



US010844410B2

(12) **United States Patent**
Pfleger et al.

(10) **Patent No.:** **US 10,844,410 B2**

(45) **Date of Patent:** **Nov. 24, 2020**

(54) **MUTANT THIOESTERASES**

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(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.

(21) Appl. No.: **16/357,635**

(22) Filed: **Mar. 19, 2019**

(65) **Prior Publication Data**

US 2019/0284588 A1 Sep. 19, 2019

Related U.S. Application Data

(60) Provisional application No. 62/644,954, filed on Mar. 19, 2018.

(51) **Int. Cl.**

C12N 9/16 (2006.01)
C12N 15/52 (2006.01)
C12N 15/00 (2006.01)
C12P 7/64 (2006.01)
C12N 15/70 (2006.01)

(52) **U.S. Cl.**

CPC **C12P 7/6409** (2013.01); **C12N 9/16** (2013.01); **C12N 15/52** (2013.01); **C12N 15/70** (2013.01); **C12Y 301/02** (2013.01)

(58) **Field of Classification Search**

CPC C12N 9/16; C12N 15/52; C12N 15/00
See application file for complete search history.

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(57) **ABSTRACT**

Mutant thioesterases having enhanced medium chain substrate activity, polynucleotides encoding and configured to express the mutant thioesterases in a transformed host cell, host cells transformed to contain the polynucleotides, and methods of using same.

24 Claims, 21 Drawing Sheets
(20 of 21 Drawing Sheet(s) Filed in Color)
Specification includes a Sequence Listing.

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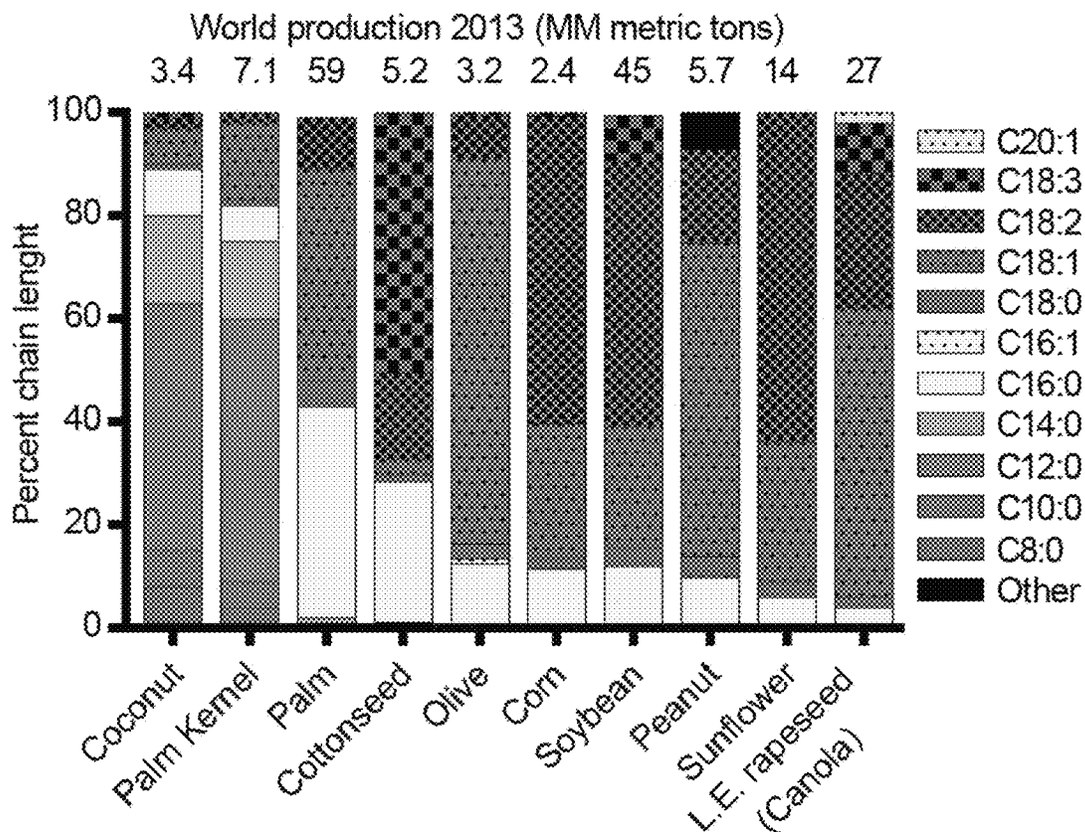


FIG. 1

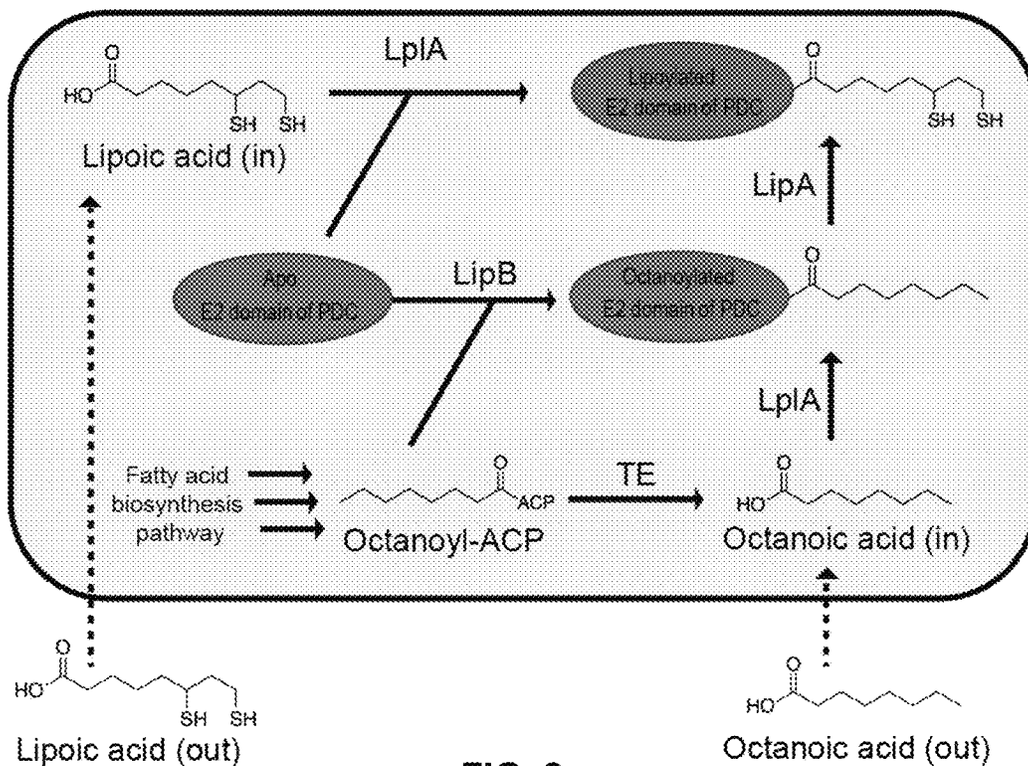


FIG. 2

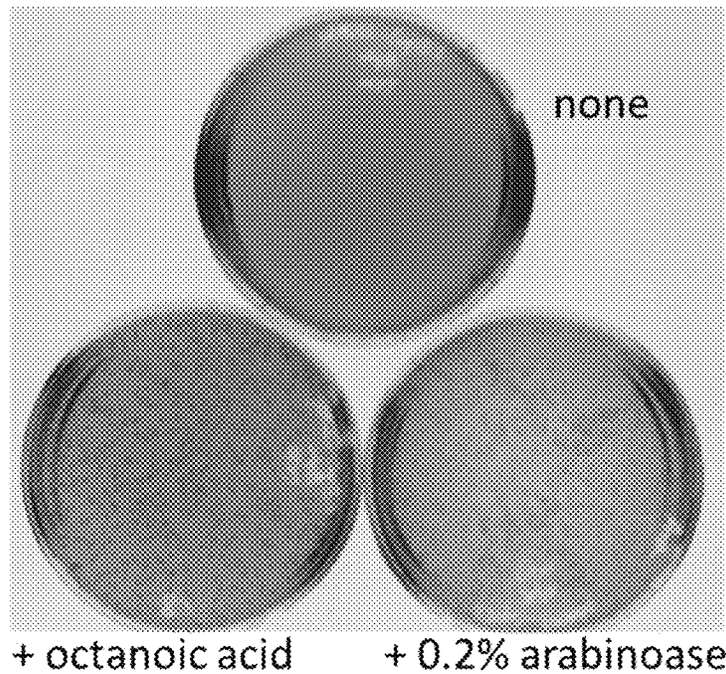
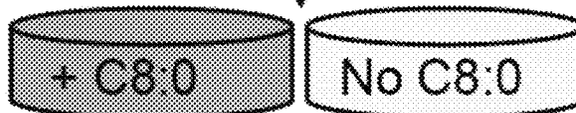


FIG. 4

Strain: *E. coli* K12 MG1655 Δ lipB
 Plasmid: pBTRCK-CpFatB1.2
 Minimal media plates
 0.2% glucose + IPTG (varies)

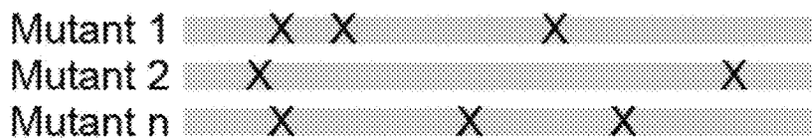


Induction μ M IPTG	Media + C8:0	Media No C8:0
Empty plasmid	2 days	No growth
0	2 days	No growth
10	2 days	4 days
20	2 days	4 days
30	2 days	3 days
50	2 days	3 days

FIG. 5

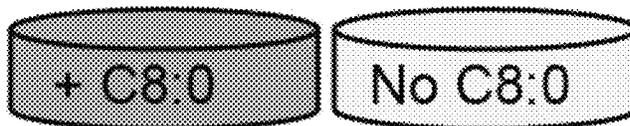
Random mutagenesis

CpFatB1.2 Coding sequence



Minimal media plates

0.2% glucose + 20 μM IPTG



plasmid	Media + 50 μM C8:0	Media No C8:0
Empty plasmid	2 days	No growth
CpFatB1.2	2 days	4 days
CpFatB1.2 variants	2 days	3 days

FIG. 6

Mutant	Residue changes
CpFatB1.2	n/a
CpFatB1.2-M1	A59S, K296R
CpFatB1.2-M2	M29T, T117S, Q163L
CpFatB1.2-M3	A59S, K296R
CpFatB1.2-M4	ΔA_{5-4} (new start codon at M19), N28S, I65M
CpFatB1.2-M5	W17R, T204S
CpFatB1.2-M6	L251M, L265I,
CpFatB1.2-M7	n/a
CpFatB1.2-M8	K15E, S207T
CpFatB1.2-M9	R261S
CpFatB1.2-M10	n/a

FIG. 7A

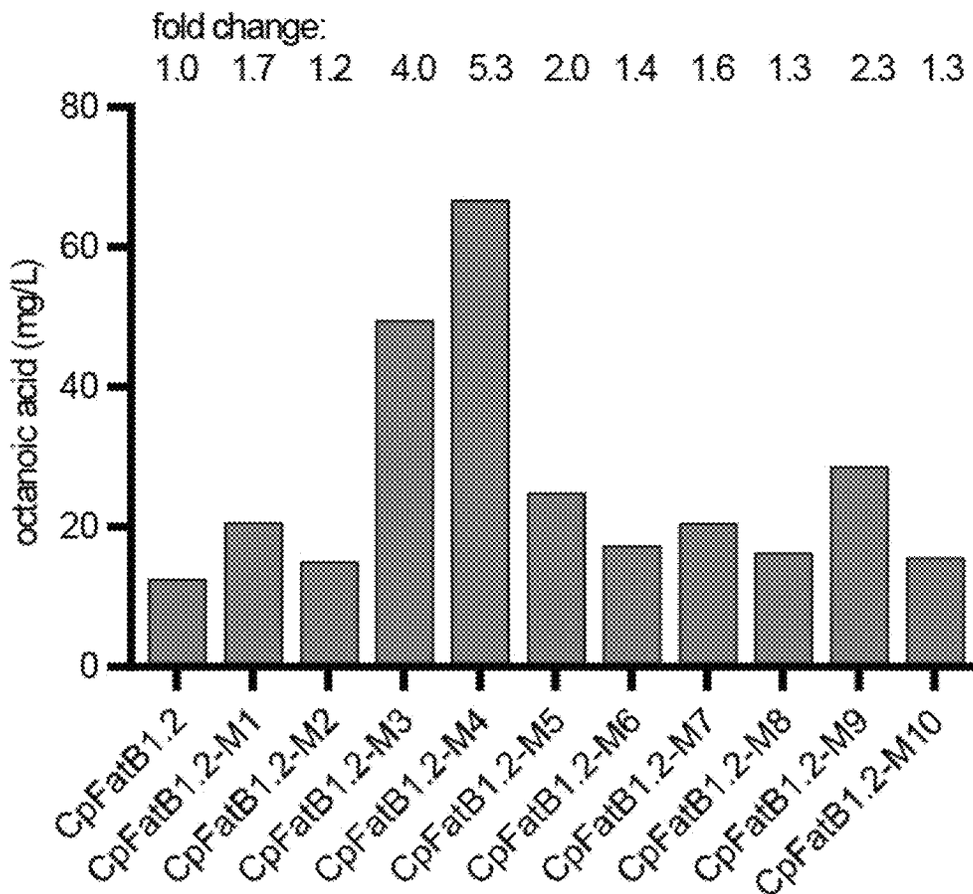


FIG. 7B

mutant	Residue changes
CpFatB1.2	n/a
CpFatB1.2-M3	A59S, K296R
CpFatB1.2-M4	ΔA_{1-18} (new start codon at M19), N28S, I65M
CpFatB1.2-M12	M136V
CpFatB1.2-M16	K296R
CpFatB1.2-M20	W17STOP, D293V
CpFatB1.2-M40	D293V
CpFatB1.2-M47	N146K, D293V, N309D
CpFatB1.2-M66	D293V
CpFatB1.2-M71	M136I
CpFatB1.2-M73	V268I, R279H, D293V
CpFatB1.2-M76	K23E, L86Q
CpFatB1.2-M79	V9M, K15E, E236A

