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A synthetic biology and green bioprocess approach to recreate agarwood sesquiterpenoid mixtures

Agarwood trees are used as incense and perfumes because of their fragrant oxygenated terpenes. Increasing consumer demand is endangering these trees. This work characterizes agarwood chemical diversity, uses synthetic biology to metabolically engineer green algae to produce terpenes and presents a green bioprocess to generate diverse oxyfunctionalized sesquiterpene mixtures like those from agarwood, but using CO_2 as an input. The process could be a sustainable source of fragrances and relieve some pressure on the endangered plants.

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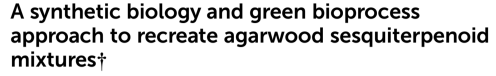


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Sergio Gutiérrez, Da Sebastian Overmans, Da Gordon B. Wellman, Da Vasilios G. Samaras, Db Claudia Oviedo, Dc,d Martin Gede, Dc,e Gyorgy Szekely Dc,d,e and Kyle J. Lauersen D*

Certain endangered Thymelaeaceous trees are major sources of the fragrant and highly valued resinous agarwood, comprised of hundreds of oxygenated sesquiterpenoids (STPs). Despite growing pressure on natural agarwood sources, the chemical complexity of STPs severely limits their synthetic production. Here, we catalogued the chemical diversity in 58 agarwood samples by two-dimensional gas chromatography—mass spectrometry and used synthetic biology to produce subsets of the complex STP mixtures found in agarwood. We improved STP yields in the unicellular alga *Chlamydomonas reinhardtii* by 25-fold through combinatorial engineering to biosynthesise nine macrocyclic STP backbones found in agarwood. A bioprocess following green-chemistry principles was developed that exploits 'milking' of STPs without cell lysis, solvent—solvent STP extraction, solvent—STP nanofiltration, and bulk STP oxy-functionalisation to obtain terpene mixtures like those of agarwood. This green bioprocess allows total solvent recycling and continuous production. Our synthetic biology and green bioprocess approach enables a more sustainable synthesis of complex, fragrant terpenes as an alternative to their exploitation from natural sources.

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Introduction

Agarwood is the common name given to the fragrant resinous woods derived from trees in the Thymeleaceae family, including *Aquilaria*, *Enkleia*, *Aetoxylon*, *Gonystylus*, *Wikstroemia*, and *Gyrinops*. Indigenous to Southeast Asian countries, these trees are prized for the complex fragrances of their resins that originate from multiple oxygenated and non-oxygenated sesquiterpenoids. These resins form after physical wounding of the tree, and their unique chemical profile develops over periods up to several decades. The use of agarwood pro-

The chemical composition of agarwood is species-specific and often unique to individual plants, depending on certain abiotic factors. ^{10,11} The main fragrant compounds in agarwood are sesquiterpenoids (STPs), an ecologically important class of 15-carbon atom terpenoid compounds that play a crucial role in cloud formation. ¹² These STPs originate from at least seven distinct primary chemical skeletons: agarofuran, agarospirane/vetispirane, cadinene, eremophilane/valencene, eudesmane/selinene, guaianes, and prezizanes. ^{3,10,13} Both abiotic and biotic factors govern their creation *in situ* where oxygenation of STPs to derivatives occurs, further shaping their individual scent profiles. The tree itself serves only as the vehicle and

ducts for incense, perfumes, medicines and ornamentation has been recorded across cultures and centuries.^{3–5} Common applications are burning wood chips as incense and steam-distillate oils as perfume ingredients.^{4,6} In particular, agarwood products were valued at \$US 44 billion within the global perfume market.⁷ Demand for agarwood has rapidly outpaced the supply of slow-growing natural sources, leading to the inclusion of 13 out of 20 *Aquilaria* species in the Appendix II of the Convention on International Trade in Endangered Species of Wild Fauna and Flora (CITES) as potentially threatened with extinction.⁸ Despite global efforts to regulate trade and develop alternative production modes through silviculture or tissue culture, the clandestine sourcing of wild trees remains an issue that further pressures their existence.^{4,9}

^aBioengineering Program, Biological, Environmental Sciences and Engineering Division (BESE), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia. E-mail: kyle.lauersen@kaust.edu.sa

^bAnalytical Chemistry Core Lab (ACL), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia

^cAdvanced Membranes and Porous Materials Center (AMPM), Physical Science and Engineering Division (PSE), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia

^dEnvironmental Science and Engineering Program, Biological, Environmental Sciences and Engineering Division (BESE), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia

^eChemical Engineering Program, Physical Science and Engineering Division (PSE), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia † Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d3gc03708h

Paper

matrix for chemical biosynthesis and resin accumulation, such that STP biosynthesis by metabolic engineering could be an alternative means for production of fragrant chemicals. ^{4,14} However, biochemical pathways leading to agarwood STPs are complex, poorly understood, and can be non-enzymatic in nature. So far, only guaiene STP synthases (STPSs) have been identified from *Aquilaria crassna* or *A. sinensis* and enzymatic routes to other primary STPs skeletons found in agarwood are not yet identified. ^{11,14–16}

Although the biochemical pathways to agarwood STPs are not elucidated, the diversity of natural product chemistry found in other organisms can provide alternative starting points to recreate agarwood STP chemical complexity. The primary carbon rings of agarofuran, cadinene, valencene, and selinane mirror STPs biosynthesised by valencene, ¹⁷ valerianol, ¹⁸ cadinol, ¹⁹ and aristolochene ²⁰ synthases, whereas zizaene synthase ²¹ produces a prezizane-like STP. This existing repertoire of known STP synthases from plant and fungal species can produce STP mixtures that largely replicate the primary STP skeletons in agarwood.

Here, we aimed to reproduce some of the chemical complexity of agarwood STPs using synthetic biology and a bioprocess following the principles of green chemistry. To inform the engineering approach, we first catalogued the chemical complexity of STPs in 58 agarwood samples using high-resolution, two-dimensional (2D) gas chromatography-time-of-flight mass spectrometry (GCxGC-TOF/MS). We then engineered unicellular algae as a sustainable cell chassis to produce nine distinct STP chemical skeletons widely found in agarwood. A lowenergy 'microbial-milking' technique was exploited to extract the STPs from growing algal cells, and energy-efficient solvent nanofiltration was used to concentrate the extracted STPs. Finally, we used green chemical reactions on these STP skeletons to generate a complex mixture of 103 oxygenated STPs typically found in agarwood. This green bioprocess employs an autotrophic host that produced chemicals from carbon dioxide as an input and a low-energy, gentle STP-harvesting technique that enables solvent recycling. Our process represents a sustainable alternative source of fragrant terpene mixtures compared to the harvest of agarwood trees.

