Bioderivatization as a concept for renewable production of chemicals that are toxic or poorly soluble in the liquid phase

Pachara Sattayawat*, Ian Sofian Yunusa*, and Patrik R. Jones*†,1

*Department of Life Sciences, Imperial College London, SW7 2AZ London, United Kingdom

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Bio-based production technologies may complement or replace petroleum-based production of chemicals, but they face a number of technical challenges, including product toxicity and/or water insolvency. Plants and microorganisms naturally biosynthesize chemicals that often are converted into derivatives with reduced toxicity or enhanced solubility. Inspired by this principle, we propose a bioderivatization strategy for biotechnological chemicals production, defined as purposeful biochemical derivatization of intended target molecules. As proof of principle, the effects of hydrophobic (e.g., esterification) and hydrophilic (e.g., glycosylation) bioderivatization strategies on the biosynthesis of a relatively toxic and poorly soluble chemical, 1-octanol, were evaluated in Escherichia coli and Synechocystis sp. PCC 6803. The 1-octanol pathway was first optimized to reach product titer at which the host displayed symptoms of toxicity. Solvent overlay used to capture volatile products partially masked product toxicity. Regardless of whether solvent overlay was used, most strains with bioderivatization had a higher molar product titer and product yield, as well as improved cellular growth and glucose consumption, compared with strains without bioderivatization. The positive effect on bioproduction was observed with both the hydrophobic and hydrophilic strategies. Interestingly, in several combinations of genotype/induction strength, bioderivatization had a positive effect on productivity without any apparent effect on growth. We attribute this to enhanced product solubility in the aqueous or solvent fraction of the bioreactor liquid phase (depending on the derivative and medium used), with consequent enhanced product removal. Overall, under most conditions, a benefit of bioproduction was observed, and the bioderivatization strategy could be considered for other similar chemicals as well.

bioderivatization | 1-octanol | toxicity | solubility | bioproduction

Microbial biotechnology offers an attractive method for renewable production of chemicals that replace those currently sourced from fossil fuel feedstocks (e.g., monomers for plastic or textile polymer synthesis) (1) or nature (e.g., vanillin) (2, 3). A critical factor determining success with such a process is the compatibility between the engineered metabolism and its microbial host (4). If the target molecule or its metabolic intermediates are toxic to the host organism, then the maximum potential for cost-effective production is likely not achieved. For lower-value chemicals, economics really matter in the face of competition from fossil fuels (5). Nonetheless, implementation of strategies to enhance product tolerance has been found to improve productivity (6).

In nature, many organisms naturally synthesize very toxic molecules (7) but yet have survived throughout evolution and perhaps even prospered because of this. If we look closer, however, in many cases these chemicals accumulate as chemical derivatives—for example, glucosides that are synthesized by plants (8, 9). In some cases, these detoxification mechanisms are so effective that chemicals (e.g., cyanogenic glucosides) that without derivatization would certainly kill the plant itself can accumulate up to 30% of dry weight (10). Other examples are esters synthesized by yeasts using native alcohol acyltransferases (AATs). It has been argued that ester synthesis is also a detoxification mechanism to convert more toxic metabolites into those that are less harmful (11). If this process is so successful, why not also adopt it for biotechnology?

Herein we propose and define bioderivatization as the purposeful in vivo transformation of chemicals into chemical derivatives by modification of functional groups. Often these functional groups (e.g., hydroxyl groups, –OH) are central to rendering a chemical toxic toward cells (12). At the same time, bioderivatization may also radically change the chemical properties (e.g., water solubility) of the target chemical and/or protect the molecule from further conversion (e.g., oxidation). Bioderivatization could also open new opportunities for strategic product/process separation that is more cost-efficient. Once the derivative has been isolated outside of the biological host, it would need to be converted back to its original form, unless the particular derivative in question is also an attractive product. If the original form of the derivatized products are desired, the final end-products (i.e., aglycons or acids and alcohols) can be obtained through either hydrogenation or hydrolysis following isolation from the bioreactor. The general bioderivatization concept is illustrated in Fig. 1.

There are several instances where bioderivatization was implemented without a rationale (e.g., vanillin glucoside biosynthesis as opposed to synthesizing plain vanillin) (2) or occurred by chance through interactions with native metabolism in the biotechnological

Significance

Microorganisms can be rationally engineered to convert CO2 and H2O into chemicals, replacing those made from fossil fuels today. Sometimes such chemicals are poorly soluble in water or negatively affect the growth of the microorganism, resulting in cost-inefficient manufacturing. In nature, this problem is often solved by converting incompatible chemicals into those more compatible with the host and/or environment. Inspired by this, we propose a similar strategy for engineered biotechnology, whereby biochemical conversion inside the microorganism is followed by chemical reversal once outside. The principle was demonstrated with 1-octanol by implementing two different conversion methods in two different species, showing enhanced bioproduction in most cases. The approach may stimulate commercialization of sustainable and renewable production of chemicals.