FIG. 8A

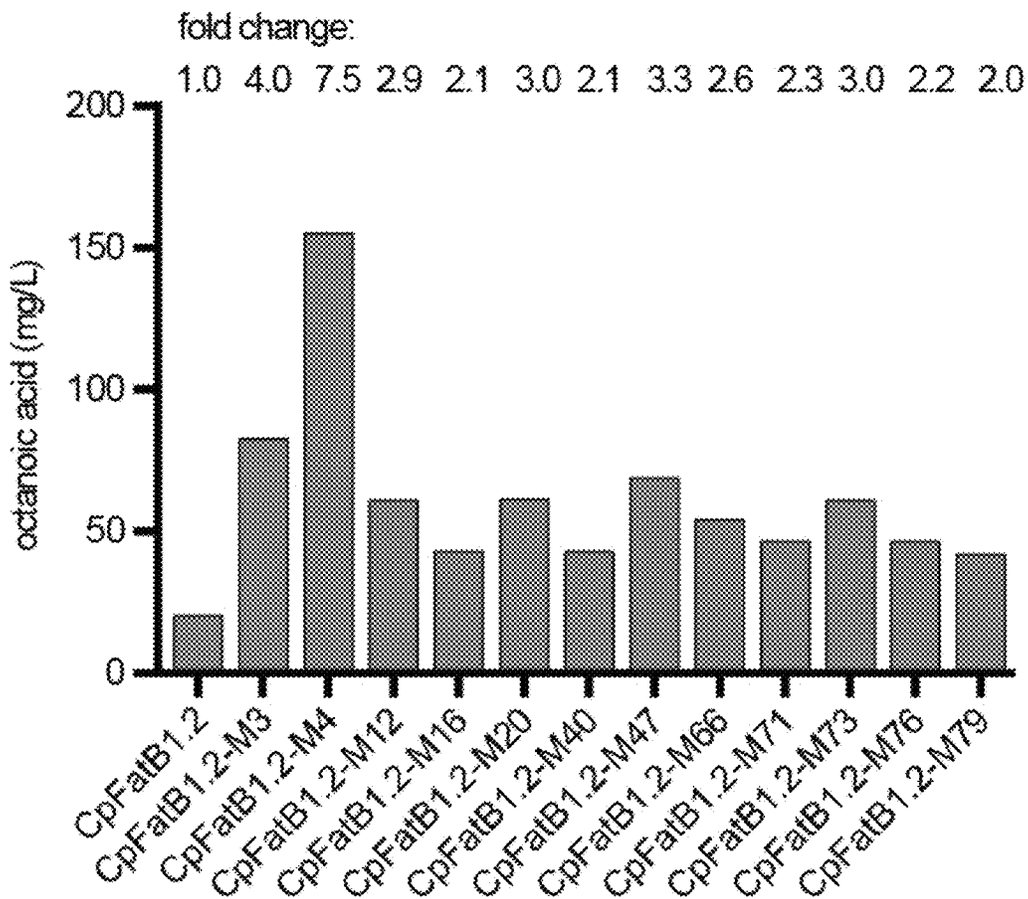


FIG. 8B

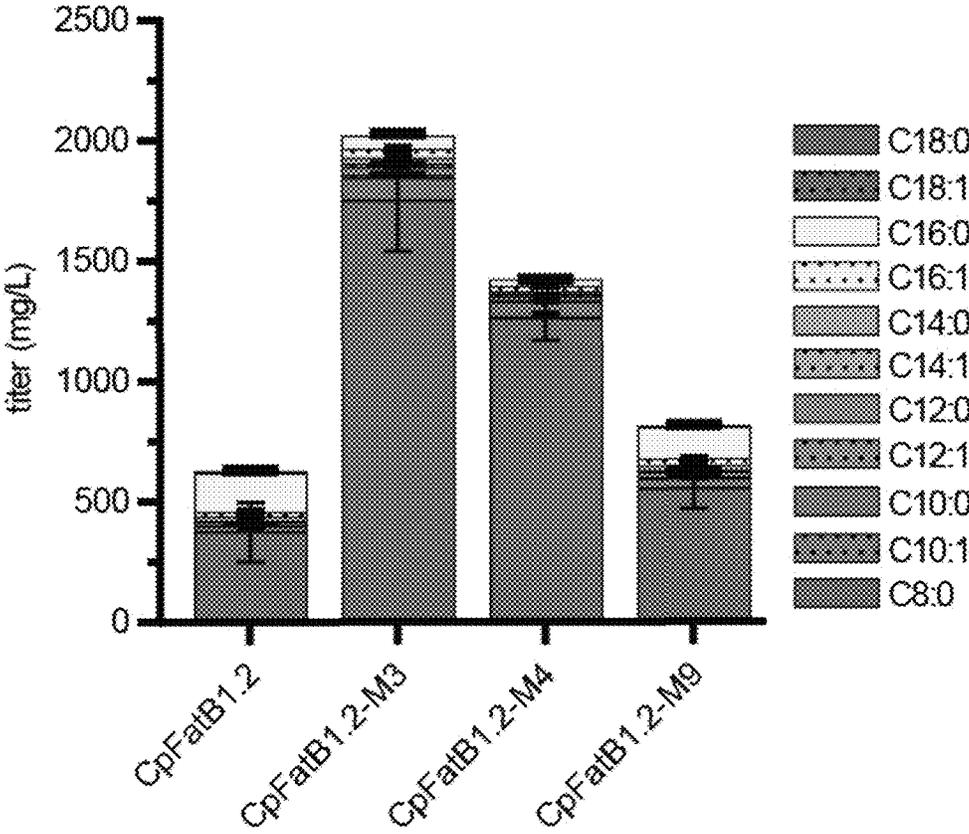


FIG. 9

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      1      10      20      30      40      50      60      70      80
      +-----+-----+-----+-----+-----+-----+-----+-----+
CpFatB1.2 MLLTAITTVFVAPEKRWTMFDRKSKRPNMLMDSFGLERVVQDGLVFRQSF SIRSYEICADRTASIE TVMNHVQETSLNQCKSIGL...
CpFatB1.2-M4 YAAHRDYHRVRCAGKRWTMFDRKSKRPNMLMDSFGLERVVQDGLVFRQSF SIRSYEICADRTASIE TVMNHVQETSLNQCKSIGL...
CpFatB1.2-M4-287 *****MFDRKSKRPNMLMDSFGLERVVQDGLVFRQSF SIRSYEICADRTASIE TVMNHVQETSLNQCKSIGL...
CpFatB1.2-M4-288 *****MLMDSFGLERVVQDGLVFRQSF SIRSYEICADRTASIE TVMNHVQETSLNQCKSIGL...
CpFatB1.2-M4-289 *****MDSFGLERVVQDGLVFRQSF SIRSYEICADRTASIE TVMNHVQETSLNQCKSIGL...
CpFatB1.2-M4-290 *****mETVMNHVQETSLNQCKSIGL...
CpFatB1.2-M4-291 *****MNHVQETSLNQCKSIGL...

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CpFatB1.2 ... (Positions 1-85 of SEQ ID NO:4)
CpFatB1.2-M4 ... (Positions 1-85 of SEQ ID NO:12)
CpFatB1.2-M4-287 ... (Positions 1-67 of SEQ ID NO:14)
CpFatB1.2-M4-288 ... (Positions 11-67 of SEQ ID NO:14)
CpFatB1.2-M4-289 ... (Positions 13-67 of SEQ ID NO:14)
CpFatB1.2-M4-290 ... (Positions 47-67 of SEQ ID NO:14)
CpFatB1.2-M4-291 ... (Positions 51-67 of SEQ ID NO:14)

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FIG. 10A

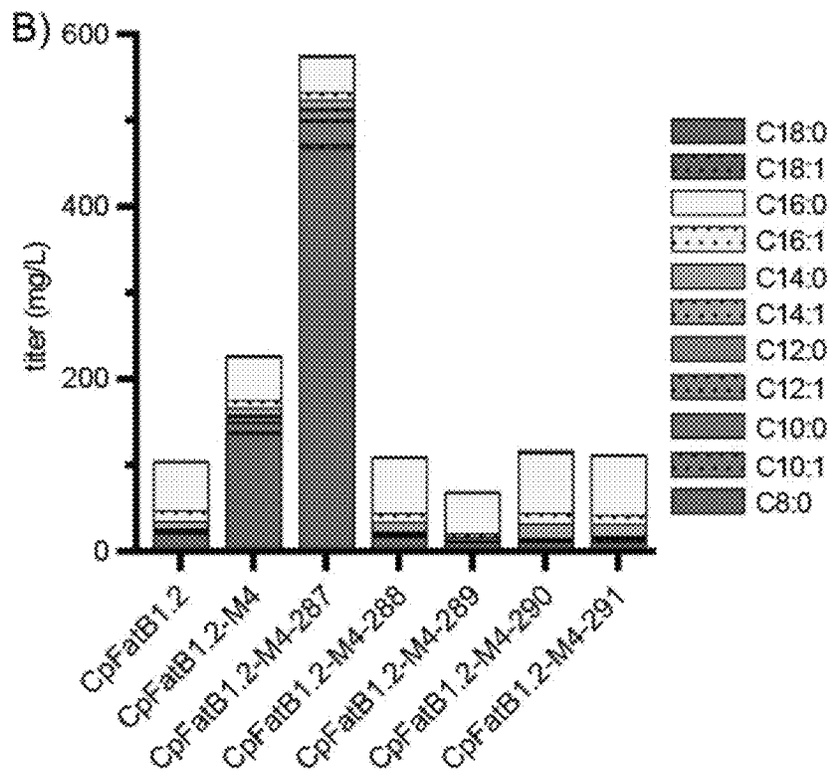


FIG. 10B

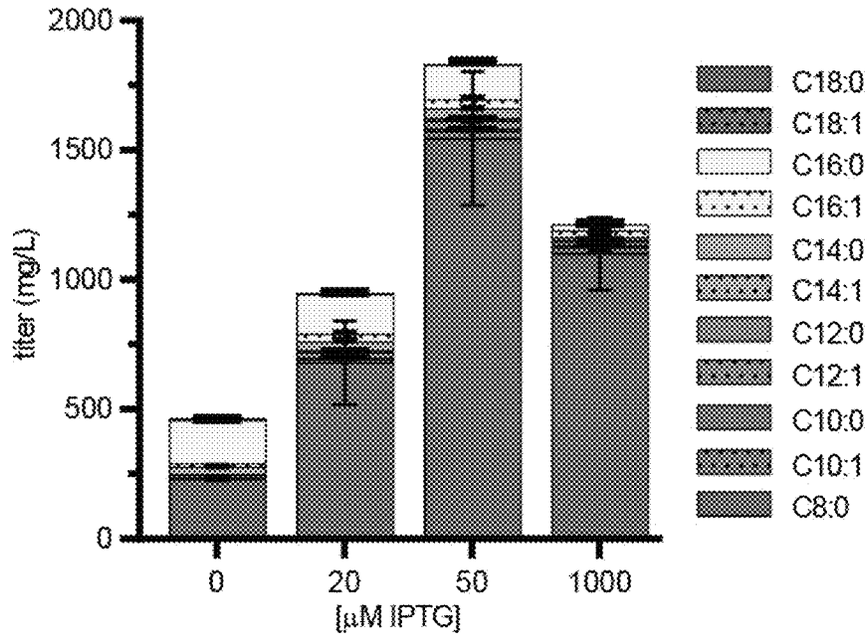


FIG. 11

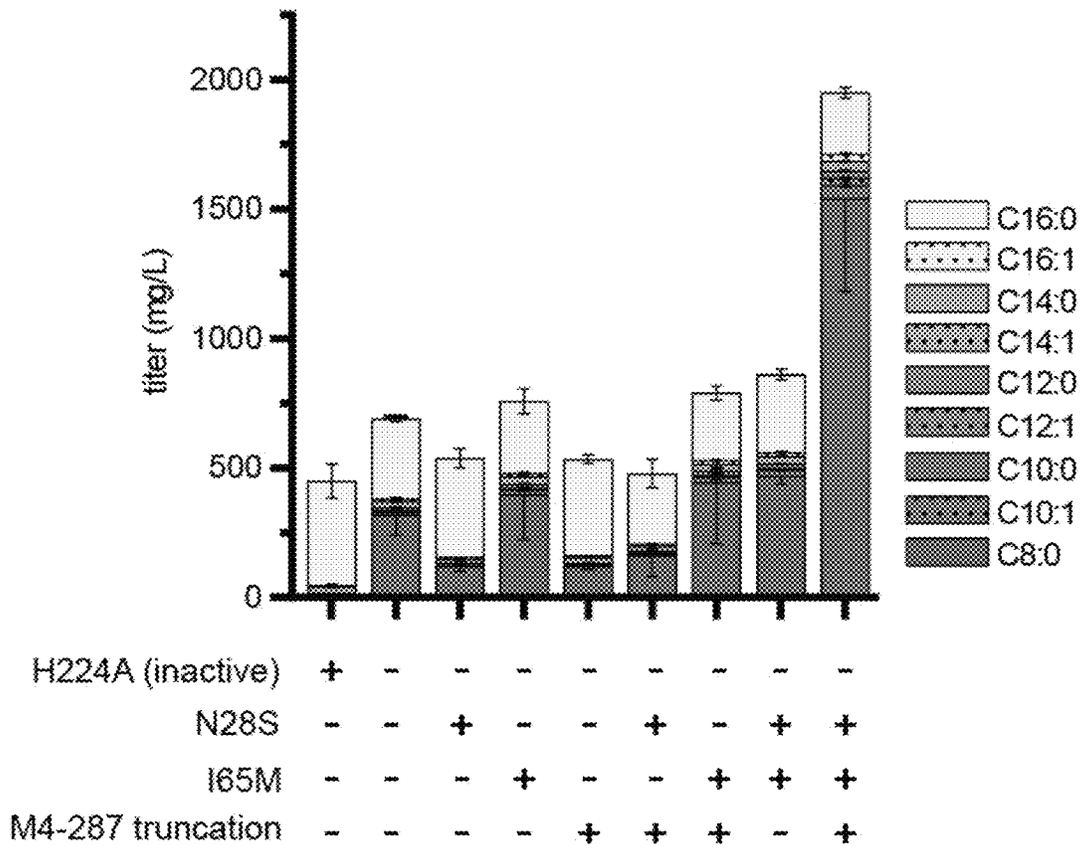


FIG. 12

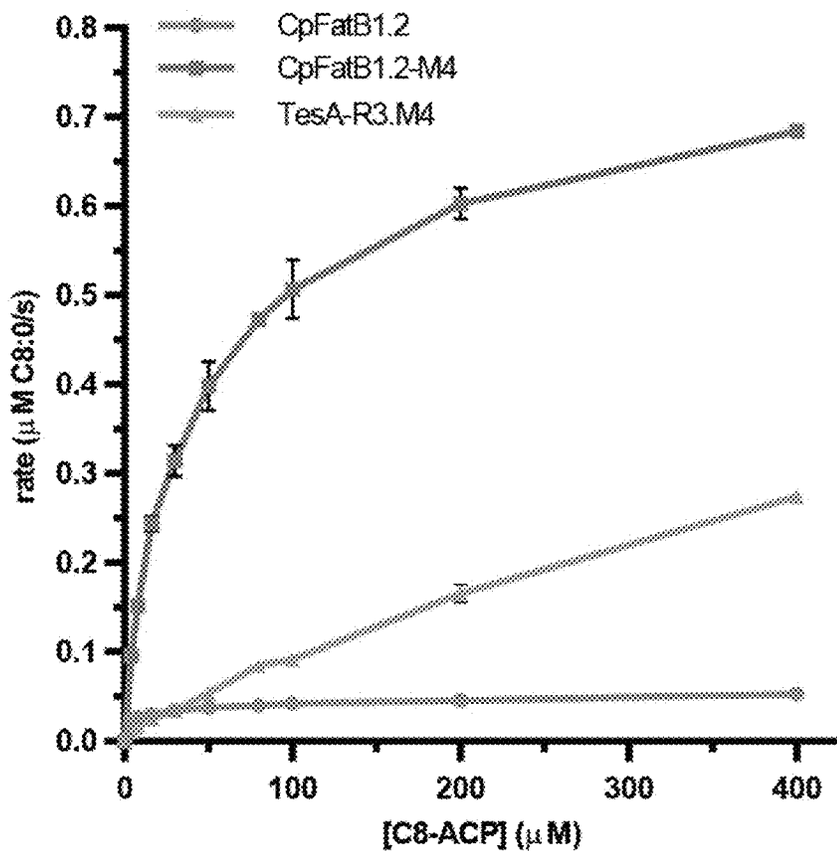


FIG. 13C

Mutant	k_{cat} (min^{-1})	K_m (μM)	k_{cat}/K_m ($\mu\text{M}^{-1} \text{min}^{-1}$)
CpFatB1.2	68.6 ± 2.1	5.92 ± 1.1	11.6 ± 2.2
CpFatB1.2-M4	1076 ± 21	36.8 ± 2.3	29.2 ± 1.9
TesA-R3.M4	1028 ± 66	606 ± 56	1.70 ± 0.2

