Transporter-mediated biofuel secretion

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Engineering microorganisms to produce biofuels is currently among the most promising strategies in renewable energy. However, harvesting these organisms for extracting biofuels is energy- and cost-intensive, limiting the commercial feasibility of large-scale production. Here, we demonstrate the use of a class of transport proteins of pharmacological interest to circumvent the need to harvest biomass during biofuel production. We show that membrane-embedded transporters, better known to efflux lipids and drugs, can be used to mediate the secretion of intracellularly synthesized model isoprenoid biofuel compounds to the extracellular milieu. Transporter-mediated biofuel secretion sustainably maintained an approximate three- to fivefold boost in biofuel production in our Escherichia coli test system. Because the transporters used in this study belong to the ubiquitous ATP-binding cassette protein family, we propose their use as “plug-and-play” biofuel-secreting systems in a variety of bacteria, cyanobacteria, diatoms, yeast, and algae used for biofuel production. This investigation showcases the potential of expressing desired membrane transport proteins in cell factories to achieve the export or import of substances of economic, environmental, or therapeutic importance.

ABC transporters | biofuel milking | green fuel | multidrug efflux

The rising cost of transportation fuel and concerns over the stability of supply have coincided with the rising era of synthetic genomics and metabolic engineering in the last few years. This scenario has led to the development of engineered microorganisms as attractive sources for the production of green fuels (1). The first-generation biofuel, ethanol, derived from crop-fed microbes, has limitations in terms of its energy density, hygroscopic properties, and the use of cultivable land (2). As a result, microbes with more economic nutritional requirements are now being heavily investigated for the production of advanced biofuel molecules, with properties comparable to petroleum-based fuels (2). Several algae, cyanobacteria, yeast, and bacteria have been investigated as potential production hosts (2). Potential biofuel compounds include derivatives of alcohols, fatty acids, polyketides, and isoprenoids (2).

A major problem that hinders successful commercialization of microbiofuel production is the energy and costs incurred while harvesting cells and extracting the fuel, because most biofuel molecules are synthesized and stored inside cells (3). The most common approach taken to abate this problem today includes lytic, metabolic, enzymatic, or electrical degradation of the cell membranes and walls to leak out biofuel molecules (3–5). An alternative approach is to use membrane transport protein pumps that can export/secrete intracellular biofuels to the extracellular milieu (“secretion” is used interchangeably with “export/efflux” in this paper, as the exported molecules are produced within the same cell and collected outside the cell) (5–7).

Analogous to obtaining milk from cows without killing or weakening them, the term “milking” is often used to describe the biofuel secretion strategy (6). Biofuel secretion from healthy living cells is expected to allow the cells to replenish their intracellular biofuel content, resulting in a sustainable system with overall higher yields compared with nonsecreting systems. The increase in yield could possibly allow liquid biofuel molecules to overcome the phase separation threshold to ultimately appear as a layer above the culture medium, enabling easy recovery. Biofuel secretion is also predicted to reverse the cellular toxicity that is associated with the intracellular accumulation of certain types of biofuel molecules.

To our knowledge, the only report concerning the use of membrane transport proteins for achieving biofuel secretion is from Dunlop et al. (8). They expressed transporters belonging to the RND (resistance-nodulation-cell division) family in bacteria and observed an ~1.5-fold increase in the production of limonene (8). However, a major drawback of this strategy is that RND transporters are large, tripartite protein complexes, spanning two membranes and crossing the periplasmic space; this configuration is exclusively found in gram-negative bacteria and some similar cyanobacteria (9). This tripartite structure limits the translation of RND secretion systems to other economically viable biofuel production hosts such as algae and yeasts.

Unlike RND proteins, transporters belonging to the ATP-binding cassette (ABC) protein family are widely found in all five kingdoms of life (10). They share a conserved structural architecture and specifically import or export a wide variety of molecules and ions across cellular membranes (11). We exclusively deal with exporters in this study. ABC exporters (particularly of the ABCB and ABCG subfamilies) are known to export a diverse range of extremely hydrophobic molecules such as lipids, drugs, and steroids (12); human P-glycoprotein (P-gp/ABCBI) is a well-known example of high pharmacological interest owing to its role in multidrug efflux (13). The functional conservation and diverse substrate poly-specificity among ABC exporters are exemplified by the studies in which a bacterial ABC exporter, LmrA, was found capable of substituting the drug transport function of P-gp in human cells (14). This finding clearly depicts the cross-species translation potential of these transporters.

The broad poly-specificity of ABC transporters in general led us to hypothesize that they could be used to recognize certain biofuel molecules and achieve their secretion out of the cell. There is considerable evidence that a wide range of extremely “greasy” molecules can be transported (12). In plants, for example, transporters from the ABCG subfamily export cuticular waxes (15). In this study, we tested our hypothesis using an isoprenoid-producing Escherichia coli model system. We report the secretion of four different long-chain isoprenoid compounds mediated by different bacterial ABC transporters.

Results

Model Test System. To test our hypothesis that ABC transporters (we exclusively refer to exporters) can secrete biofuel molecules, we established a simple model test system in E. coli. Two different plasmids were cotransformed into E. coli BL21 (DE3) cells. These plasmids have different origins of replication and antibiotic resistance markers, so that they could be stably maintained together in each cell. The first plasmid was used for the constitutive
production of isoprenoids, whereas the second was used for the inducible expression of ABC transporters.

pAC184-based plasmids have been previously engineered to carry operons responsible for the constitutive production of various isoprenoids via the mevalonate pathway (MEV) pathway (16). We used these plasmids for the production of four isoprenoids (17), zeaxanthin, canthaxanthin, β-carotene, and lycopene, in *E. coli* (Fig. 1). These selected isoprenoids are long-chain C40 carotenoids and were chosen as model biofuel compounds for several reasons: (i) they are readily detectable using colorimetric methods (each of these compounds has a unique characteristic absorbance spectrum in the visible region, allowing easy detection and quantification using absorbance spectrophotometry; Fig. S1); (ii) they lack an indigenous background in noncarotenogenic *E. coli*; and (iii) they have a similar overall chemical composition to squalene and botryococcenes (Fig. 1), which have been validated to be hydrocracked to yield gasoline and other useful products (18).

