Engineering of *Ralstonia eutropha* H16 for Autotrophic and Heterotrophic Production of Methyl Ketones

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**Running title:** Autotrophic methyl ketone production in *R. eutropha*

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Abstract

*Ralstonia eutropha* is a facultatively chemolithoautotrophic bacterium able to grow with organic substrates or H₂ and CO₂ under aerobic conditions. Under conditions of nutrient imbalance, *R. eutropha* produces copious amounts of poly[(R)-3-hydroxybutyrate] (PHB). Its ability to utilize CO₂ as a sole carbon source renders it an interesting new candidate host for the production of renewable liquid transportation fuels. We engineered *R. eutropha* for the production of fatty acid-derived, diesel-range methyl ketones. Modifications engineered in *R. eutropha* included overexpression of a cytoplasmic version of the TesA thioesterase, which led to a substantial (>150-fold) increase in fatty acid titer under certain conditions. In addition, deletion of two putative β-oxidation operons and heterologous expression of three genes (acyl-CoA oxidase from *Micrococcus luteus* and *fadB* and *fadM* from *E. coli*) led to the production of 50 to 65 mg/l of diesel-range methyl ketones under heterotrophic growth conditions and 50 to 180 mg/l under chemolithoautotrophic growth conditions (with CO₂ and H₂ as the sole carbon source and electron donor, respectively). Induction of the methyl ketone pathway diverted substantial carbon flux away from PHB biosynthesis and appeared to enhance carbon flux through the pathway for biosynthesis of fatty acids, which are the precursors of methyl ketones.
Introduction

*R. eutropha* is an industrially relevant, facultatively chemolithoautotrophic bacterium able to grow with organic substrates or H₂ and CO₂ under aerobic conditions. When experiencing nutrient limitation, *R. eutropha* directs most of its reduced carbon flux to the synthesis of poly[(R)-3-hydroxybutyrate] (PHB), a biopolymer stored in intracellular granules. When cultivated with H₂ and CO₂, *R. eutropha* is able to form up to 61 g/l of PHB in 48 h, representing about 70% of total cell weight (1). Mutants defective in PHB production secrete large amounts of pyruvate into the growth medium when cultured chemolithoautotrophically, suggesting that these mutants maintain a comparable carbon flux in the presence or absence of PHB biosynthesis (2, 3).

Biosynthesis of PHB under nutrient-limiting conditions in *R. eutropha* has been industrially exploited in the production of biodegradable plastic for more than a decade (e.g., by ICI/Zeneca, Monsanto, and Metabolix). Only recently has *R. eutropha* been recognized as a candidate for biofuel production from CO₂, potentially producing carbon-neutral biofuels from non-photosynthetic sources. Li and coworkers engineered a *R. eutropha* strain for electromicrobial conversion of CO₂ to C₄ and C₅ alcohols (4). In their system, electricity powered the electrochemical reduction of CO₂ at a cathode to produce formate, which was then converted to isobutanol and 3-methyl-1-butanol by an engineered *R. eutropha* strain. To meet future transportation fuel demands, higher energy-density, diesel-compatible biofuels such as medium-chain hydrocarbons will also need to be developed.

Medium-chain methyl ketones are promising targets as diesel fuel blending agents (5). Recently, Goh and co-workers demonstrated the production of over 380 mg/l of C₁₁ to C₁₅ methyl ketones from glucose (0.2%) in an engineered *E. coli* DH1 strain, which represented a
700-fold increase in titer relative to a fatty acid-overproducing strain (5). The major features of this E. coli strain were a re-engineered β-oxidation pathway designed to overproduce β-ketoacyl-CoA s and overexpression of the native thioesterase FadM. Subsequently, Park and co-workers (6) described engineering of an E. coli MG1655 strain that was able to produce 14 mg/l of methyl ketones (2% glucose) when overexpressing a β-ketoacyl-ACP thioesterase (shmks2) and a β-keto acid decarboxylase (shmks1) from Solanum habrochaites (wild tomato) and when adhE, ldhA, poxB, and pta were deleted from the chromosome. Batch incubations of the best strain under optimized growth conditions with 5% glucose resulted in a methyl ketone titer of ca. 500 mg/l.

We report here for the first time the autotrophic production of methyl ketones by engineered bacteria (R. eutropha H16). Wild-type R. eutropha does not produce detectable levels of methyl ketones natively and intracellular concentrations of fatty acids (methyl ketone precursors) are very low in this bacterium. In this study, methyl ketone-producing strains were initially developed under heterotrophic conditions and the best performing strain was then tested under autotrophic conditions. R. eutropha engineered with a modified β-oxidation pathway produced up to 180 mg/l of methyl ketones from CO₂ as the sole carbon source.

