos has commercialized Isobionics® Santalol oil, which, in terms of composition and odorous properties, is very close to the natural one, but it is produced on a biotechnological basis from renewable raw materials (cornstarch).⁴ Alternatively, a new category of synthetic sandalwood-scented fragrances (1) like Brahmanol®, Firsantol®, Sandalore®, Ebanol®, and many others was developed.⁵ These compounds are efficiently synthesized starting from α -pinene, which is an abundant and renewable natural source (Scheme 1B). As a result, their production can be considered more environmentally sustainable than the sandalwood oil extraction from endangered *Santalum* trees.

The carbon skeleton of alcohols 1 (C_{13} – C_{14}) is constructed from campholenic aldehyde 2 (obtained from α-pinene oxide) through well-established C=C double bond forming reactions (crossed aldol condensation, Claisen rearrangement and Mannich methylene homologation, see ESI†). However, since the campholenic ring is a key structural element needed for the perception of sandalwood odour, the reduction of the carbonyl precursors, preserving the ring unsaturation, is undoubtedly the most challenging and delicate step of the overall synthesis. Currently, these reductive steps are accomplished either by hydrosilylation, hydrogen transfer or hydrogenation reactions, mostly catalysed by precious transition metals.5,6 Although these methodologies were preferred to the metal hydride-based procedures, mainly for practicality reasons, they are not yet fully chemoselective, leading to possible overreduced products and to C=C double bond positional isomerization.

[†] Electronic supplementary information (ESI) available: Preparation and characterization of products, Mosher's ester analysis, optimization of multienzymatic protocols, olfactory evaluation; copies of ¹H and ¹³C NMR spectra. See DOI: https://doi.org/10.1039/d4gc00746h

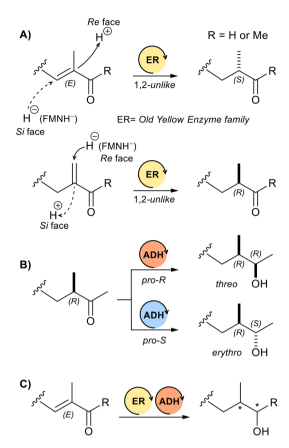
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Scheme 1 (A) α - and β -santalol are the major components of Indian sandalwood oil (around 65-75% w/w) and are responsible for of its characteristic scent. (B) Partial retrosynthesis of selected commercially available campholenic sandalwood fragrances.

In addition, preliminary olfactive evaluations on selected sandalwood fragrances have revealed significant differences in scent and odour thresholds among the stereoisomers.7 The correlation between stereochemistry and odour perception is not unusual among chiral odorant molecules,8 and, in some cases, even odourless enantiomers have been identified.9 Nevertheless, most commercial fragrances are still manufactured relying on non-stereoselective methodologies, resulting in a substantial waste of reagents, solvents and energy employed for the synthesis of all isomers that do not contribute to the scent of the fragrance.

To the best of our knowledge, biocatalytic reductions based on ene-reductases (ERs) and alcohol dehydrogenases (ADHs) enzymatic activities have never been applied to sandalwood fragrances synthesis. 10 Indeed, based on safety, environmental but also on stereo- and chemoselectivity issues, such catalytic approach might significantly improve the sustainability of the productive process.

In more detail, the ERs catalyse the stereospecific reduction of C=C double bonds conjugated with electron-withdrawing groups (typical EWGs are -COH, -COR, -CN, -NO₂ or -CO₂R), usually high stereoselectivities and conversions are achieved. 11 Especially, many of the ERs belonging to the old yellow enzymes family (OYEs) are known to catalyse the reduction of (E)- α -methyl enones or enals with (S)stereoselectivity. 12,13 Conversely, the reduction of their α -methylene regioisomers affords the (R) enantiomers (Scheme 2A).13 Thus, the stereochemical course of the



Scheme 2 (A) Stereoselectivity of flavin mononucleotide (FMN) cofactor dependent ERs like the old yellow enzymes (OYEs): (S) enantioselectivity is observed with α -methyl enals or enones, while for α -methylene isomers is (R). (B) The selection of an ADH with pro-R or pro-S stereoselectivity enables the access to either the erythro or to the threo alcohols. (C) ADH and ER enzymes can be combined in a one-pot two-step cascade process.