Results

Chemical characterisation of agarwood and distillates

The *Aquilaria* sp. wood and distillate samples investigated here originate from various Southeast Asian countries (Fig. 1a and ESI Table 1†). Wood samples (n=36) exhibit a range of colours, with darker tones (14 to 18 based on Gardner colour scale) indicating higher resinous content (Fig. 1b, ESI Table 1, and ESI Fig. 1, 2†), while the distillate oils (n=22) exhibit varying amber-brown shades (Gardner scale 6 to 18) (Fig. 1c, ESI Table 1 and ESI Fig. 1, 2†). Representative 2D GCxGC-TOF/MS analyses of wood (sample 34) and distillate (sample 39) are presented in Fig. 1d and e. Comprehensive data for all samples can be found in ESI File 1.† Wood samples and distillate oils

contain a combination of monoterpenes, sesquiterpenoids (STPs), functionalised sesquiterpenoids (fSTPs), and carboxylic acids. Each sample was unique in its chemical profile, but 298 STPs and fSTPs were common across agarwood samples and 338 STPs and fSTPs were common to all distillate samples (ESI Tables 2 and 3†). Various STP skeleton types were found within the samples, including: agarofurans (α -agarofuran), agarospiranes (isoagarospirol), eudesmanes (isolongifolene, eudesmol, copaene, selinene, maaliene), eremophilanes (valencene, aristolochene, nootkatone, aristolene), guaianes (α -guaiene, β -guaiene, δ -guaiene, rotundone, viridiflorol, ledol, gurjunene, arodadendrene, guaiol, cedrol, patchoulene), cadinanes (τ -cadinol) and prezizanes (prezizaene, zizaene). Additional skeleton structures were identified, including patchoulol, caryophyllene, santalol, and bisabolol (ESI Tables 2–4, ESI Files 1–3†).

The identified sesquiterpenoid skeletons, spanning from the aromatic agarofurans to the complex prezizanes, offer a detailed insight into agarwood products and their chemistry. Compounds such as α -guaiene, δ -guaiene, and isolongifolene, expected from earlier research, were confirmed, indicating the reliability of our analytical approach. Although, many compounds were identified across all agarwood and its distillates, each sample had a distinct chemical profile. These detailed observations increase our understanding of the natural chemical complexity of agarwood samples.

Optimising algal metabolism to biosynthesise heterologous STPs

Chlamydomonas reinhardtii is an emerging algal host for metabolic engineering and heterologous terpene production. ^{22,23} Enhanced heterologous terpenoid yields have been demonstrated from this green alga following metabolic engineering. ^{24–26} These include the knockdown of native squalene synthase (CrSQS-k.d.) which increased farnesyl pyrophosphate pools in the cytoplasm and led to improvements in heterologous bisabolene or patchoulol yields when their synthases were expressed. ^{24,26} Disruption of carotenoid homeostasis by introduction of a beta-carotene ketolase to biosynthesise ketocarotenoids in the plastid led to pronounced increases of heterologous isoprene biosynthesis²⁵ (Fig. 2a, right panel). Here, the effects of these two interventions alone or in combination were quantified in the alga using the fragrant STP patchoulol as a reporter of biosynthetic activity (Fig. 2b).

When expressed and localised in the algal cytoplasm, the patchoulol synthase from *Pogostemon cablin* (*PcPS*) produced 5.4 ± 2.1 fg per cell patchoulol. Biosynthesis of ketocarotenoids and perturbation of carotenoid homeostasis through *CrBKT* overexpression led to higher patchoulol levels of 41.2 ± 5.9 fg per cell, representing an 8-fold increase over *PcPS* expression alone. Suppressing endogenous *CrSQS* through artificial micro-RNA in a strain expressing *PcPS* further increased patchoulol yield to 71.7 ± 7.8 fg per cell (a 12-fold increase over a strain expressing *PcPS* alone). The combination of *CrSQS*-k.d. and *CrBKT* overexpression in a strain expressing *PcPS* gave patchoulol yields of 132.7 ± 41.1 fg per cell ($2215 \mu g L^{-1}$ culture, ESI Fig. 4, ESI Table $5\dagger$). This strategy resulted in a

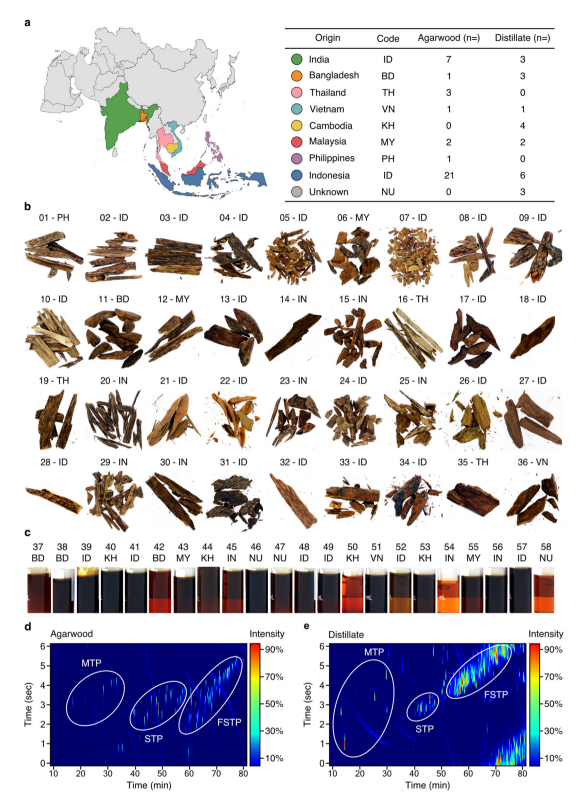


Fig. 1 Comprehensive chemical characterisation of agarwood samples and distillates. (a) Left: map of Asia, colour-coded to represent different origins of agarwood samples. Right: countries, their ISO 366 codes, and the total samples analysed from each, further categorised into agarwood wood chips ("bakhour") and steam distillate oils ("oudh"). (b and c) Photographs of agarwood (b) and distillate samples (c). (d) 2D chromatogram of wood-extract sample (34) with three primary chemical clusters circled. (e) 2D chromatogram for distillate sample (39). MTP: monoterpenoids; STP: non-functionalised sesquiterpenoids; fSTP: functionalised sesquiterpenoids. 2D chromatograms for each sample shown are found in ESI Table 1.† ESI Tables 2 and 3† provide summaries of common STPs identified in agarwoods and distillates, respectively. ESI Files 1-3† contain the supporting data for GC-TOF/MS, GC-MS/FID, and GCxGC-TOF/MS, respectively.