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Competing interest statement: The authors are inventors on the UK patent application entitled “Bio-based production of toxic chemicals.” The subject is a method of producing a derivative of 1-octanol, as well as microorganisms and expression vectors for use in said method.

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†To whom correspondence may be addressed. Email: p.jones@imperial.ac.uk.

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host (e.g., geraniol synthesis resulting in geranyl acetate formation through the action of native AATs) (13). To our knowledge, however, the impact of this process on biotechnological objectives has not yet been systematically studied.

We assumed that only toxic or labile products, or those that are expensive to separate (e.g., organic acids) (14), were likely to benefit from the strategy. For example, shorter-chain alcohols, such as l-butanol and ethanol, have relatively low toxicity and effective production systems already in place. Thus, in the present work, the hypothesis that bioderivatization offers benefits for a microbial biotechnological process was tested using 1-octanol (15, 16) as the model product using two different model organisms (17). A stainless steel AAT (SAAT) gene (saat) from Frageria x ananassa (18) and an alcohol O-acetyltransferase (ATF1) gene (atf) from Saccharomyces cerevisiae (19) were chemically synthesized from Integrated DNA Technologies (IDT) and codon-optimized for E. coli. Five genes encoding glycosyl transferases and one gene encoding sucrose synthase (SUS) were also synthesized from IDT for octyl glucoside production.

Plasmids used for gene expression were constructed using Bioart Biopolymer Standard for Idempotent Cloning (BASIC) (19) or traditional restriction enzyme ligase-based cloning. The strains and plasmid construction methods are described in detail in SI Appendix. The UniProtKB accession numbers for all of the proteins used in this study are listed in SI Appendix, Table S3, and the primers used for PCR analysis are listed in SI Appendix, Table S4. All linkers used for BASIC are listed in SI Appendix, Table S5. The amino acid sequence alignments of the C8-prefering thioesterases are shown in SI Appendix, Fig. S1.

Materials and Methods

Strains and Plasmids. E. coli DH5α (Thermo Fisher Scientific) was used to propagate all the plasmids used in this work. Two strains of E. coli (E. coli C43 [DE3; Lucigen] and BW25113 [Keio collection]) and one species of cyanobacteria (Synechocystis sp. PCC 6803) were used as hosts for 1-octanol and octyl acetate production. All E. coli and cyanobacterial strains used in this study are listed in SI Appendix, Tables S1 and S2, respectively. The genes encoding thioesterases Tes3, phosphopantetheinyl transferase Sfp, and carboxylic acid reductase (CAR) were obtained from plasmid pET-TPC3 (15), whereas the chloramphenicol acetyltransferase (CAT) gene (cat) was amplified from plasmid pACYP-Ctet-fpr (17). A stainless steel AAT (SAAT) gene (saat) from Frageria x ananassa (18) and an alcohol O-acetyltransferase (ATF1) gene (atf) from Saccharomyces cerevisiae (19) were chemically synthesized from Integrated DNA Technologies (IDT) and codon-optimized for E. coli. Five genes encoding glycosyl transferases and one gene encoding sucrose synthase (SUS) were also synthesized from IDT for octyl glucoside production.

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Evaluating the Effect of 1-Octanol, Octyl Acetate, and Octyl Glucoside on Growth and Liquid-Phase Partitioning. E. coli strain C43 (DE3) was cultivated in 10 mL of lysogenic broth (LB broth; Sigma-Aldrich) overnight at 37 °C and 180 rpm. The overnight culture was washed twice with fresh M9 minimal medium (47.8 mM Na2HPO4, 2 mM KH2PO4, 8.55 mM NaCl, 18.69 mM NH4Cl, 2 mM MgSO4, 0.1 mM CaCl2, and 2% [wt/vol] glucose) and resuspended in the same medium to an initial OD730 of 0.1. The liquid cultures were spiked with various concentrations of 1-octanol (0 to 50 mM), octyl acetate (0 to 200 mM), and octyl glucoside (0 to 50 mM) and transferred (200 μL) into a well in a 96-well microtiter plate. The plate was incubated in a Tecan Infinite M200 Pro Spectrophotometer at 37 °C with continuous shaking at 432 rpm, and OD730 was measured every hour for 15 h. The specific growth rates were calculated for each treatment.