reduction proceeds for these enzymes with unlike stereotopicity (proton addition from Re enantioface to give (S) products for α,β -unsaturated substrates and from Si face to give (R) products for α-methylene substrates, Scheme 2A).14 Only very recently, deazaflavin (F420) dependent ene-reductases (FDRs) have been found to be stereodivergent (like stereotopicity) to most common flavin mononucleotide (FMN) dependent ERs, including the OYEs.15

ADHs, on the other hand, catalyse the reduction of carbonyl groups of ketones or aldehydes. These redox enzymes typically give excellent results in terms of conversion and stereoselectivity, and, as opposed to the ER enzymes, it is not rare to find ADHs with opposite stereoselectivity (Scheme 2B). 16

Lastly, the combination of ER and ADH activities in a cascade process has proven to be highly effective for the reduction of several prochiral enones and enals.¹⁷ Chiral alcohols with up to two stereogenic centres are usually obtained in good yields and with a high optical purity (Scheme 2C).

In the following we show how the application of biocatalytic methods might significantly impact on the sustainability of sandalwood fragrances 1a-d.

Results and discussion

Epoxidation of α -pinene and synthesis of campholenic aldehyde

To begin, two literature procedures for the conversion of (-)- α -pinene ([α]_D = -37.1, c = 1, CHCl₃, ee = 82% by chiral GC, see ESI†) into pinene oxide 3 were evaluated, conversion and selectivity data are reported in Table 1. However, epoxide 3, due to its high ring-strain energy, underwent rearrangement already during the reaction, regardless of the method applied, affording a mixture of constitutional isomers: mainly campholenic aldehyde 2 and other terpenes¹⁸ (pinocamphone and trans-pinocarveol, detected by GC-MS). Only the non-campholenic terpenes must be considered as side-products of the reaction, since 2 is the product of the next step (Meinwald rearrangement). However, the product distribution was significantly correlated to the reaction temperature. Indeed, although the epoxidation with O2/acetaldehyde in the presence of a catalytic amount of N-hydroxyphtalimide (NHPI) at 60 °C is very appealing for its high atom economy, 19 it yielded quantities of other terpenes considerably higher (28% vs. 10% by GC-MS) compared to the epoxidation with m-chloroperbenzoic acid (MCPBA),²⁰ which instead was carried out at a lower temperature (5-10 °C). Nonetheless, the use of peracids has two main drawbacks: (a) safety concerns and (b) non-negligible production of carboxylic acid by-products (in this case m-chloro-

Surprisingly, the Payne epoxidation²¹ has never been applied to α -pinene. This procedure involves the addition of hydrogen peroxide (H₂O₂) to a nitrile (RCN) to produce a peroxycarboximic acid (RC(OOH)NH). Then, the acid oxidizes the olefin, resulting in the formation of epoxide and amide byproduct (RCONH₂).

Three different nitriles were tested (MeCN, PhCN and CCl₃CN), conversions and chemoselectivity are reported in

Table 1. Both MeCN and PhCN gave unsatisfactory conversions and required high reaction temperatures, that, as seen before, favour the rearrangement of pinene oxide into pinocamphone or carveol. 22,23 In contrast, the higher reactivity of CCl₃CN allowed to form 3 at room temperature with great benefit to the selectivity (86% by GC-MS). Even in this reaction, as seen for the MCPBA oxidation, we produced a stoichiometric amount of by-product (trichloroacetamide), quantitatively isolated by simple precipitation in hexane during the reaction work-up. But in this case, by heating the amide in presence of dehydrating phosphoric anhydride P₄O₁₀ in refluxing xylene,²⁴ we could recover over 65% of the initial nitrile by distillation (Scheme 3, for more details see ESI†). The recovered nitrile was of sufficient purity (96% by quantitative ¹³C-NMR in presence of Cr(acac)3, see ESI†) to be reused for further epoxidations (Scheme 3).

Although H_2O_2 is considered a "green" oxidant, we explored the possibility of generating and using it *in situ* in the Payne oxidation through a telescopic approach. Currently, H_2O_2 is produced by the anthraquinone process, which requires a Pd

Scheme 3 Reaction conditions: for steps (i) and (ii) see Table 1; (iii) 0.8 eq. P_4O_{10} , refluxing xylene.