pET19b-derived plasmids are widely used to heterologously express proteins in *E. coli* in an isopropyl-β-D-1-thiogalactopyranoside (IPTG)-inducible manner. We have previously used pET19b to overexpress bacterial ABC transporters for X-ray crystal structure determination (19). We chose to express a panel of 19 different bacterial ABC transporters, many of which are homologs of the ABC exporter, MsbA (Fig. 1; Table S1). MsbA is known to export the lipid A-core moiety of LPS (lipopolysaccharide) from the inner leaflet to the outer leaflet of the inner membrane in *E. coli* (Fig. 24) (20). Subsequently, various lipid transfer proteins and transporters participate to move the lipid A-core (assembled into LPS) to the outer membrane during LPS biogenesis (Fig. 24) (21). MsbA is also known to be capable of exporting a broad range of chemotherapeutic drugs and antibiotics, likely using a mechanism that is similar to lipid A export (22, 23).

Carotenoid-producing *E. coli* generally accumulate the product carotenoids in the inner membrane, likely in the inner leaflet (24). This localization is probably because the carotenoid biosynthetic pathway enzymes encoded by the pAC184-derived plasmids are membrane-localized (24, 25), and in part, because these carotenoids are virtually insoluble in aqueous environments like the cytoplasm, periplasm, and growth medium. The inner membrane localization of carotenoids is akin to that of lipid A (Fig. 2A and B), making carotenoids amenable to be exported by MsbA and probably sharing the same alternating access transport mechanism that is conserved among these types of ABC transporters (11, 26). However, with regard to secretion, the aqueous insolubility of carotenoids would prevent them from exiting the membranes in ordinary culture medium.

To circumvent the insolubility issue, the double transformant *E. coli* liquid cultures were grown with an overlay of decane, which is a biocompatible organic solvent (Fig. 2). The use of such two-phase culture systems is common for collecting hydrophobic secreted products from cells (8, 27, 28). Our hypothesis was that the MsbA-exported carotenoids would partition from the membranes and periplasmic space into the decane phase (Fig. 2B). We also hypothesized that the hydrophobicity of the carotenoid would determine the speed of partitioning into decane (Fig. 2B), thus affecting the time taken to detect the secreted carotenoid. Testing this hypothesis was made possible through our selection of carotenoids that span a range of increasing hydrophobicities (decreasing polarity), from zeaxanthin to lycopene (Fig. 1).

### Isoprenoid Secretion from Cells in Culture Medium

In our carotenoid-producing double transformants, the expression of 3 of our battery of 19 transporters was not detectable on a Western blot; these were excluded from further analysis (Fig. S2). The screens for carotenoid secretion involved three different measurements for each sample: the concentration of secreted carotenoid (in the decane phase; Figs. S3A and B and S4A and B), the concentration of produced carotenoid (in cells, extracted with acetone for 15 min at 55 °C; Figs. S3 C and D and S4 C and D), and cell viability (optical density of the cell culture; Figs. S3E and S4E). A positive hit was defined as a transporter that, when expressed, led to an increase in the secreted amount of carotenoid compared with the nonexpressing (empty vector) control, without reductions in the amount of produced carotenoid and cell viability.

In our screens for zeaxanthin and canthaxanthin secretion, the best hits were *Salmonella enterica* ser. *typhimurium* MsbA (containing a substitution I89T, abbreviated as StMsbA*) for zeaxanthin and *Escherichia coli* MsbA (abbreviated as EcoMsbA) for canthaxanthin (Figs. S3 and S4). Other expressed ortholog transporters showed comparatively poor secretion/negative hits (Figs. S3 and S4), suggesting that zeaxanthin and canthaxanthin

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### Table S1

<table>
<thead>
<tr>
<th>No.</th>
<th>ABC transporters</th>
<th>Isoprenoids/Biofuel compounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>MsbA – Vbrío chloae*</td>
<td>Zeaxanthin [logP&lt;sub&gt;oct&lt;/sub&gt;] 8.35</td>
</tr>
<tr>
<td>4.</td>
<td>MsbA – Helicobacter pylori*</td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td>MsbA – <em>Pasteurella multocida</em></td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>MsbA – <em>Pseudomonas aeruginosa</em></td>
<td></td>
</tr>
<tr>
<td>7.</td>
<td>MsbA – <em>Klebsiella pneumoniae</em></td>
<td></td>
</tr>
<tr>
<td>8.</td>
<td>MsbA – <em>Escherichia coli</em> (abbrev. EcoMsbA)</td>
<td></td>
</tr>
<tr>
<td>9.</td>
<td>MsbA – <em>Haemophilus influenzae</em></td>
<td></td>
</tr>
<tr>
<td>10.</td>
<td>Cyd – <em>Haemophilus influenzae</em></td>
<td></td>
</tr>
<tr>
<td>11.</td>
<td>CydD – <em>Haemophilus influenzae</em></td>
<td></td>
</tr>
<tr>
<td>12.</td>
<td>Putative ABC – <em>Haemophilus influenzae</em></td>
<td></td>
</tr>
<tr>
<td>13.</td>
<td>Putative ABC – <em>Haemophilus influenzae</em></td>
<td></td>
</tr>
<tr>
<td>14.</td>
<td>Putative ABC – Bacillus subtilis*</td>
<td></td>
</tr>
<tr>
<td>15.</td>
<td>Putative ABC – Aggregatibacter actinomycetemcomitans*</td>
<td></td>
</tr>
<tr>
<td>16.</td>
<td>MsbA – <em>Salmonella enterica</em> ser. Paratyphi A</td>
<td></td>
</tr>
<tr>
<td>17.</td>
<td>Putative ABC – <em>Enterococcus faecalis</em></td>
<td></td>
</tr>
<tr>
<td>18.</td>
<td>Putative ABC – <em>Streptococcus pneumoniae</em></td>
<td></td>
</tr>
<tr>
<td>19.</td>
<td>Putative ABC – <em>Chlorobium tepidum</em></td>
<td></td>
</tr>
</tbody>
</table>