FIG. 13D

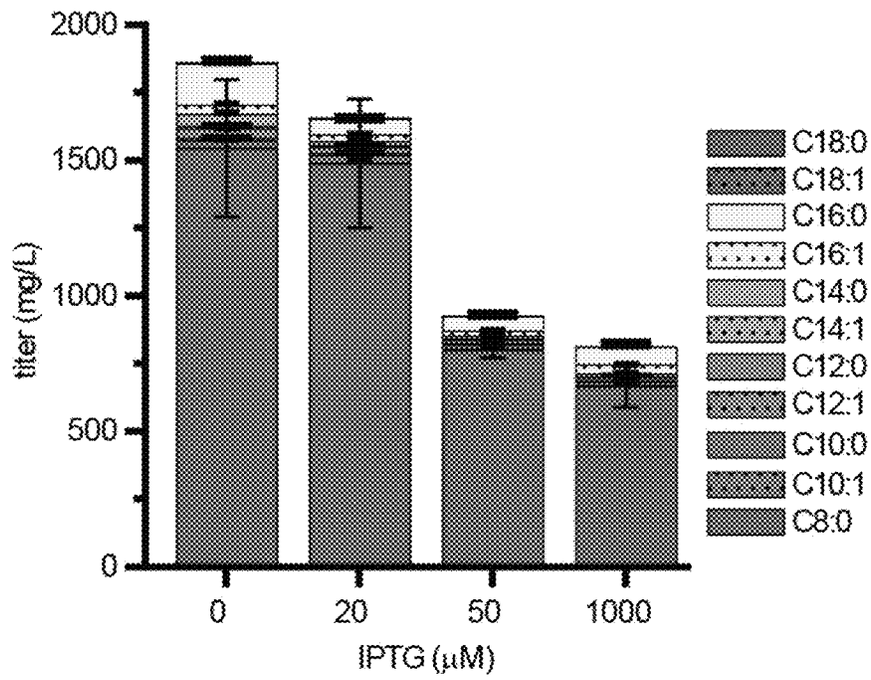


FIG. 14A

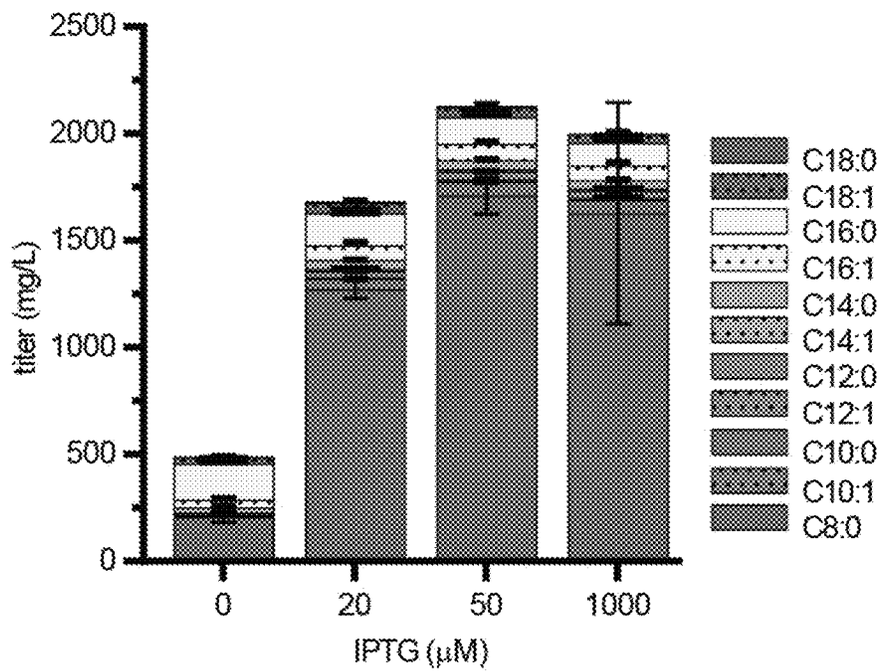


FIG. 14B

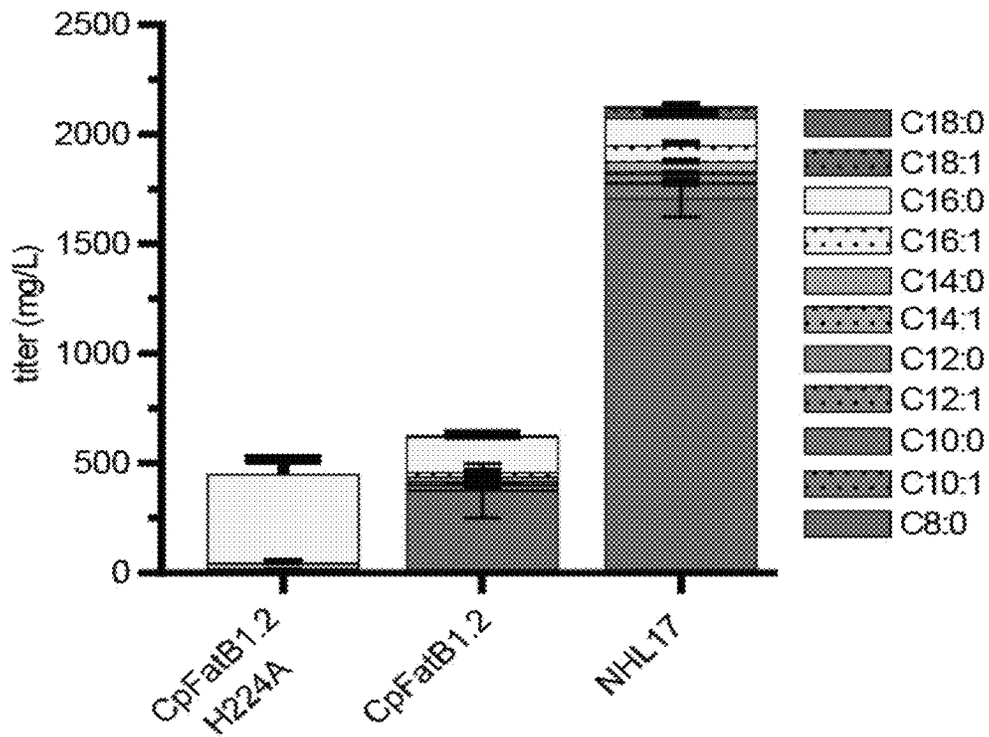


FIG. 15A

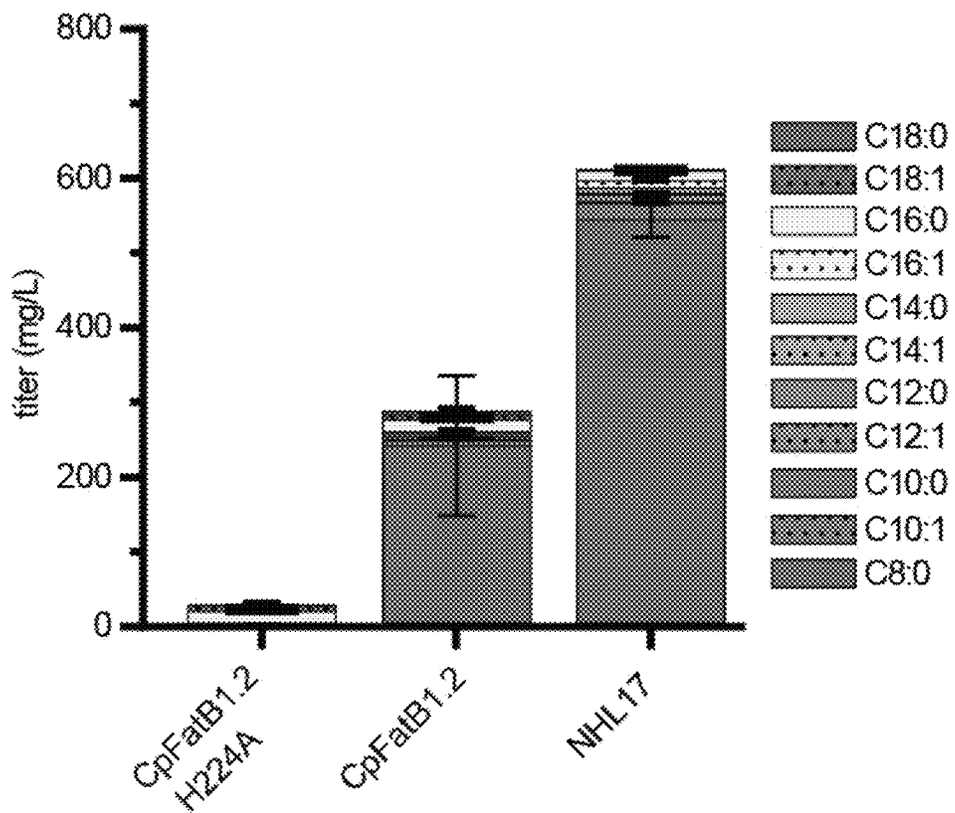


FIG. 15B

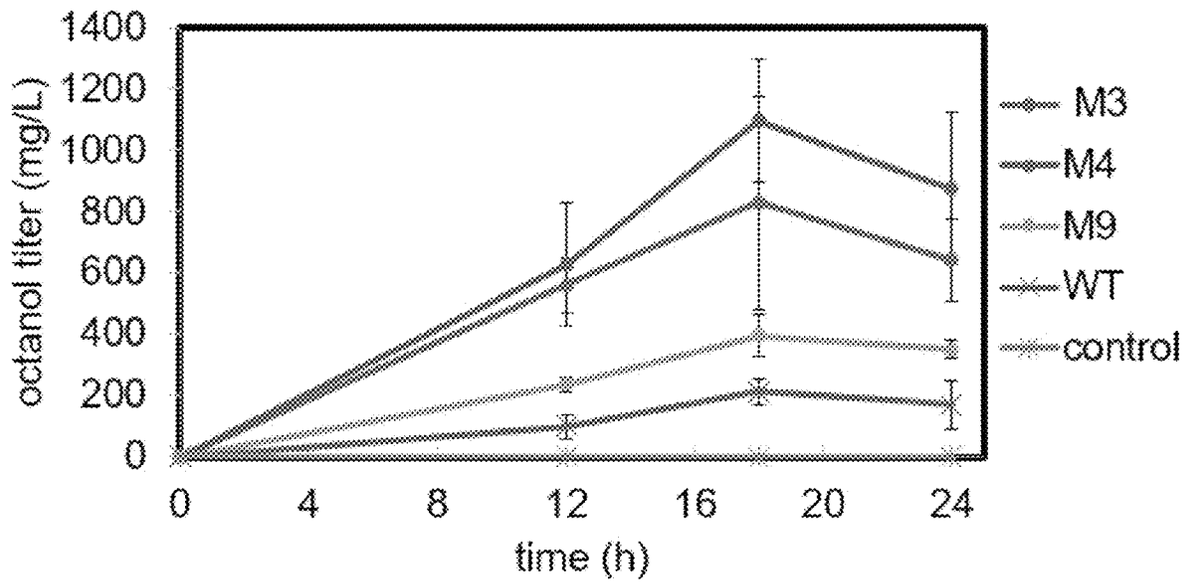


FIG. 16

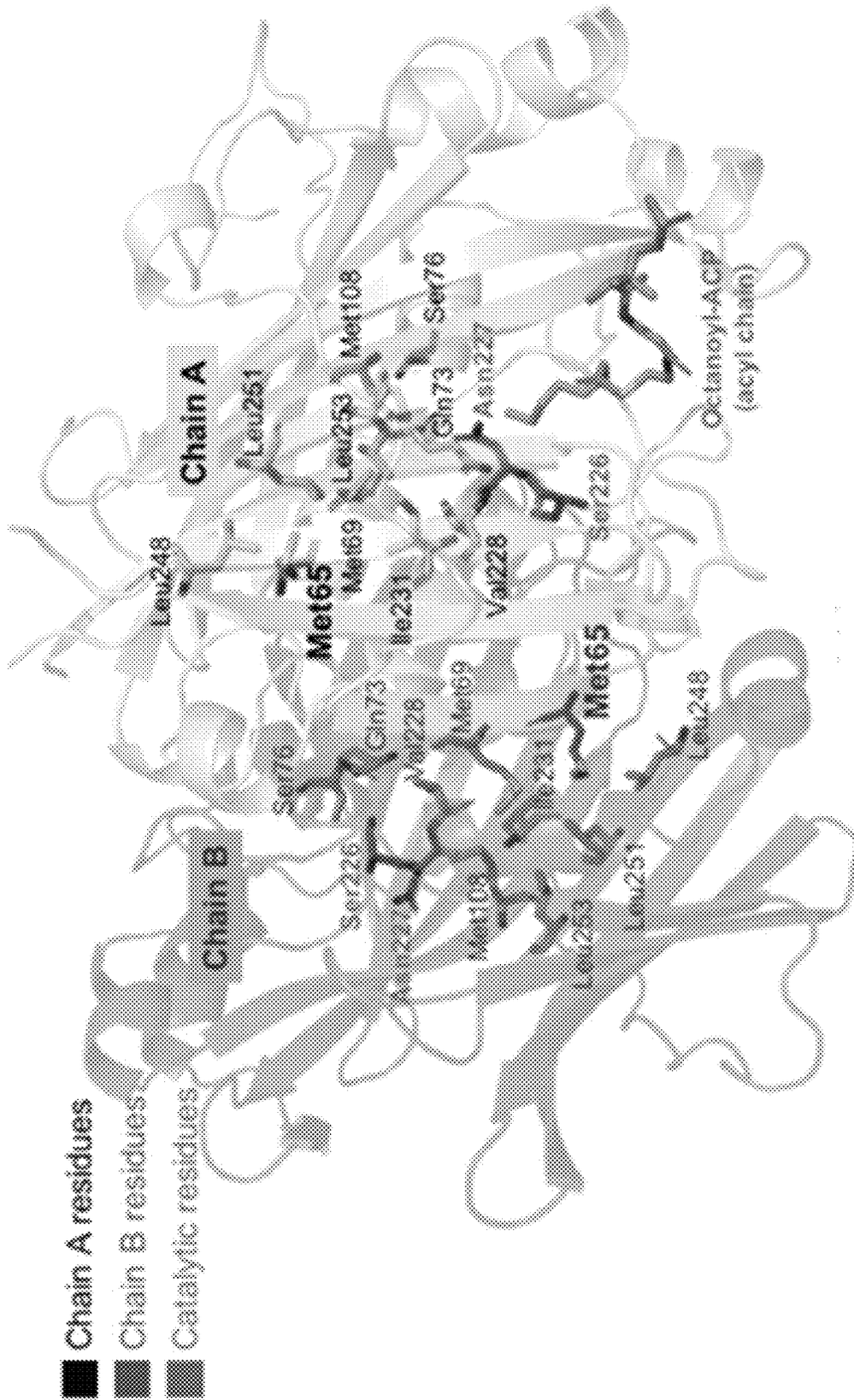


FIG. 17A

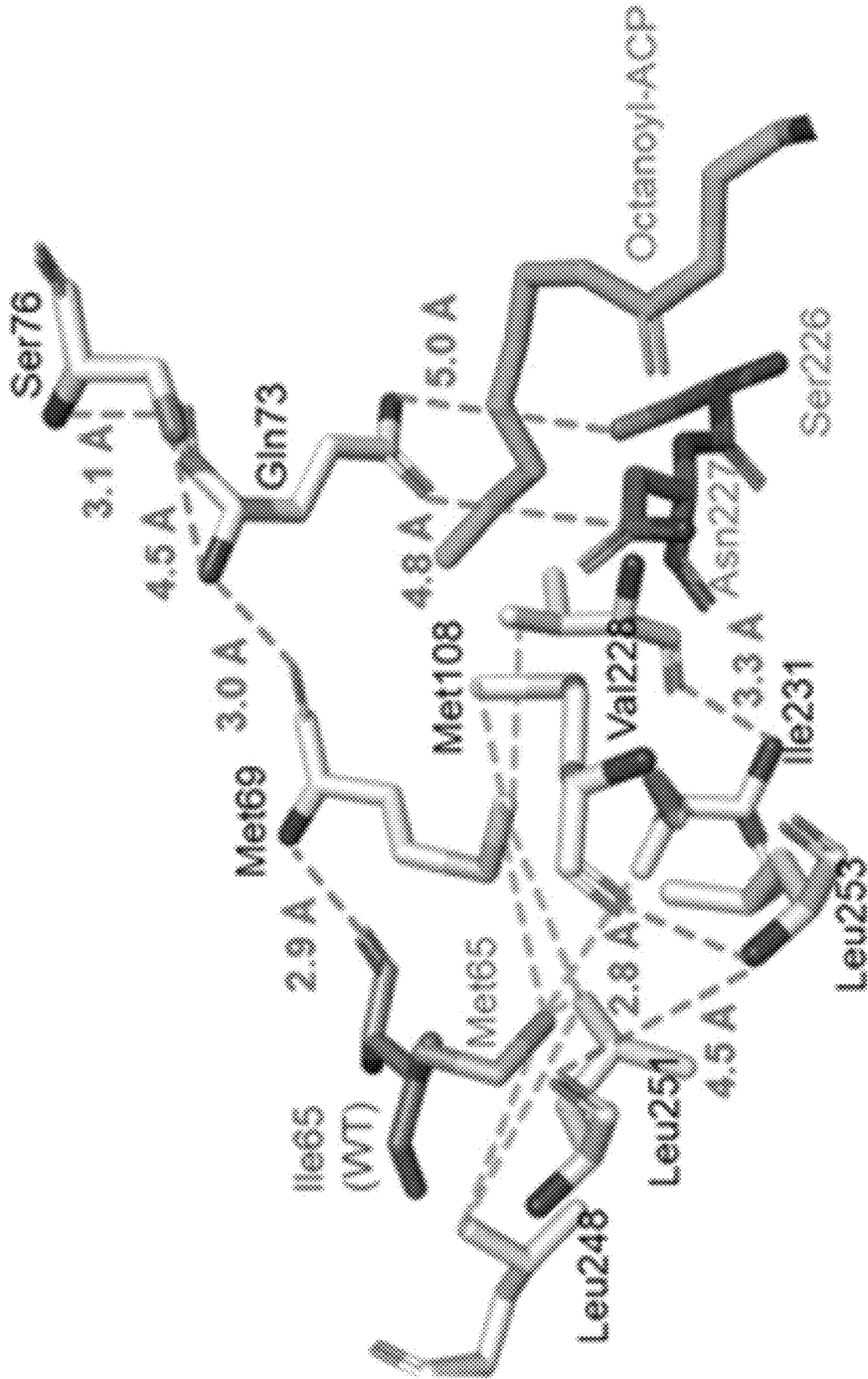
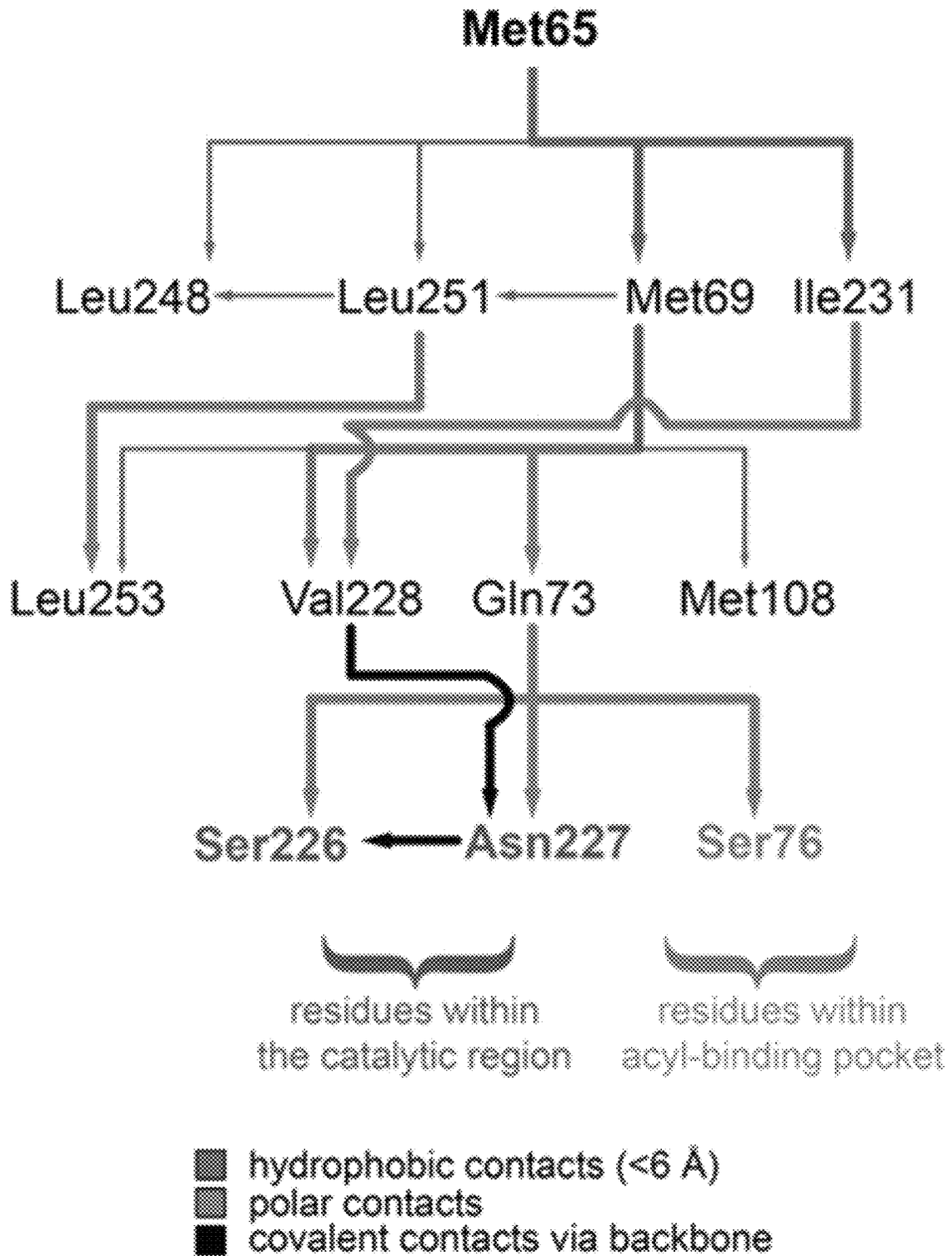