Table 1 Screening of different epoxidation methods of α -pinene

		Prod. distrib. ^a (%)		
Reaction conditions	Conv. a (%)	3	2	Other
O ₂ , ^b 3 eq. CH ₃ CHO, 0.1 eq. NHPI, MeCN, 60 °C (ref. 19)	62	55	17	28
1.3 eq. MCPBA, 1.0 eq. NaHCO ₃ , CH ₂ Cl ₂ , 5–10 °C (ref. 20)	91	82	8	10
1.1 eq. H ₂ O ₂ , ^c 1.8 eq. MeCN, MeOH, pH = 9 KHCO ₃ buffer, rt (ref. 23)		_	_	
1.1 eq. H ₂ O ₂ , c 13.3 MeCN, pH = 7 KP _i , 60 °C (ref. 22)	18	64	7	29
1.5 eq. H_2O_2 , 6.6 eq. PhCN, pH = 7 KP; 60 °C (ref. 21)	_	_	_	
1.2 eq. H_2O_2 , c 1.2 eq. CCl_3CN , pH = 7 KP _i , CH ₂ Cl ₂ , rt (ref. 21)	100	86	5	9
(i) O_2 , cat. GOX, 3.0 eq. glucose, pH = 7 KP _i , 30 °C; (ref. 27) (ii) 1.5 eq. CCl ₃ CN, CH ₂ Cl ₂ , rt (ref. 21)	65	68	17	15

 $[^]a$ Not isolated yield, by GC-MS. b Bubbling. c 30% w/w.

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catalysed hydrogenation to regenerate the starting 2-alkyl anthrahydroquinone.25 Alternatively, H2O2 could be generated by a biocatalytic approach, avoiding the use of H₂ and precious transition metals. For instance, glucose oxidase (GOX) catalyses the reduction of molecular oxygen to H₂O₂, ²⁶ using glucose as sacrificial substrate (Scheme 3). More recently, H₂O₂ was also bio-produced by oxidation of choline to trimethylglycine using choline oxidase enzymes and used in situ for limonene oxidation.27

In our case we succeeded to generate a dilute solution of H_2O_2 ($\approx 1.2\%$ w/w) by oxidation of glucose to gluconolactone (around 70% by 13C-NMR, see ESI†) in a phosphate buffer by bubbling O2 in presence of GOX from Aspergillus niger. Then, the solution was submitted to Payne oxidation of pinene, however both conversion and selectivity were not as good as those achieved using directly commercial H₂O₂ (65% vs. 100%, Table 1), in which case the oxidation was easily scaled-up on a multi-g scale, and on (+)- α -pinene ([α]_D = +40.8, c = 1, CHCl₃, 92% ee by chiral GC, ESI†) as well. Finally, the oxidation reaction mixture was submitted to the Meinwald rearrangement catalysed by ZnBr₂ (7% w/w) in toluene to give the campholenic aldehyde 2 in a quantitative yield. 18,28

Stereoselective one-pot two-step bioreduction

With the aldehyde 2 in our hands (purity 91% by GC-MS) we prepared the substrates for the bioreductions: the α,β -unsaturated carbonyl compounds **4a–c** and **6** (Fig. 1A), and the α-methylene aldehydes 5a-b and ketones 5c-d (Fig. 1B). However, concerning their syntheses, we avoided as much as possible column chromatography purifications, more details on the synthetic routes are available in the ESI.†

The reduction of the conjugated C=C double bond of substrates was catalysed by recombinant OYE2 from S. cerevisiae. This enzyme was selected from a panel of ERs from different

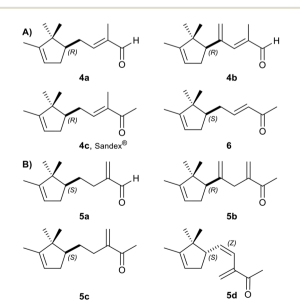


Fig. 1 Substrates submitted to bioreduction: (A) α , β -unsaturated carbonyl compounds; (B) α -methylene isomers.