*Against transporters indicates mutation(s) in protein sequence, which are detailed in the Table S1. The octanol/water partition coefficients ([logP<sub>oct</sub>] from www.chemicalize.org) of the carotenoids show increasing hydrophobicity (decreasing polarity) from zeaxanthin to lycopene. Structural comparisons of the carotenoids with validated biofuel compounds, squalene and botryococcene (18, 30), are shown.

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Fig. 1. A battery of 19 bacterial ABC transporters were tested for the secretion of four isoprenoid carotenoid molecules that were chosen as model biofuel compounds in this study. *Against transporters indicates mutation(s) in protein sequence, which are detailed in the Table S1. The octanol/water partition coefficients ([logP<sub>oct</sub>] from www.chemicalize.org) of the carotenoids show increasing hydrophobicity (decreasing polarity) from zeaxanthin to lycopene. Structural comparisons of the carotenoids with validated biofuel compounds, squalene and botryococcene (18, 30), are shown.
secretion by StMsbA* and EcoMsbA, respectively, were unlikely to be a result of secondary effects like lipid A cotransport or the expression of unrelated pumps/proteins. The failure of many transporters in these screens, particularly the conserved MsbA-homologs, suggests that even minor differences in protein sequence have notable effects on biofuel secretion.

We could detect zeaxanthin secreted by StMsbA* in the decane phase after 24 h of aerobic incubation at 30 °C, and the secreted amounts further increased over the next 48 h (Fig. 3A and B). By 72 h, there was a highly significant (P < 0.003) ~2.4-fold increase in secreted zeaxanthin mediated by StMsbA* compared with the nonexpressing control (Fig. 3B). Notably, there were no reductions in produced zeaxanthin amounts per cell (Fig. 3C) or cell viabilities (Fig. S5A), suggesting that the increase in secreted zeaxanthin was not due to cell death. An ~4.4-fold, significant (P < 0.03) increase in secreted canthaxanthin was not due to cell death. An ~4.4-fold increase in secreted canthaxanthin was not due to cell death. An ~4.4-fold increase in secreted canthaxanthin was not due to cell death.

In this paper, the process of biofuel export by MsbA across the IM, combined with the partitioning of biofuels into the organic phase, is collectively termed biofuel “secretion.”

**Isoprenoid Secretion from Washed Cells in Buffer.** In our isoprenoid secretion experiments described above, the double transformant *E. coli* cells were grown in decane-overlayed LB medium overnight and IPTG-induced. *E. coli* cells were grown in decane-overlayed LB medium overnight and IPTG-induced. (A) Physiologically, the lipid A-core moiety of LPS is exported from the inner leaflet to the outer leaflet of the inner membrane (IM) by the multidrug ABC exporter, MsbA, in *E. coli* (20). A network of different transporters and lipid transfer proteins (all dark blue symbols) subsequently participate in moving the lipid A-core (assembled into LPS) through the periplasmic space (PS) to the outer membrane (OM), during LPS biogenesis (21). (B) Carotenoids that are C40 isoprenoid compounds (Fig. 1) were selected as model biofuel compounds. The IM localization of biosynthesized carotenoids in *E. coli* resembles that of lipid A, making them accessible to MsbA for export. In the absence of specific transfer proteins for the “foreign” carotenoids in *E. coli*, the time taken for carotenoids to partition through the PS and OM into the organic phase increases with an increase in carotenoid hydrophobicity. Eventually, a threshold in hydrophobicity is reached where the partitioning is negligible. In this paper, the process of biofuel export by MsbA across the IM, combined with the partitioning of biofuels into the organic phase, is collectively termed biofuel “secretion.”

**Isoprenoid Hydrophobicity Affects Secretion Detection Times.** Zeaxanthin and canthaxanthin are “foreign” to the heterologous host, *E. coli*, which is unlikely to harbor specific carotenoid-binding/transfer proteins. As a result, we hypothesized that the partitioning of free carotenoid molecules from the membranes and periplasm into decane should primarily depend on their chemical polarity and hydrophobicity (Figs. 1 and 2). Canthaxanthin (detected in decane after 96 h) is moderately more hydrophobic than zeaxanthin (detected in decane after 24 h; Fig. 1), which according to our hypothesis, should be responsible for the longer incubation time required.

To further test this hypothesis, we attempted to measure the secretion of more hydrophobic carotenoids: β-carotene and lycopene (Fig. 1). From our ATPase measurements (described later), we had an indication that EcoMsbA could interact with β-carotene. In our secretion experiments using intact cells in culture medium, we found a highly significant (P < 0.008) ~4.4-fold increase in secreted β-carotene from EcoMsbA-expressing cells versus nonexpressing control (Fig. 5A and B), without any