FIG. 17B



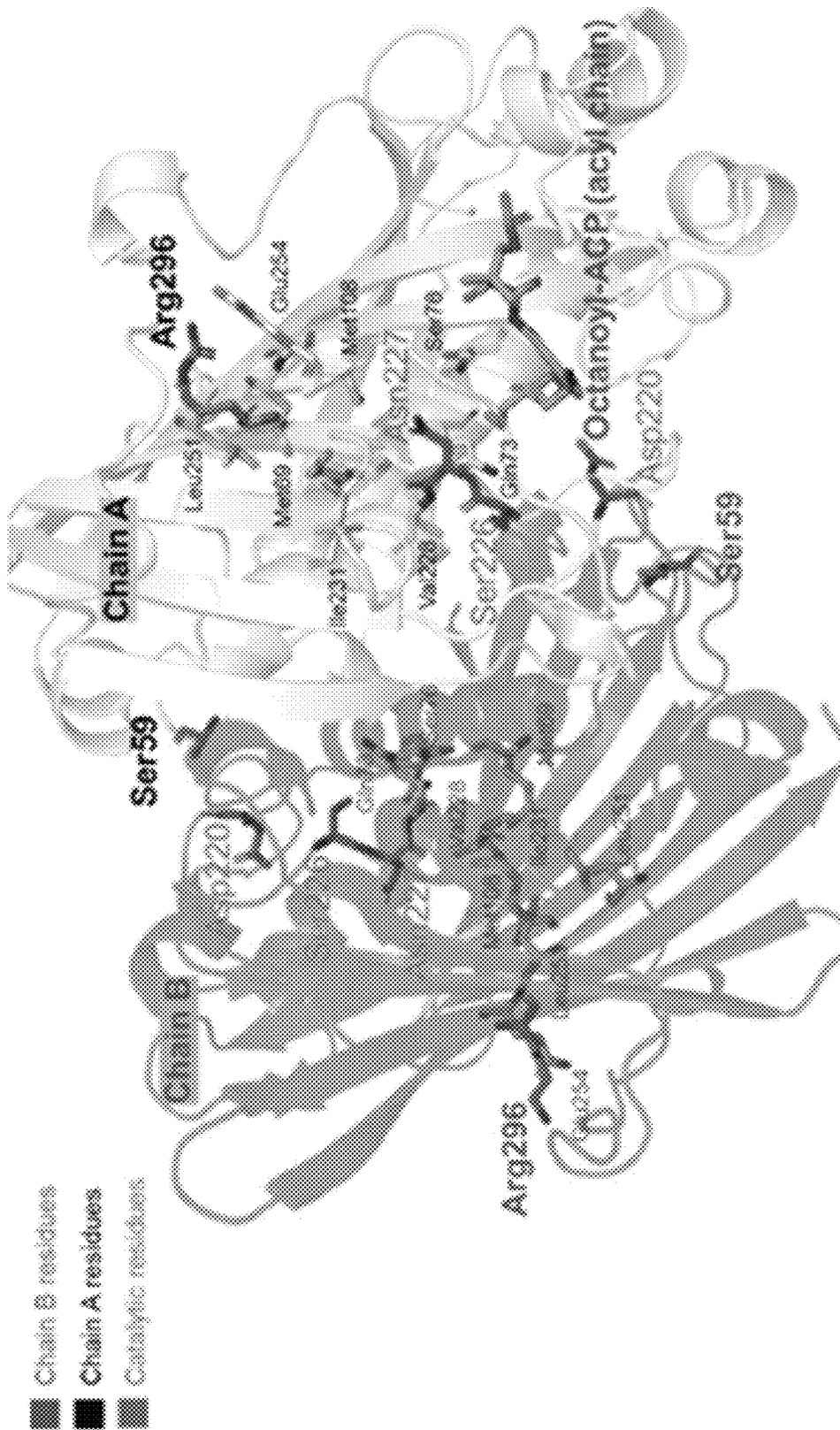


FIG. 18A

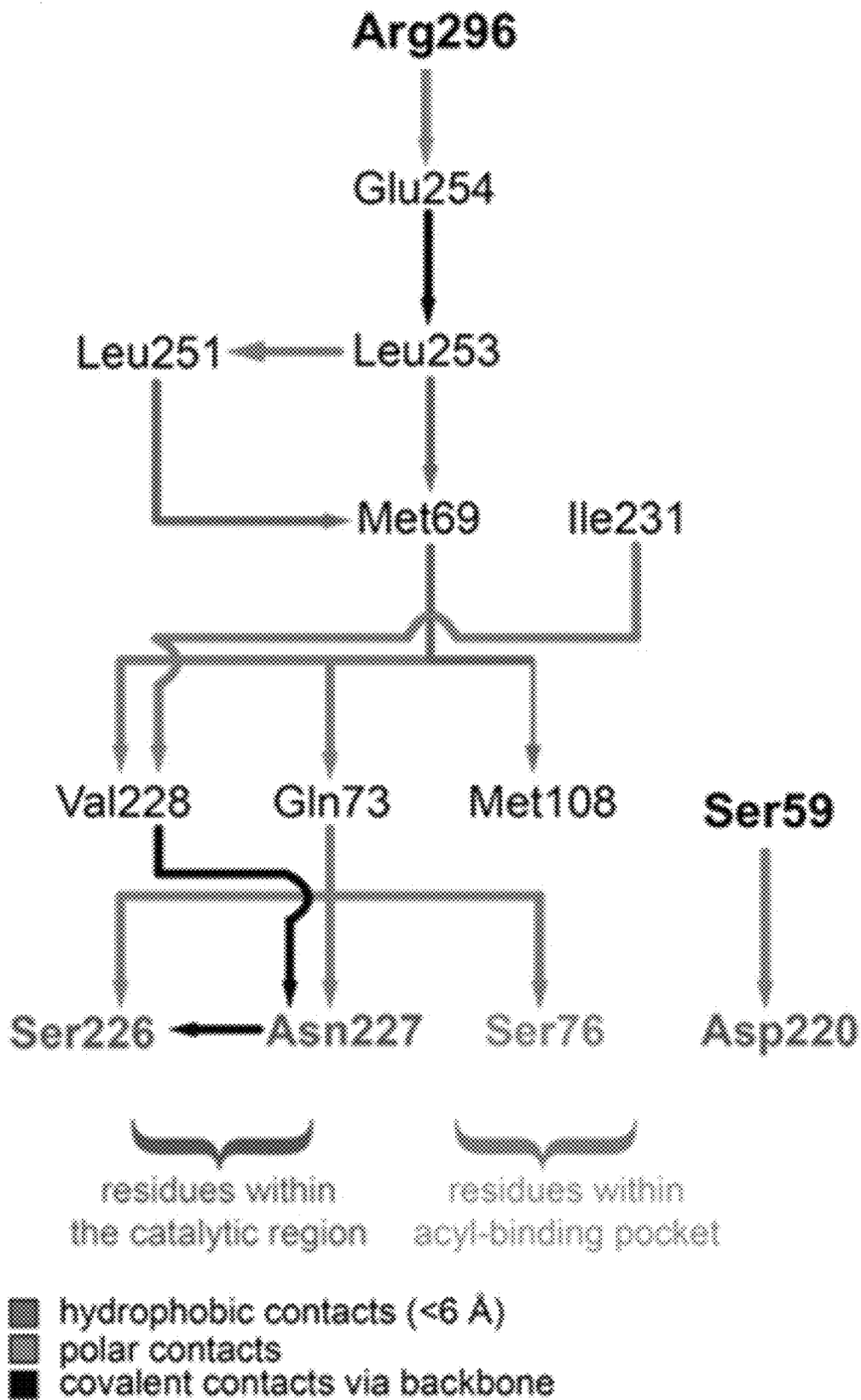


FIG. 18C

MUTANT THIOESTERASES

FEDERAL FUNDING STATEMENT

This invention was made with government support under 5
CBET1149678 awarded by the National Science Founda-
tion. The government has certain rights in the invention.

FIELD OF THE INVENTION

The invention is directed to mutant thioesterases having
enhanced medium chain substrate activity, polynucleotides
encoding and configured to express the mutant thioesterases
in a transformed host cell, host cells transformed to contain
the polynucleotides, and related methods.

BACKGROUND

Free fatty acids (FFAs) are energy-rich molecules capable
of serving as precursors for the production of liquid trans-
portation fuels and high-value oleochemicals. Fuel prop-
erties are dictated by the aliphatic chain length and degree of
saturation of the FFA precursors. Medium-chain (C6-C12)
FFA feedstocks can be converted to hydrocarbons with fuel
properties comparable to gasoline, diesel, or jet fuel (Choi et
al. 2013 and Lee et al. 2008). Fuels derived from microbially
produced FFAs would facilitate reduction of the carbon
footprint and, unlike bioethanol, avoid expensive and labo-
rious infrastructure and engine remodeling (Howard et al.
2013).

Escherichia coli is a popular microbial host for FFA
production because of its established type II fatty acid
biosynthesis (FAB) pathway, short doubling time, and
genetic tractability. The *E. coli* FAB pathway is initiated by
the ATP-dependent carboxylation of acetyl-CoA to malonyl-
CoA. Subsequently, CoA is exchanged with acyl carrier
protein (ACP), the recognition tag of FAB, producing mal-
onyl-ACP. Malonyl-ACP and acetyl-CoA are condensed to
yield acetoacetyl-ACP. The alkyl chain of the β -ketoacyl-
ACP is successively extended by two carbon atoms that
originate from additional malonyl-ACP. This cycle is termi-
nated by the acyl-ACP thioesterase, which hydrolyzes the
thioester bond to generate the FFA and ACP. The specificity
of the acyl-ACP thioesterase controls the terminal aliphatic
chain length and chemical properties of the FFA product
composition. Regulation of the FFA chain length produced
through the FAB pathway has typically been achieved by the
overexpression of the two native *E. coli* thioesterases (TesA
and TesB), or heterologous expression of various plant and
bacterial thioesterases, which exhibit a wide range of sub-
strate specificities (Choi et al. 2013, Steen et al. 2010, Zhang
et al. 2011, Lu et al. 2008, Voelker et al. 1994, Dormann et
al. 1995).

Several of these thioesterases have been evolved to fur-
ther diversify the gamut of attainable FFA compositions.
Despite this diversification, very few thioesterases are spe-
cific towards a unique aliphatic chain length. Of these
studied thioesterases, ‘TesA (a cytosolic TesA that lacks the
N-terminal signal peptide and whose crystal structure has
been elucidated) produces one of the highest FFA titers
(Steen et al. 2010, Choi et al. 2013, Cho et al. 1993, Lo et
al. 2005). In spite of these clear advantages, ‘TesA has broad
substrate specificity that necessitates costly downstream
separation (Steen et al. 2010, Choi et al. 2013).

The carbon chain length of fatty acids is economically
significant because the natural occurrence of certain types of
fatty acids, such as medium-chain fatty acids (carbon chain

of 6 to 12 carbon atoms) in general and C8 carbon chain
length fatty acids in particular, is notably less than long-
chain fatty acids (carbon chain longer than 12 carbon
atoms). There are currently only two notable sources for C8
fatty acids, coconut and palm kernel, and C8 fatty acids are
only a minor fraction of the fatty acids made by these
sources. C8 fatty acids and related C8 compounds are
important in light of their use in cosmetics, plastics, and
other oleochemical products.

10 Tools and methods for producing high amounts C8 fatty
acids and products derived therefrom are needed.

SUMMARY OF THE INVENTION

15 One aspect of the invention is directed to an unnatural,
mutated protein. The protein can comprise an amino acid
sequence at least about 80% identical to positions 28-317 of
SEQ ID NO:4. The amino acid sequence can comprise one
or more of: a residue other than asparagine at a position
corresponding to position 28 of SEQ ID NO:4; a residue
other than methionine at a position corresponding to position
29 of SEQ ID NO:4; a residue other than alanine at a
position corresponding to position 59 of SEQ ID NO:4; a
residue other than isoleucine at a position corresponding to
position 65 of SEQ ID NO:4; a residue other than leucine at
a position corresponding to position 86 of SEQ ID NO:4; a
residue other than threonine at a position corresponding to
position 117 of SEQ ID NO:4; a residue other than methio-
nine at a position corresponding to position 136 of SEQ ID
NO:4; a residue other than asparagine at a position corre-
sponding to position 146 of SEQ ID NO:4; a residue other
than glutamine at a position corresponding to position 163 of
SEQ ID NO:4; a residue other than threonine at a position
corresponding to position 204 of SEQ ID NO:4; a residue
other than serine at a position corresponding to position 207
of SEQ ID NO:4; a residue other than glutamate at a position
corresponding to position 236 of SEQ ID NO:4; a residue
other than leucine at a position corresponding to position
251 of SEQ ID NO:4; a residue other than arginine at a
position corresponding to position 261 of SEQ ID NO:4; a
residue other than leucine at a position corresponding to
position 265 of SEQ ID NO:4; a residue other than valine at
a position corresponding to position 268 of SEQ ID NO:4;
a residue other than arginine at a position corresponding to
position 279 of SEQ ID NO:4; a residue other than aspartate
at a position corresponding to position 293 of SEQ ID NO:4;
a residue other than lysine at a position corresponding to
position 296 of SEQ ID NO:4; and a residue other than
asparagine at a position corresponding to position 309 of
SEQ ID NO:4; and/or the protein lacks an N-terminal
portion having an amino acid sequence identical to positions
1-94 of SEQ ID NO:2. The protein preferably exhibits
thioesterase activity.

The protein in some versions can comprise one or more
of: a serine or a conservative variant of serine at the position
corresponding to position 28 of SEQ ID NO:4; a threonine
or a conservative variant of threonine at the position corre-
sponding to position 29 of SEQ ID NO:4; a serine or a
conservative variant of serine at the position corresponding
to position 59 of SEQ ID NO:4; a methionine or a conser-
vative variant of methionine at the position corresponding to
position 65 of SEQ ID NO:4; a glutamine or a conservative
variant of glutamine at the position corresponding to posi-
tion 86 of SEQ ID NO:4; a serine or a conservative variant
of serine at the position corresponding to position 117 of
SEQ ID NO:4; a valine, an isoleucine, or a conservative
variant of valine or isoleucine at the position corresponding

to position 136 of SEQ ID NO:4; a lysine or a conservative variant of lysine at the position corresponding to position 146 of SEQ ID NO:4; a leucine or a conservative variant of leucine at the position corresponding to position 163 of SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 204 of SEQ ID NO:4; a threonine or a conservative variant of threonine at the position corresponding to position 207 of SEQ ID NO:4; an alanine or a conservative variant of alanine at the position corresponding to position 236 of SEQ ID NO:4; a methionine or a conservative variant of methionine at the position corresponding to position 251 of SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 261 of SEQ ID NO:4; an isoleucine or a conservative variant of isoleucine at the position corresponding to position 265 of SEQ ID NO:4; an isoleucine or a conservative variant of isoleucine at the position corresponding to position 268 of SEQ ID NO:4; a histidine or a conservative variant of histidine at the position corresponding to position 279 of SEQ ID NO:4; a valine or a conservative variant of valine at the position corresponding to position 293 of SEQ ID NO:4; an arginine or a conservative variant of arginine at the position corresponding to position 296 of SEQ ID NO:4; and an aspartate or a conservative variant of aspartate at the position corresponding to position 309 of SEQ ID NO:4.

In some versions, protein further comprises a sequence corresponding to positions 1-27 of SEQ ID NO:4, wherein the amino acid sequence comprises one or more of: a residue other than valine at a position corresponding to position 9 of SEQ ID NO:4; a residue other than lysine at a position corresponding to position 15 of SEQ ID NO:4; a residue other than tryptophan at a position corresponding to position 17 of SEQ ID NO:4; and a residue other than lysine at a position corresponding to position 23 of SEQ ID NO:4. The protein in some versions can comprise one or more of: a methionine or a conservative variant of methionine at the position corresponding to position 9 of SEQ ID NO:4; a glutamate or a conservative variant of glutamate at the position corresponding to position 15 of SEQ ID NO:4; an arginine or a conservative variant of arginine at the position corresponding to position 17 of SEQ ID NO:4; and a glutamate or a conservative variant of glutamate at the position corresponding to position 23 of SEQ ID NO:4.