sources for its optimal balance between selectivity and activity (for the screening see ESI†). For the regeneration of the reduced nicotinamide adenine dinucleotide phosphate (NADPH) cofactor, glucose dehydrogenase enzyme (GDH) from B. megaterium together with an excess of glucose, as sacrificial co-substrate, 29 was used (Scheme 4). In most cases, the conversions were nearly quantitative (by GC-MS), usually within 10-12 hours, and occasionally, supplemental quantities of enzyme and cofactor was necessary to ensure the complete consumption of all starting material. Then, the commercially available Evo440 ADH (from evoxx technologies GmbH), known for its high activity on a broad range of substrates, 17f-i was added to the reaction mixture containing the saturated aldehydes (I, R = H), yielding the primary alcohols 1a-b. After product removal from the reaction mixture by absorption on hydrophobic polyaromatic resins, 17b,29,30 followed by column chromatography and/or bulb-to-bulb distillation purification procedures, each diastereoisomer of Brahmanol® and Firsantol® fragrances was isolated with a very high purity (>98% by GC-MS). This purification process ensured the elimination of all off-odours and contaminants typically arise during the biotransformation, rendering the fragrances suitable for the olfactory evaluation. For the reduction of ketone intermediates (I, R = Me), either a pro-R or a pro-S ADH, respectively Evo270 or Evo440, was employed, following the same work-up described above, affording secondary alcohols

In summary, the reaction conditions together with yields, diastereomeric ratios drs (measured by 13C-NMR) and optical rotatory values of alcohols 1a-d are reported in Scheme 4. In addition, since our attempts to determine the enantiomeric excess of either the carbonyl intermediates I or of the alcohols 1 by chiral GC or HPLC failed, we opted to quantify indirectly the stereoselectivity of OYE2 (Scheme 4) by a mass balance approach taking into account the initial ee of starting pinene, which was reasonably retained throughout the synthesis (see ESI†) and the dr of alcohols **1a-b** (by ¹³C-NMR) and of ketone intermediates I (by GC-MS or by ¹³C-NMR).

Interestingly, even though Brahmanol® and Firsantol® differ only by a methylene group at C(4) position, the stereoselectivity of OYE2 in reducing the Firsantol® precursors 4b-5b was significantly higher than that of Brahmanol® homologues 4a-5a (Scheme 4), regardless of the type of substrate (methylene or α,β -unsaturated). A similar enhancement of selectivity was observed for the reduction of 5d (Ebanol® precursor) compared to the reduction of substrates 4c-5c (Sandalore® precursors), which differ from 5d solely by the absence of the C=C unsaturation. Clearly, increasing the steric hindrance of substrates by adding a methylene group or by stiffening their structure proved beneficial to the stereoselectivity.

Lastly, in Fig. 2 we show, as representative example, the stacked ¹³C-NMR expansion spectra relative to the C(3) signal of each diastereoisomer of Sandalore® (1c), including the commercial mixture; integration of this signal allowed the determination of dr.

96:4, +106.0

>99(R)

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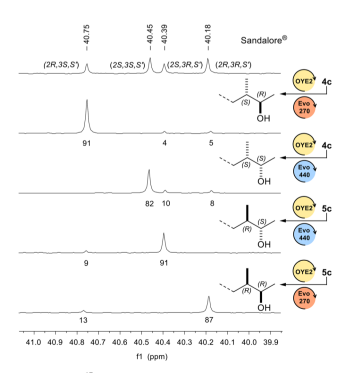
Scheme 4 Biocatalytic reduction of substrates 4a-c and 5a-d. Reaction conditions: substrate (3 mmol), pH = 7 KP, buffer (15 mL), i-PrOH cosolvent (1% v/v), 30 °C, 150 rpm; enzymes: for primary alcohols and secondary (2S) alcohols: (i) OYE2 (6 mL, ≈3 mg mL⁻¹), GDH (≈200 U), NADP+ (20 mg), glucose (4.2 g); (ii) after 10-12 h Evo440 (30-40 mg) and NADP+ (10 mg) were added. For (2R) alcohols: same reaction conditions but in step (ii) Evo270 (30-40 mg) was used instead of Evo440. ^a Isolated yield after column chromatography followed by bulb-to-bulb distillation. ^b By ratio of 13 C-NMR signal integrations: C(1) for 1a-b, C(3) for 1c; C(2) for 1d. c In CHCl₃, $c \approx 1$. d Estimated by mass balance taking in account the initial ee of pinene, see ESI.†

(2S,3R,S')-1c

72% from **5c**

91:9, +30.3

94(R):6(S)



ÓН

(2R,S')-1a

82% from 5a

87:13, +17.0 vs +17.3^[7b]

95(R):5(S)

(2R,R')-**1b**

80% from 5b

91:9, -57.6 vs -60.8^[7c]

>99(R)

Fig. 2 Stacked ¹³C-NMR spectra relative to the C(3) signal of each diastereoisomer of 1c (CDCl₃, 100 MHz).