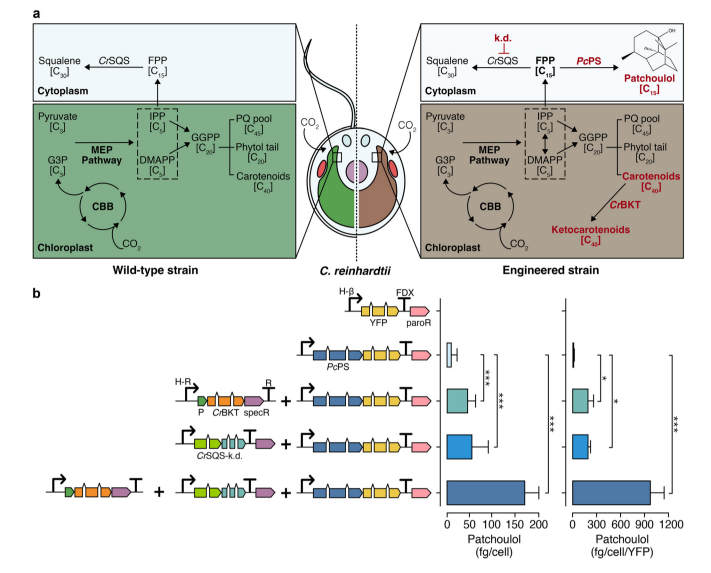


Fig. 2 Native and engineered terpenoid pathway in *C. reinhardtii* for enhanced sesquiterpenoid biosynthesis. (a) Left: illustration of the endogenous terpenoid biosynthetic pathway in wild type *C. reinhardtii*. Right: representation of an engineered *C. reinhardtii* strain characterised by absent flagella, cell-wall deficiency, and modifications of its terpenoid biosynthesis pathway known to improve heterologous STP production. Engineered changes are highlighted in red. (b) Left: depiction of algal-optimised transgenes including mVenus (yellow fluorescent protein, YFP), PcPS-mVenus fusion, CrBKT, and CrSQS-k.d., transformed into C. reinhardtii. Right: yield of patchoulol from each engineered construct combination. Abbreviations: k.d.: knockdown, PcPS: P. cablin patchoulol synthase; CrBKT: C. reinhardtii β -carotene ketolase; CrSQS: C. reinhardtii squalene synthase amiRNA expression construct, FPP: farnesyl pyrophosphate, G3P: glyceraldehyde β -phosphate, β -carotene ketolase; β -carotene

25-fold improvement in patchoulol yields over PcPS alone and approximately a 2-3-fold enhancement over each individual strategy (ESI Table 5†), one-way ANOVA (F(6, 2) = 158, p < 0.05). These yields were higher than any previous single engineering event for patchoulol production to date under the growth conditions tested. No statistically significant growth changes were detected in any strain grown photoheterotrophically with acetate (F(3, 20) = 0.035, p = 0.99) or photoautotrophically with CO₂ (F(3, 20) = 0.265, p = 0.84) (ESI Fig. 5†). A strain with both

*Cr*BKT overexpression and *Cr*SQS-k.d. was used in all subsequent STP production experiments.

Biosynthesis of diverse STPs in C. reinhardtii

Transformation and expression of 21 STPS isoforms in various fusions to the yellow fluorescent protein (YFP) reporter were performed, along with a YFP reporter alone control (23 plasmids in total, Fig. 3a and ESI Table 6†). Due to non-homologous DNA integration in the algal nuclear genome, random

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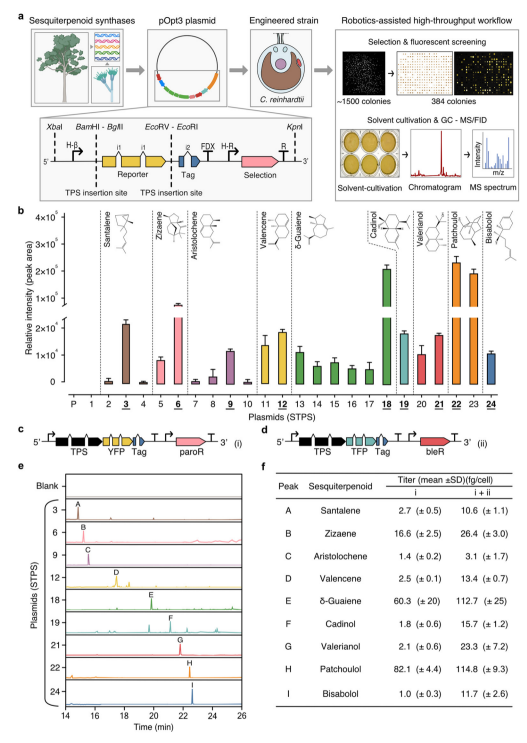


Fig. 3 Biosynthesis of diverse STPs in C. reinhardtii through sesquiterpenoid synthase isoforms expression. (a) The research workflow depicts the progression from selecting sesquiterpenoid synthases across various species for expression in C. reinhardtii. Isoform integration location into the pOpt3 in fusion to fluorescent reporter and robotics-assisted colony screening. Top-right box: depicted is a transformant plate one week after transformation, organised transformant colonies and YFP signal indicating synthase-reporter fusion expression. Brightest colonies were screened for expression in replica-plate assays with dodecane solvent overlay, which could be separated from the culture by gravity and directly used for analysis in GC-MS/FID. (b) Isoform performance was quantified from the most highly expressed transformants per STP (n = 25). Those chosen for further analysis are underlined (c). An illustrated vector structure of the STPS-YFP with paromomycin resistance and (d) -mTFP1 with bleomycin resistance. (e) GC-MS/FID chromatograms for each target STP. Annotated peaks include: A - santalene, B - zizaene, C - aristolochene, D - valencene, E - δ -guaiene, F - cadinol, G - valerianol, H - patchoulol, I bisabolol. (f) STP titres of single and double transformations. Abbreviations: YFP: mVenus yellow fluorescent protein; TFP: mTFP1 cyan fluorescent protein; paroR and bleR: paromomycin and bleomycin resistance, respectively; STPS: terpene synthase; Tag: strepII epitope. Chevrons i1/i2: intron addition to codon-optimised transgenes to enable strong algal expression. For a comprehensive list of plasmids, refer to ESI Table 5.† Detailed calculations of productivity and cell concentrations are in ESI Tables 6-8.† ESI Files 5 and 6† provide supporting data for GC-MS/FID and genetic constructs, respectively.

Table 1 Sesquiterpenoid synthase isoforms

Isoform number	Origin	Sesquiterpenoid	UniProt	Ref
1	Santalum album	Santalene	E3W202	54
2	Clausena lansium	Santalene	E5LLI1	54
3	Cinnamomum camphora	Santalene	A0A7S6HRZ4	55
4	Chrysopogon zizanioides	Zizaene	AJQ30127	37
5	Penicillium expansus	Aristolochene	C0LW28	56
6	Penicillium camemberti	Aristolochene	A0A0G4NXL3	57
7	Penicillium chrysogenum	Aristolochene	A0A167QKS4	58
8	Penicillium roqueforti	Aristolochene	Q03471	59
9	Citrus sinensis	Valencene	D0UZK2	60
10	Callitropsis nootkatensis	Valencene	S4SC87	61
11	Aquilaria crassna	δ-Guaiene	D0VMR6	15
12	Aquilaria crassna	δ-Guaiene	D0VMR8	15
13	Aquilaria crassna	δ-Guaiene	D0VMR7	15
14	Aquilaria sinensis	δ-Guaiene	K9MQ67	62
15	Aquilaria sinensis	δ-Guaiene	K9MNV6	62
16	Aquilaria sinensis	δ-Guaiene	K9MPP8	62
17	Lavandula angustifolia	τ-Cadinol	U3LW50	63
18	Camellia sinensis	Valerianol	A0A167V661	18
19	Camellia hiemalis	Valerianol	A0A348AUV5	18
20	Pogostemon cablin	Patchoulol	$Q49SP3^a$	64
21	Cynara cardunculus	Bisabolol	XP_024994640	65

^a Mutation C415F and H454A.