**Materials and Methods**

**Bacterial strains, media, and cultivation.** Strains used in this study are listed in Table 1. E. coli strains (S17-lambda pir, JM1, JM2, JM3), R. eutropha strains (H16 wild type, JM7, JM8 (Re2303), JM9, JM16), and plasmids (pJM3, pJM9, pJM20) along with their associated information (annotated GenBank-format sequence files), have been deposited in the public version of the JBEI Registry (https://public-registry.jbei.org; entries JPUB_000923-934) and are physically available from the authors and/or addgene (http://www.addgene.org) upon request.
E. coli strains were propagated at 37°C in Lysogeny Broth (LB). Where necessary, medium was solidified with 1.5% (w/v) agar and supplemented with 50 µg/ml kanamycin. *R.* eutropha* H16* strains were grown at 30°C in LB medium supplemented with 300 µg/ml kanamycin and 15 µg/ml gentamycin, where required, or modified minimal medium (7) containing either 2% fructose and 0.1% NH₄Cl (nitrogen-sufficient conditions) or 1% fructose and 0.01% NH₄Cl (nitrogen-deficient growth conditions). For studies of heterologous gene expression, *R.* eutropha strains were grown in 15 ml minimal medium in 30-ml glass tubes with 200 rpm agitation at 30°C. Medium was supplemented as necessary with 300 µg/ml kanamycin and 15 µg/ml gentamycin. L-Arabinose was added at a final concentration of 0.2% (w/v) to cultures requiring gene induction (48 h post-inoculation). All liquid cultures were inoculated with single colonies originating from freshly transformed strains, and were grown for up to 120 h before being harvested for analysis. The first samples for methyl ketones were taken 48 h post-induction.

Plasmids and primers. Plasmids and primers used in this study are summarized in Table 1 and Table S1 (provided in the supplemental material), respectively.

**Plasmid construction for heterologous expression in *R.* eutropha.** Cloning of genes originating from *E.* coli and *Micrococcus luteus* was accomplished by ligase-dependent cloning techniques described elsewhere. All plasmids constructed in this study contain the backbone from the broad-host range vector pBBR1-MCS2 (8), including the functions required for plasmid mobilization. Target genes were amplified using primers listed in Table S1 (provided in the supplemental material). Proper plasmid construction was confirmed by sequencing before transformation into *E.* coli S17-lambda pir (9) (Table 1). Plasmids were subsequently transferred into the desired *R.* eutropha strain by conjugation. Following conjugation, plasmid DNA was
extracted from *R. eutropha* and frozen. Before each expression experiment, *R. eutropha* strains were transformed by electroporation to preclude *E. coli* contamination. Notably, transformation efficiencies were at least 150-fold higher when the plasmid to be introduced into H16 wild-type by electroporation was prepared from H16 wild-type instead of from *E. coli* strains. For electroporation of *R. eutropha*, a single colony was inoculated into LB medium and grown to mid-exponential growth phase. One milliliter of the culture was pelleted and washed with 1 ml sorbitol (pH 7.4). The cell pellet was resuspended in 35 µl sorbitol (pH 7.4) and 1-2 µg of plasmid DNA was added. The cell suspension was transferred in a 1-mm cuvette and electroporation was executed with the following parameters: 0.55 kV, 25 µF, and 200 Ω. Immediately following electroporation, 400 µl of SOC medium was added and the cells were transferred into a 5-ml culture tube. Electroporated cells were recovered for 2 h before being plated onto selective medium.

**Strain construction for heterologous expression in *R. eutropha***. Genome modifications in *R. eutropha* were performed by homologous recombination as described by Brigham and co-workers (10). Briefly, in-frame deletion constructs were created by amplifying the 500-bp regions flanking the target gene. Subsequently the flanking regions were fused via a 20-bp complementary tag that was added to the 5’- end of each inner primer. The fusion product was inserted into the BamHI-digested suicide vector pGY46. pGY46 was previously used for deletion of *phaC1*, so the *phaC1* flanking regions had to be removed by BamHI restriction digestion. The mobilizing strain, *E. coli* S17-lambda pir (9), was transformed with this plasmid. Single-crossover events were selected for on LB plates containing kanamycin and confirmed by colony PCR using two primer combinations: (1) primer X-F and primer 3-O and (2) primer X-R and primer 5-O, where primer X-F and primer X-R will bind upstream and downstream of the
flanking regions, respectively. Resolution of the integrated vector by a second crossover event was performed with positive strains. These strains were grown in LB medium without selection and plated onto solid LB medium containing 10% sucrose. Deletion events were verified by the PCR using primer X-F and primer X-R.

**Quantification of fructose.** Culture medium of *R. eutropha* wild-type and mutants was collected at specified time points. The supernatant was separated from cells by centrifugation and cell debris was removed by filtration (0.45-µm pore size). An Agilent 1200 series binary-pump liquid chromatography system equipped with refractive index and diode array detectors was used to measure fructose in the supernatant. Isocratic separation of fructose was achieved with a Bio-Rad Aminex HPX-87H column (Richmond, CA) at 60°C using a mobile phase of 4 mM H₂SO₄ flowing at a rate of 0.6 ml per min.