In some versions, the protein comprises at least one of: a residue other than asparagine at the position corresponding to position 28 of SEQ ID NO:4; and a residue other than isoleucine at the position corresponding to position 65 of SEQ ID NO:4. The protein in some versions can comprise: a residue other than asparagine at the position corresponding to position 28 of SEQ ID NO:4; and a residue other than isoleucine at the position corresponding to position 65 of SEQ ID NO:4. The protein in some versions can comprise at least one of: a serine or a conservative variant of serine at the position corresponding to position 28 of SEQ ID NO:4; and a methionine or a conservative variant of methionine at the position corresponding to position 65 of SEQ ID NO:4. The protein in some versions can lack an N-terminal portion having an amino acid sequence identical to positions 1-18 of SEQ ID NO:4.

In some versions, the protein comprises at least one of: a residue other than alanine at the position corresponding to

position 59 of SEQ ID NO:4; and a residue other than lysine at the position corresponding to position 296 of SEQ ID NO:4. The protein in some versions can comprise: a residue other than alanine at the position corresponding to position 59 of SEQ ID NO:4; and a residue other than lysine at the position corresponding to position 296 of SEQ ID NO:4. The protein in some versions can comprise at least one of: a serine or a conservative variant of serine at the position corresponding to position 59 of SEQ ID NO:4; and an arginine or a conservative variant of arginine at the position corresponding to position 296 of SEQ ID NO:4. The protein in some versions can comprise: a serine or a conservative variant of serine at the position corresponding to position 59 of SEQ ID NO:4; and an arginine or a conservative variant of arginine at the position corresponding to position 296 of SEQ ID NO:4.

In some versions, the protein comprises a residue other than aspartate at the position corresponding to position 293 of SEQ ID NO:4. The protein in some versions can comprise a valine or a conservative variant of valine at the position corresponding to position 293 of SEQ ID NO:4.

In some versions, the protein comprises a residue other than lysine at the position corresponding to position 296 of SEQ ID NO:4. The protein in some versions can comprise an arginine or a conservative variant of arginine at the position corresponding to position 296 of SEQ ID NO:4.

Another aspect of the invention is directed to polynucleotides encoding the proteins of the invention.

Another aspect of the invention is directed to host cells comprising the polynucleotides of the invention.

Another aspect of the invention is directed to methods of producing a fatty acid derivative. The methods can comprise cultivating the host cell of the invention under conditions that permit production of the fatty acid derivative. In some versions, the fatty acid derivative comprises a C8 fatty acid derivative. In some versions, the C8 fatty acid derivative comprises octanoic acid.

The objects and advantages of the invention will appear more fully from the following detailed description of the preferred embodiment of the invention made in conjunction with the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

The patent or application file contains at least one drawing executed in color. Copies of this patent or patent application publication with color drawing(s) will be provided by the Office upon request and payment of the necessary fee.

FIG. 1. Plant oils and fats as a percentage of their chain-length composition. Coconut plant oil and palm kernel oil are main sources of medium-chain fatty acids (MCFA) (C8-C12).

Octanoic acid is a minor component in both of these sources.

FIG. 2. Synthesis of lipoyl group of pyruvate dehydrogenase. Pyruvate dehydrogenase complex (PDC) requires a lipoylated E2 domain (green) for *E. coli* to grow in aerobic conditions. This can be achieved via direct incorporation of lipoyl acid or octanoic acid (C8:0) from the media through LplA and LipA (only for C8:0) or the LipB-mediated octanoylation of Apo-E2 domain (red) followed by insertion of sulfur via LipA. In the absence of lipoyl acid and octanoic acid from the media, a Δ lipB strain can be rescued by the action of an octanyl-ACP thioesterase, which provides with intracellular C8:0 that can be used to lipoylate E2 domain of PDC.

FIG. 3A. CpFatB1 truncation sites with arrows indicating the position M1 for each new truncation.

FIG. 3B. Free fatty acid production from CpFatB1 truncations shown in FIG. 3A. Each truncation was cloned into p\DeltafadD) *E. coli* strain and grown in LB 0.4% glycerol and 1 mM IPTG. CpFatB1.2 was chosen as a baseline thioesterase for mutagenesis and selection in Δ lipB strain.

FIG. 4. Rescue of *E. coli* lipB null mutant expressing CpFatB1.2. *E. coli* lipB null mutant containing pBAD33-CpFatB1.2 can grow on MOPS minimal media-agarose containing 0.2% arabinose compared without the addition of the pBAD33-CpFatB1.2. The positive control was the plate with the addition of 50 μ M octanoic acid. This shows that it is possible to use a thioesterase as an alternative to LipB in minimal media.

FIG. 5. Selection of optimal conditions for lipB screen with CpFatB1.2. A low copy plasmid (pBTRCK-CpFatB1.2) containing CpFatB1.2 under an IPTG-inducible promoter was transformed into *E. coli* Δ lipB strain and plated into MOPS minimal media plates containing different concentrations of IPTG. Growth in these plates was monitored for 4 days. Plates with 20 μ M IPTG was selected for selection of mutants because *E. coli* Δ lipB carrying CpFatB1.2 took an extra day to grow (4 days), giving room for improved mutants to be selected the day before (3 days).

FIG. 6. Screening for improved mutants of CpFatB1.2. A library of pBTRCK-CpFatB1.2 mutants was created by random mutagenesis of the coding sequence CpFatB1.2 at a low mutation rate. This library was transformed into *E. coli* Δ lipB strain and plated into MOPS minimal media plates containing 20 μ M IPTG. 90 putative mutant colonies were selected after 3 days for further characterization.

FIG. 7A. Mutations in CpFatB1.2 mutants 1-10. Sequence identity of CpFatB1.2 mutants 1-10 shows a mixture of CpFatB1.2 and CpFatB1.2 variants. The top three improved mutants (M3, M4 and M9) (see FIG. 7B) contained variations from CpFatB1.2. CpFatB1.2-M4 contains an early stop codon.

FIG. 7B. Octanoic acid production from CpFatB1.2 mutants 1-10. Plasmids from 10 of the 90 putative mutants were isolated and transformed into the RL08ara (Δ fadD) strain and grown in LB supplemented with 0.4% glycerol and 20 μ M IPTG. Octanoic acid production for first ten mutants relative to baseline CpFatB1.2 identified two mutants with several fold improvement over CpFatB1.2.

FIG. 8A. Mutations in additional CpFatB1.2.

FIG. 8B. Octanoic acid production from the additional CpFatB1.2 mutants presented in FIG. 8A.

FIG. 9. Free fatty acid production from mutants CpFatB1.2-M3, CpFatB1.2-M4, and CpFatB1.2-M9 under high expression conditions. Mutants CpFatB1.2-M3, CpFatB1.2-M4, and CpFatB1.2-M9 were subcloned into high copy plasmid p\DeltafadD) strain and grown in MOPS media (Kim et al. 2015) enriched with tryptone and yeast extract and 1 mM IPTG. Data shows several fold improvements over CpFatB1.2 in high expression conditions, consistent with the data shown in FIG. 7B for low expression conditions.

FIG. 10A. Generation of CpFatB1.2-M4 truncation variants CpFatB1.2-M4-287, CpFatB1.2-M4-288, CpFatB1.2-M4-289, CpFatB1.2-M4-290, and CpFatB1.2-M4-291. Characterization of CpFatB1.2-M4. Alignment of CpFatB1.2 with CpFatB1.2-M4 (Δ A54, N28S, I65M) shows

a frame shift which that rose from nucleotide A54 deletion, producing a stop codon. Given that the enzyme is active, we proposed that it was being translated from a different in-frame methionine from a non-specific ribosome binding site (RBS). We identified five methionines (highlighted in green) in frame with CpFatB1.2 and cloned in-frame CpFatB1.2-M4 variants (CpFatB1.2-M4-287, CpFatB1.2-M4-288, CpFatB1.2-M4-289, CpFatB1.2-M4-290, and CpFatB1.2-M4-291) with the methionines as start codons.

FIG. 10B. Free fatty acid production from the CpFatB1.2-M4 truncation variants. Isolated plasmids of each variant in FIG. 10A were transformed into the RL08ara (Δ fadD) strain and grown in LB 0.4% glycerol and 1 mM IPTG. Only variant CpFatB1.2-M4-287 showed high level of octanoic acid production characteristic of its parent CpFatB1.2-M4 sequence. Moreover, increased production observed in variant CpFatB1.2-M4-287 over CpFatB1.2-M4 suggests that the hypothesis of CpFatB1.2-M4 being translated from a non-specific RBS was correct. This places CpFatB1.2-M4-287 as having a ~20-fold increased production under low expression conditions tested.

FIG. 11. Free fatty acid production from CpFatB1.2-M4-287 at different expression levels. CpFatB1.2-M4-287 was cloned into high copy plasmid p\DeltafadD) strain, grown in MOPS media (Kim et al. 2015) enriched with tryptone and yeast extract, and induced under different concentrations of IPTG. Under these conditions, 50 μ M IPTG gave maximum octanoic acid production. At 1 mM IPTG, there was a growth defect that can be seen in the drastic decrease in C16 species.

FIG. 12. Analysis of CpFatB1.2-M4-287 mutations. We made every single and double mutant combination contained in CpFatB1.2-M4-287 (N28S, I65M, 287-truncation) in the CpFatB1.2 base thioesterase and studied FFA production under the same conditions described for FIG. 10B. H224A is a catalytically inactive version of CpFatB1.2, which was used as a negative control. I65M was observed to have a minor contribution to activity as a single mutant. No significant additive effects were observed with double mutants. Only the triple mutant was able to outperform CpFatB1.2 drastically.

FIG. 13A. Scheme for octanoyl-ACP assay. Holo-ACP generated by thioesterase activity reacts with DTNB forming TNB compound with absorbance at 412 nm.

FIG. 13B. Verification of octanoyl-ACP synthesis. Shown are HPLC traces of apo-ACP and holo-ACP mixture as purified from *E. coli* (top); holo-ACP after incubation of *E. coli* mixture with Sfp (center); and octanoyl-ACP produced after incubation of holo-ACP with AasS (bottom).

FIG. 13C. Activity of CpFatB1.2, CpFatB1.2-M4 and TesA-R3.M4 as a function of octanoyl-ACP concentration.

FIG. 13D. Kinetic parameters for each enzyme in FIG. 13C based on non-linear least squares fit of the curve.

FIG. 14A. Free fatty acid production from CpFatB1.2-M4-287 expressed from a plasmid. CpFatB1.2-M4-287 was cloned into a pBTRCK plasmid with a strong RBS, transformed into the RL08ara (Δ fadD) strain, grown in MOPS media (Kim et al. 2015) enriched with tryptone and yeast extract, and induced with different concentrations of IPTG. Maximum production was achieved without induction with IPTG.

FIG. 14B. Free fatty acid production from CpFatB1.2-M4-287 expressed from the chromosome. Strain NHL17 (*Escherichia coli* K12 MG1655 Δ araBAD Δ fadD::trc-CpFatB1.2-M4-287) was created to contain a single copy of a gene expressing CpFatB1.2-M4-287 and was tested for free fatty acid production under the same conditions described

for FIG. 14A. NHL17, with a single copy of CpFatB1.2-M4-287 in the chromosome, was capable of making titers higher than that achieved from plasmids.

FIGS. 15A and 15B. Summary of highest titer achieved with CpFatB1.2 in high copy plasmid at maximum induction (CpFatB1.2) versus CpFatB1.2-M4-287 from a single copy in the chromosome (NHL17). FIG. 15A shows results from culturing in MOPS media enriched with tryptone and yeast extract. FIG. 15B shows results from culturing in MOPS minimal media with phosphate limitation.

FIG. 16. Octanol production from *E. coli* strains co-expressing CpFatB1.2 variants, an acyl-CoA synthetase, and a hybrid acyl-CoA reductase/aldehyde reductase.

FIGS. 17A-17C. Effect of I65M mutation on CpFatB1.2-M4-287 mutant activity. FIG. 17A shows a structural overview of CpFatB1.2-M4-287 mutant. The mutated residues are shown in bold. The catalytic residues are marked in pink. The docked configuration of the substrate (blue) has been shown in one of the two (chain A) identical binding pockets present in two chains. FIG. 17B shows key residues that connect residue 65 to the acyl binding pocket. All the hydrophobic (pink dashes) and polar contacts (green dashes) have been overlaid on the CpFatB1.2-M4-287 mutant model. The polar distances have been labeled in green. FIG. 17C shows a contact map showing the trace of hydrophobic, polar, or covalent contacts from residue 65 to the catalytic region. The map terminates upon reaching one or more residues from the catalytic region (red) or the acyl binding-pocket (gray). Edge thickness correlates to importance of interactions.

FIGS. 18A-18C. Effect of A59S and K296R mutations on CpFatB1.2-M3 mutant activity. FIG. 18A shows a structural overview of CpFatB1.2-M3 mutant. The mutated residues are shown in bold. The catalytic residues are marked in pink. The docked configuration of the substrate (blue) has been shown in one of the two (chain A) identical binding pockets present in two chains. FIG. 18B shows key residues that connect residues 59 and 296 to the acyl binding pocket. All the hydrophobic (pink dashes) and polar contacts (green dashes) have been overlaid on the CpFatB1.2-M3 mutant model. The polar distances have been labeled in green. The enhanced salt bridge formation between R296 and E254 has been shown inset. FIG. 18C shows a contact map showing the trace of hydrophobic (pink), polar (green), or covalent contacts (black) from residues 59 and 296 to the catalytic region. Edge thickness correlates to importance of interactions.

DETAILED DESCRIPTION OF THE INVENTION

One aspect of the invention is directed to mutant thioesterases. In some versions, the mutant thioesterases have enhanced activity with C8 substrates.

The mutant thioesterases may comprise an amino acid sequence at least about 30%, least about 40%, at least about 50%, at least about 60%, at least about 70%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 97%, or at least about 99%, identical to positions 41-317, positions 40-317, positions 39-317, positions 38-317, positions 37-317, positions 36-317, positions 35-317, positions 34-317, positions 33-317, positions 32-317, positions 31-317, positions 30-317, positions 29-317, positions 28-317, positions 27-317, positions 26-317, positions 25-317, positions 24-317, positions 23-317, positions 22-317, positions 21-317, positions 20-317, positions

19-317, positions 18-317, positions 17-317, positions 16-317, positions 15-317, positions 14-317, positions 13-317, positions 12-317, positions 11-317, positions 10-317, positions 9-317, positions 8-317, positions 7-317, positions 6-317, positions 5-317, positions 4-317, positions 3-317, positions 2-317, or positions 1-317 of SEQ ID NO:4.

The mutant thioesterases may have one or more substitutions at positions corresponding to particular positions of SEQ ID NO:4. For example, the mutant thioesterases may comprise one or more of: a residue other than valine at a position corresponding to position 9 of SEQ ID NO:4; a residue other than lysine at a position corresponding to position 15 of SEQ ID NO:4; a residue other than tryptophan at a position corresponding to position 17 of SEQ ID NO:4; a residue other than lysine at a position corresponding to position 23 of SEQ ID NO:4; a residue other than asparagine at a position corresponding to position 28 of SEQ ID NO:4; a residue other than methionine at a position corresponding to position 29 of SEQ ID NO:4; a residue other than alanine at a position corresponding to position 59 of SEQ ID NO:4; a residue other than isoleucine at a position corresponding to position 65 of SEQ ID NO:4; a residue other than leucine at a position corresponding to position 86 of SEQ ID NO:4; a residue other than threonine at a position corresponding to position 117 of SEQ ID NO:4; a residue other than methionine at a position corresponding to position 136 of SEQ ID NO:4; a residue other than asparagine at a position corresponding to position 146 of SEQ ID NO:4; a residue other than glutamine at a position corresponding to position 163 of SEQ ID NO:4; a residue other than threonine at a position corresponding to position 204 of SEQ ID NO:4; a residue other than serine at a position corresponding to position 207 of SEQ ID NO:4; a residue other than glutamate at a position corresponding to position 236 of SEQ ID NO:4; a residue other than leucine at a position corresponding to position 251 of SEQ ID NO:4; a residue other than arginine at a position corresponding to position 261 of SEQ ID NO:4; a residue other than leucine at a position corresponding to position 265 of SEQ ID NO:4; a residue other than valine at a position corresponding to position 268 of SEQ ID NO:4; a residue other than arginine at a position corresponding to position 279 of SEQ ID NO:4; a residue other than aspartate at a position corresponding to position 293 of SEQ ID NO:4; a residue other than lysine at a position corresponding to position 296 of SEQ ID NO:4; and a residue other than asparagine at a position corresponding to position 309 of SEQ ID NO:4. The mutant thioesterases may have any one or more of the above-referenced substitutions in any combination.