position effects change expression dynamics across a population of transformants and large numbers of colonies must be screened to identify robust over-expressing transformants. Here, high-throughput robotics assisted handling was coupled to plate-level fluorescent protein expression screening for at least 384 transformants per plasmid. This screening allowed identification individual transformants for each target plasmid with robust STPS-FP fusion expression. The STPSs selected for expression were chosen based on their reported ability to make primary STPs skeletons commonly found in our analysis of agarwood samples. These STPSs biosynthesise nine distinct STP products (Fig. 3b). Santalene, zizaene, aristolochene, valencene, δ-guaiene, cadinol, valerianol, patchoulol, and bisabolol, were detectable as new GC-MS/FID peaks in dodecane overlays obtained from transgenic strains compared to overlays from control strains (Fig. 3b, e and ESI File 4†). Each STPS isoform varied in the level of sesquiterpenoids they biosynthesised (Fig. 3b). Zizaene synthase had higher activity when YFP was placed at the N-terminal (plasmid 8, Fig. 3b), similar to reports of SUMO-tagged variants in E. coli.21

The optimal isoform with highest target-compound production was identified for each STPS (Fig. 3b). Subsequently, a second transformation was performed with the same STPS (Fig. 3c) fused to the monomeric teal (cyan) fluorescent protein 1 (mTFP1) reporter with a different selection marker (Fig. 3d). Secondary transformation was used to increase heterologous STPS enzyme titter in the cell to catalyse increased target STP biosynthesis. STP yield improvements were observed in all strains after secondary transformation (Fig. 3f), with an average 5.3-fold improvement in titre for each product (ESI Tables 7–9†). Additional STP peaks were also identified in these extracts by GCxGC-TOF/MS, representing known side products of the individual synthases (ESI Table 10†).

A green bioprocess for functionalised STPs

To achieve sustainable production of complex STP mixtures present in agarwood, we developed a bioprocess design that minimised waste and could be operated at room temperature with only non-hazardous reagents and, therefore, follows the principles of green chemistry (Fig. 4a). STP-producing algal cells were subjected to culture-solvent two-phase cultivations with a fluorocarbon underlay that allowed simple gravity separation of the fluorocarbon liquids containing STPs. Algal cultivation was followed by liquid-liquid extraction of STPs from the fluorocarbon solutions with ethanol, yielding individual STP-ethanol solutions, which were combined to form a STP mixture in ethanol (Fig. 4a and b). Low-energy, pressure-driven, organic-solvent nanofiltration (OSN) was then applied at room temperature, using only the pressure derived from a gas cylinder to drive ethanol through a nanofiltration membrane (Fig. 4a). OSN enabled concentration of STPs in ethanol by 53 times without evaporative loss (Fig. 4b).

The hydroboration-oxidation technique is commonly used in organic synthesis for alkenes conversion to alcohols. Compared to halogenation, it results in consistent stereochemistry and minimal waste generation and is within the principles of green chemistry. The non-functionalised cyclic primary STP structures produced by algae were then subjected to hydroboration reactions to cause bulk hydroxy-functionalisation of double-bond locations within the STP backbone (Fig. 4a). Reactions used either borane (BH₃) or 9-borabicyclo [3.3.1]nonane (9-BBN) reagents and subsequent treatment with hydrogen peroxide (H_2O_2) to furnish the hydroxy-functionalised STP backbones, thereby mimicking natural chemical-oxygenation processes (Fig. 4a). The nine STPs observed in chromatograms were successfully hydroxylated, or otherwise oxyfunctionalized, to yield α -santalol/ β -santol, cedrenol, 12-nor-

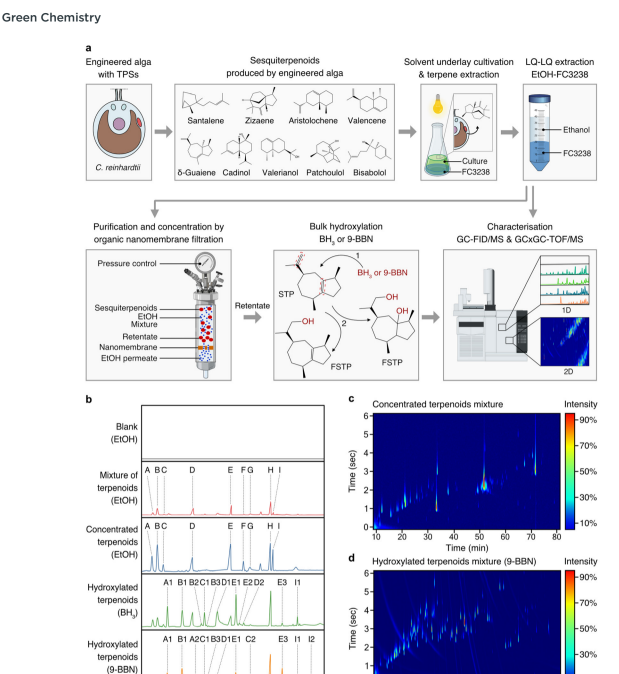


Fig. 4 A green bioprocess workflow for the generation of agarwood-related sesquiterpenoids. (a) STPs biosynthesised by engineered algae can be milked from the growing culture using heavy perfluorinated solvent (FC3283) underlays. STP-containing fluorocarbons can be separated from culture and STPs rapidly transferred to ethanol through liquid-liquid extraction from fluorocarbons. STPs can be concentrated in ethanol at room temperature in gaspressure-driven nanofiltration cells wherein a nanoporous membrane retains STPs and allows permeation of ethanol. Concentrated STPs in ethanol can be subjected to hydroxylation reactions using BH₃ or 9-BBN to increase molecular species diversity. Analysis throughout the process can be performed by GC-MS and GCxGC-TOF/MS. (b) One-dimensional (1D) chromatograms of analysed process fractions, with STP skeletons labelled A to I and their respective derivatives after functionalisation: A: santalene, A1: santalol, A2: bergamotene, B: zizaene, B1: cedrenol, B3: 12-nor-preziza-7(15)-en-2-one, B3: khusimol, C: aristolochene, C1: aristolone, C2: dihydrokaranone, D: valencene, D1: nootkatol, D2: nootkatone, E: δ-quaiene, E1: quaiol, E2: globulol, E3: quaia-1(10),11-dien-15,2-olide, F: cadinol, G: valerianol, H: patchoulol, I: bisabolol, I1: bisabolol oxide B, I2: bisabolol oxide A. (c) Two-dimensional (2D) chromatogram depicting the starting mixed STPs. (d) 2D chromatogram of STPs post-hydroxylation. Abbreviations: BH3: borane; 9-BBN: 9-borabicyclo [3.3.1]nonane; nfSTP: non-functionalised sesquiterpenoid; fSTP: functionalised sesquiterpenoid; H₂O₂: hydrogen peroxide; GC-MS/FID: gas chromatography with flame-ionisation detection and mass spectrometry; GCxGC-TOF/MS: 2D gas chromatography with time-of-flight mass spectrometry; EtOH: ethanol; 1: step one; anti-Markovnikov hydroboration of the STP, BH₃ adds boron to the alkene in the less substituted carbon. 2: step two, a three-part oxidation process: initiation by H2O2 ion donation to the trialkyl borane, rearrangement leading to a hydroxide ion removal, and oxidation completion by reaction with aqueous NaOH, forming alcohol and sodium borate. ESI Tables 9 and 10† provide detailed GCxGC-TOF/MS data for panels (c) and (d), ESI File 4† contains supporting raw data for (b). ESI File 7† contains supporting raw data for GCxGC-TOF/MS.