**Quantification of pyruvate.** Pyruvate in filtered culture medium was quantified using a commercially available fluorescence-based assay kit (K609-100, BioVision, Mountain View, CA) according to the manufacturer's instructions.

**Extraction and quantification of PHB.** PHB was extracted and quantified according to a method described by Brandl and co-workers (11). One milliliter of chloroform was added to 5-20 mg of lyophilized cells, followed by the addition of 0.85 ml methanol and 0.15 ml concentrated sulfuric acid. The mixture was heated for 2.5 h at 100°C. After allowing the solution to cool down to room temperature, samples were stored on ice and 0.5 ml of water was added. Phases were separated by centrifugation at 2000 x g for 5 min. Subsequently, the organic phase was removed and transferred to a new tube and dried with 200 mg of anhydrous Na₂SO₄. Before analysis by gas chromatography/flame ionization detection (GC/FID), samples were filtered through a 0.2-µm PVDF syringe filter. GC/FID analysis (1 µl injection) was performed.
with Thermo Scientific model Focus GC, which was equipped with a DB-WAX column (15-m length, 0.32-mm inner diameter, 0.25-µm film thickness; J&W Scientific). The GC oven temperature was programmed to hold at 80°C for 5 min before increasing to 170°C at 20°C/min. The temperature of the injection port was 250°C. Ultra high-purity helium served as the carrier gas flowing at 1 ml/min. External standard quantification with 3-hydroxybutyrate methyl ester standards was performed.

**Extraction of fatty acids from bacterial cultures.** Fatty acids were hexane-extracted from cell pellets of *R. eutropha* using modifications of methods described previously (5). All solid materials that came in contact with the hexane extracts during the extraction process were solvent-cleaned. 15-ml cultures of *R. eutropha* wild-type and mutants were grown in 30-ml glass tubes with PTFE-lined screw caps and sacrificed at specified time points. Tubes were centrifuged for 20 min at 4,000 rpm and 20°C. Following centrifugation, the supernatant was decanted and the pellet was resuspended in 100 µl reagent-grade water by vortex mixing. Subsequently, 1 ml of high-purity methanol and 4 ml of high-purity hexane were added to the cells and the mixture was homogenized by vortex mixing and sonication. To determine sample-specific recovery, the hexane was amended with perdeuterated alkane standards (C₁₀D₂₂ and C₂₄D₅₀). Sonication was performed for 15 min in an ice water bath. After sonication, samples were allowed to equilibrate for 10 min at 20°C before being subjected to another centrifugation step at 3,500 rpm for 15 min. After centrifugation, the hexane layer was transferred into a 10-ml conical vial and concentrated to 100 µl under a gentle stream of N₂. Fifty microliters of the concentrated hexane extract were removed, derivatized with ethereal diazomethane to generate fatty acid methyl esters (FAME) (5), and concentrated to 50 µl for GC/mass spectrometry (GC/MS) analysis.
Extraction of methyl ketones from bacterial cultures. Methyl ketones from \textit{R. eutropha} cultures were extracted either using the above-described method for extraction of fatty acids or by means of a decane overlay (5). 50-100 µl of the decane overlay was used for GC/MS analysis directly or was diluted in decane before injection.

Analysis of fatty acids and methyl ketones by GC/MS. Fatty acids (derivatized as methyl esters) and methyl ketones extracted from \textit{R. eutropha} were quantified using electron-ionization GC/MS as described previously (5).

Chemolithoautotrophic fermentation. Pre-cultures for chemolithoautotrophic fermentation were grown by inoculating a single colony of strain JM13 (\Delta beta ox background; Table 1) in 5 ml of LB supplemented with 10 µg/ml gentamycin and 300 µg/ml kanamycin and incubated at 30°C for 24 h with shaking. The culture was diluted 1:100 into 50 ml of minimal medium (12) supplemented with 10 g/l fructose, 2 g/l ammonium sulfate, 10 µg/ml gentamycin, and 300 µg/ml kanamycin. Cultures were incubated in 250-ml baffled flasks at 30°C with shaking (250 rpm) for 72 h. Cells were harvested via centrifugation and washed once in minimal medium absent of any exogenous carbon source.