Determination of the absolute stereochemical configuration

(2R,3R,S')-1c

70% from 5c

87:13, +27.0

94(R):6(S)

The configurations initially assigned on the basis of the typical stereochemical courses commonly observed in the reductions catalysed by OYE2 (unlike stereotopicity), 12,17 Evo270 (pro-S), and Evo440 (pro-R) enzymes, 17h,i were confirmed by a combination of experiments on Sandalore® stereoisomers. Firstly, the absolute configuration at C(2) stereogenic centre was determined by analysis of ¹H and ¹³C-NMR spectra of esters obtained by condensation of α-methoxyα-trifluoromethylphenylacetic acid (MTPA) with 1c alcohols.³¹ In Fig. 3A is shown the expanded region of ¹³C-NMR spectra relative to C(1) signal of both (R)-MTPA and (S)-MTPA esters of the couple of alcohols 1c obtained from the bioreduction of 4c with OYE2 and Evo440, and with OYE2 and Evo270, respectively.

According to Mosher's model, when the C(1) methyl group lies closer to the shielding region of phenyl ring, its ¹H and ¹³C chemical shifts are up-field, conversely if it is distanced from the aromatic ring its protons are less deshielded, resulting in a downfield chemical shift. The differential chemical shift $\Delta \delta^{S,R}$ ($\Delta \delta^{S,R} = \delta_{\text{MTPA}(S)} - \delta_{\text{MTPA}(R)}$) resulted +0.56 ppm for Evo270 and −0.29 ppm for Evo440, confirming the stereoselectivities of both ADHs. Then, we assigned the absolute configuration at C(3) centre knowing the C(2)-C(3) relative stereochemistry (syn or anti), which was attributed by ¹H-NMR ana**Green Chemistry** Paper

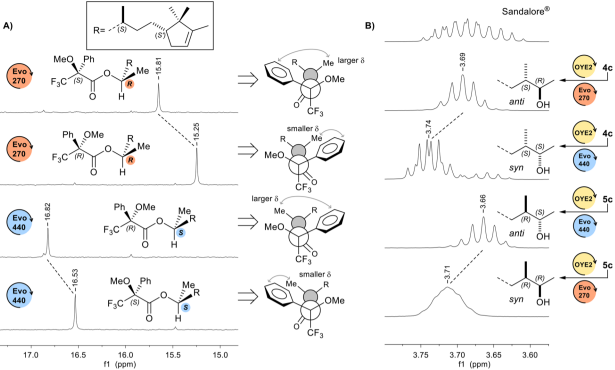


Fig. 3 (A) ¹³C-NMR expansion spectra relative to C(1) signal of (R)-MTPA and (S)-MTPA esters of alcohols 1c obtained from the bioreduction of 4c (CDCl₃, 100 MHz). (B) ¹H-NMR expansion spectra relative to the CHOH signal of Sandalore® and its diastereoisomers (CDCl₃, 400 MHz).

lysis of the chemical shift and multiplicity of CHOH signal. Indeed, it is known that for syn diastereoisomers this signal is a doublet of quartet (J = 6.3 and 4.4 Hz) downfield compared to that of its syn isomers ($\delta = 3.74$ and 3.71 ppm for syn alcohols vs. 3.69 and 3.66 ppm for the couple of anti diastereoisomers, in CDCl₃), for which instead it appears as a quintet (Fig. 3B).32

Notably, only by isolating ketone (Z)-7**d**, obtained by repeating the reduction without ADH, it was possible to confirm the (R) stereoselectivity of OYE2 in reducing α -methylene- β , γ -unsaturated substrates like dienone 5d (Fig. 4).

The measurement of Nuclear Overhauser Effect (NOE) through 2D NOESY experiment, 33 between H(6) and C(3)CH3 protons allowed to determinate their average distance, which turned out to be crucial for the assignment of the relative configuration, and therefore of the absolute configuration as well. Indeed, a conformational study, conducted by density functional theory (DFT) computational chemistry (model chemistry: B3LYP/ 6-31+g(d,p), and SCRF = SMD, solvent = CHCl₃, for more details see ESI \dagger), ³⁴ revealed that the diastereoisomer with (Z,3R,S') configuration predominantly exists as an equilibrium mixture of Conf1 (66%) and Conf2 (34%) conformers (Fig. 4), resulting in a weighted interproton distance of 3.7 Å, not so much different from that experimentally determined (3.1 Å). Conversely, the C(3) epimer is mainly populated by only one conformer Conf1 (99.9%), in which the H(6)-C(3)CH₃ distance is much longer (4.6 Å) than the experimental one, approaching the typical detection limit (5 Å) of NOESY experiments.