16

20

Time (min)

10%

40

Time (min)

60

Fig. 5 Summary of synthesis of agarwood-like sesquiterpenoids *via* an integrated algal-chemical methodology. Schematic representation of sesquiterpenoids (STPs) analogous to agarwood, synthesized through a bioengineered algal pathway and characterized by GC–MS. The precursor, farnesyl pyrophosphate (FPP), undergoes enzymatic conversion by various STP synthase isoforms (numerical annotations) to yield distinct STPs labeled A–I. Subsequent hydroxy-boron oxidation produces functionalized derivatives, denoted by additional numbers or asterisks. A: santalene, A1: santalol, A2: bergamotene, B: zizaene, B1: cedrenol, B3: 12-nor-preziza-7(15)-en-2-one, B3: khusimol, C: aristolochene, C1: aristolone, C2: dihydrokaranone, D: valencene, D1: nootkatol, D2: nootkatone, E: δ-guaiene, E1: guaiol, E2: globulol, E3: guaia-1(10),11-dien-15,2-olide, F: cadinol, G: valerianol, G*: eudesmol, G**: jinkoheremol, H: patchoulol, H*: pogostol, and I: bisabolol, I1: bisabolol oxide B, I2: bisabolol oxide A. ESI,† including mass spectra and raw data from GC-MS/FID, are in ESI Tables 2, 3, 11–13, and ESI File 5.†

preziza-7(15)-en-2-one, khusimol, aristolone, dihydrokaranone, nootkatol, nootkatone, guaiol, globulol, and bisabolol oxide (Fig. 4b and ESI Tables 10 and 11†). This mixture was clear and did not have the amber colour associated with traditional agarwood distillates which are largely derived from other chemicals in the resin (Fig. 1). Using GCxGC–TOF/MS quantification, STP diversity increased in the 9-BBN reaction: from an initial 42 to 103 chemical species (Fig. 4c, d and ESI Tables 10 and 11†). A total of 41 agarwood-STPs derivatives matched compounds typically found in natural agarwood terpene profiles, including α -santalol/ β -santol, cedrenol, guaiol, and bisabolol oxide B (Fig. 5 and ESI Tables 2, 3, 10–13†). Overlap of chemical species in this mixture and agarwood or its steam distillate are compared in figure in ESI Table 12.†

Discussion

Agarwoods have captivated human interest for centuries due to their complex and diverse fragrances.²⁸ However, the

increasing demand for their resinous wood has placed an unprecedented pressure on their natural sources, particularly older trees native to Southeast Asian forests, and the requirement of tissue wounding to promote resin formation further increases this pressure. ^{3,8} In this sample set, we observed that each agarwood and distillate sample contained hundreds of unique STPs and share a subset of these chemicals in varying abundances (ESI Files 1–3 and ESI Tables 2–4†). There was no obvious correlation between agarwood prices, origin locations, colour, or sample type, indicating that their market value is subjective, largely based on perceived fragrance or rarity (ESI Table 1, ESI Fig. 2, 7†).

We sought to develop a sustainable approach using metabolically engineered algae to heterologously biosynthesise the STP backbones found in agarwood, followed by oxy-functionalisation using green chemistry as an alternative to tree sources of STPs. Our goal was to generate not only a microbial production alternative to agarwood STPs, but also to develop a sustainable process that generates minimal waste and uses only gentle reagents within the principles of green chemistry

and bioprocess design. 29,30 We opted for a microalgal host as our bioengineering platform due to its ability to thrive on the nutrients found in wastewaters and CO_2 as a sole carbon source. 31 This not only enhances the sustainability of the process but also potentially facilitates the conversion of wasted materials into valuable resources, aligning with the principles of circular economy. Our downstream bioprocess uses only low-energy methods for STP extraction and concentration; that is, microbial-milking of STPs during cultivation, liquid-liquid STP transfer to ethanol, and nanofiltration for STP concentration. Using boron and $\mathrm{H}_2\mathrm{O}_2$, we generated complex, functionalised STP mixtures that resemble those found in agarwood samples in nature.

Green Chemistry

Endogenous isoprenoids in C. reinhardtii are supplied exclusively by the methyl-D-erythritol phosphate (MEP) pathway localised to the plastid (Fig. 2a).³² That algae rely on this pathway for essential photosynthetic carotenoids implies an inherent capacity to channel carbon fixed by photosynthesis to isoprenoids, which could be tapped for heterologous terpene biosynthesis. The first reports of micrograms of heterologous terpenes per litre of algal culture²⁷ have now been superseded by hundreds of milligrams per litre, 25,33 owing to improved transgene design, 34,35 enhanced metabolic channelling, and high-cell-density cultivation strategies.^{26,36} Here, we demonstrate for the first time a combinatorial modification of two metabolic pathways simultaneously in C. reinhardtii by perturbing plastid carotenoid and cytoplasmic sterol biosynthesis in the same strain (Fig. 2). This combination improved target STP production by up to 25-fold over wild-type levels and enabled us to use this strain to produce other STPs of interest. Improvements in transgene optimisation and DNA synthesis capacity together with improved robotic handling and fluorescent screening methods of transformants has facilitated STP isoform benchmarking directly in alga (Fig. 3). This is a key step as isoform-performance testing is a role previously conducted only in fermentative microbes. Screening directly in alga enabled nuances of expression in this host to be ruled out directly when choosing a target isoform.

Terpene extraction from engineered microbes is a challenge in any bioprocess design. Microbes lack the specialised tissues of multicellular organisms to accumulate hydrophobic terpene products, necessitating a physical hydrophobic contactor be used as sink for terpene extraction during cultivation. This can be achieved in numerous ways: using resinous particles,³⁷ membrane contactors,38 and through biocompatible solventculture two-phase milking.^{17,39} We previously found that perfluorinated heat-transfer fluids (namely fluorocarbons) can be used as heavy solvents that sit underneath microbial cultures as an underlay for terpene milking.³⁹ We also found that accumulated terpenes in the fluorocarbons rapidly partitioned into ethanol in subsequent liquid-liquid extractions.³⁹ We leveraged this property to first milk STP skeletons from algal cultures and subsequently pool the ethanol extracts to make a starting mixture of STPs (Fig. 4). This extraction method is advantageous for its simplicity, operating at standard atmospheric conditions, and not requiring complex machinery to

yield STPs in ethanol. Although not yet demonstrated at larger scales, the process presented here could readily be incorporated into tubular photobioreactor or even raceway pond algal production facilities.