Chemoautotrophic cultures were grown in a 1-liter bioreactor. Reactors contained 700 ml of minimal medium lacking an organic carbon source and were supplemented with 5 g/l ammonium sulfate and 2 g/l arabinose (for induction of the PBAD promoter). Reactors were inoculated to a starting optical density of 1.0 (9.5 x 10^8 cfu/ml). Decane (66.67 ml) was added to the reactor following inoculation. A blend of hydrogen:oxygen:carbon dioxide (80:4:16) was introduced into the reactor at a flow rate of 0.1 l/min. The reactor was incubated at 30°C with constant stirring (200 rpm). Samples of the culture medium as well as the decane layer were removed periodically using sterile glass Pasteur pipettes and were analyzed as described above.
Detection of CO₂ in batch cultures by the Respiration Activity MOnitoring System (RAMOS). General set-up and operation of the RAMOS system (Kuhner, Switzerland) has been described previously (13). Experiments were performed as follows: eight 250-ml RAMOS flasks were filled with 40 ml of fructose minimal medium and equilibrated overnight. Following equilibration, a sample of the culture medium was taken for fructose analysis. Subsequently, four of the flasks were inoculated with 4 ml of either a pre-adapted JM13 (Δbeta ox background; Table 1) or JM15 (Δbeta ox ΔphaCAB background) culture and incubated at 30°C and 100 rpm. Sixty hours post-inoculation, half of the RAMOS cultures were induced with 0.2% L-arabinose. All cultures were overlain with 2.5 ml of decane at this point. Cultures were sampled for determination of optical density and fructose every 24 h post-inoculation. Decane was sampled for methyl ketone analysis every 24 h post-induction. The experiment was terminated 156 h post-inoculation and all culture cell pellets were collected for PHB analysis. During fermentation, a CO₂ measuring cycle was continuously repeated; the measuring cycle consisted of a 240-min measuring phase and a 10-min purging phase.

Results

Expression of ‘tesA in R. eutropha led to enhanced production of fatty acids and methyl ketones under heterotrophic growth conditions. Overproduction of fatty acids in E. coli was previously achieved by expressing a cytoplasmically directed version of the tesA thioesterase (‘tesA) (14). Overexpression of ‘tesA in a ΔfadE mutant, deficient in β-oxidation, was shown to produce 1.1 g/l of free fatty acids in E. coli (14). Based on these findings, and the fact that methyl ketones derive from fatty acids, we attempted to build fatty acid-overproducing host strains for the production of methyl ketones in R. eutropha. Several strains heterologously expressing ‘tesA were tested for fatty acid production: H16 wild-type (strain JM5; Table 1), a
ΔfadE double mutant (strain JM17; Table 1), as well as Δbeta ox and Δbeta ox ΔphaCAB strains (JM12 and JM14, respectively; Table 1). These strains were cultivated for up to 96 h either in the presence or absence of a decane overlay.

Expression of ‘tesA in R. eutropha wild-type (strain JM5) and a fadE double-mutant host (JM17) led to the production of 10 mg/l of fatty acids when the strains were cultivated in LB medium in the absence of a decane overlay. The fatty acids were detected in the cell pellets and no fatty acids were found in the supernatant of these cultures. Trace amounts (55 µg/l) of fatty acids were detected in a control strain expressing green fluorescent protein (GFP) (strain JM4).

Thus, expression of ‘tesA in H16 wild-type led to a 180-fold improvement in fatty acid production when compared to the GFP control strain. Myristate and palmitate (14:0 and 16:0, respectively) were the major fatty acids produced by these strains (i.e., JM5 and JM17), with lesser amounts of palmitoleate and oleate (16:1 and 18:1, respectively). Notably, in-frame deletions of both fadE copies in R. eutropha did not increase fatty acid production, unlike results reported for E. coli (14). To ensure that β-oxidation was in fact eliminated, we tested ‘tesA expression in a Δbeta ox mutant background (strain Re2303) that had been previously shown to be incapable of metabolizing oleic acid (10). Additionally, we created a Δbeta ox ΔphaCAB mutant strain (strain JM9) that was deficient in PHB synthesis, postulating that acetyl-CoA not being used for PHB biosynthesis might instead feed into fatty acid biosynthesis. Fatty acid production of ‘tesA-overexpressing strains JM12 (Δbeta ox mutant) and JM14 (Δbeta ox ΔphaCAB mutant) was then compared to production under uninduced conditions and in wild-type H16. Under minimal medium conditions (2% fructose), Δbeta ox mutant strains expressing ‘tesA (strain JM12) produced only 0.001 mg/l of total fatty acids, which is comparable to fatty acid production when ‘tesA expression was not induced (Fig. 1) and in wild-type H16 (data not
shown). In contrast, expression of ‘tesA in a Δbeta ox ΔphaCAB mutant strain (JM14) produced 0.017 mg/l of total fatty acids. Thus, deletion of the PHB biosynthesis pathway contributed to a 17-fold increase in fatty acid titer when compared to the uninduced strain or wild-type H16.

Myristate and palmitate (14:0 and 16:0, respectively) were the only fatty acids produced under minimal medium conditions in strain JM14 (Δbeta ox ΔphaCAB mutant). Palmitate and stearate (16:0 and 18:0, respectively) were the two fatty acids detected in the cell pellets of strain JM12 (Δbeta ox mutant). While testing strains JM12 and JM14 for fatty acid overproduction, we observed that both strains also synthesized medium-chain methyl ketones (Fig. 1); ca. 0.5 mg/l methyl ketones were detected in the cell pellets of these strains.