Production of 1-Octanol, Octyl Acetate, and Octyl Glucoside in Engineered E. coli. Overnight cultures were grown in LB media (10 mL) containing appropriate antibiotics with final concentrations as follows: kanamycin 50 μg/mL, spectinomycin 50 μg/mL, and carbenicillin 100 μg/mL. The overnight cultures were washed twice with fresh M9 minimal medium and resuspended in 2 mL of M9 minimal medium before inoculation in 25 mL of M9 minimal medium with a starting OD600 of ~0.1 in 100 mL Erlenmeyer flask with appropriate antibiotics. For 1-octanol and octyl acetate production experiments, the liquid cultures were cultivated at 37 °C and 180 rpm for 4 h and then induced with various concentrations of isopropyl-β-D-thiogalactopyranoside (IPTG; 0.02, 0.05, 0.2, and 0.5 mM). After induction, 10% (vol/vol) hexadecane solvent overlay (Sigma-Aldrich) was added, and the liquid cultures were cultivated at 30 °C and 150 rpm. OD600 was measured every 24 h after inoculation for 48 h unless stated otherwise. Samples from the liquid cultures and hexadecane solvent overlay were collected every 24 h after inoculation for HPLC and GC-MS analysis, respectively. For octyl glucoside production, the cultures were cultivated at 37 °C and 180 rpm for 4 h and induced with 0.5 mM IPTG before continuing the incubation at 30 °C and 150 rpm for 48 h. Samples were centrifuged, and the supernatants were analyzed by HPLC. Sucrose (15 or 100 mM) was supplemented to the culture medium at the time of induction when the function of AtSUS1 was assessed. Solvent overlay was used when the production of 1-octanol and octyl glucoside was compared.

Production of 1-Octanol and Octyl Acetate in Engineered Cyanobacteria. All cyanobacterial strains were cultivated in BG11 medium without cobalt (hereinafter BG11-Co; SI Appendix, Text), as cobalt was used as an inducer. The preculture was grown in 5 mL of BG11-Co containing appropriate antibiotic(s) (final concentrations: kanamycin, 10 μg/mL; spectinomycin, 10 μg/mL; carbenicillin, 100 μg/mL; and ampicillin, 20 μg/mL) and 1% (vol/vol) CO2 in an Algaetron AG230 (Photon Systems Instruments). When the OD730 reached 3 to 4, the liquid preculture was transferred into an
autoclaved 100-mL Erlenmeyer flask covered with aluminum foil. The OD_{730} was adjusted to 0.2 by adding BG11-Co to a final volume of 25 mL, and antibiotics were added accordingly. The liquid culture was induced on day 2 with 15 μM nickel and 625 nM cobalt, and the OD_{730} was monitored for 20 d in the presence or absence of a 30% (vol/vol) hexadecane solvent overlay. On day 20, the liquid culture was transferred into fresh BG11-Co medium containing antibiotics with initial OD_{730} = 0.2. When the OD_{730} reached 1 to 1.5, the liquid culture was induced with 15 μM nickel and 625 nM cobalt, and hexadecane solvent overlay 30% (vol/vol) was added. The hexadecane solvent overlay was sampled at 16 d after induction. The GC-MS analysis. In a separate experiment, production cultures were inoculated directly from the initial preculture and cultured for 10 d in the presence of 30% (vol/vol) hexadecane solvent overlay, with induction of protein expression on day 2, as described above. The toxicity of the different products to *Synechocystis* sp. PCC 6803 were evaluated as described in SI Appendix, Fig. S2.

Quantification and Analysis of 1-Octanol, Octyl Acetate, Octyl Glucoside, and Other Metabolites.

**GC-MS.** Hexadecane solvent overlay was used to capture 1-octanol and octyl acetate from the liquid culture. Alternatively, 0.1 volume of hexadecane was used to extract samples of the aqueous phase from mock or production cultures. One hundred microliters of hexadecane overlay or extract was transfered into an insert in a 2-mL screw top GC vial (Agilent). Samples were analyzed using an Agilent 7890B gas chromatograph with an HP-5 ms column, a 7683 autosampler, and a 5977E MSD system. Then 1 μL of sample was injected using a pulsed split ratio of 10:1 and split flow at 10 min/mL. The GC was programmed with an initial temperature of 70 °C for 30 s, followed by a first ramp at 30 °C/min to 250 °C before ramping up to 300 °C with a final hold for 2 min at 40 °C/min. Target products were identified by comparing mass spectra and retention times with external standards. Serial dilutions of 1-octanol ≥99%; Sigma-Aldrich) and octyl acetate (≥99%; ACROS Organics) standards were used to quantify the concentrations of 1-octanol and octyl acetate in the sample.