**Fig. 2.** Schematic representation of the biofuel secretion system. To collect and detect hydrophobic biofuel compounds secreted from *E. coli* cells (golden circles), a two-phase culture system was adopted. A biocompatible organic solvent phase was layered over *E. coli* liquid cultures. (A) Physiologically, the lipid A-core moiety of LPS is exported from the inner leaflet to the outer leaflet of the inner membrane (IM) by the multidrug ABC exporter, MsbA, in *E. coli* (20). A network of different transporters and lipid transfer proteins (all dark blue symbols) subsequently participate in moving the lipid A-core (assembled into LPS) through the periplasmic space (PS) to the outer membrane (OM), during LPS biogenesis (21). (B) Carotenoids that are C40 isoprenoid compounds (Fig. 1) were selected as model biofuel compounds. The IM localization of biosynthesized carotenoids in *E. coli* resembles that of lipid A, making them accessible to MsbA for export. In the absence of specific transfer proteins for the “foreign” carotenoids in *E. coli*, the time taken for carotenoids to partition through the PS and OM into the organic phase increases with an increase in carotenoid hydrophobicity. Eventually, a threshold in hydrophobicity is reached where the partitioning is negligible. In this paper, the process of biofuel export by MsbA across the IM, combined with the partitioning of biofuels into the organic phase, is collectively termed biofuel “secretion.”
significant reductions in the produced amount (Fig. 5C) and cell viability (Fig. S5C). However, \( \beta \)-carotene took 120–144 h to be secreted, which is 1–2 d longer than it took for canthaxanthin.

Lycopene is one of the most hydrophobic carotenoids and is slightly more hydrophobic than \( \beta \)-carotene (Fig. 1). Yoon et al. (28) reported earlier that very little lycopene could be recovered from \( E. \ coli \) despite culturing in two-phase systems with decane, unless the outer membrane was removed through spheroplast formation. We reproduced this experimental finding by making spheroplasts out of our double transformant \( E. \ coli \) cells (Fig. S6A), using the same and well-established osmolytic method (28, 29). Our transporter screen in spheroplasts revealed three hits.

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**Fig. 3.** MsbA-mediated isoprenoid secretion from cells in culture medium. (A) Zeaxanthin-producing \( E. \ coli \) BL21(DE3), overexpressing StMsbA* against nonexpressing control, were grown in 10 mL LB medium containing 25 \( \mu \)g/mL chloramphenicol, 50 \( \mu \)g/mL carbenicillin, and 1 mM IPTG, gently overlayed with 5 mL decane, at 30 °C, with aeration at 250 rpm for 72 h, with aliquots of the decane and culture phases taken every 24 h. The image represents the 48-h time point. The absorbencies of the decane phases at the absorbance maxima (\( \lambda_{\text{max}} \)) – 450 nm are shown for comparison (\( n = 3 \) biological repeats, i.e., performed with independently cultured cells ± SEM). (B) Secreted zeaxanthin concentrations in the decane phase were measured spectrophotometrically using a standard curve (\( n = 3 \) biological repeats ± SEM; \( P \) from two-tailed, paired Student t tests). (C) Cells from the culture phase were harvested by centrifugation at 16,000 \( \times \) g for 1 min. Supernatant was removed, and the cell pellets were treated with acetone for 15 min at 55 °C with thorough agitation. The cell debris was collected by a second centrifugation, and the produced zeaxanthin concentration, now in the acetone phase, was analyzed spectrophotometrically. These values were normalized against the bacterial cell number, estimated using absorbance (OD) at 595 nm of 1 = 8 \times 10^{11} \text{ cells/L} (\( n = 3 \) biological repeats ± SEM; \( P \) from two-tailed, paired Student t tests). (D–F) Canthaxanthin overproducing \( E. \ coli \) BL21 (DE3) cells, overexpressing \( E. \ coli \) MsbA, or nonoverexpressing controls were grown and subjected to canthaxanthin secretion and production analyses as described in A–C (canthaxanthin \( \lambda_{\text{max}} = 460 \) nm). Data obtained at the 96-h time point are shown (\( n = 4 \) biological repeats ± SEM; \( P \) from two-tailed, paired Student t tests).

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**Fig. 4.** MsbA-mediated isoprenoid secretion from washed cells in buffer. (A) Zeaxanthin- or (B) canthaxanthin-producing cells containing the pET19b-derived plasmid for StMsbA* or EcoMsbA overexpression or the pET19b empty vector control were grown in 5 mL LB medium containing 25 \( \mu \)g/mL chloramphenicol and 50 \( \mu \)g/mL carbenicillin at 37 °C, with aeration at 200 rpm to OD_{595} 0.5–0.6; 1 mM IPTG was then added, and the cultures were grown for a further 4 h under the same conditions. Cells were then harvested at 3,000 \( \times \) g, 10 min, and 4 °C. Pellets were washed with ice cold 50 mM KPi buffer, pH 7.0, containing 5 mM MgSO_{4}. After three similar washes, cells were resuspended in 5 mL wash buffer containing 0.5% glucose, gently overlayed with 2 mL decane, and incubated at 30 °C, with aeration at 250 rpm. Secreted and produced zeaxanthin/canthaxanthin amounts and cell viability were measured as described in the legend to Fig. 3. Fold change for the values obtained from transporter-expressing cells compared with control cells are presented, where a fold change of 1 represents identical values (zeaxanthin: \( n = 4 \) biological repeats ± SEM; canthaxanthin: \( n = 5 \) biological repeats ± SEM; Student t tests as in legend to Fig. 3).
for lycopene secretion, out of which StMsbA* resulted in a highly significant ($P < 0.004$) ~4.3-fold increase in secreted lycopene (Fig. S6B).

These results strongly support our hypothesis: the time taken for secreted isoprenoids to appear in the decane phase in two-phase *E. coli* culture systems increases with the increase in hydrophobicity of the isoprenoid. In other words, isoprenoid secretion out of cells is not synchronous with isoprenoid export across the inner membrane; the former is dependent on the partitioning of isoprenoids between the membranes, periplasmic space, and decane, and the latter (transporter-mediated export) is dependent on the transporter’s kinetics per se.