Re-engineering of β-oxidation in *R. eutropha* produced methyl ketones under heterotrophic growth conditions. We observed that *R. eutropha* produced methyl ketones when ‘tesA was overexpressed in both a β-oxidation-deficient host (strain JM12) and a β-oxidation- and PHB-deficient host (strain JM14). To further exploit this ability and increase methyl ketone titers, we modified *R. eutropha* based on a strategy described for the production of methyl ketones from glucose in *E. coli* (5). This strategy involved the following: (a) replacing the native β-oxidation pathway in *R. eutropha* with an overexpressed, heterologous, truncated version of β-oxidation in which the soluble and high-specific-activity acyl-CoA oxidase from *M. luteus* replaces FadE (acyl-CoA dehydrogenase), the *E. coli* version of FadB (which converts *trans*-2-enoyl-CoAs to β-ketoacyl-CoAs) is overexpressed, and *fadA* is deleted, all of which was designed to facilitate the overproduction of β-ketoacyl-CoAs, and (b) overexpressing the *E. coli* thioesterase FadM, thereby converting β-ketoacyl-CoAs to β-keto acids, which can then spontaneously decarboxylate to form methyl ketones.
Of the four genetic backgrounds hosting the heterologous methyl ketone pathway, the highest methyl ketone titer was observed in strain JM13 (Δbeta ox mutant; Table 1) at 96 h post-induction (Fig. 2). This strain produced >50 mg/l of total methyl ketones under heterotrophic growth conditions with fructose as the carbon source. The Δbeta ox ΔphaCAB mutant strain (strain JM15; Table 1) reached similar concentrations, whereas the wild-type (strain JM6) and the ΔphaCAB (strain JM11) hosts produced significantly lower concentrations of methyl ketones (<2 mg/l). Thus, deletion of two putative β-oxidation operons in *R. eutropha*, both of which included fadA, was crucial for higher production of methyl ketones in this microorganism. Notably, almost all of the methyl ketone production occurred during a period of insignificant growth (48 to 96 h post induction; Fig. 2). Methyl ketones were not detectable in any of these strains when the methyl ketone pathway was not induced (Fig. 2). In accordance with the fatty acid distributions observed in *R. eutropha* Δbeta ox ΔphaCAB mutants (JM14), the major methyl ketones detected in this host strain expressing the methyl ketone pathway (JM15) were 13:0 and 15:0 methyl ketones (derived, respectively, from 14:0 and 16:0 fatty acids) (Fig. 3). Minor amounts of 15:1 methyl ketones were observed (derived from 16:1 fatty acids). Unexpectedly, 13:0 and 15:0 methyl ketones were also the major methyl ketones and 15:1 the minor methyl ketone detected in Δbeta ox mutant strains (JM13). This is at least partially in disagreement with the observed fatty acid distribution of this mutant strain (JM12), as the major fatty acids observed were palmitate and stearate (16:0 and 18:0 fatty acids). As shown in Fig. 3, wild-type (strain JM6) and ΔphaCAB (strain JM11) mutant backgrounds showed similar methyl ketone distributions. In these two low-titer strains, 35% of total methyl ketones were represented by 13:0 and 65% by 15:0 methyl ketones. The proportions of 13:0 and 15:1 methyl ketones were higher in the better producing strains (with β-oxidation knockouts).
The methyl ketone pathway diverts carbon flux from the PHB pathway. Biosynthesis of PHB from acetyl coenzyme A (acetyl-CoA) in *R. eutropha* is enabled by enzymes encoded by the *phaCAB* operon. *R. eutropha* produces PHB under conditions of nutrient imbalance when an excess of carbon is available and one or more essential growth factor(s), such as nitrogen or O₂, is limited. We wanted to re-route carbon (specifically, acetyl-CoA) flux from PHB biosynthesis to fatty acid/methyl ketone biosynthesis by deleting the *phaCAB* operon. However, as reported earlier, deleting PHB biosynthesis in the wild-type H16 and ∆beta ox mutant strains expressing the methyl ketone pathway did not lead to increased methyl ketone production (Fig. 2). While deletion of PHB biosynthesis did not lead to improved methyl ketone production under the conditions tested, assessments of PHB production indicated that induction of the methyl ketone pathway did divert substantial carbon flux away from PHB biosynthesis (in strains with intact *phaCAB*). Specifically, we found that, when the methyl ketone pathway was induced, the percentage of fructose carbon allocated to PHB decreased from 9.4% to 2.1% in the wild-type background (strain JM6) and from 9.4% to 1.3% in the ∆beta ox mutant background (strain JM13) (Table 2). Under both induced and uninduced conditions, PHB production was negligible in ∆*phaCAB* control strains tested, namely, strains JM11 and JM15 (Table 2).