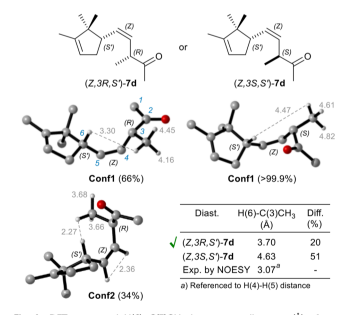


Fig. 4 DFT computed H(6)-C(3)CH₃ interproton distances (Å) of most stable conformers of C(3) epimers, compared to the experimental distance determined by quantitative NOE, for clarity some hydrogen atoms are omitted. In round brackets is reported the Boltzman conformer population.

Comparison of OYE2 chemoselectivity with other methodologies

Besides the stereoselective reduction of prochiral enones or enals (4-5),35 OYE2 has demonstrated its superiority in terms

Table 2 Reduction of 6 with different methodologies

		Prod. distrib. ^b (%)		
Reaction conditions a	Conv. ^b (%)	8	9	Other
OYE2, (cond. of Scheme 4, without ADH)	96	>99(87) ^d	_	
0.15 eq. Pd/C, e NH ₄ HCO ₂ , MeOH, 30 °C (ref. 36a)	>99	95	4	1
0.2 eq. RANEY®-Ni, pyridine, EtOH, rt	90	28	72	_
5 eq. Mg, MeOH, rt (ref. 36 <i>b</i>)	<i>f</i>	_	_	_
0.25 eq. Lindlar cat., H ₂ , MeOH, 50 °C	15	84	_	16
0.1 eq. Wilkinson cat., H_2 , g CH_2Cl_2 , 36 o C (ref. 36 c)	_	_	_	_
3.0 eq. NaBH ₄ , 5.0 eq. NiCl ₂ ·6H ₂ O, MeOH/H ₂ O, rt (ref. 36 <i>d</i>)	93	97	2	1

^a Catalyst/substrate ratio is in w/w. ^b Not isolated yield, by GC-MS. ^c Allylic alcohol, saturated alcohol, positional isomers. ^d Isolated yield without column chromatography. ^e 5% w/w. ^f Decomposition. ^g Pressure 1 atm.

of chemoselectivity compared to other catalytic procedures.^{5b} Indeed, in the synthesis of substrate 5c (Sandalore® precursor) we faced the problem of hydrogenating the α,β -unsaturated ketone 6 to ketone 8 preserving the campholenic double bond. Various reductive methodologies, known for their high chemoselectivity toward the reduction of conjugated C=C double bonds were tested, 36 the results of this screening are shown in Table 2. However, unlike in the case of OYE2 catalysed reduction, together with 8 a non-negligible quantity of overreduced product (9) and other products was consistently present, regardless of the method applied. In addition, the GC-MS analysis of a commercially available Sandalore® sample revealed the presence of small amounts of completely saturated alcohols (around 3% by GC-MS, ESI†). This result highlights that even in the current industrial process, achieving complete chemoselectivity remains an unresolved issue.5

Olfactory evaluation of Sandalore® stereoisomers

The olfactory evaluation of Sandalore® revealed that the isomer with (S) configuration at C(2) stereogenic centre and (R) configuration at C(3) centre fits more effectively into the human olfactory receptors compared to the other isomers, since a stronger and more pleasant sandalwood odour was perceived (Table 3). These results agree with earlier evaluations carried out (detailed olfactory assessments are reported into ESI,† in Scheme 4 the most appealing stereoisomers are high-

lighted in grey) and confirm the postulated olfactophore model for the sandalwood fragrances. The latter consists of a bulky lipophilic moiety (such as the campholenic ring or the norbornane of santalol) separated by a spacer from the hydroxyl polar group.³⁷ According to this model, the correct orientation of the osmophoric hydroxy group and the methyl substituent within the odour receptor is crucial for the perception of sandalwood scent, in contrast to the stereochemical configuration of campholenic ring that seems not to have influence.