**In Vitro Detection of Isoprenoid-ABC Transporter Interactions.** To substantiate our in vivo results on isoprenoid secretion mediated by ABC transporters, we attempted to obtain in vitro evidence of substrate-like interactions between isoprenoid molecules and the ABC transporter MsbA. Many compounds (substrates, inhibitors, or modulators) that interact with ABC transporters modulate (stimulate/inhibit) their ATP hydrolysis (ATPase) activities in a concentration-dependent manner (22). We tested isoprenoid compounds for their ability to modulate the ATPase activity of EcoMsbA.

Organic solvents such as decane, hexane, acetone, ethanol, or methanol readily denature pure detergent-solubilized proteins. Thus, the poor solubility of isoprenoids in aqueous or polar aprotic solvents such as DMSO that are routinely used for ATPase studies presented a problem for this assay. We observed that 1 mM aprotic solvents such as DMSO that are routinely used for ATPase production readily denature pure detergent-solubilized proteins. Thus, the poor solubility of isoprenoids in methanol-water solutions were used as background controls. The background-subtracted ATPase activity of EcoMsbA revealed a β-carotene concentration-dependent stimulation-inhibition pattern that is characteristic of many known substrates of MsbA and other ABC transporters (Fig. 6A). Furthermore, the concentration range across which stimulation was observed (0.5–2 μM) is consistent with the lower micromolar affinity constants observed for almost all substrates of ABC transporters (Fig. 6B) (22). These data strongly suggest substrate-like interactions between isoprenoids and MsbA and provide excellent complementation to our in vivo data demonstrating isoprenoid secretion by MsbA.

To further ensure that the ATPase modulation observed was not due to methanol present in β-carotene solutions, we resorted to test substrate-like interactions between EcoMsbA and other terpenoid compounds that could be dissolved in routine solvents like DMSO. We observed that essential oils, such as spearmint oil, for example, could be readily mixed with DMSO at 0.1%. The DMSO background-subtracted ATPase activity of EcoMsbA showed a similar stimulation-inhibition pattern as was observed for β-carotene (Fig. 6B), lending further in vitro evidence for substrate-like interactions between oily model biofuel compounds and ABC transporters.

**Discussion**

Harvesting large quantities of cells and chemically extracting biofuels are major hindrances to successful commercialization of biofuel production. The use of ABC transporters to facilitate the export of specific biofuel molecules from cells, abating the need for harvest and extraction, could enable easy extracellular recovery and sustained intracellular production. To our knowledge, this study is the first demonstration of using ABC transporters to mediate the secretion of isoprenoid biofuel compounds from intact, healthy, biofuel-producing cells. Concurrent with biofuel secretion, cells rapidly replenish their intracellular biofuel reservoirs allowing production to be sustained over a longer period. In our *E. coli* model test system, the secretion process was sustained for at least 6 d without the need to replenish the growth medium or culture. Thus, for the same quantity of biofuel produced...

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**Fig. 5.** EcoMsbA secretes β-carotene. The assay was performed, and the results were analyzed and are presented in a manner identical to the description in legend to Fig. 3. (A and B) Secreted β-carotene was quantified (A-absorbance at λ$_{max}$ = 453 nm) in the decane phase after 144-h incubation at 30 °C, with aeration at 250 rpm, and was significantly higher for EcoMsbA-expressing cells compared with nonexpressing control ($n = 3$ biological repeats ± SEM; Student t test as in the legend to Fig. 3). Control was indistinguishable from background, probably due to the relatively lower expression of genomic MsbA and negligible passive diffusion of β-carotene, but was nevertheless quantified for uniformity. (C) No significant differences ($P > 0.11$) were observed in the intracellular produced amounts of β-carotene between EcoMsbA-expressing and nonexpressing cells.

**Fig. 6.** In vitro substrate-like interactions between biofuel compounds and MsbA. ATP hydrolysis (ATPase) activity of purified EcoMsbA was measured by the colorimetric detection of Pi using the absorbance of malachite green-ammonium molybdate at 595 nm, as established elsewhere (31). Briefly, detergent-purified MsbA was incubated with substrates on ice for 15 min, before the addition of 2.5 mM ATP, incubation at 30 °C for 5 min, and subsequent addition of the colorimetric reagent. Substrates included (A) β-carotene in methanol-water (see text for main details) or (B) spearmint oil in DMSO at the indicated concentrations controlled against equal volumes of the diluents, methanol-water, or DMSO. Control values were subtracted, and substrate-stimulated ATPase activities of EcoMsbA are presented ($n = 4$ independent experiments ± SEM for both experiments).
conventionally, we have a dramatic reduction in biomass scale and significant gain in the ease of recovering the biofuel.

The possibility of extending this study for liquid alkanes/alkenes could abate the requirement of a two-phase system necessary to collect the secreted isoprenoids tested in this study. These test carotenoids are solids in their pure state and insoluble in water; hence, the need for decane in our assays (Fig. 2). This strategy, whereas attractive for the commercial production of carotenoids, is, however, not commercially feasible for biofuel production on a large scale. Instead, isoprenoids that are liquids under standard conditions, such as squalene produced by microbes, animals, and plants, and botryococcenes produced by the *Botryococcus braunii* Race B, may be more attractive (18, 30). Most of these oils have proven to be direct substitutes for crude oil in the production of gasoline (18). If these oils can be secreted in large enough quantities, they could potentially overcome the partition threshold to generate an oil layer above the culture medium that can be easily harvested.

Our model system coupling a biofuel synthetic pathway with an ABC transporter system has the potential to become plug-and-play, which is important for translating to other organisms like cyanobacteria, diatoms, algae, and yeast. The promiscuity of ABC transporters throughout all kingdoms of life significantly increases the probability of their successful heterologous expression in the production host. The poly-specificity of these proteins also provides confidence in obtaining a hit when screening a small number of ABC transporters that may be preselected based on expression in the desired host. Large-scale production of advanced biofuels, which includes harvesting, extraction, and refining, is currently pursued by several biotech companies. We envision that our plug-and-play secretion system would add significant value when applied to these commercial ventures.