As the observed methyl ketone titers cannot account for the magnitude of the carbon flux diverted from PHB by induction of the methyl ketone pathway, we hypothesized that the carbon may be diverted to mineralization of fructose to CO₂. To test this hypothesis, a RAMOS study was conducted with ∆beta ox (strain JM13) and ∆beta ox ∆*phaCAB* (strain JM15) hosts bearing the methyl ketone pathway. Over the entire incubation period, CO₂, fructose, and methyl ketone concentrations were determined in all samples and O₂ was confirmed to be present. Methyl ketone concentrations in these shake flask cultures were comparable to those previously
determined in 30-ml glass tubes (data not shown). As before, we observed a diversion of carbon flux from PHB biosynthesis under induced conditions (Table 3). Specifically, the Δbeta ox mutant (strain JM13) had 7% less fructose carbon utilized for PHB synthesis under induced conditions (Table 3). However, CO₂ constituted less than 1% of fructose carbon consumed for both strains under induced conditions and thus could only account for a small portion of the carbon diverted from PHB. Similarly, methyl ketones accounted for a relatively small portion of fructose carbon (2.2%) (Table 3). Additionally, we have determined that pyruvate accounts for a negligible portion of the carbon diverted from PHB biosynthesis; no pyruvate was detected in the culture medium after incubation of strain JM13 (see Fig. S1 in the supplemental material). Also, acetate was not detected in these samples at concentrations greater than 5 μM. In summary, the fate of the diverted PHB carbon is not fully accounted for and merits further investigation.

Engineered R. eutropha produces methyl ketones autotrophically from hydrogen and carbon dioxide. The ultimate goal of this study was to produce methyl ketones from CO₂ as the sole carbon source. We chose the Δbeta ox mutant expressing the methyl ketone pathway (strain JM13) to investigate autotrophic production, as this strain had the best titer under heterotrophic conditions.

Chemolithoautotrophic cultures were inoculated with pre-adapted cells into minimal medium without an organic carbon source at an initial optical density of 1. L-Arabinose and methyl ketone concentrations as well as optical density were determined every 24 h post-induction. L-Arabinose concentrations remained constant over the entire fermentation period (< 1.5% relative standard deviation for L-arabinose concentrations from the time of induction until the end of incubation in multiple experiments), confirming that R. eutropha was unable to utilize this sugar and thus, CO₂ was the only available carbon source present in the culture. Fig. 4 shows
a time course of methyl ketone production under chemolithoautotrophic conditions. The methyl ketone concentration increased from undetectable levels at induction to ~50 - 70 mg/l at 48 h post induction. In other experiments conducted under autotrophic conditions, methyl ketone titers were as high as 180 mg/l (at OD values comparable to those shown in Fig. 4; see Fig. S2 in the supplemental material). The composition of methyl ketones produced under chemolithoautotrophic growth conditions was very similar to that observed under heterotrophic growth conditions (data not shown). Thus, *R. eutropha* engineered to overproduce methyl ketones is able to generate titers under chemolithoautotrophic conditions that are comparable to (or greater than) those observed under heterotrophic conditions.

**Discussion**

Production of fatty acid-based methyl ketones from glucose was first shown in an engineered *E. coli* DH1 strain at titers of 380 mg/l (5). Our data revealed that similar genetic modifications introduced into *R. eutropha* H16 led to autotrophic production of methyl ketones at concentrations of ~50 to 180 mg/l. Considering the relatively high energetic demands of carbon fixation, it is perhaps surprising that production of methyl ketones by *R. eutropha* strain JM13 under autotrophic conditions was comparable to that under heterotrophic conditions (Figs. 2 and 4). However, the lower titers of methyl ketones under heterotrophic growth conditions can be partially attributed to the low optical densities of the cultures (Fig. 2). Under heterotrophic growth conditions these cultures reached optical densities of ~1, whereas the autotrophic cultures reached optical densities of ~2.

A key feature of this engineered methyl ketone pathway in both *R. eutropha* and *E. coli* is the enhancement of early steps of β-oxidation (here catalyzed by acyl-CoA oxidase and FadB), allowing for production of β-ketoacyl-CoAs, and elimination of the final (FadA-catalyzed)
reaction of \( \beta \)-oxidation, allowing \( \beta \)-ketoacyl-CoAs to be diverted (\emph{via} FadM and spontaneous
decarboxylation) to methyl ketones rather than acyl-CoAs that are two carbons shorter than the
acyl-CoAs entering the cycle. In this context, one can rationalize the large improvement in
methyl ketone titers in \textit{R. eutropha} strains with native \( \beta \)-oxidation knocked out (strains JM13 and
JM15 vs. JM6 and JM11; Fig. 2): the deleted \( \beta \)-oxidation operons include \textit{fadA} (5), which must
be removed to truncate \( \beta \)-oxidation at \( \beta \)-ketoacyl-CoA.