**HPLC.** An Agilent 1200 series HPLC instrument equipped with different columns and a reflective index detector was used to determine the concentrations of octyl glucoside, glucose, fructose, sucrose and fermentation products in the *E. coli* sample every 24 h. One milliliter of liquid cultures was sampled at 24 h and centrifuged at 17,000 g for 15 min to separate the aqeous and hexadecane layers. The supernatant was transferred into a 2-mL HPLC vial. For octyl glucoside detection, samples (20 μL) were analyzed with a Zorbax XDB-C18 column (Agilent). The flow rate was set at 1 mL/min with a column temperature of 30 °C (20). For the analysis of sugars (glucose, sucrose, and fructose) from samples supplemented with sucrose, an Aminex HPX-87P column (Bio-Rad) was used to analyze the samples (20 μL) with the flow rate of 0.6 mL/min, and the column temperature was set at 85 °C. For glucose and other fermentation products when sucrose was not supplemented, samples (100 μL) were analyzed with an Aminex HPX-87H column (Bio-Rad), and the flow rate and column temperature were set at 0.6 mL/min and 60 °C, respectively. Serial dilutions of glucose (Sigma-Aldrich), sucrose (Sigma-Aldrich), fructose (Sigma-Aldrich), sodium acetate (Sigma-Aldrich), sodium lactate (Sigma-Aldrich), and absolute ethanol (VWR) were used to determine the amounts of these compounds in the sample.

**Statistical Treatment and Data.** Three biological replicates were used for each treatment and/or condition. Unless stated otherwise, symbols or bar graphs represent the mean ± SD from three biological replicates. Individual data from line graphs are presented in SI Appendix, Fig. S3. Because all samples were collected from cultures that most likely were reasonably homogenous, normality was assumed in all cases, as discussed by Fay and Gerow (21). With selected data, indicated in the text or figure legends with *p* values obtained, a two-sided Student’s t test was used, with asterisks indicating significance (*p* ≤ 0.05; **p** ≤ 0.01; ***p** ≤ 0.005).

The data supporting the findings of this study are available within the paper and its Supplemental Information files. All other data that support the findings of this study are available from the corresponding author upon reasonable request.

**Results and Discussion**

**Comparing the Toxicity of 1-Octanol, Octyl Acetate, and Octyl Glucoside.** In previous studies (15, 16), we observed that 1-octanol was toxic, resulting in reduced growth and also genetic instability in cyanobacteria (16). Alcohols are known to compromise the integrity of cell membranes (22), thereby causing cellular toxicity that is most commonly observed as a growth defect. With 1-octanol having only a single hydroxyl-group as a functional group, we initially considered two different types of derivatization: O-glucosylation and esterification. For ester synthesis, the most valuable organic acid moiety that could be recovered following hydrolysis of the bioderivative would make the most sense economically; however, although acetate is not a particularly valuable end product, the biosynthesis of acetate esters is easiest to implement and thus served as the starting point.

Before commencing with metabolic engineering, we evaluated the tolerance of the *E. coli* host strain to 1-octanol and its two proposed derivatives (octyl glucoside and octyl acetate) in 96-well microtiter plates. The cells were unable to grow when the concentration of 1-octanol in the liquid culture was >0.75 mM (Fig. 2 and SI Appendix, Fig. S4A). In contrast, growth was observed at all tested concentrations of octyl acetate (0 to 50 mM) (Fig. 2 and SI Appendix, Fig. S4B) and above (SI Appendix, Fig. S5), and up to 2.5 mM for octyl glucoside (Fig. 2 and SI Appendix, Fig. S4C). In previous studies, alcohols displayed varying (sometimes more, sometimes less) toxicity relative to its corresponding esters, depending on the specific product and derivative in question (12); for example, butyl acetate was found to be more toxic than 1-butanol. To further complicate matters, the apparent toxicity of externally applied substances is likely influenced by environmental factors, including varying types of solvent overlay, and the toxicity of internally accumulated 1-octanol, octanal, and octanoic acid it remains unknown. Nevertheless, at least in the case of externally applied chemicals, both the corresponding acetyl ester and glucoside were less toxic than 1-octanol under the tested conditions. Based on this experiment, both derivatives were pursued in vivo as a model system, although clearly a “one size fits all” generalization is not possible.

**Selection of AAT for Octyl Acetate Biosynthesis.** Octyl acetate is naturally found in wild strawberry (*Fragaria vesca*) (23). For use as a food flavor additive, it is also synthesized by a direct esterification reaction between acetic acid and octyl alcohol, catalyzed by acids, ion-exchange resins, or ionic liquids (24). Until now, we have found reports of microbial in vivo production of octyl acetate using a renewable substrate.

The TPC3 pathway was extended with an AAT under the assumption that native acetyl-CoA was not limiting (Fig. 3A). Three
AAT enzymes—CAT (25), SAAT (18), and ATF1 (26)—were selected based on the literature. CAT has previously been used in *E. coli* for ester biosynthesis (27), while both SAAT and ATF1 have been reported to use 1-octanol as a substrate (18, 27, 28). Before the in vivo production of 1-octanol and octyl acetate, a mock experiment with spiked 1-octanol (1 or 3.84 mM) and octyl acetate (1 or 2.9 mM) was carried out to investigate whether 1-octanol or octyl acetate remained in the aqueous or solvent phases after a 24-h incubation with solvent overlay. None of the compounds was detected in the aqueous phase at the lower concentration (Fig. 3B). The highest octyl acetate titer (0.54 ± 0.01 mM [93.82 mg/L]) at 48 h after inoculation) and yield (12.54 mmol/mmol glucose) were found in cultures of the strain expressing ATF1 (strain no. 5; *SI Appendix*, Table S1). The introduction of AAT did not result in marked changes in growth (Fig. 3C), and esterification of 1-octanol had only a small positive impact on the final product titer. There may be at least two possible explanations for this, including (i) the hexadecane solvent overlay reduced the toxicity of the products by in situ product removal (27) or (ii) the 1-octanol–producing strain reached a final titer of only 0.32 ± 0.03 mM, which is lower than the concentration that affected the growth of *E. coli*, as shown in Fig. 2.