To create alternatives to fragrant agarwood-derived STPs, synthases for the central seven classes of common STP skeletons were required. However, only a handful of δ-guaiene synthase isoforms from *Aguilaria* trees have been reported. 11,15,40 We overcame this limitation by using terpene synthases from different plants and fungi that could produce substitute chemical skeletons to those detected in agarwood (Table 1). However, the fragrances of agarwood largely comprise oxy-functionalised derivative STPs. Terpene functionalisation in metabolic engineering is traditionally conducted by the concerted action of highly regio- and stereo-specific enzymes, such as P450s.41 The chemical complexity of agarwood terpene profiles is both the product of time and likely non-enzymatic chemistry, however, some P450s may be implicated in certain classes of functionalisation observed within.⁴² These P450s have yet to be biochemically characterised and would require expression optimisation gene-by-gene to participate in single-STP functionalisation in a microbe. By starting with a mixture of STP skeletons and subjecting them to a chemical process that catalyses oxygenation reactions at sterically accessible double bonds, we could bypass biological functionalisation to obtain complex STP mixtures (Fig. 4). In practice, this procedure mimics parts of the natural maturation process of STPs subjected to abiotic factors in resins over time and successfully reproduced numerous functionalised STPs typical of agarwood samples using a green-chemical process (Fig. 4, 5 and ESI Table 12†).

Traditional methods of oxidation or hydroxylation of STPs have often involved the use of heavy-metal catalysts⁴³ (such as chromium and mercury), strong oxidising agents44 (i.e., osmium tetroxide), toxic solvents⁴⁵ (i.e., benzene), or hazardous reagents⁴⁶ (i.e., chlorine)—all of which pose serious environmental and health risks. There are few means of achieving regio-specific hydroxylation of alkene sites using green chemistry. Here, we used organoboron reagents to introduce hydroxyl groups to the most accessible double bonds of the STP skeletons and eliminated use of heavy-metal reagents and hazardous by-products associated with conventional methods (Fig. 4b-d, 5 and ESI Table 11†). This represents a green chemistry solution to increasing the chemical diversity of starting terpene backbones. Coupling this green chemical step to a photosynthetic (green) cell host, and chemical milking strategy involving complete solvent recycling and low energy OSN concentration, allows classification of this as a complete green bioprocess to fragrant STPs.

Mono- and di-terpenes are also present in agarwood (Fig. 1 and ESI File 1†). Using more complex mixtures of terpenes obtained by additional metabolic engineering will undoubtedly increase the complexity achievable in synthetically functionalised mixtures. Here, we did not express specific α -humulene, β -eudesmol¹⁴ and vetispiradiene synthases, which could broaden the complexity of the starting mixtures,

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as would expression of mono and di-terpene synthases. The combination of possibilities for these mixtures is immense, offering wide scope for additional combinations of microbial engineered terpene production and synthetic chemistry to yield fragrant mixtures. Previous hype around novel terpene production techniques created instability in natural-product markets, leading to economic challenges within legitimate supply chains. ⁴⁷ This work showcases the potential of combining synthetic biology, sustainable bioprocess design and green-chemistry as a complementary alternative source of fragrant terpenes to natural agarwood and may alleviate some of its overharvest from the wild. Our approach should be considered alongside silviculture or emerging methods like tissue

In summary, we have developed a strategy following greenbioprocess design principles that combines metabolic engineering, terpene milking, and pharmaceutical-grade chemical processing with a green chemistry-functionalization step to obtain terpene mixtures that mimic some of the natural chemical diversity found in agarwood. Our set of 103 semi-synthetic engineered sesquiterpenoids partially captures the diversity found in agarwood and it will be possible to expand the chemical diversity by simply engineering production of other terpene backbones. Our approach presents a sustainable complement to legitimate sources of fragrances from agarwood, and offers a potential solution to the current demand that has surpassed natural supply for these culturally important goods. We demonstrate the feasibility of a cyclic, synthetic-biology-mediated and chemically complemented green bioprocess to generate functionalised STP mixtures.

Materials and methods

Agarwood sample collection and processing

culture as alternatives to fragrance sourcing.

Agarwood samples were procured from the old-town market "Al Balad" in Jeddah, Saudi Arabia (21.481° N, 39.187° E) in Winter 2023. The collection includes 36 different samples of agarwood chips, dried Aquilaria spp. wood, also known as "bahkour", and 22 agarwood steam-distillate oils, known as "oudh". The ages of the plants at harvest and origins of these samples cannot be accurately traced; however, many were labelled with their country of origin. The samples varied in price at the time of purchase, reflecting their purported rarity, the density of fragrant compounds, and complexity of aromatic notes (ESI Table 1†). It was determined that the appropriate organic solvent for extraction was acetone based on preliminary analysis with different solvents, and each agarwood sample was then diluted to obtain chromatograms with clear peaks for product detection and identification. All samples were processed within 16 weeks of being procured. The wood samples were weighed (1 g) and ground into a fine powder using a combination of freeze-thaw cycles with liquid nitrogen and mechanical grinding. The homogenised samples were immersed in 5 mL of 1:1 hexane: acetone and subjected to ultrasonic agitation at 40 °C for 8 h to facilitate terpenoid

extraction. The samples were then passed through a 0.2 μm filter to obtain clear solvent extracts that were evaporated under a nitrogen stream for 20 min to concentrate the terpenoids in the samples. Concentrated samples were resuspended in 500 μL acetone and stored at −20 °C until gas chromatography–flame-ionisation detector and mass spectrometry (GC-MS/FID) and two-dimensional gas chromatography with time-of-flight mass spectrometry (GCxGC-TOF/MS) analyses. All photographs were captured with a Canon EOS RP camera using a Canon RF 24–105 mm f/4–7.1 IS STM lens (Canon, Tokyo, Japan) and ColorChecker Passport (CCPP2, Calibrite LLC, DE, USA) used for colour calibration.

Algal strain cultivation

Chlamydomonas reinhardtii strain UPN22²⁶ was used for all experiments. C. reinhardtii UPN22 is a derivative of the UVM4 strain.⁴⁸ It has been genetically enhanced to use phosphite and nitrate as a sole source of phosphorous and nitrogen, respectively, to minimise contamination and maximise cell densities in cultivation.²⁶ Strains were cultured in Tris-acetate phosphite nitrate (TAPhi-NO₃,²⁶) liquid medium with shaking at 120 rpm or on solid agar and 150 μmol m⁻² s⁻¹ light intensity from a combination of cooland warm white LED tubes (light spectra as reported in²⁶). 500 mL algal cultures were agitated with stirring in Erlenmeyer flasks under the same growth conditions.

Plasmid construction, algal transformation, and screening for sesquiterpenoid synthase expression

Heterologous expression of sesquiterpenoid synthases (STPSs) in C. reinhardtii was achieved through synthetic transgene redesign based on the amino-acid sequences of each STPS, following previously established protocols.34,49 We selected 21 STPSs from various species encoding isoforms that yield nine different STP skeletons (Table 1). Targeted STPSs for aristolochene, δ-guaiene, santalene, valencene, valerianol, zizaene, τ-cadinol, bisabolol, and patchoulol were designed; all accession numbers are listed in Table 1. Their amino-acid sequences were used to generate algal-adapted nucleotide coding sequences using the Intronserter programme.34 This programme back-translates amino-acid sequences to frequently used codons, removes unwanted restriction sites, and systematically integrates the first intron of CrRBCS2i1 at a set distance to enable expression from the C. reinhardtii nuclear genome, as previously reported. 49,50 Algal-adapted coding synthesised and were sub-cloned sequences pOptimized_3 expression plasmids35 by Genscript (Piscataway, NJ, USA). Ketocarotenoid biosynthesis in alga was achieved by transformation with the pOpt2_CrBKT_aadA plasmid,51 and knockdown of C. reinhardtii squalene synthase (Uniprot: A8IE29) was achieved using the previously reported luciferaseartificial microRNA expression plasmid pOpt2_ cCA_gLuc-TAA_i3_ami_Spec.²⁴ Both plasmids confer resistance to spectinomycin as a selectable marker. Combining both cassettes into a single plasmid was unsuccessful. Co-transformation of both plasmids and selection on spectinomycin was combined with robotics assisted colony picking and plate-level screening of 768 transformants to find those with luciferase activity (indicative of SQS knockdown) and a brown-colony phenotype from *CrBKT*-mediated ketocarotenoid biosynthesis. All plasmid constructs used are listed in ESI Table 6† and the complete annotated plasmid sequences are available in ESI File 5.†