Although fatty acids are precursors for methyl ketones, the relationship between fatty
acid production and methyl ketone production is subtle. For example, deletion of the PHB
biosynthesis pathway resulted in increased fatty acid production (Fig. 1) but comparable methyl
ketone production, either with or without the heterologous methyl ketone pathway (Figs. 2 and 1,
respectively). Nonetheless, our data strongly suggest that expression of the heterologous methyl
ketone pathway greatly increases flux through the fatty acid biosynthesis pathway. To illustrate,
in \( \Delta \beta \)-oxidation (JM13) and \( \Delta \beta \)-oxidation and \( \Delta \text{phaCAB} \) (JM15) strains expressing the methyl
ketone pathway, methyl ketones accounted for 2 to 4\% of the fructose carbon consumed.
However, in the analogous strains expressing \textit{tesA} but lacking the rest of the heterologous
methyl ketone pathway (JM12 and JM14, respectively), we found that free fatty acids accounted
for only ~0.00016\% of the fructose carbon consumed. As methyl ketones derive from fatty
acids, it appears that expression of the methyl ketone pathway substantially increases flux
through the fatty acid biosynthetic pathway (assuming that the amount of fatty acids converted to
phospholipids in strains with and without the methyl ketone pathway does not differ by several
orders of magnitude).

Contrary to our expectations, deletion of PHB biosynthesis in \textit{R. eutropha} did not
improve methyl ketone production. Although we were unsuccessful in diverting acetyl-CoA
toward methyl ketone production in PHB-minus strains, induction of the methyl ketone pathway in PHB-positive strains led to substantial diversion of carbon flux away from PHB biosynthesis (Tables 2 and 3). While we demonstrated that the fate of the diverted carbon is primarily not methyl ketones, CO₂, acetate, or pyruvate, the disposition of this carbon is not currently known. 

A better understanding of metabolic flux in methyl ketone-producing *R. eutropha* strains, and use of that information for engineering refinements, should allow us to substantially increase methyl ketone titers and enhance the viability of chemolithoautotrophic biofuel production. 

**Acknowledgments**

We thank Chris J. Brigham for providing us with the Re2303 mutant strain. We also thank the A. J. Sinskey laboratory (MIT) for general technical advice. We are grateful to Ee-Been Goh for technical advice and helpful discussions and we thank Veronica Teixeira Benites for technical assistance and strain maintenance. This work was funded by DOE's ARPA-E Electrofuels Program to Lawrence Berkeley National Laboratory under contract DE-AC02-05CH11231 and performed at the Joint BioEnergy Institute.
References


FIGURE LEGENDS

**FIG. 1.** Fatty acid and methyl ketone production in *R. eutropha* strains expressing 'tesA. Titers of fatty acids (extracted from the cell pellet and derivatized as FAME) and methyl ketones (accumulated in the decane overlay) are shown for two fructose-grown *R. eutropha* mutant strains expressing 'tesA (mutants with β-oxidation and/or phaCAB chromosomal deletions; Table 1). Asterisks represent strains that were not induced with arabinose. Bar heights represent the average of at least three independent biological replicates and error bars represent one standard deviation at 96 h post-induction.

**FIG. 2.** Heterotrophic methyl ketone production in *R. eutropha*. Methyl ketone titers are shown for four fructose-grown *R. eutropha* strains expressing the methyl ketone pathway (wild type and mutants with β-oxidation and/or phaCAB chromosomal deletions; Table 1). Asterisks represent strains that were not induced with arabinose. For methyl ketones, bar heights represent the average of at least three independent biological replicates and error bars represent one standard deviation. For optical density, datum points represent the average of at least three independent biological replicates.

**FIG. 3.** Distribution of 13:0, 15:0, and 15:1 methyl ketones produced by the *R. eutropha* strains represented in Fig. 2. Data represent the average methyl ketone distributions of three biological replicates at 96 h post-induction.

**FIG. 4.** Autotrophic methyl ketone production in *R. eutropha* strain JM13 (β-oxidation mutant) under a H₂:O₂:CO₂ atmosphere (80:4:16). Data for two biological replicates are shown.
TABLE 1. Bacterial strains and plasmids used in this study