The mutant thioesterases may comprise one or more of: a methionine or a conservative variant of methionine at the position corresponding to position 9 of SEQ ID NO:4; a glutamate or a conservative variant of glutamate at the position corresponding to position 15 of SEQ ID NO:4; an arginine or a conservative variant of arginine at the position corresponding to position 17 of SEQ ID NO:4; a glutamate or a conservative variant of glutamate at the position corresponding to position 23 of SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 28 of SEQ ID NO:4; a threonine or a conservative variant of threonine at the position corresponding to position 29 of SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 59 of SEQ ID NO:4; a methionine or a conservative variant of methionine at the position corresponding to position 65 of SEQ ID NO:4; a glutamine or a conservative variant of glutamine at the position corresponding to position 86 of

SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 117 of SEQ ID NO:4; a valine, an isoleucine, or a conservative variant of valine or isoleucine at the position corresponding to position 136 of SEQ ID NO:4; a lysine or a conservative variant of lysine at the position corresponding to position 146 of SEQ ID NO:4; a leucine or a conservative variant of leucine at the position corresponding to position 163 of SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 204 of SEQ ID NO:4; a threonine or a conservative variant of threonine at the position corresponding to position 207 of SEQ ID NO:4; an alanine or a conservative variant of alanine at the position corresponding to position 236 of SEQ ID NO:4; a methionine or a conservative variant of methionine at the position corresponding to position 251 of SEQ ID NO:4; a serine or a conservative variant of serine at the position corresponding to position 261 of SEQ ID NO:4; an isoleucine or a conservative variant of isoleucine at the position corresponding to position 265 of SEQ ID NO:4; an isoleucine or a conservative variant of isoleucine at the position corresponding to position 268 of SEQ ID NO:4; a histidine or a conservative variant of histidine at the position corresponding to position 279 of SEQ ID NO:4; a valine or a conservative variant of valine at the position corresponding to position 293 of SEQ ID NO:4; an arginine or a conservative variant of arginine at the position corresponding to position 296 of SEQ ID NO:4; and an aspartate or a conservative variant of aspartate at the position corresponding to position 309 of SEQ ID NO:4. The mutant thioesterases may have any one or more of the above-referenced residues in any combination.

Some mutant thioesterases of the invention may comprise a residue other than asparagine at the position corresponding to position 28 of SEQ ID NO:4 and/or a residue other than isoleucine at the position corresponding to position 65 of SEQ ID NO:4. These mutant thioesterases may comprise a serine or a conservative variant of serine at the position corresponding to position 28 of SEQ ID NO:4 and/or a methionine or a conservative variant of methionine at the position corresponding to position 65 of SEQ ID NO:4.

Some mutant thioesterases of the invention may comprise a residue other than alanine at the position corresponding to position 59 of SEQ ID NO:4 and/or a residue other than lysine at the position corresponding to position 296 of SEQ ID NO:4. These mutant thioesterases may comprise a residue other than alanine at the position corresponding to position 59 of SEQ ID NO:4 and/or a residue other than lysine at the position corresponding to position 296 of SEQ ID NO:4.

Some mutant thioesterases of the invention may comprise a residue other than aspartate at the position corresponding to position 293 of SEQ ID NO:4. These mutant thioesterases may comprise a valine or a conservative variant of valine at the position corresponding to position 293 of SEQ ID NO:4.

Some mutant thioesterases of the invention may comprise a residue other than lysine at the position corresponding to position 296 of SEQ ID NO:4. These mutant thioesterases may comprise an arginine or a conservative variant of arginine at the position corresponding to position 296 of SEQ ID NO:4.

Some mutant thioesterases of the invention may comprise a residue other than arginine at the position corresponding to position 261 of SEQ ID NO:4. These mutant thioesterases may comprise a serine or a conservative variant of serine at the position corresponding to position 261 of SEQ ID NO:4.

The mutant thioesterases may lack various N-terminal portions characteristic of various natural thioesterases. The

mutant thioesterases, for example, may lack an N-terminal portion having amino acid sequence identical to positions 1-10, positions 1-20, positions 1-30, positions 1-40, positions 1-50, positions 1-60, positions 1-65, positions 1-70, positions 1-75, positions 1-80, positions 1-81, positions 1-82, positions 1-83, positions 1-84, positions 1-85, positions 1-86, positions 1-87, positions 1-88, positions 1-89, positions 1-90, positions 1-91, positions 1-92, positions 1-93, or positions 1-94 of SEQ ID NO:2. The mutant thioesterases may lack an N-terminal portion having an amino acid sequence identical to positions 1-2, positions 1-3, positions 1-4, positions 1-5, positions 1-6, positions 1-7, positions 1-8, positions 1-9, positions 1-10, positions 1-11, positions 1-12, positions 1-13, positions 1-14, positions 1-15, positions 1-16, positions 1-17, or positions 1-18 of SEQ ID NO:4. The N-terminal portions are lacking at positions N-terminal (i.e., closer to the N-terminus) of the amino acid sequence at least about 30%, least about 40%, at least about 50%, at least about 60%, at least about 70%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 97%, or at least about 99%, identical to positions 41-317, positions 40-317, positions 39-317, positions 38-317, positions 37-317, positions 36-317, positions 35-317, positions 34-317, positions 33-317, positions 32-317, positions 31-317, positions 30-317, positions 29-317, positions 28-317, positions 27-317, positions 26-317, positions 25-317, positions 24-317, positions 23-317, positions 22-317, positions 21-317, positions 20-317, positions 19-317, positions 18-317, positions 17-317, positions 16-317, positions 15-317, positions 14-317, positions 13-317, positions 12-317, positions 11-317, positions 10-317, positions 9-317, positions 8-317, positions 7-317, positions 6-317, positions 5-317, positions 4-317, positions 3-317, positions 2-317, or positions 1-317 of SEQ ID NO:4.

The mutant thioesterases of the invention may be derived from a precursor thioesterase, wherein each of the mutants (or the naturally-occurring equivalents) has at least one altered property in vitro and/or in vivo, as compared to the properties of the precursor thioesterase. The altered property preferably comprises an enhancement of an aspect of thioesterase activity. The altered property may include increased thioesterase activity with medium-chain substrates, such as C8 substrates. The altered property may comprise an increase in selectivity or catalytic rate in hydrolyzing a medium-chain acyl-acyl carrier protein (ACP) substrate or a medium-chain acyl-CoA substrate to yield a free fatty acid or a free fatty acid derivative. The altered property may comprise an increase in selectivity or catalytic rate in hydrolyzing a C8-ACP substrate or a C8 acyl-CoA substrate to yield a free fatty acid or a free fatty acid derivative. An exemplary precursor thioesterase is *Cuphea palustris* FatB1 thioesterase (CpFatB1) represented by SEQ ID NO:1 (nucleotide coding sequence) and SEQ ID NO:2 (protein sequence).

Another aspect of the invention is a polynucleotide (or a gene) encoding a mutant thioesterase of the invention. Another aspect of the invention is a vector comprising the polynucleotide (or the gene) according to the invention. Vectors of the invention can be transformed into suitable host cells to produce recombinant host cells.

Another aspect of the invention is a recombinant host cell comprising a polynucleotide encoding a mutant thioesterase or a naturally-occurring equivalent thereof. In some versions, known genomic alteration or modification techniques can be employed to alter or modify the endogenous thio-

esterases of the host cell, effectuating one or more of the aforementioned mutations, such that at least one of the mutant endogenous thioesterases has at least one altered property. In other versions, the recombinant host cell is engineered to include a plasmid comprising a polynucleotide encoding a mutant thioesterase. In yet other versions, the recombinant host cell is engineered to include the polynucleotide encoding the mutant thioesterase integrated into the chromosome of the host cell.

The recombinant host cell of the invention can be selected from any cell capable of expressing a recombinant gene construct, and can be selected from a microbial, plant or animal cell. In a particular embodiment, the host cell is bacterial, cyanobacterial, fungal, yeast, algal, human or mammalian in origin. In a particular embodiment, the host cell is selected from any of Gram positive bacterial species such as Actinomycetes; Bacillaceae, including *Bacillus alkalophilus*, *Bacillus subtilis*, *Bacillus licheniformis*, *Bacillus lentus*, *Bacillus brevis*, *Bacillus stearothermophilus*, *Bacillus alkalophilus*, *Bacillus amyloliquefaciens*, *Bacillus coagulans*, *Bacillus circulans*, *Bacillus lautus*, *Bacillus megaterium*, *B. thuringiensis*; *Brevibacterium* sp., including *Brevibacterium flavum*, *Brevibacterium lactofermentum*, *Brevibacterium ammoniagenes*, *Brevibacterium butanicum*, *Brevibacterium divaricatum*, *Brevibacterium healii*, *Brevibacterium ketoglutamicum*, *Brevibacterium ketosoreduc-tum*, *Brevibacterium lactofermentum*, *Brevibacterium lin-ens*, *Brevibacterium paraffinolyticum*; *Corynebacterium* spp. such as *C. glutamicum* and *C. melassecola*, *Coryne-bacterium herculis*, *Corynebacterium lilium*, *Corynebacte-rtium acetoacidophilum*, *Corynebacterium acetoglutami-cum*, *Corynebacterium acetophilum*, *Corynebacterium ammoniagenes*, *Corynebacterium fujiokense*, *Corynebacte-rium nitrilophilus*; or lactic acid bacterial species including *Lactococcus* spp. such as *Lactococcus lactis*; *Lactobacillus* spp. including *Lactobacillus reuteri*; *Leuconostoc* spp.; *Pediococcus* spp.; *Serratia* spp. such as *Serratia marc-escens*; *Streptomyces* species, such as *Streptomyces lividans*, *Streptomyces murinus*, *S. coelicolor* and *Streptococcus* spp. Alternatively, strains of a Gram negative bacterial species belonging to Enterobacteriaceae including *E. coli*, *Cellulo-monas* spp.; or to Pseudomonadaceae including *Pseudomo-nas aeruginosa*, *Pseudomonas alcaligenes*, *Pseudomonas fluorescens*, *Pseudomonas putida*, *Pseudomonas syringae* and *Burkholderia cepacia*, *Salmonella* sp., *Stenotrophomo-nas* spp., and *Stenotrophomonas maltophilia*. Oleaginous microorganisms such as *Rhodococcus* spp., *Rhodococcus opacus*, *Ralstonia* spp., and *Acetivobacter* spp. are useful as well. Furthermore, yeasts and filamentous fungal strains can be useful host cells, including *Absidia* spp.; *Acremonium* spp.; *Agaricus* spp.; *Anaeromyces* spp.; *Aspergillus* spp., including *A. aculeatus*, *A. awamori*, *A. flavus*, *A. foetidus*, *A. fumaricus*, *A. fumigatus*, *A. nidulans*, *A. niger*, *A. oryzae*, *A. terreus*; *A. tubingensis* and *A. versicolor*; *Aeurobasidium* spp.; *Cephalosporium* spp.; *Chaetomium* spp.; *Coprinus* spp.; *Dactyllum* spp.; *Fusarium* spp., including *F. conglo-merans*, *F. decemcellulare*, *F. javanicum*, *F. lini*, *F. oxyspo-rum* and *F. solani*; *Gliocladium* spp.; *Kluyveromyces* sp.; *Hansenula* sp.; *Humicola* spp., including *H. insolens* and *H. lanuginosa*; *Hypocrea* spp.; *Mucor* spp.; *Neurospora* spp., including *N. crassa* and *N. sitophila*; *Neocallimastix* spp.; *Orpinomyces* spp.; *Penicillium* spp.; *Phanerochaete* spp.; *Phlebia* spp.; *Pichia* sp.; *Piromyces* spp.; *Rhizopus* spp.; *Rhizomucor* species such as *Rhizomucor miehei*; *Saccaro-myces* species such as *S. cerevisiae*, *S. pastorianus*, *S. eubayanus*, and *S. fragilis*; *Schizophyllum* spp.; *Schizosac-charomyces* such as, for example, *S. pombe* species; *chyta-*

lidium sp., *Sulpholobus* sp., *Thermoplasma* sp., *Thermomyces* sp.; *Trametes* spp.; *Trichoderma* spp., including *T. reesei*, *T. reesei (longibrachiatum)* and *T. viride*; *Yarrowinia* sp.; and *Zygorhynchus* spp and in particular include oleaginous yeast just *Phafia* spp., *Rhorosporidium toruloides* Y4, *Rhodotorula Glutinis* and *Candida* 107.

In some versions of the invention, a recombinant host cell is provided, which expresses or overexpresses a gene encoding the mutant thioesterase, and which also expresses (or overexpresses) one or more genes encoding one or more enzymes that utilize, as substrates, reaction products of the mutant thioesterase (e.g., fatty acids, fatty acyl-CoAs, fatty acyl-phosphate esters, fatty aldehydes, fatty esters, or fatty alcohols) or reaction products of one or more other enzymes that are parts of a metabolic pathway, including reaction products of the mutant thioesterase (e.g., fatty acids) as precursors and/or substrates.

In one embodiment of the invention, a recombinant host cell is provided, which expresses or overexpresses a gene encoding a mutant thioesterase and which also expresses (or overexpresses) one or more genes encoding one or more enzymes that react with a substrate that is necessary as a precursor to a reaction in a fatty acid biosynthetic pathway. In a particular embodiment, the recombinant host cell includes a gene that encodes a thioesterase and a gene that encodes an enzyme that reacts with a substrate that is necessary as a precursor to a reaction in a fatty acid synthetic pathway, which comprises the overexpression or modifica-tion of a gene selected from *pdh*, *panK*, *aceEF*, *fabH*, *fabD*, *fabG*, *acpP*, *fadR*, *accABCD*, *fabI*, *fabA*, *fabB*, *fabF*, and/or any homologs thereof.

In some versions of the invention, the recombinant host cell comprises a gene (or a polynucleotide) that encodes a mutant thioesterase and also comprises the attenuation or deletion of a gene that reduces carbon flux, or a gene that competes for substrates, cofactors, or energy requirements within a fatty acid biosynthetic pathway. In a particular embodiment, the attenuated gene comprises at least one of *fadE*, *gpsA*, *ldhA*, *pflB*, *adhE*, *pta*, *poxB*, *ackA*, *ackB*, *plsB*, *ldh*, *glta*, *sfa*, and/or any homologs thereof.

In some versions of the invention, a recombinant host cell comprises a gene (or a polynucleotide) encoding a mutant thioesterase and a heterologously-introduced exogenous gene encoding at least one fatty acid derivative enzyme. In certain embodiments, the exogenous gene or polynucleotide encodes, for example, an acyl-CoA synthase, a wax/ester synthase, an alcohol acyltransferase, an alcohol dehydroge-nase, an acyl-CoA reductase, an acyl-ACP reductase, a fatty-alcohol-forming acyl-CoA reductase, an alcohol O-acyltransferase, an aldehyde deformylating oxygenase, a fatty-acid O-methyltransferase, a carboxylic acid reductase, a decarboxylase, an aldehyde reductase, a fatty alcohol acetyl transferase, an acyl condensing enzyme, an amino-transferase, and/or a decarboxylase.

In some versions of the invention, a gene encoding the mutant thioesterase and/or a fatty acid derivative enzyme, for example, an acyl-CoA synthase, a wax/ester synthase, an alcohol acyltransferase, an alcohol dehydrogenase, an acyl-CoA reductase, an acyl-ACP reductase, a fatty-alcohol-forming acyl-CoA reductase, an alcohol O-acyltransferase, an aldehyde deformylating oxygenase, a fatty-acid O-methyltransferase, a carboxylic acid reductase, a decarboxylase, an aldehyde reductase, a fatty alcohol acetyl transferase, an acyl condensing enzyme, an aminotransferase, a polyhy-droxalkanoate (PHA) synthase, and/or a decarboxylase, is overexpressed.

In some versions of the invention, genes encoding mutant thioesterases, fatty acid derivative enzymes and/or other recombinantly expressed genes in a recombinant host cell are modified to optimize at least one codon for expression in the recombinant host cell.

In some versions of the invention, the recombinant host cell comprises at least one gene encoding a mutant thioesterase and a gene encoding an acyl-CoA synthase. The acyl-CoA synthase gene can be any of fadD, fadK, BH3103, yhfL, pfl-4354, EAV15023, fadD1, fadD2, RPC_4074, fadDD35, fadDD22, faa3p, or the gene encoding the protein ZP_01644857. Other examples of acyl-CoA synthase genes include fadDD35 from *M. tuberculosis* HR7Rv [NP_217021], yhfL from *B. subtilis* [NP_388908], fadD1 from *P. aeruginosa* PAO1 [NP_251989], the gene encoding the protein ZP_01644857 from *Stenotrophomonas maltophilia* R551-3, or faa3p from *Saccharomyces cerevisiae* [NP_012257]. Other examples are described elsewhere herein.

In some versions of the invention, a recombinant host cell is provided comprising at least one gene or polynucleotide encoding a mutant thioesterase (and a gene or polynucleotide encoding an ester synthase, such as an ester synthase gene obtained from *Acinetobacter* spp., *Alcanivorax borkumensis*, *Arabidopsis thaliana*, *Saccharomyces cerevisiae*, *Homo sapiens*, *Simmondsia chinensis*, *Mortierella alpina*, *Cryptococcus curvatus*, *Alcanivorax jadensis*, *Alcanivorax borkumensis*, *Acinetobacter* sp. HO1-N, or *Rhodococcus opacus*. Examples of ester synthase genes include wax/dgat, encoding a bifunctional ester synthase/acyl-CoA: diacylglycerol acyltransferase from *Simmondsia chinensis*, *Acinetobacter* sp. strain ADPJ, *Alcanivorax borkumensis*, *Pseudomonas aeruginosa*, *Fundibacter jadensis*, *Arabidopsis thaliana*, or *Alkaligenes eutrophus*. The gene encoding the ester synthase may be overexpressed.