The glass-bead protocol was used to transform the nuclear genome of C. reinhardtii with a plasmid DNA.52 Each plasmid was linearised using restriction enzymes (XbaI + KpnI, Thermo Scientific FastDigest), and 10 µg of DNA was used for each transformation. Following an ~8 h recovery period in liquid TAPhi-NO₃ medium under low light, algal cells were plated on a selective medium with paromomycin (10 μg mL⁻¹), spectinomycin (200 μg mL⁻¹) or zeocin (15 μg mL⁻¹) antibiotics either individual or combinations of selection agents relative to each target plasmid. Plates were illuminated continuously for ~7 d before colony picking. A PIXL robot (Singer Instruments, Watchet, UK) transferred up to 384 colonies per transformation event to TAPhi-NO₃ agar plates. After an additional 3 d, a ROTOR robot (Singer Instruments) was used to replicate colonies onto new medium and plates containing amido black (150 µg mL⁻¹) for fluorescence screening as previously described.³⁵

All algal-optimised STPSs were expressed as fusions with mVenus (yellow fluorescent protein) or the monomeric teal (cyan) fluorescent protein 1 (mTFP1). Transgene expression of each STPS was determined by fluorescence imaging at the agar-plate level, as previously described. 27,35 Chlorophyll fluorescence was observed with 2 s of 475/20 nm excitation and 640/160 nm emission to show colony presence/absence on amido black-containing plates. Cyan-green fluorescence was captured with 420/20 nm excitation and 480/20 nm emission filters using 2.5 min exposure. Yellow fluorescent signals were captured with 504/10 nm excitation and 530/10 nm emission filters with 30-s exposures. Transformants displaying strong fluorescent-protein signals were selected and inoculated into 12-well plates containing 2 mL of liquid TAPhi-NO3 medium and grown with shaking at 160 rpm. Predicted molecular masses of the expressed heterologous STPS-FP fusions were verified using sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) in-gel fluorescence against the fusion-protein fluorescent reporter.

Heterologous sesquiterpenoid biosynthesis analyses

For each transformant, biosynthesis of heterologous STPs was assessed by gas chromatography-mass spectrometry (GC-MS/FID). Four individual transformants containing each plasmid with the highest fluorescence signals were selected, and solvent-culture two-phase living extractions were performed using a 10% v/v dodecane-culture overlay in 6-well plates, as previously reported. ^{17,53} Cultivations were performed in biological triplicate for 6 d in 4.5 mL TAPhi-NO₃ media with 500 μ L dodecane overlay. Phases were separated, culture samples were taken for cell-density analysis by flow cytometry as previously described, ²⁶ and dodecane was spun at 20 000 × g for 3 min, then 150 μ L clarified solvent was transferred into amber GC vials in triplicate prior to analysis.

Samples were analysed as previously described 26,38 using an Agilent 7890A gas chromatograph equipped with a mass spectrometer and a flame-ionisation detector (GC-MS/FID). The gas chromatograph comprises a 5975C inert MSD with a triple-axis detector and a DB-5MS column (30 m × 0.25 mm i. d., 0.25 µm film thickness). The injector, interface, and ionsource temperature profiles were set to 250 °C, 250 °C, and 220 °C, respectively. In splitless mode, 1 µL of the sample was injected using an autosampler (G4513A, Agilent). The column flow was constant at 1 mL min⁻¹, with helium as carrier gas. The initial GC oven temperature was set to 80 °C for 1 min, increased to 120 °C at 10 °C min⁻¹, raised to 160 °C at 3 °C min⁻¹, and to 240 °C at 10 °C min⁻¹, holding for 3 min. After a 12 min solvent delay, mass spectra were recorded using a scanning range of $50-750 \, m/z$ at 20 scans per second. Chromatograms were analysed with MassHunter Workstation software version B.08.00 (Agilent), and STPs were identified using the National Institute of Standards and Technology (NIST) library (Gaithersburg, MD, USA). Further identification was conducted with purified standard calibration curves ranging from concentrations of 1-1200 µM in dodecane of δ-guaiene (CAT#B942760), patchoulol (CAT#P206200), santalene (CAT#S15065), valerianol (CAT#V914000, Toronto Research Chemicals, ON, Canada), bisabolol (CAT#95426), valencene (CAT#06808), cedrene (CAT#22133, Sigma-Aldrich, MO, USA) (ESI Fig. 6†). For compound identification, retention-time acquisition, internal digital-library calibration, and method development, we used a set of 12 microampules containing a standard terpene mixture, which covered 98 terpenes at 1 mM in methanol (CAT# MSITPN101, MetaSci, ON, Canada, ESI Table 4†).

Two-dimensional Gas chromatography time-of-flight mass spectrometry (GCxGC-TOF/MS) analysis

Comprehensive two-dimensional gas chromatography time-offlight mass spectrometry (GCxGC-TOF/MS) analysis of agarwood and distillate extracts in acetone was performed using an Agilent 7890B gas-chromatography system equipped with a Zoex ZX1 cryogenic thermal modulator and a JEOL TOF MS (AccuTOF GCx-plus, JEOL, Japan). The GCxGC system featured a normal (non-polar × mid-polar) two-dimensional column configuration, comprising a first-dimension with a 30 m nonpolar HP-5MS UI capillary column (5%-phenyl-methylpolysiloxane) and a second-dimension column with a 2 m mid-polar BPX-50 capillary column (50% phenyl polysilphenylene-siloxane). We used helium (99.999%) as the carrier gas at a constant flow rate of 0.8 mL min⁻¹. The GCxGC-TOF/MS injector temperature was maintained at 300 °C with a 10:1 split ratio. The oven temperature was initially held at 80 °C for 1 min, then increased to 325 °C at 2 °C min⁻¹. The modulation period was set at 6 s with a pulse time of 0.35 ms. The mass spectrometer operated in electron ionisation (EI+) mode at 70 eV. Both the transfer-line and ion-source temperatures were maintained at 250 °C. The detector voltage of TOF was set to 2500 Volts, and data were acquired at a rate of 50 Hz. Mass

spectra were obtained within a mass-to-charge ratio (m/z) from 50 to 600.