Supporting Information

Doshi et al. 10.1073/pnas.1301358110

SI Materials and Methods

Bacterial Strains, Plasmids, and Culture Conditions. pAC184-derived plasmids for carotenoid production, i.e., pAC-ZEAXipi (zeaxanthin), pAC-CANTHIpI (canthaxanthin), pAC-BETAipi (β-carotene), and pAC-LYClyкопene (lycopene) (1) were kind gifts from Francis Cunningham (University of Maryland, College Park, MD) and Juergen Polle (City University of New York, Brooklyn, NY). Nineteen different ATP-binding cassette (ABC) transporters were cloned into pET19b from PCR products using the restriction enzymes NdeI and BamHI or XhoI. All 19 transporters were sequence verified to confirm their identities. *Escherichia coli* Top10 cells were used to host single plasmids for transformation and plasmid preparation purposes, whereas BL21 (DE3) cells were used for double transformations, production of carotenoids from pAC184-derived plasmids, and the expression of transporters from pET19b-derived plasmids. All transformations were performed using the heat-shock method. Cultures were grown in Luria-Bertani (LB) medium containing 25 μg/mL chloramphenicol (for pAC plasmids) and 50 μg/mL carbenicillin (for pET plasmids); 1 mM isopropyl-β-D-1-thiogalactopyranoside (IPTG) was used to induce expression of the transporter from pET plasmids.

Two-Phase Culture System. To detect the secreted carotenoids, a two-phase culture system was adopted for growing *E. coli* (2–6). Double transformant colonies (containing pAC184- and pET19b-derived plasmids) were picked into fresh LB medium containing the two antibiotics and grown overnight at 37 °C, with aeration at 200 rpm. Glycerol stocks were made from these overnight cultures using 25% (vol/vol) sterile glycerol. For secretion experiments, overnight cultures of the double transformants were set up from glycerol stocks in LB medium containing the two antibiotics growing at 37 °C, with aeration at 200 rpm. For transporter screens (Figs. S3 and S4), 0.5 mM overnight cultures were used to inoculate 5 mL LB medium containing the two antibiotics and 1 mM IPTG and were gently overlayed with 1 mL decane (>99% purity; Fisher/Acros Organics). For thorough analyses of transporter-mediated carotenoid secretion (Figs. 3 and 5), 1 mL overnight cultures were used to inoculate 10 mL LB medium with the antibiotics and 1 mM IPTG and were gently overlayed with 5 mL decane. Two-phase cultures were incubated at 30 °C, with aeration at 250 rpm under mild-absent light conditions for 24–144 h, depending on the carotenoid and experiment.

Carotenoid Secretion from Washed Cells in Buffer. Overnight cultures (0.5 mL) of double transformants were inoculated into 5 mL LB medium with the two antibiotics and aerobically grown at 37 °C, 200 rpm for, for 2–3 h until the OD₅₉₅ reached 0.5–0.6 (log phase); 1 mM IPTG was added to all cultures that were further grown aerobically at 37 °C, 200 rpm, for 4 h. Cells were collected by centrifugation at 3,000 × g, 10 min, and 4 °C, and washed with 10 mL of ice cold 50 mM KPi (potassium phosphate) buffer containing 5 mM MgSO₄. After two more similar washes, the cells were resuspended in 5 mL of the wash buffer supplemented with 0.5% (wt/vol) sterile glycerol and were gently overlayed with 2 mL decane. These two-phase cell suspensions were incubated at 30 °C, with aeration at 250 rpm for 24–96 h, depending on the carotenoid.

Analytical Measurements. At any given time point, aliquots were drawn from the two-phase cultures sufficient to make three different spectrophotometric measurements; secreted carotenoid, produced carotenoid, and cell viability. Secreted carotenoid amounts were measured in the decane phase, after removing any membrane debris by centrifugation at 16,000 × g for 1 min. Pure decane (stored in the absence of light to avoid noisy spectra) was used to blank the spectrophotometer (Beckman DU 800), and the concentrations of carotenoids in the samples were extrapolated from standard curves generated from analytical standard carotenoids dissolved in decane (zeaxanthin, Sigma Aldrich; canthaxanthin, Santa Cruz Biotech; β-carotene, MP Biomedicals; lycopene, Toronto Research Chemicals). Some carotenoid solutions needed to be heated with stirring to allow the carotenoids to dissolve completely. All spectrophotometric measurements involving organic solvents were made in quartz microcuvettes with a path length of 1 cm. The molar extinction coefficients (ε) in decane were calculated as the averages of all data points used to derive the respective standard curves, using the formula A = εcLs, where A is absorbance; ε is molar extinction coefficient; L is path length; and c is concentration. εs of carotenoids are known to be highly solvent dependent (7). In the absence of published ε in decane, we compared our calculated ε to the published values obtained in other solvents (8). The extent of the agreement depended on the hydrophobicity of the carotenoid; lycopene (ε_decane = 159 × 10⁵ M⁻¹ cm⁻¹; ε_hexane = 186 × 10⁵ M⁻¹ cm⁻¹) and β-carotene (ε_decane = 132 × 10⁵ M⁻¹ cm⁻¹; ε_hexane = 139 × 10³ M⁻³ cm⁻³) agreed the best, canthaxanthin (ε_decane = 987 × 10² M⁻² cm⁻¹; ε_petroleum ether = 124 × 10⁵ M⁻¹ cm⁻¹) agreed moderately, and zeaxanthin’s coefficient (ε_decane = 298 × 10¹ M⁻¹ cm⁻¹; ε_petroleum ether = 134 × 10³ M⁻¹ cm⁻¹) was found to be considerably lower in decane. These comparisons suggested low solubility of the polar carotenoids zeaxanthin and canthaxanthin in decane, and therefore their absolute quantities must be interpreted with caution. Produced carotenoid was measured from the aqueous fractions of the two-phase cultures. Cells were collected from a 200 mM aliquot by centrifugation at 16,000 × g for 1 min, and the supernatant was removed carefully using a pipette. Cell pellets were treated with 200 mM acetone and were incubated at 55 °C for 15 min with vigorous intermittent vortices to extract the intracellular carotenoid content. The cell debris was then removed by centrifugation at 16,000 × g for 2 min, and absorbance of the supernatant was measured, after blanking the spectrophotometer with acetone (certain pure carotenoid standards were insoluble in acetone at the required concentrations to generate ε_acetone, so ε_decane was used instead, for uniformity with secreted concentrations). Finally, cell viability was assessed from the absorbance of 200 μL of the aqueous culture phase, using a 595-nm filter on a plate reader (Beckman DTX 880). Cell number was calculated using the relation OD₅₉₅ of 1 = 8 × 10⁻¹³ cells/L and was used to calculate the produced carotenoid amounts, presented as milligram per cell × 10⁻¹³.