In some versions of the invention, the recombinant host cell comprises at least one gene encoding a fatty aldehyde biosynthetic enzyme. A fatty aldehyde biosynthetic gene can be, for example, a carboxylic acid reductase gene (e.g., a car gene).

In some versions of the invention, the recombinant host cell comprises at least one fatty alcohol production gene. Fatty alcohol production genes include, for example, fatty acyl-CoA reductases such as acrl or the fatty acyl-CoA reductase from *Marinobacter aquaeolei* VT8 (Robert M. Willis, Bradley D. Wahlen, Lance C. Seefeldt, and Brett M. Barney. Characterization of a Fatty Acyl-CoA Reductase from *Marinobacter aquaeolei* VT8: A Bacterial Enzyme Catalyzing the Reduction of Fatty Acyl-CoA to Fatty Alcohol. *Biochemistry* 2011 50 (48), 10550-10558). Other fatty alcohol production genes are described in, for example, PCT Publication Nos. 2008/119082 and 2007/136762, the disclosures of which are herein incorporated by reference. Other examples are provided elsewhere herein.

In some versions of the invention, the recombinant host cell comprises a gene encoding a mutant thioesterase and a gene encoding at least one olefin producing gene. The gene may be a terminal olefin producing gene or an internal olefin producing gene. As examples of terminal olefin producing genes, those described in PCT Publication No. 2009/085278, including orf880, are appropriate. As examples of internal olefin producing genes, those described in PCT Publication No. 2008/147781 A2 are appropriate. The disclosures of PCT Publication Nos. 2009/085278 and 2008/147781 A2 are herein incorporated by reference.

In some versions of the invention, a recombinant host cell is provided comprising at least one gene or polynucleotide

encoding a mutant thioesterase, and at least one of (a) a gene or polynucleotide encoding a fatty acid derivative enzyme and (b) a gene or polynucleotide encoding an acyl-CoA dehydrogenase enzyme that is attenuated. Preferably that gene encoding a fatty acid derivative enzyme that is attenuated or deleted is endogenous to the host cell, encoding, for example, an acyl-CoA synthase, a wax/ester synthase, an alcohol acyltransferase, an alcohol dehydrogenase, an acyl-CoA reductase, an acyl-ACP reductase, a fatty-alcohol-forming acyl-CoA reductase, an alcohol O-acyltransferase, an aldehyde deformylating oxygenase, a fatty-acid O-methyltransferase, a carboxylic acid reductase, a decarboxylase, an aldehyde reductase, a fatty alcohol acetyl transferase, an acyl condensing enzyme, an aminotransferase, and/or a decarbonylase. In one embodiment, the attenuated gene encodes an acyl-CoA synthase or an ester synthase.

In some versions of the invention, the recombinant host cell has an endogenous gene encoding an acyl-CoA dehydrogenase enzyme that is deleted or attenuated.

In some versions of the invention, a method is provided wherein the recombinant host cell according to the invention is cultured under conditions that permit expression or overexpression of a mutant thioesterases of the invention. The mutant thioesterase can be recovered, and more preferably substantially purified, after the host cell is harvested and/or lysed.

In some versions of the invention, a method is provided wherein the recombinant host cell according to the invention is cultivated under conditions that permit production of fatty acid derivatives. In a preferred embodiment, the fatty acid derivative can be recovered, and more preferably the fatty acid derivative is substantially purified. In a particularly preferred embodiment, the fatty acid derivative composition is substantially purified from other components produced during cultivation by centrifugation.

In some versions of the invention, a method is provided for producing a fatty acid derivative, comprising cultivating a recombinant host cell of the invention under conditions suitable to ensure expression or overexpression of a mutant thioesterase, and recovering the fatty acid derivative that is produced.

In some versions of the invention, a method is provided for extracellularly producing a fatty acid derivative in vitro, comprising cultivating a recombinant host cell under conditions suitable for expression or overexpression of a mutant thioesterase of the invention, harvesting the cells, and lysing the cells, such that the thioesterase enzyme that is produced can be recovered and used to produce fatty acid derivatives in vitro. In an exemplary embodiment, the mutant thioesterase is substantially purified. In another exemplary embodiment, the mutant thioesterase is not purified from the cell lysate. The purified mutant thioesterase enzyme or the cell lysate comprising such an enzyme can then be subject to suitable thioesterase substrates under conditions that allow the production of fatty acid derivatives extracellularly. Techniques for introducing substrates to enzymes are well known in the art. A non-limiting example is adding the substrate(s) in a solution form to the enzyme solution or the cell lysate, and allowing the mixture to incubate. Another non-limiting example involves incubating the substrate(s) and enzyme solution or cell lysate by either attaching the substrate(s) or the enzyme to a solid medium (e.g., beads, resins, plates, etc.) and passing the enzyme solution/lysate or the substrate(s), respectively, through the solid medium in a speed that allows for sufficient contact between the substrate(s) and the enzyme.

In some versions of the invention, a method is provided for producing a fatty acid derivative, which comprises cultivating a recombinant host cell under conditions suitable to ensure expression of the mutant thioesterase, and recovering the fatty acid derivative that is secreted or released extracellularly. Accordingly, the fatty acid derivative product is recovered from, for example, the supernatant of a cultivation broth wherein the host cell is cultured.

In some versions of the invention, a method is provided for obtaining a fatty acid derivative composition extracellularly by cultivating a recombinant host cell that has been transformed with a polynucleotide encoding a mutant thioesterase, cultivating under conditions that permit production of a fatty acid derivative, a major or minor portion of which is secreted or released extracellularly, and recovering the fatty acid derivative that is produced. In an exemplary embodiment, the fatty acid derivative is produced within the cell, but a portion of it is released by the host cell. Accordingly, the method further comprises harvesting the cells, lysing the cells, and recovering the fatty acid derivative.

In some versions of the invention, a method of producing fatty acid derivatives is provided comprising: transforming the host cell with a polynucleotide sequence encoding a mutant thioesterase, such that the production of fatty acid derivatives in the host cell is altered relative to a cell that has not been transformed with the mutant thioesterase gene.

In some versions of the invention, a method of producing fatty acid derivatives is provided comprising: providing a polynucleotide sequence comprising a gene encoding a mutant thioesterase; transforming a suitable host cell under conditions wherein said polynucleotide sequence is incorporated into said chromosome of said cell and said gene is expressible within said host cell; cultivating the transformed host cell under conditions suitable for said host cell to express said gene and produce a mutant thioesterase protein; and recovering the fatty acid derivatives.

In any of the embodiments above, derivatives of a certain carbon chain length can be recovered at a greater proportional yield, in comparison with the production of such fatty acid derivatives of the same carbon chain length in the same host cell in the absence of the mutant thioesterase. In a particular embodiment, the fatty acid derivatives that are recovered at an increased or decreased yield comprise a primary chain length of a C8 fatty acyl chain. The fatty acid derivatives that are recovered at an increased or decreased yield in the composition can be selected from all types of fatty acid derivatives, including, for example, hydrocarbons, fatty acids, fatty esters, fatty aldehydes, fatty alcohols terminal olefins, internal olefins, alkanes, diols, fatty amines, dicarboxylic acids, polyhydroxyalkanoates, or ketones, or combinations thereof.

Alternatively, in any of the embodiments above, a particular fatty acid derivative can be produced at an increased or decreased proportional or percentage yield relative to the other fatty acid derivatives, when compared to the proportional or percentage yield of that particular fatty acid derivative in the same host cell in the absence of the mutant thioesterase.

In some versions of the invention, fatty acid derivative compositions are provided that are produced by the host cells of the invention. Such compositions can comprise hydrocarbons, esters, alcohols, ketones, aldehydes, fatty acids, dicarboxylic acids, internal olefins, terminal olefins, polyhydroxyalkanoates, and/or combinations thereof. Such compositions are useful in applications in the chemical industry, for example in the production of surfactants and

detergents, or as a biofuel and a substitute for petroleum, heating oil, kerosene, diesel, jet fuel or gasoline.

In a particular version, the fatty acid derivative composition comprises C8 (i.e., a carbon chain length of 8 carbons) fatty esters, C8 fatty acids, C8 fatty aldehydes, C8 fatty alcohols, or polyhydroxyalkanoates with C8 side chains.

In a particular version, the fatty acid derivatives of the invention comprise straight chain fatty acid derivatives, branched chain fatty acid derivatives, and/or cyclic moieties. In a particular embodiment, the fatty acid derivatives are unsaturated (e.g., monounsaturated) or saturated.

In some versions of the invention, the fatty acid derivative composition includes octanoic acid.

Another aspect of the invention is directed to a method of screening thioesterase mutants for C8 thioesterase activity. The method comprises, introducing a gene encoding a mutant thioesterase in a microorganism lacking lipB, incubating the microorganism in a medium devoid of lipoic acid and octanoic acid, and recovering the microorganism after growth in the medium. As shown in the examples, the method can be used to screen for thioesterases having enhanced C8 thioesterase activity from a library of mutant thioesterases by recovering microorganisms capable of faster growth in the medium. The incubating preferably comprises incubating the microorganism under conditions (i.e., temperature, etc.) suitable for growth when lipoic acid and/or octanoic acid is supplied exogenously. The microorganism is preferably *E. coli*.

Throughout the specification, a reference may be made using an abbreviation of a gene name or a polypeptide name, but it is understood that such an abbreviated gene or polypeptide name represents the genus of genes or polypeptides, respectively. Such gene names include all genes encoding the same polypeptide and homologous polypeptides having the same physiological function. Polypeptide names include all polypeptides that have the same activity (e.g., that catalyze the same fundamental chemical reaction).

Unless otherwise indicated, the accession numbers referenced herein are derived from the NCBI database (National Center for Biotechnology Information) maintained by the National Institute of Health, U.S.A.

EC numbers are established by the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology (NC-IUBMB) (available at www.chem.qmul.ac.uk/iubmb/enzyme/). The EC numbers referenced herein are derived from the KEGG Ligand database, maintained by the Kyoto Encyclopedia of Genes and Genomics, sponsored in part by the University of Tokyo.

As used herein, the term "alcohol dehydrogenase" (EC 1.1.1.*) is a polypeptide capable of catalyzing the conversion of a fatty aldehydes to an alcohol (e.g., a fatty alcohol). Additionally, one of ordinary skill in the art will appreciate that some alcohol dehydrogenases will catalyze other reactions as well. For example, some alcohol dehydrogenases will accept other substrates in addition to fatty aldehydes. Such non-specific alcohol dehydrogenases are, therefore, also included in this definition. Polynucleotide sequences encoding alcohol dehydrogenases are known in the art, and such dehydrogenases are publicly available.

The term "altered property" refers to a modification in one or more properties of a mutant polynucleotide or mutant protein with reference to a precursor polynucleotide or precursor protein. In one embodiment, the altered property is a changed preference for particular substrates, as reflected in, for example, a changed preference for particular acyl-CoA or acyl-acyl carrier protein substrates such as C8 acyl-CoA or acyl-acyl carrier protein substrates

The term "alignment" refers to a method of comparing two or more polynucleotides or polypeptide sequences for the purpose of determining their relationship to each other. Alignments are typically performed by computer programs that apply various algorithms, however it is also possible to perform an alignment by hand. Alignment programs typically iterate through potential alignments of sequences and score the alignments using substitution tables, employing a variety of strategies to reach a potential optimal alignment score. Commonly-used alignment algorithms include, but are not limited to, CLUSTALW, (see, Thompson J. D., Higgins D. G., Gibson T. J., CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice, *Nucleic Acids Research* 22: 4673-4680, 1994); CLUSTALV, (see, Larkin M. A., et al., CLUSTALW2, ClustalW and ClustalX version 2, *Bioinformatics* 23(21): 2947-2948, 2007); Jotun-Hein, Muscle et al., MUSCLE: a multiple sequence alignment method with reduced time and space complexity, *BMC Bioinformatics* 5: 113, 2004); Mafft, Kalign, ProbCons, and T-Coffee (see Notredame et al., T-Coffee: A novel method for multiple sequence alignments, *Journal of Molecular Biology* 302: 205-217, 2000). Exemplary programs that implement one or more of the above algorithms include, but are not limited to MegAlign from DNASTar (DNASTar, Inc. 3801 Regent St. Madison, Wis. 53705), MUSCLE, T-Coffee, CLUSTALX, CLUSTALV, JalView, Phylip, and Discovery Studio from Accelrys (Accelrys, Inc., 10188 Telesis Ct, Suite 100, San Diego, Calif. 92121). In a non-limiting example, MegAlign is used to implement the CLUSTALW alignment algorithm with the following parameters: Gap Penalty 10, Gap Length Penalty 0.20, Delay Divergent Seqs (30%) DNA Transition Weight 0.50, Protein Weight matrix Gonnet Series, DNA Weight Matrix IUB.

The term "carbon chain length" is defined herein as the number of carbon atoms in a carbon chain of a thioesterase substrate or a fatty acid derivative. The carbon chain length of a particular molecule is marked as CX, wherein the "X" refers to the number of carbons in the carbon chain. "Long-chain" (e.g., long-chain fatty acid, fatty acyl-ACP, or fatty acyl-CoA) refers to molecules having a carbon chain longer than 12 carbons. "Medium-chain" (e.g., medium-chain fatty acid, fatty acyl-ACP, or fatty acyl-CoA) refers to molecules having a carbon chain of 6 to 12 carbons. "Short-chain" (e.g., short chain fatty acid, fatty acyl-ACP, or fatty acyl-CoA) refers to molecules having a carbon chain fewer than 6 carbons.

The term "carbon source" means a substrate or compound suitable to be used as a source of carbon for prokaryotic or simple eukaryotic cell growth. Carbon sources can be in various forms, including, but not limited to polymers, carbohydrates, acids, alcohols, aldehydes, ketones, amino acids, peptides, gases (e.g., CO and CO₂), and the like. These include, for example, various monosaccharides such as glucose, fructose, mannose and galactose; oligosaccharides such as fructo-oligosaccharide and galacto-oligosaccharide; polysaccharides such as xylose, and arabinose; disaccharides such as sucrose, maltose and turanose; cellulosic material such as methyl cellulose and sodium carboxymethyl cellulose; saturated or unsaturated fatty acid esters such as succinate, lactate and acetate; alcohols such as ethanol, etc., or mixtures thereof. The carbon source can additionally be a product of photosynthesis, including, but not limited to glucose. Glycerol can be an effective carbon source as well. Suitable carbon sources can be generated from any number of natural and renewable sources, includ-

ing particularly biomass from agricultural, municipal and industrial waste, so long as the material can be used as a component of a cultivation to provide a carbon source. Biomass sources include corn stover, sugarcane, switchgrass, animal matter, or waste materials.

The term "chromosomal integration" means the process whereby an incoming sequence is introduced into the chromosome of a host cell. The homologous regions of the transforming DNA align with homologous regions of the chromosome. Then, the sequence between the homology boxes can be replaced by the incoming sequence in a double crossover (i.e., homologous recombination). In some embodiments of the present invention, homologous sections of an inactivating chromosomal segment of a DNA construct align with the flanking homologous regions of the indigenous chromosomal region of the microbial chromosome. Subsequently, the indigenous chromosomal region is deleted by the DNA construct in a double crossover.

The term "conditions that permit product production" refers to any cultivation conditions that allow a production host to produce a desired product, such as acyl-CoA or fatty acid derivatives including, for example, fatty acids, hydrocarbons, fatty alcohols, waxes, polyhydroxyalkanoates, or fatty esters. Cultivation conditions usually comprise many parameters. Exemplary conditions include, but are not limited to, temperature ranges, levels of aeration, pH ranges, and media composition (e.g., solvents and solutes). Each of these conditions, individually and in combination, allows the production host to grow. Exemplary media include broths or gels. Generally, a suitable medium includes a carbon source, such as glucose, fructose, cellulose, or the like, which can be metabolized by the microorganism directly. In addition, enzymes can be used in the medium to facilitate the mobilization (e.g., the depolymerization of starch or cellulose to fermentable sugars) and subsequent metabolism of the carbon source. To determine if the culture conditions are suitable for product production, the production host can be cultured for about 4, 8, 12, 24, 36, 48, or 72 hours. During culturing or after culturing, samples can be obtained and analyzed to determine if the culture conditions permit product production. For example, the production hosts in the sample or the medium in which the production hosts were grown can be tested for the presence of the desired product. When testing for the presence of a product, assays, such as, but not limited to, TLC, HPLC, GC/FID, GC/MS, LC/MS, MS, as well as those provided in the examples herein, can be used.

The term "consensus sequence" or "canonical sequence" refers to an archetypical amino acid sequence against which all variants of a particular protein or sequence of interest are compared. Either term also refers to a sequence that sets forth the nucleotides that are most often present in a polynucleotide sequence of interest. For each position of a protein, the consensus sequence gives the amino acid that is most abundant in that position in the sequence alignment.

The term "conservative substitutions" or "conserved substitutions" refers to, for example, a substitution of an amino acid with a conservative variant.