Bio- vs. chemo-reduction process

Finally, we conducted a comparison between the Takasago reductive process for the stereoselective synthesis of the (S,S)-enantiomer of Brahmanol® and our bioreductive process, since both synthetic routes started from the common intermediate 4a. The Table 4 are reported the key technological and environmental factors and metrics for those two processes, including the simplified environmental factor (sEF), the complete environmental factor (cEF), and the EcoScale semi-quantitative analysis score.

Regarding the chemical reduction, in the first step the α,β -unsaturated aldehyde **4a** was transformed into the allylic alcohol intermediate by treatment with the very harsh reducing agent LiAlH₄ (0 °C in Et₂O).^{39,44} According to the patent, the allylic alcohol was then hydrogenated in presence of a Ru-

Table 3 Olfactory evaluation of Sandalore® diastereoisomers

Isomer	Odour description
(2R,3S,S')-1c	Prominent amber-woody essence with a slightly creamy and tobacco notes. After few minutes the impact seems to be stronger (8/10) than the commercial Sandalore®. ³⁸ Very tenacious, it holds the scent consistently over time (7/10).
(2S,3S,S')-1c	Unpleasant head impact $(6/10)$. After a few minutes, although remaining unpleasant, the intensity of the note drops significantly compared to $1c-(2R,3S,S')$, $(3/10)$.
(2S,3R,S')-1c	Impact 8/10. Creamy appearance reminiscent of sandalwood. Impressively tenacious even over time 9/10 (the most appealing of all diastereoisomers).
(2R,3R,S')- 1c	Delivering a notable impact at $7/10$, it shares an olfactory resemblance to $1c-(2R,3S,S')$, albeit slightly creamier and less dry. However, its tenacity falls short in comparison, scoring approximately $3/10$ tenacity.

Table 4 Technological and environmental factors comparison: chemovs. bio-reduction

$$R = (S)$$

$$Cat. = Ru_2Cl_4[(R)-Tol-BINAP]_2 NEt_3$$

$$R = (S)$$

$$Aa O O O H$$

$$Cat. = Ru_2Cl_4[(R)-Tol-BINAP]_2 NEt_3$$

$$R = (S)$$

$$Aa O O O O O O O O$$

	Chemic	Bioreduction ^b	
Factors	LiAlH ₄ ³⁹	${\rm H_2}^{7b}$	OYE2 + Evo440
T (°C)	0 to rt	80	30
P (atm)	1	48	1
Substrate conc.	0.4	1.3	0.2
[M]			
Time (h)	3	24	24
Catalytic	No	Yes	Yes
Solvent ⁴⁰	Et_2O	MeOH	Water
	hazardous	problematic	recommended
Technical set-up	Standard	High pressure	Standard
Work-up	Complex ^c	Very simple ^d	Simple ^e
Energy usage	Medium	Very high	Low
Resource		Ru	_
depletion			
Conversion (%)	f	>99	>99
Yield (%)	75 (84 and 85)		84
Chemoselec. (%)		f	
Stereoselectivity	88(S):12(R)		99(S):1(R)
Purification	Distil	Distillation ×2	
Waste	Non-biodegradable		Biodegradable
Safety			<u>(1)</u>
sEF	0.3	0.4	9.0
cEF	86	4	181
EcoScale	69/100	46/100	82/100
Average EcoScale		57	82

^a Starting from pinene with 96% *ee.* ^b Starting from pinene with 82% *ee.* ^c Ice quenching, acidification, extraction, drying, concentration under vacuum. ^d Concentration under vacuum. ^e Extraction, drying, concentration under vacuum. ^f Not reported.

based Noyori catalyst (48 atm, 80 °C, in MeOH).^{7b} Although the overall yield and the stereoselectivity were acceptable, they were significantly lower than those achieved in the one-pot bioreductive process (98% *ee vs.* 76% *ee*).

Undeniably, both the sEF and cEF metrics reveal that the bioreduction generates a significantly higher amount of waste per kg of product compared to the chemical reductions, with a difference of one order of magnitude, irrespective of the environmental factor considered (Table 4). This result is not surprising, considering that the transition metal catalysed hydrogenations, among all catalytic transformations, are those operating at highest substrate concentrations with perfect atom-economy.

However, these metrics are strictly related to the yield and to the reagents concentration, but do not take in account other important parameters such as: (i) selectivity (chemo- and/or stereoselectivity), (ii) ease of workup/purification, (iii) safety, (iv) energy consumption, (v) technical set-up (pot economy and/or requirement for high pressure equipment), (vi) nature of waste (biodegradable vs. non-biodegradable, presence of heavy metals), (vii) resource depletion; all of which have an impact on the environmental sustainability of the process.