Spheroplast Formation. For lycopene secretion experiments, spheroplasts were made out of healthy BL21 (DE3) cells harboring pAC184- and pET19b-derived plasmids, using a method described before (2, 9, 10). Overnight cultures of the double transformants were collected by centrifugation at 3,000 × g, 15 min, for 4 °C. The supernatant was removed, and the cell pellets were washed once with 10 mM KPi buffer, pH 7.0. After collecting the cells again by centrifugation, the pellets were resuspended in 10 mL of a 0.5 M sucrose solution in 10 mM KPi buffer, pH 7.0. Lysozyme (chicken egg white) was then added to a final concentration of 50 μg/mL, and the samples were incubated in a 37 °C water bath for 2 h. After the incubation, 10 mM KPi buffer, pH 7.0, was added to dilute the samples 1:1. Subsequently, 10 mM
K-EDTA, pH 7.0, was added, and the samples were incubated at room temperature for 15 min. The spheroplasts were collected by centrifugation at 500 × g, 15 min, for 4 °C. The supernatant was gently removed, and the spheroplasts were resuspended in 10 mL LB medium containing the two antibiotics and 1 mM IPTG and were gently overlayed with 5 mL decane. These two-phase spheroplast cultures were grown at 30 °C, with aeration at 250 rpm under mild-absent light conditions.

**ATP Hydrolysis (ATPase) Measurements.** ATPase activity measurements on purified *E. coli* MsbA (EcoMsbA) were performed using a colorimetric method described elsewhere (11, 12); 0.5 μg of detergent-purified EcoMsbA (13) was used for each measurement, and 100 mM potassium-Hepes buffer, pH 7.0, containing 5 mM MgSO₄ was used as the assay buffer. EcoMsbA diluted in the assay buffer was mixed with a range of concentrations of the substrates to be tested (or equal volumes of their solvents as controls) and incubated on ice for 15 min. The substrates were β-carotene (predominantly trans; MP Biomedicals) (prepared by first making a 1 mM stock in methanol and then diluting 50 μL of it in 1 mL water to produce a 50 μM working stock in 1:20 methanol:water) and spermidine oil (100%, Now Essential Oils) [prepared fresh as a working stock of 0.1% (vol/vol) in anhydrous DMSO]. After incubation on ice, 2.5 mM ATP (dissolved in the assay buffer) was added to each reaction simultaneously using a multichannel pipette. The reactions were incubated at 30 °C for 5 min, followed by protein denaturation at 80 °C for 15 s. A malachite green-aminommonium molybdate solution, freshly activated using 1:100 10% Triton X-100, was added to each reaction and incubated for 5 min at room temperature with mild shaking. The absorbance was read on a plate reader (Beckman DTX 880) using a 595-nm filter, and the concentration of Pi was extrapolated from a Pi standard curve. ATPase values are presented as nmol of Pi liberated from hydrolyzed ATP per minute per milligram of transporter.