Gas chromatography data analysis

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The analysis of terpenoid extracts followed a procedure similar to previously reported methods, ^{26,38,39} and qualitative analyses primarily relied on the retention index and match factor. The GC-MS/FID data were processed using the MassHunter Workstation software version B.08.00 (Agilent Technologies, USA). The identification of compounds was assisted by the NIST Mass Spectral Library Version 2.3 (National Institute of Standards and Technology, Gaithersburg, MD, USA). The mass-spectral data, derived from the GCxGC-TOF/MS analysis, were evaluated with GC ImageTM Version 2.9 software (Lincoln, NE, USA) and referenced against the NIST2020/EPA/NIH EI Mass Spectral Library. The spectral data were cross-referenced with library spectra to identify potential chemical structures, facilitated by calculating the match factor and probability, thereby generating a list of probable compound matches. The match factor directly compares the unknown mass spectrum peaks with those in the library spectra, indicating their similarity. In contrast, the probability determines the relative likelihood of the list of hits being accurate, assuming that the unknown spectrum is present in the library. Using these metrics provides the relative assuredness of matching chemical structures within the library spectra.

Sesquiterpenoid production and concentration

Transformants of best-performing STPS isoforms that accumulated the most of each heterologous sesquiterpenoid in screening conditions, were subjected to solvent-culture two-phase cultivations at 300 mL scale in TAPhi-NO3 medium using FC-3283 as a solvent underlay phase, as previously described.³⁹ FC-3283 is a perfluorinated amine that is inert and denser than water, forming an underlay to the culture and accumulating heterologous terpene products from the algae. After 6 days of cultivation, gravity and gentle centrifugation separated the fluorocarbon and cultures. After centrifugation to clarify, fluorocarbons were subjected to liquid-liquid extraction to partition accumulated STPs into an equal volume of 96% ethanol. The mixture was shaken for 16 h at room temperature at 200 rpm. Next, samples were again centrifuged gently to further separate the phases. FC-3283, after ethanol extraction, can be reused on algal cultures and is effectively recycled in this process. 500 µL aliquots from each phase were sampled and stored in separate GC vials at -20 °C for subsequent analysis and separation performance quantification.

For each of the nine STPs biosynthesised by the algae, 20 mL 96% ethanol-containing STPs were generated. The ethanol fractions were pooled and subjected to organic solvent nanofiltration (OSN) in a dead-end cell to concentrate the terpenes in ethanol without evaporative losses. A Duramem solvent-resistant membrane (Evonik, Germany) with a nominal molecular weight cut-off value of 300 g mol⁻¹ suitable for STP retention and chemical compatibility with ethanol was selected. The 200 mL ethanol-sesquiterpenoid mixture was

loaded into the OSN chamber containing a 16 mm membrane disc and subjected to 20 bar pressure delivered by $\rm CO_2$ as an inert gas, to drive the ethanol phase through the nanofiltration membrane at 2.84 L m⁻² h⁻¹ flux. While ethanol permeated the membrane, STPs were kept in the retentate due to their molecular weight and consequently concentrated in the ethanol. The permeate ethanol is suitable for recycling and can be reused in subsequent liquid–liquid extraction processes.

Bulk hydroxylation of sesquiterpenoid backbones

To selectively introduce hydroxyl groups at the double bonds present in the terpenes, hydroboration-oxidation reactions were performed using two distinct organoboron reagents: borane-tetrahydrofuran complex (BH3·THF) (CAT#176192, Sigma-Aldrich) and 9-borabicyclo[3.3.1]nonane (151076, Sigma-Aldrich). Three different stoichiometric ratios (0.5, 1.0, and 2.0) were explored for each reagent to investigate different degrees of hydroxylation within the STP mixture. The sterically hindered 9-BBN was anticipated to selectively react with the least sterically demanding sites of the terpenes, while the borane-tetrahydrofuran complex was expected to facilitate hydroxylation even at the more challenging endocyclic positions. To remove residual ethanol and water content from the concentrated terpene mixture, a rotary evaporator was used under reduced pressure at 40 °C, ensuring effective removal without compromising the integrity of the terpenes. To ensure the complete removal of water, molecular sieves (CAT#105734, Sigma-Aldrich) were added to the terpene-THF solvent mixture. The reactions were carried out in anhydrous tetrahydrofuran (THF, Sigma-Aldrich) to enable removal by evaporation after completion. Hydrogen peroxide (H2O2, 30%, VWR Chemicals) and sodium hydroxide aqueous solution (NaOH, Sigma-Aldrich) were used to convert the organoboron intermediates into non-toxic, water-soluble boric acid after the reaction. All subsequent terpene derivatives were extracted from the aqueous mixture with ethyl acetate.

Data analysis

To evaluate the impact of engineered modifications on STP biosynthesis and growth characteristics in C. reinhardtii strains, one-way analysis of variance (ANOVA) was performed to compare mean STP (patchoulol) titres and growth rates among different transformants and the parental control strain. This statistical approach allowed for the simultaneous analysis of differences between multiple groups, to provide a robust assessment of the experimental manipulations. ANOVAs were performed separately for STP titres and growth rates under mixotrophic and phototrophic conditions. Post-hoc pairwise comparisons using Tukey's HSD were conducted to identify which groups were statistically significantly different. A oneway ANOVA was used to assess the effects of differences in STP biosynthesis among tested strategies (single or double transformation), followed by a post hoc Tukey's HSD test for specific pairwise data-set comparisons. Mean production values (in fg cell⁻¹ and mg L⁻¹) for each STP were compared, considering

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standard deviations to evaluate data variability. Mean values were considered statistically significantly different at a level of p < 0.05. For data analyses, JMP v.16 (SAS Institute Inc., NC., USA) and Rstudio 3.6.2 (Posit Software, Boston, USA) were used. Data visualisation was done using JMP v.16 and GraphPad Prism v.10 (GraphPad Software, MA, USA). Image adjustments involved ColourChecker calibration (Calibrite LLC, DE, USA) paired with Adobe Lightroom (Adobe Inc., CA, USA) for colour accuracy. The Gardner Colour Scale was used for agarwood sample assessment. Images were cropped and organised in Affinity Photo v.1.10.6 (Serif Ltd, WB, UK). Analysis of images and assignment of colour values according to the Gardner scale was conducted using ImageJ software (NIH, USA). Diagrams and illustrations were made using Affinity Designer v.1.10.6 (Serif Ltd, WB, UK), chemical structures with ChemDraw v.20.1 (PerkinElmer, MA, USA) and all visual elements were harmonised in Affinity Publisher v.1.10.6 (Serif Ltd, WB, UK).

Data availability

Data supporting the findings of this work are available within the paper and its ESI.† Source GCxGC-TOF/MS, GC-MS/FID for the findings of this study and genetic data can be accessed in DRYAD (https://doi.org/10.5061/dryad.h70rxwdq5).

Author contributions

S. G., S. O., G. B. W., and K. J. L. conceptualised and planned the project. S. G. and K. J. L. designed the genes and plasmids for this study. S. G. and S. O. undertook in situ extraction experiments and analysed the GC-FID/MS results. S. G. and V. G. S. conducted experiments and interpreted the GCxGC-TOF/MS data. C. O. executed the nanofiltration experiments and M. G. conducted the hydroxy-boron oxidation experiments in the lab of G. S., who also participated in experimental design. S. G., S. O., G. B. W., and K. J. L. analysed the data and wrote the manuscript. K. J. L. supervised, managed and was responsible for funding acquisition. All authors reviewed and discussed the manuscript.

Conflicts of interest

The authors declare no competing financial interest with regards to this work.

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