To more comprehensively evaluate the impact of bioderivatization, we hypothesized that it was important to exceed the titer at which the underivatized product affected the growth of the host, at least in the absence of solvent overlay. Thus, the next task was to improve flux through the 1-octanol pathway.

### Identification of Limiting Factor(s) for 1-Octanol Pathway Flux

The titer of 1-octanol and octyl acetate in the previous experiment were low, indicating that one or more reactions in the introduced pathway prevented efficient pathway flux. In cyanobacteria, the availability of octanoic acid was found to limit the biosynthesis of 1-octanol (16). To evaluate whether the supply of octanoic acid was sufficient for the production of 1-octanol and octyl acetate, AHR, aldehyde reductase. (B) Plasmids used to generate strains 1 to 5 as listed in *SI Appendix*, Table S1. (C) Production of octyl acetate by introducing AAT enzymes in *E. coli* C43 (DE3). (D) Chromatogram of overlay sampled from Tes3-Sfp-CAR and Tes3-Sfp-CAR + ATF1 at 48 h. Peak identification: (1) 1-octanol; (2) octyl acetate. (E) Biomass accumulation of strains in C. AAT enzymes were able to convert 1-octanol and acetyl-CoA into octyl acetate; however, SAAT and ATF1 were more effective than CAT (*P = 0.003* and 0.0002, respectively) (Fig. 3B). The highest octyl acetate titer (0.54 ± 0.01 mM [93.82 mg/L]) at 48 h after inoculation) and yield (12.54 mmol/mmol glucose) were found in cultures of the strain expressing ATF1 (strain no. 5; *SI Appendix*, Table S1). The introduction of AAT did not result in marked changes in growth (Fig. 3C), and esterification of 1-octanol only had a small positive impact on the final product titer. There may be at least two possible explanations for this, including (i) the hexadecane solvent overlay reduced the toxicity of the products by in situ product removal (27) or (ii) the 1-octanol–producing strain reached a final titer of only 0.32 ± 0.03 mM, which is lower than the concentration that affected the growth of *E. coli*, as shown in Fig. 2.

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also limiting in *E. coli*, octanoic acid was added externally to the strains expressing the first-generation 1-octanol pathway with (strain no. 4) or without (strain no. 2) coexpression of SAAT (SI Appendix, Table S1). Substantially more of each product was observed in cultures to which octanoic acid was added, confirming that the supply of acid was indeed limiting the 1-octanol pathway (Fig. 4A). As all the 1-octanol was converted into the corresponding ester in acetyl-CoA and the AAT activity were not limiting (Fig. 4C and D at 48 h; $P$ = 0.027). This is in line with a previous report by Wilbanks and Trinh (12). The 1-octanol–producing strain (Tes3-Sfp-CAR) showed noticeably lower cell density (Fig. 4C, D, G, and H) and glucose consumption (Fig. 4E and F), an effect that was largely alleviated by esterification (Tes3-Sfp-CAR + SAAT), thereby providing early insight into the main hypothesis of the work. The differences between the alcohol- and ester-forming strains were reduced in the absence of externally added octanoic acid (Fig. 4C, D, G, and H), supporting the idea that further pathway optimization was essential to evaluate the effect of bioderivatization under conditions that were more likely to be relevant for application.

A similar substrate feeding experiment was also carried out in the absence of the hexadecane solvent overlay. However, this precluded quantification of the products, since both 1-octanol and octyl acetate are volatile. Thus, growth and glucose consumption were instead evaluated as indicators of cellular and metabolic activity. The experiment indicated that octanoic acid feeding reduced the growth of *E. coli* even without conversion to 1-octanol and octyl acetate (compare the negative control in Fig. 4C and D at 48 h; $P$ = 0.027). This is in line with a previous report by Wilbanks and Trinh (12). The 1-octanol–producing strain (Tes3-Sfp-CAR) showed noticeably lower cell density (Fig. 4C, D, G, and H) and glucose consumption (Fig. 4E and F), an effect that was largely alleviated by esterification (Tes3-Sfp-CAR + SAAT), thereby providing early insight into the main hypothesis of the work. The differences between the alcohol- and ester-forming strains were reduced in the absence of externally added octanoic acid (Fig. 4C, D, G, and H), supporting the idea that further pathway optimization was essential to evaluate the effect of bioderivatization under conditions that were more likely to be relevant for application.