**Fig. S1.** The absorbance spectra of all carotenoids used in this study. Analytical standard carotenoids, zeaxanthin, canthaxanthin, β-carotene, and lycopene, were dissolved in decane and tested for absorbance in the visible region of the spectrum at the indicated concentrations, blanked against decane. Each carotenoid revealed a characteristic absorbance spectrum between 380 and 540 nm with the following absorbance maxima (λₘₐₓ): zeaxanthin, 450 nm; canthaxanthin, 460 nm; β-carotene, 453 nm; lycopene, 474 nm. The molar extinction coefficients (ε) obtained in decane at the λₘₐₓ were as follows: zeaxanthin, 298 × 10⁶ M⁻¹ cm⁻¹; canthaxanthin, 987 × 10⁶ M⁻¹ cm⁻¹; β-carotene, 132 × 10⁶ M⁻¹ cm⁻¹; lycopene, 159 × 10⁶ M⁻¹ cm⁻¹ (see SI Materials and Methods for further details).
Fig. S2. The expression of 19 ABC transporters used in this study. *E. coli* BL21 (DE3) cells harboring pac184-based plasmid for producing the carotenoid and pET19b-derived plasmid for expressing the ABC transporter were alkaline lysed and loaded onto a reducing 10% SDS/PAGE gel. Western blot analysis using the anti-6xHis primary antibody (Santa Cruz Biotech) showed that apart from transporter numbers 4, 6, and 12, the expression of the rest was detectable, and the proteins migrated close to their predicted sizes (for transporter names and sequences 1–19; Table S1). Cells harboring pET19b empty vector control were devoid of a positive signal.
The constitutive production of zeaxanthin in *E. coli* was achieved through the use of the plasmid pAC-ZEAXipi (1). Double transformant cells producing zeaxanthin and expressing the panel of transporters were grown in two-phase culture systems as described in the legend to Fig. 3, with the following exceptions: 1 mL decane was overlayed over 5 mL cell culture, and the incubation period was 24 h. The relative amounts of zeaxanthin in the samples were estimated by comparing the absorbance values in the visible region of the spectrum ($\lambda_{\text{max}} - 450$ nm). The raw absorbance values are presented here. (A and B) Secreted zeaxanthin was measured in the decane fractions of the samples. Debris was removed using centrifugation at 16,000 $\times$ g for 1 min before making absorbance measurements. (C and D) Produced intracellular zeaxanthin amounts were measured using 1-mL cell pellets that were treated with acetone for 15 min at 55 °C, after which cell debris was removed by centrifugation as above. (E) Cell viability was assessed by measuring the absorbance cultures at 595 nm. Transporter number 2, i.e., *Salmonella enterica ser. typhimurium* MsbA, containing mutation I89T (abbreviated StMsbA*), was the best hit in this screen for zeaxanthin secretion.
Fig. S4. Transporter secretion screen for canthaxanthin. The constitutive production of canthaxanthin in *E. coli* was achieved through the use of the plasmid pAC-CANTHipi (1). The screen was performed in an identical manner described in the legend to Fig. S3, except the incubation period was 48 h. As suggested by the relative peak absorbance values at $\lambda_{\text{max}}$ = 460 nm pertaining to (A and B) secreted and (C and D) produced canthaxanthin and (E) the viability of cell cultures, transporter number 8, i.e., *E. coli* MsbA (abbreviated EcoMsbA), was the best hit for canthaxanthin secretion.
Fig. S5. Transporter expression facilitates cell viability. In double transformant E. coli BL21 (DE3) cells overexpressing StMsbA* or EcoMsbA, and producing (A) zeaxanthin (n = 3 biological repeats ± SEM), (B) canthaxanthin (n = 4 biological repeats ± SEM), or (C) β-carotene (n = 3 biological repeats ± SEM), the cell viability measured through absorbance at 595 nm was found to be significantly higher compared with the carotenoid-producing cells containing the non-transporter-expressing empty vector pET19b (P from two-tailed, paired Student t tests). The time points for measurements were 24–72 h for zeaxanthin-, 96 h for canthaxanthin-, and 144 h for β-carotene–producing cells.
Lycopene secretion from spheroplasts. The production of lycopene in *E. coli* was achieved using the plasmid pAC-LYC (1). (A) Lycopene-producing *E. coli* BL21 (DE3) cells expressing StMsbA* were grown in two-phase culture systems with decane. However, as observed by others (2), we were unable to achieve any extraction of lycopene into the decane phase unless we used *E. coli* spheroplasts instead of whole cells. Spheroplasts were made using a well-established osmolytic method (2, 9, 10). However, the time taken for lycopene to be detected in the decane phase varied between 24 and 72 h, probably due to the variability in spheroplast formation and the proportion of revertants. (B) StMsbA*-mediated secretion, as extrapolated from *λ*max = 474 nm, was found to result in a significant ~4.3-fold increase in lycopene in the decane phase compared with the nonexpressing control (*P* < 0.004; two-tailed paired Student *t* test, *n* = 3 biological repeats ± SEM). Unlike healthy whole cells, spheroplasts did not actively divide during the incubation, as monitored at OD$_{595}$.

Table S1. Nineteen ABC transporters bearing homology with MsbA were selected to screen for the secretion of isoprenoid compounds in this study

<table>
<thead>
<tr>
<th>No.</th>
<th>ABC transporter</th>
<th>NCBI accession no.</th>
<th>Mutations</th>
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<tr>
<td>1</td>
<td>MsbA–<em>Vibrio cholerae</em></td>
<td>ZP_01948624</td>
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<td>MsbA–<em>Shewanella sp.</em> MR-1/MR-4</td>
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<td>Δ(M1-H54), G147R, S354N, Q391K, A491V, H506N, I549T.</td>
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<td>Q352P.</td>
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<td>E480D.</td>
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<td>L495, Y416C.</td>
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<td>Putative ABC–<em>Chlorobium tepidum</em></td>
<td>NP_661163</td>
<td>T450A, D524G.</td>
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</table>

Among the 19 transporters, some had mutations that were created for the purpose of solving X-ray crystal structures.

*The protein sequence of this transporter has similar (≈97%) identity with two sequences in the NCBI database: MsbA–*Shewanella oneidensis* MR-1 (NP_718380.1) and *Shewanella sp.* MR-4 (YP_734550.1). Because this transporter was not found to be a positive hit for biofuel secretion, we did not investigate its identity any further. The *N*- and *C*-terminal sequences of the transporter are given for the reference of the reader: *N* terminus, MTASPKDEMWTVFKRL-LAYLKPMMGFLLSVVAFLVVGLVDAAFISFGIDKFG555TPAISNGIALPNTQGFHADNVLLAPVVLIMFLSRLGFANFVYSGYSMSARLMDMRQOOGVHVYSYMUKK-ENTGNJLSKVTDFDQARASGALISVRDGYTVIGMLMFYNSKWSLCLVIQMPMLGMVMTISVRFKVSQOJTMGDSVAATEOMIGKHSGNVLAFGGOETETARFKAINDRR-HQNMLIAAQISQPLMVWGSFAL; and *C* terminus, PESDTGYTVKRAKGFLRDNVSFGYEQQERRALDKIFEVSQGQTALVRGGSGKSTIAPSLVTRFYTGALSGDILDDVSI-YDYSKLNSLRVLVSQVYTLFDNTIAISAYAYFPGEVREQIIEATLALAHMEFIELPDGLDQTVGENVLLSGGQRQIAIARMLRDAVPVLIIDEATSALDTESEKAIQQGLDN- LRQNRTSVMHRLSTIESAQDLILVUGRIVERGTHKSLLELGGMYAKLYQMQFGS.

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