In this context, a semi-quantitative analysis based on the EcoScale approach could offer valuable insights for a more comprehensive and fair comparison of the two routes beyond what the cEF and the sEF had provided. Using this approach, a maximum of points is arbitrarily assigned to the ideal process (100/100), where all technological and environmental factors are optimized. For each parameter that deviates from the ideal process, penalty points are assigned accordingly. The higher the score, the more sustainable is the methodology applied.

Remarkably, the biocatalytic process scored very well, 82/100 on EcoScale, while both chemo-reductions had a significantly higher number of point penalties, 69/100 and 46/100 (detailed calculations are reported in the ESI†). The main penalties were ascribable to a lower stereoselectivity, to the use of hazardous and highly flammable reagents (LiAlH $_4$ and H $_2$) and solvents (Et $_2$ O and MeOH), to the need of a highly energy-consuming reaction set-up (cooling or heating, high-pressure), and to the production of non-biodegradable wastes.

In addition, our multi-enzymatic process is carried out in a single vessel, in accord with the principle of pot economy. Such set-up simplifies reaction work-up and product isolation, both in terms of energy consumption and quantity of solvents used. On the contrary, in the Takasago process the incompatibility between the two reductive steps prevents their combination in a more appealing and sustainable one-pot synthesis, and even the telescoped transformation results impracticable. To this regard biocatalysis maintains its primacy over other methodologies, including also the more recent and efficient homogenous metal-based stereoselective catalysts. Indeed, to our knowledge, these catalysts have never been combined in a one-pot two hydrogenations-step capable of stereoselectively reducing enals or enones into the corresponding alcohols. 35

Lastly, the bioreduction is undoubtedly a safer process, as clearly indicated by the higher number of hazards and risks typically associated with the hydrogenations and the hydridebased reductions.

Experimental

A solution of substrate (3.0 mmol) in *i*-PrOH (0.8 mL) was added to a KP $_i$ buffer solution (pH 7.0, 50 mM, 15 mL) containing OYE2 (\approx 3 mg mL $^{-1}$, 5 mL), GDH (200 U), glucose (4.2 g), NADP $^+$ (20 mg). The mixture was incubated for 10–12 hours in an orbital shaker (150 rpm, 30 °C). The reaction was monitored by TLC until complete conversion. Eventually, more enzymes were added to increase the conversion. After 10–12 hours ADH (Evo270 or Evo440, 30–40 mg) and NADP $^+$ (10 mg) were added to the reaction mixture. After 10–12 hours XAD-1180 resins (5.0 g) were added to the reaction

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mixture and left to shaker for 30 minutes. The mixture was filtered into a porous filter (porosity 0) and the resins were washed several times with EtOAc (15 mL \times 4). The combined organic phase was washed with water (10 mL), dried over Na₂SO₄, and the solvent was removed under reduced pressure. The crude material was submitted to silica gel column chromatography purification and then distilled by bulb-to-bulb apparatus affording alcohol 1.

Conclusions

Some of the most important commercial sandalwood fragrances were prepared without employing transition metal catalysed hydrogenations, and even the use of metal hydride based carbonyl reductions was very limited (preparation of 4b and 5b substrates). Such reductive methodologies were proficiently replaced by bioreductions, combing ER and ADH enzymatic activities in a one-pot two-step process. The yield and especially the chemo- and the stereoselectivity achieved through biocatalysis compare favourably with the actual industrial procedures.^{5a} In addition, the most powerful and appealing stereoisomers were synthesized, with good diastereomeric ratios, by stereoselective reduction of methylene precursors 5.

A fair comparison of the technical, safety and environmental factors, including the EcoScale semi-quantitative analysis, applied to the stereoselective synthesis of Brahamanol® clearly showed the superior eco-friendliness of the bioreductive approach with respect to the traditional chemo reductions. Lastly, such development should open an appealing perspective in the field of fragrance chemistry.

Author contributions

F. G. G. conceived and designed the experiments. M. C. C., D. M., L. D. M., F. P., F. G. G. and D. F. performed experiments and analysed data. F. G. G. prepared the manuscript, all authors equally contributed to the revision.

Conflicts of interest

There are no conflicts to declare.

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