The first-generation 1-octanol pathway was also evaluated in two different *E. coli* strain backgrounds: *E. coli* B strain C43 (DE3) and K-12 strain BW25113. As the latter strain (0.74 mM, 0.58 mg/L, 27 mmol/mol glucose) displayed greater alcohol yield ($P$ = 0.042) than the former (0.35 mM, 45.44 mg/L, 10.15 mmol/mol glucose) (SI Appendix, Fig. S7), the *E. coli* BW25113 strain background was used in the subsequent experiments.

**Fig. 4.** (A and B) Identification of limiting substrates in 1-octanol and octyl acetate production by substrate feeding and effect of bioderivatization on *E. coli* growth and glucose consumption without overlay. *E. coli* C43 (DE3) Tes3-Sfp-CAR (A) and Tes3-Sfp-CAR + SAAT (B) cultivated with 10% (vol/vol) hexadecane overlay. Different concentrations of octanoic acid were added to the cultures. (C–F) Growth and metabolism of *E. coli* C43 (DE3) fed with 0 mM (C and E) and 1.5 mM (D and F) octanoic acid in the absence of solvent overlay. Three strains were tested: negative control (gray triangles), Tes3-Sfp-CAR (black circles), and Tes3-Sfp-CAR + SAAT (red squares). (G and H) Photographs of liquid cultures taken at 48 h when supplemented with 0 mM octanoic acid (G) and 1.5 mM octanoic acid (H) in the absence of solvent overlay.
377.37 mg/L (44.20 mmol/mol glucose) at the higher IPTG concentration of 0.2 mM (Fig. 5C). With these improvements, the internally produced 1-octanol now exceeded the titer (0.75 mM) at which the underivatized product affected the growth of the strain in the absence of solvent overlay.

Lozada et al. (31) reported that ‘CpFatB1-4’ was most effective when the expression level was low, as the strain expressing it showed a growth defect at higher expression levels. This was also observed in our study (SI Appendix, Fig. S8A). Since the same promoter was also used in the second AAT-encoding plasmid, we speculated that an imbalance in pathway enzyme activities may compromise the final outcome. Thus, the second-best thioesterase (‘CaFatB3-5’) was also used in further production experiments, given that growth of the ‘CaFatB3-5’ strain was not greatly influenced by the IPTG concentration (SI Appendix, Fig. S8B).

Following optimization of the 1-octanol pathway, the effect of esterification on growth and glucose consumption of the biocatalytic host was evaluated. Strains harboring ‘CpFatB1-4’, with or without ATFI, were induced at the IPTG concentration found to be optimal for ‘CpFatB1-4’ (0.05 mM) and grown in the absence of a hexadecane overlay. The presence of ATFI enhanced both growth and glucose consumption (Fig. 5E and F), as was also observed in the previous octanoic acid feeding experiments (Fig. 4) with P = 0.006 and 0.003 when comparing growth and glucose consumption at 48 h, respectively.

**Effect of Bioderivatization on Production with Enhanced Pathway Flux.** Would bioderivatization also influence C8 productivity now that pathway flux was up to 8 times greater? To answer this question, the effect of ATFI in the improved (high-flux) ‘CpFatB1-4’ and ‘CaFatB3-5’ strains was evaluated in the presence of a hexadecane solvent overlay at three different IPTG levels. The results were complex depending on the conditions, but a number of interesting observations were made. Cellular growth and glucose consumption were positively influenced by ATFI in some of the induction/thioesterase combinations (e.g., ‘CaFatB3-5’ at 0.2 and 0.5 mM IPTG) but not in others (e.g., none of the strains at 0.05 mM IPTG) (SI Appendix, Figs. S9 and S10). Microtiter well growth at 0.2 mM IPTG also indicated enhanced growth in the first 24 h (SI Appendix, Fig. S11). In contrast, no difference in growth was observed in response to esterification at low IPTG induction, as was also found with the first-generation low-flux strains (Fig. 3), despite the fact that a substantial impact on growth was observed in the absence of solvent overlay (Fig. 5E and F). Thus, the solvent overlay partially mitigated the “growth and metabolism” defect caused by the pathway and/or its product, at least when compared with the same strain cultured in the absence of solvent overlay, that is, 0.05 mM IPTG with the ‘CpFatB1-4’ strain.

At the lowest protein expression inducer level (0.05 mM IPTG), esterification had no impact on yield (Fig. 6) or titer (SI Appendix, Fig. S12) for both strains. In contrast, at the higher IPTG levels, almost all IPTG/strain/time combinations showed both improved titer and yield when ATFI was coexpressed. Interestingly, this means that all combinations of IPTG dosage and strain sampled at 24 and 48 h except one (‘CpFatB3-5’, 0.2 mM, 48 h), had increased C8 productivity, even with the variable impacts on growth or glucose consumption. In other words, the effect of esterification under these conditions was partially independent of any effect on cellular activity, and thus the effect of esterification on product toxicity was not the sole reason for the improved productivity under these conditions.