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Relevant genotype or description</th>
<th>Source or reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>S17-lambda pir</td>
<td><em>thi pro recA hsdR [RP4-2Tc::Mu-Km::tn7]lambda pir Tp' Sm'</em></td>
<td>[9]</td>
</tr>
<tr>
<td>JM1</td>
<td>S17 with pJM3</td>
<td>This study</td>
</tr>
<tr>
<td>JM2</td>
<td>S17 with pJM9</td>
<td>This study</td>
</tr>
<tr>
<td>JM3</td>
<td>S17 with pJM20</td>
<td>This study</td>
</tr>
<tr>
<td>E. coli</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R. eutropha</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H16 DSM28</td>
<td>H16 wild type</td>
<td>DSM</td>
</tr>
<tr>
<td>JM4</td>
<td>H16 with pJM3</td>
<td>This study</td>
</tr>
<tr>
<td>JM5</td>
<td>H16 with pJM9</td>
<td>This study</td>
</tr>
<tr>
<td>JM6</td>
<td>H16 with pJM20</td>
<td>This study</td>
</tr>
<tr>
<td>JM7</td>
<td>H16 ΔphaCAB (mutant is deficient in PHB biosynthesis)</td>
<td>This study</td>
</tr>
<tr>
<td>Re2303 (JM8)</td>
<td>H16 Δ(H16_A0459-0464, H16_A1526-1531); (Δbeta ox); mutant is deficient in native β-oxidation</td>
<td>[10]</td>
</tr>
<tr>
<td>JM9</td>
<td>H16 Δ(H16_A0459-0464, H16_A1526-1531)ΔphaCAB; (Δbeta oxΔphaCAB); mutant is deficient in native β-oxidation and PHB biosynthesis</td>
<td>This study</td>
</tr>
<tr>
<td>JM10</td>
<td>JM7 with pJM9</td>
<td>This study</td>
</tr>
<tr>
<td>JM11</td>
<td>JM7 with pJM20</td>
<td>This study</td>
</tr>
<tr>
<td>JM12</td>
<td>Re2303 with pJM9</td>
<td>This study</td>
</tr>
<tr>
<td>JM13</td>
<td>Re2303 with pJM20</td>
<td>This study</td>
</tr>
<tr>
<td>JM14</td>
<td>JM9 with pJM9</td>
<td>This study</td>
</tr>
<tr>
<td>JM15</td>
<td>JM9 with pJM20</td>
<td>This study</td>
</tr>
<tr>
<td>JM16</td>
<td>H16 Δ(H16_A0460, H16_A1530); mutant carries deletions in both copies of <em>fadE</em></td>
<td>This study</td>
</tr>
<tr>
<td>JM17</td>
<td>JM16 with pJM9</td>
<td>This study</td>
</tr>
<tr>
<td>Plasmids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pJM3</td>
<td>Km'; broad host range vector pBBR1-MCS2 containing <em>gfp</em> under control of BAD promoter</td>
<td>This study</td>
</tr>
<tr>
<td>pJM9</td>
<td>Km'; broad host range vector pBBR1-MCS2 containing <em>tesA</em> thioesterase under control of BAD promoter</td>
<td>This study</td>
</tr>
<tr>
<td>pJM20</td>
<td>Km'; broad host range vector pBBR1-MCS2 containing entire MK pathway (<em>tesA, fadB, Mlut_11700 and fadM</em>) under control of BAD promoter</td>
<td>This study</td>
</tr>
</tbody>
</table>

* Numbers with the prefix "H16_A" represent *R. eutropha* locus tags

* For the sake of brevity, "beta ox" is used in text and figures to refer to host strain Re2303.
For the sake of brevity, "beta ox\DeltaphaCAB" is used in text and figures to refer to host strain JM9.
TABLE 2. PHB production in *R. eutropha* strains bearing the methyl ketone pathway under induced and uninduced heterotrophic conditions (30-ml glass tube experiments).

<table>
<thead>
<tr>
<th>Strain</th>
<th>PHB yields (% fructose C consumed)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Induced</td>
</tr>
<tr>
<td>WT (JM6)</td>
<td>2.1</td>
</tr>
<tr>
<td>ΔphaCAB (JM11)</td>
<td>0</td>
</tr>
<tr>
<td>Δbeta ox (JM13)</td>
<td>1.3</td>
</tr>
<tr>
<td>Δbeta ox ΔphaCAB (JM15)</td>
<td>0</td>
</tr>
</tbody>
</table>

*a* All strains carried the MK pathway (Table 1)

*b* Data represent a pooled sample of five independent cultures
TABLE 3. PHB, CO₂, and methyl ketone production in *R. eutropha* strains bearing the methyl ketone pathway under induced and uninduced heterotrophic conditions (RAMOS experiments).

<table>
<thead>
<tr>
<th>Strain&lt;sup&gt;a,b&lt;/sup&gt;</th>
<th>Carbon (% fructose C consumed)</th>
<th>PHB</th>
<th>CO₂</th>
<th>MK</th>
</tr>
</thead>
<tbody>
<tr>
<td>∆beta ox (JM13)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>4 ± 0.73&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0.1</td>
<td>2.2 ± 0.05&lt;sup&gt;e&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>∆beta ox ∆phaCAB (JM15)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.3 ± 0.03&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0.5</td>
<td>3.5 ± 0.66&lt;sup&gt;e&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>∆beta ox&lt;sup&gt;d&lt;/sup&gt;</td>
<td>11 ± 0.77&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>∆beta ox ∆phaCAB&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.1</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> All strains carried the MK pathway (Table 1)
<sup>b</sup> Data represent an average of two independent cultures
<sup>c</sup> Induced conditions
<sup>d</sup> Uninduced conditions
<sup>e</sup> Mean ± SD (SD were not calculated for values near the detection limit)
Relative distribution of methyl ketones (as % of total)

<table>
<thead>
<tr>
<th>Strain</th>
<th>WT</th>
<th>ΔphaCAB</th>
<th>Δbeta ox</th>
<th>ΔphaCAB</th>
</tr>
</thead>
<tbody>
<tr>
<td>JM6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JM11</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JM13</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JM15</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- WT
- ΔphaCAB
- Δbeta ox
- ΔphaCAB

15:1
15:0
13:0