Most likely, the additional positive effect of bioderivatization on productivity can be explained by esterification enhancing product removal, either by enhancing the compatibility with native
efflux transporter(s) or, more likely, by enhancing product solubility in the solvent overlay and thereby facilitating the transfer away from the cell. In turn, this would reduce the local product concentration, with consequences for both toxicity and pathway thermodynamics (due to a reduction in the actual free energy change of the entire pathway including efflux). The lack of effect of bioderivatization on C8 productivity at the low (0.05 mM) IPTG induction level might be explained by an imbalance between pathway catalysts complicated by the contrasting impact of IPTG on the two different thioesterases. This difference is illustrated by the change in the product titer ratio between strains with and without ATF1 in response to the concentration of IPTG used for induction (SI Appendix, Figs. S13 and S14).

To ensure that the measurements from the solvent phase were representative of the total bioproduction system, the distribution of products between the liquid and solvent phases was quantified with a 1-octanol- and octyl acetate-producing strain. Similar to the mock experiments (SI Appendix, Fig. S6), the majority of 1-octanol (84%; mole 1-octanol in solvent phase/mole 1-octanol in aqueous and solvent phases) and all the octyl acetate were found in the solvent phase (SI Appendix, Fig. S15).

Transfer of the Bioderivatization Concept to a Different Organism. We recently reported the introduction of 1-octanol biosynthesis into cyanobacteria (16). The use of solvent overlay was found to be important, as the cyanobacterium (Synechocystis sp. PCC 6803) was even more sensitive than E. coli to 1-octanol. Given the positive impact of esterification on C8 biosynthesis in E. coli, we wondered whether similar benefits could also be observed in Synechocystis sp. PCC 6803. External addition of 1-octanol and octyl acetate indicated reduced sensitivity to the ester compared with 1-octanol (SI Appendix, Fig. S2), although Synechocystis sp. PCC 6803 was clearly more sensitive than E. coli to octyl acetate.

Heterologous expression of SAAT in combination with the CaFatB3-5 thioesterase, Sfp, and CAR enabled complete conversion of 1-octanol into octyl acetate in this species as well. Strains expressing ATF1 could not be obtained despite repeated transformation attempts. Cultivation of the two strains in the absence of a solvent overlay resulted in marked differences in growth and appearance by day 10 (Fig. 7 A and B), but by day 20 the 1-octanol strain had caught up, and both cultures were vibrant green. The day 20 culture was then used as a preculture for a new culture with fresh medium that contained hexadecane solvent overlay.

In the presence of solvent overlay, the addition of SAAT resulted in enhanced growth (Fig. 7 C and D), C8 product titer (Fig. 7E), and yield (Fig. 7F) after 16 d of cultivation and induction of protein expression on day 2. Surprisingly, none of the 1-octanol-producing strains lost the ability to accumulate its product in this study, in contrast to what we observed earlier (16). The initial no-induction precultures of the same strains were also used to inoculate cultures that were immediately provided a solvent overlay and then induced for protein expression and cultured for 10 d. Such cultures accumulated 1.6 ± 0.4 mM 1-octanol and 2.4 ± 0.5 mM octyl acetate of each respective product. Similar to what was found with E. coli (SI Appendix, Fig. S15), the majority of the 1-octanol (96%; mole 1-octanol in solvent phase/mole 1-octanol in aqueous and solvent phases) produced by the strain lacking AAT accumulated in the solvent phase (Fig. 7G).

Effect of Conjugate Type: Hydrophilic Instead of Hydrophobic Bioderivatization. Other conjugation types are also possible—for example, glycosylation. This requires changes in metabolic engineering and has implications for both cellular efflux and product separation. The simplest glycosylation to implement is O-glucosylation, thereby resulting in the formation of octyl glucoside, a nonionic alkyl glucoside used as a surfactant (32). Octyl glucoside is much more soluble in water than the alcohol—5.5 g/L (33) vs. 0.5 g/L (34), respectively—and will require a different choice of downstream processing for separation compared with esters.

The CaFatB3-5 thioesterase 1-octanol pathway was extended overexpression of a glycosyltransferase (GT) (Fig. 3A). Five
candidates were selected based on their reported activity toward longer-chain alcohols (35, 36). By combining a glycosyltransferase from Medicago truncatula (MH2) (37) with the 1-octanol pathway, 0.73 mM (214 mg/L) octyl glucoside was produced after 48 h (Fig. 8B). Therefore, MH2 was selected for further investigations. The octyl glucoside titers were low in comparison with the octyl acetate strain (Fig. 3). Insufficient glucosylation may be due to a limitation in the UDP-glucose pool, enhanced regeneration may be important to achieve high glucoside productivity. SUS catalyzes the reversible conversion of sucrose and UDP into UDP-glucose and has been shown to enhance glucoside production (40). Thus, the SUS from Arabidopsis thaliana (AtSUS1) with the 1-octanol pathway, carrying only the 1-octanol pathway. Similarly, the product titers were also improved by bioderivatization (≤ 0.0009 and 0.008 at 24 and 48 h, respectively) (SI Appendix, Fig. S17).

The combined analysis suggests that bioderivatization has the potential to enhance at least some biotechnological production systems in which product toxicity and/or insufficient product solubility places excessive limits on bio-based production of valuable chemicals. In most circumstances, the growth defect caused by the target product (or pathway/parts) was overcome either by adding solvent overlay or by bioderivatization. This is most clearly shown in SI Appendix, Fig. S2 or by comparing Fig. 5E with SI Appendix, Fig. S94. In several instances, bioderivatization enhanced productivity (Figs. 6 and 8) without also enhancing growth (e.g., SI Appendix, Figs. S9, S11, and S18).

The increased water solubility of octyl-β-glucoside relative to 1-octanol, along with the small difference in toxicity between the two compounds, support the notion that enhanced product solubility also plays an important role—that is, bioderivatization does not improve productivity solely by mitigating product toxicity. Regardless, and most importantly, the addition of solvent overlay alone did not result in the same level of productivity enhancement—the ultimate objective of the biotechnological application—as bioderivatization (e.g., Fig. 7). Thus, even if solvent overlay can help overcome the growth defect, bioderivatization achieves the same effect and also improves productivity. In addition, as mentioned in the introductory paragraph, solvent overlay is not always going to be a suitable universal solution.

Fig. 7. Effect of bioderivatization on product titer, yield, and cellular growth of engineered Synechocystis sp. PCC 6803. (A) Biomass accumulation of Δaas-PnrsB-Sfp-CAR-Pcoa-CpFatB1-4 and Δaas-PnrsB-Sfp-CAR-Pcoa-CpFatB1-4-SAAT cultivated in the absence of overlay for 20 d. All strains were induced to express recombinant proteins on day 2 with 625 nM cobalt and 15 μM nickel. After 20 d of cultivation in the absence of overlay, each culture was used to inoculate fresh cultures with overlay and induced to express recombinant proteins as above. (B and C) Photographs of the 1-octanol strain (−AAT) and octyl acetate strain (+AAT) on day 10 when cultivated in the absence (B) or presence (C) of 30% (vol/vol) hexadecane solvent overlay. (D–F) Biomass accumulation (D), product titer (E), and product yield (F) in solvent overlay cultures sampled on day 16. (G) Localization of 1-octanol from a 1-octanol–producing strain with solvent overlay sampled on day 10. Student’s t test analyses on all data shown in D–F were statistically significant (P ≤ 0.01).
In future studies, it would be interesting to evaluate also how this approach may influence product:process separation, as illustrated in Fig. 1. For example, esters are attractive in this respect, as they have lower water solubility than their corresponding alcohols, which is likely to reduce the energetic cost of the product separation process from the aqueous media (41). The postbiology separation strategy and capability for cellular excretion will also influence whether to opt for a more hydrophobic or a more hydrophilic derivative. At least in the case of 1-octanol, the host cells evaluated in this work were able to excrete both conjugate types, but this might not apply for all target chemicals.

Conclusions

In this work, two different strategies for bioderivatization were implemented, and the effect on bioproduction of the toxic chemical 1-octanol was systematically evaluated in two different species. In *E. coli*, the conversion of 1-octanol into octyl acetate resulted in enhanced product titer and yield in most cases and also enhanced growth and glucose consumption in one-half of the cases. Clear and more consistent positive effects on both growth and productivity were observed when the hydrophobic bioderivatization strategy was transferred to cyanobacteria or when a hydrophilic bioderivatization strategy was implemented. The impact of esterification on bioproduction was influenced by the presence or absence of solvent overlay, as well as by the species and induction level, while the positive effect of glucosylation could be observed under both environmental conditions. We assumed that any improvements in productivity that were accompanied by improvements in cellular activity were caused by reduced product toxicity. Interestingly, closer inspection of the data suggested independent effects of bioderivatization on cellular activity (i.e., growth and glucose consumption) with or without an effect on productivity, or...
on productivity without any effect on cellular activity. The latter effect is likely related to enhanced product removal caused by the enhanced product solubility of derivatives, with esters more soluble in solvent overlay and glucosides more soluble in the aqueous phase. The results presented in this paper support the idea that engineered bioderivatization that mimics the evolved metabolism of many specialized metabolite accumulating species may offer benefits for biotechnology, even with entirely synthetic metabolism.

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