"Conservative variant" refers to residues that are functionally similar to a given residue. Amino acids within the following groups are conservative variants of one another: glycine, alanine, serine, and proline (very small); alanine, isoleucine, leucine, methionine, phenylalanine, valine, proline, and glycine (hydrophobic); alanine, valine, leucine, isoleucine, methionine (aliphatic-like); cysteine, serine, threonine, asparagine, tyrosine, and glutamine (polar); phenylalanine, tryptophan, tyrosine (aromatic); lysine, arginine,

and histidine (basic); aspartate and glutamate (acidic); alanine and glycine; asparagine and glutamine; arginine and lysine; isoleucine, leucine, methionine, and valine; and serine and threonine.

The terms “corresponds to” or “corresponding to” refer to an amino acid residue or position in a first protein sequence being positionally equivalent to an amino acid residue or position in a second reference protein sequence by virtue of the fact that the residue or position in the first protein sequence aligns to the residue or position in the reference sequence using bioinformatic techniques, for example, using the methods described herein for preparing a sequence alignment. The corresponding residue in the first protein sequence is then assigned the position number in the second reference protein sequence.

The term “deletion,” when used in the context of an amino acid sequence, means a deletion in or a removal of one or more residues from the amino acid sequence of a precursor protein, resulting in a mutant protein having at least one less amino acid residue as compared to the precursor protein. The term can also be used in the context of a nucleotide sequence, which means a deletion in or removal of a nucleotide from the polynucleotide sequence of a precursor polynucleotide.

The term “DNA construct” and “transforming DNA” (wherein “transforming” is used as an adjective) are used interchangeably herein to refer to a DNA used to introduce sequences into a host cell or organism. Typically a DNA construct is generated in vitro by PCR or other suitable technique(s) known to those in the art. In certain embodiments, the DNA construct comprises a sequence of interest (e.g., an incoming sequence). In some embodiments, the sequence is operably linked to additional elements such as control elements (e.g., promoters, etc.). A DNA construct can further comprise a selectable marker. It can also comprise an incoming sequence flanked by homology targeting sequences. In a further embodiment, the DNA construct comprises other non-homologous sequences, added to the ends (e.g., stuffer sequences or flanks). In some embodiments, the ends of the incoming sequence are closed such that the DNA construct forms a closed circle. The transforming sequences may be wildtype, mutant or modified. In some embodiments, the DNA construct comprises sequences homologous to the host cell chromosome. In other embodiments, the DNA construct comprises non-homologous sequences. Once the DNA construct is assembled in vitro it may be used to: 1) insert heterologous sequences into a desired target sequence of a host cell; 2) mutagenize a region of the host cell chromosome (i.e., replace an endogenous sequence with a heterologous sequence); 3) delete target genes; and/or (4) introduce a replicating plasmid into the host. A polynucleotide is said to “encode” an RNA or a polypeptide if, in its native state or when manipulated by methods known to those of skill in the art, it can be transcribed and/or translated to produce the RNA, the polypeptide, or a fragment thereof. The antisense strand of such a polynucleotide is also said to encode the RNA or polypeptide sequences. As is known in the art, a DNA can be transcribed by an RNA polymerase to produce an RNA, and an RNA can be reverse transcribed by reverse transcriptase to produce a DNA. Thus a DNA can encode an RNA, and vice versa.

An “ester synthase” is a peptide capable of catalyzing a biochemical reaction to producing esters. For example, an ester synthase is a peptide that is capable of participating in converting a thioester to a fatty ester. In certain embodiments, an ester synthase converts a thioester, acyl-CoA, to a

fatty ester. In an alternate embodiment, an ester synthase uses a thioester and an alcohol as substrates to produce a fatty ester. Ester synthases are capable of using short and long chain acyl-CoAs as substrates. In addition, ester synthases are capable of using short and long chain alcohols as substrates. Non-limiting examples of ester synthases include wax synthases, wax-ester synthases, acyl-CoA: alcohol transacylases, acyltransferases, fatty acyl-coenzyme A: fatty alcohol acyltransferases, fatty acyl-ACP transacylases, fatty-acid O-methyltransferases (EC 2.1.1.15), alcohol O-acyltransferases such as ATF (Rodriguez G M, Tashiro Y, Atsumi S. Expanding ester biosynthesis in *Escherichia coli*. *Nat Chem Biol*. 2014 April; 10(4):259-65), and alcohol acetyltransferases. An ester synthase that converts an acyl-CoA thioester to a wax is called a wax synthase. Exemplary ester synthases include those classified under the enzyme classification number EC 2.3.1.75. The term “ester synthase” does not comprise enzymes that also have thioesterase activity. The ones that have both ester synthase activity and thioesterase activity are categorized as thioesterases herein.

The term “expressed genes” refers to genes that are transcribed into messenger RNA (mRNA) and then translated into protein, as well as genes that are transcribed into types of RNA, such as transfer RNA (tRNA), ribosomal RNA (rRNA), and regulatory RNA, which are not translated into protein.

The terms “expression cassette” or “expression vector” refer to a polynucleotide construct generated recombinantly or synthetically, with a series of specified elements that permit transcription of a particular polynucleotide in a target cell. A recombinant expression cassette can be incorporated into a plasmid, chromosome, mitochondrial DNA, plasmid DNA, virus, or polynucleotide fragment. Typically, the recombinant expression cassette portion of an expression vector includes, among other sequences, a polynucleotide sequence to be transcribed and a promoter. In particular embodiments, expression vectors have the ability to incorporate and express heterologous polynucleotide fragments in a host cell. Many prokaryotic and eukaryotic expression vectors are commercially available. Selection of appropriate expression vectors is within the knowledge of those of skill in the art. The term “expression cassette” is also used interchangeably herein with “DNA construct,” and their grammatical equivalents.

The term “fatty acid derivative,” as used herein, refers to a composition that is derived from a metabolic pathway, which pathway includes a thioesterase reaction. Thus, fatty acid derivative products can be products that are, or are derived from, fatty acid fatty thioester, or fatty esters that are directly or indirectly products of a thioesterase reaction. Fatty acid derivatives thus include, for example, products that are, or that are derived from, fatty acids that are the direct or indirect reaction product of a thioesterase, and/or a fatty ester that is a direct or indirect reaction product of a thioesterase. Exemplary fatty acid derivatives include, for example, short and long chain alcohols, hydrocarbons, and fatty alcohols and esters, including waxes, fatty acid esters, and/or fatty esters. Specific non-limiting examples of fatty acid derivatives include fatty acids, fatty acid methyl esters, fatty acid ethyl esters, fatty alcohols, fatty alkyl-acetates, fatty aldehydes, fatty amines, fatty amides, fatty sulfates, fatty ethers, ketones, alkanes, internal olefins, terminal olefins, dicarboxylic acids, polyhydroxyalkanoates, diols and terminal and/or internal fatty acids.

The term “fatty acid derivative enzymes” refers to, collectively and individually, enzymes that may be expressed or

overexpressed in the production of fatty acid derivatives. These enzymes may be parts of a fatty acid biosynthetic pathway. Non-limiting examples of fatty acid derivative synthases include fatty acid synthases, thioesterases, acyl-CoA synthases, acyl-CoA reductases, wax/ester synthases, alcohol dehydrogenases, alcohol acyltransferases, fatty alcohol acetyl transferases, fatty alcohol-forming acyl-CoA reductase, fatty-alcohol-forming acyl-CoA reductases, fatty acid decarboxylases, alcohol O-acyltransferases, carboxylic acid reductases, fatty alcohol acetyl transferases, aldehyde deformylating oxygenases, aldehyde reductases, decarboxylases, acyl condensing enzymes, aminotransferases, decarboxylases, fatty-acid O-methyltransferases, carboxylic acid reductases, decarboxylases, and ester synthases.

Fatty acid derivative enzymes convert substrates into fatty acid derivatives. In certain circumstances, a suitable substrate may be a first fatty acid derivative, which is converted by a fatty acid derivative enzyme into a different, second fatty acid derivative.

The term "fatty alcohol" refers to an alcohol having the formula ROH. In certain embodiments, a fatty alcohol is an alcohol made from a fatty acid or fatty acid derivative. In one embodiment, the R group is at least about 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 carbons in length. R can be straight or branched chain. The branched chains may have one or more points of branching. In addition, the branched chains may include cyclic branches, such as cyclopropane or epoxide moieties. Furthermore, R can be saturated or unsaturated. If unsaturated, R can have one or more points of unsaturation. In one embodiment, the fatty alcohol is produced biosynthetically. Fatty alcohols have many uses. For example, fatty alcohols can be used to produce specialty chemicals. Specifically, fatty alcohols can be used as bio-fuels; as solvents for fats, waxes, gums, and resins; in pharmaceutical salves, emollients and lotions; as lubricating-oil additives; in detergents and emulsifiers; as textile antistatic and finishing agents; as plasticizers; as nonionic surfactants; and in cosmetics, for example as thickeners.

The term "fatty alcohol forming peptides" refers to peptides capable of catalyzing the conversion of acyl-CoA to fatty alcohol, including fatty alcohol forming acyl-CoA reductase (FAR, EC 1.1.1.*), acyl-CoA reductase (EC 1.2.1.50), long-chain acyl-(acyl-carrier-protein) reductase (EC1.2.1.80), or alcohol dehydrogenase (EC 1.1.1.1). Additionally, one of ordinary skill in the art will appreciate that some fatty alcohol forming peptides will catalyze other reactions as well. For example, some acyl-CoA reductase peptides will accept substrates other than acyl-CoA such as acyl-ACP. Such non-specific peptides are, therefore, also included. Polynucleotide sequences encoding fatty alcohol forming peptides are known in the art and such peptides are publicly available.

The term "fatty aldehyde" refers to an aldehyde having the formula RCHO characterized by an unsaturated carbonyl group (C=O). In certain embodiments, a fatty aldehyde is an aldehyde made from a fatty acid or fatty acid derivative. In one embodiment, the R group is at least about 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 carbons in length. R can be straight or branched chain. The branched chains may have one or more points of branching. In addition, the branched chains can be cyclic branches. Furthermore, R can be saturated or unsaturated. If unsaturated, R can have one or more points of unsaturation. In one embodiment, the fatty aldehyde is produced biosynthetically. Fatty aldehydes have many uses. For example, fatty aldehydes can be used to produce specialty chemicals. Specifically, fatty aldehydes can be used to produce poly-

mers, resins, dyes, flavorings, plasticizers, perfumes, pharmaceuticals, and other chemicals. Some are used as solvents, preservatives, or disinfectants. Some natural and synthetic compounds, such as vitamins and hormones, are also aldehydes.

The terms "fatty aldehyde biosynthetic polypeptide," "carboxylic acid reductase," and "CAR" are used interchangeably herein.

The term "fatty ester" refers to an ester having greater than 5 carbon atoms. In certain embodiments, a fatty ester is an ester made from a fatty acid, for example a fatty acid ester. In one embodiment, a fatty ester contains an A side (i.e., the carbon chain attached to the carboxylate oxygen) and a B side (i.e., the carbon chain comprising the parent carboxylate). In a particular embodiment, when a fatty ester is derived from the fatty acid biosynthetic pathway, the A side is contributed by an alcohol, and the B side is contributed by a fatty acid. Any alcohol can be used to form the A side of the fatty esters. For example, the alcohol can be derived from the fatty acid biosynthetic pathway. Alternatively, the alcohol can be produced through non-fatty acid biosynthetic pathways. Moreover, the alcohol can be provided exogenously. For example, the alcohol can be supplied to the cultivation broth in instances where the fatty ester is produced by an organism. Alternatively, a carboxylic acid, such as a fatty acid or acetic acid, can be supplied exogenously in instances where the fatty ester is produced by an organism that can also produce alcohol. The carbon chains comprising the A side or B side can be of any length. In one embodiment, the A side of the ester is at least about 1, 2, 3, 4, 5, 6, 7, 8, 10, 12, 14, 16, 18, or 20 carbons in length. The B side of the ester is at least about 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, or 26 carbons in length. The A side and/or the B side can be straight or branched chain. The branched chains may have one or more points of branching. In addition, the branched chains may include cyclic branches, such as cyclopropane or epoxide moieties. Furthermore, the A side and/or B side can be saturated or unsaturated. If unsaturated, the A side and/or B side can have one or more points of unsaturation. In one embodiment, the fatty ester is produced biosynthetically. In this embodiment, first the fatty acid is "activated." Non-limiting examples of activated fatty acids are acyl-CoA, acyl ACP, acyl-AMP, and acyl phosphate. Acyl-CoA can be a direct product of fatty acid biosynthesis or degradation. In addition, acyl-CoA can be synthesized from a free fatty acid, a CoA, and an adenosine nucleotide triphosphate (ATP). An example of an enzyme that produces acyl-CoA is an acyl-CoA synthase. After the fatty acid is activated, it can be readily transferred to a recipient nucleophile. Exemplary nucleophiles are alcohols, thiols, amines, or phosphates. In another embodiment, the fatty ester can be derived from a fatty acyl-thioester and an alcohol. In one embodiment, the fatty ester is a wax. The wax can be derived from a long chain fatty alcohol and a long chain fatty acid. In another embodiment, the fatty ester is a fatty acid thioester, for example fatty acyl coenzyme A (acyl-CoA). In other embodiments, the fatty ester is a fatty acyl pantothenate, an acyl acyl carrier protein (acyl-ACP), a fatty acyl enzyme ester, or a fatty phosphate ester. An ester can be formed from an acyl enzyme ester intermediate through the alcoholysis of the ester bond to form a new ester and the free enzyme. Fatty esters have many uses. For example, fatty esters can be used as, or as a component of, a biofuel or a surfactant.

"Gene" refers to a polynucleotide (e.g., a DNA segment), which encodes a polypeptide, and may include regions

preceding and following the coding regions as well as intervening sequences (introns) between individual coding segments (exons).

The term "homologous genes" refers to a pair of genes from different but related species, which correspond to each other and which are identical or similar to each other. The term encompasses genes that are separated by the speciation process during the development of new species) (e.g., orthologous genes), as well as genes that have been separated by genetic duplication (e.g., paralogous genes).

The term "endogenous protein" refers to a protein that is native to or naturally occurring in a cell. "Endogenous polynucleotide" refers to a polynucleotide that is in the cell and was not introduced into the cell using recombinant engineering techniques. For example, a gene that was present in the cell when the cell was originally isolated from nature. A gene is still considered endogenous if the control sequences, such as a promoter or enhancer sequences that activate transcription or translation, have been altered through recombinant techniques. Conversely, the term "heterologous" is also used herein, and refers to a protein or a polynucleotide that does not naturally occur in a host cell.

The term "homologous recombination" refers to the exchange of DNA fragments between two DNA molecules or paired chromosomes at sites of identical or nearly identical nucleotide sequences. In certain embodiments, chromosomal integration is homologous recombination.

The term "homologous sequences" as used herein refers to a polynucleotide or polypeptide sequence having, for example, about 100%, about 99% or more, about 98% or more, about 97% or more, about 96% or more, about 95% or more, about 94% or more, about 93% or more, about 92% or more, about 91% or more, about 90% or more, about 88% or more, about 85% or more, about 80% or more, about 75% or more, about 70% or more, about 65% or more, about 60% or more, about 55% or more, about 50% or more, about 45% or more, or about 40% or more sequence identity to another polynucleotide or polypeptide sequence when optimally aligned for comparison. In particular embodiments, homologous sequences can retain the same type and/or level of a particular activity of interest. In some embodiments, homologous sequences have between 85% and 100% sequence identity, whereas in other embodiments there is between 90% and 100% sequence identity. In particular embodiments, there is 95% and 100% sequence identity.

"Homology" refers to sequence similarity or sequence identity. Homology is determined using standard techniques known in the art (see, e.g., Smith and Waterman, *Adv. Appl. Math.*, 2:482, 1981; Needleman and Wunsch, *J. Mol. Biol.*, 48:443, 1970; Pearson and Lipman, *Proc. Natl. Acad. Sci. USA* 85:2444, 1988; programs such as GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package (Genetics Computer Group, Madison, Wis.); and Devereux et al., *Nucl. Acid Res.*, 12:387-395, 1984). A non-limiting example includes the use of the BLAST program (Altschul et al., Gapped BLAST and PSI-BLAST: a new generation of protein database search programs, *Nucleic Acids Res.* 25:3389-3402, 1997) to identify sequences that can be said to be "homologous." A recent version such as version 2.2.16, 2.2.17, 2.2.18, 2.2.19, or the latest version, including sub-programs such as blastp for protein-protein comparisons, blastn for nucleotide-nucleotide comparisons, tblastn for protein-nucleotide comparisons, or blastx for nucleotide-protein comparisons, and with parameters as follows: Maximum number of sequences returned 10,000 or 100,000; E-value (expectation value) of 1e-2 or 1e-5, word size 3, scoring matrix BLOSUM62, gap

cost existence 11, gap cost extension 1, may be suitable. An E-value of 1e-5, for example, indicates that the chance of a homologous match occurring at random is about 1 in 10,000, thereby marking a high confidence of true homology.

The term "host strain" or "host cell" refers to a suitable host for an expression vector comprising a DNA of the present invention. The host may comprise any organism, without limitation, capable of containing and expressing the nucleic acids or genes disclosed herein. The host may be prokaryotic or eukaryotic, single-celled or multicellular, including mammalian cells, plant cells, fungi, etc. Examples of single-celled hosts include cells of *Escherichia*, *Salmonella*, *Bacillus*, *Clostridium*, *Streptomyces*, *Staphylococcus*, *Neisseria*, *Lactobacillus*, *Shigella*, and *Mycoplasma*. Suitable *E. coli* strains (among a great many others) include BL21(DE3), C600, DH5 α F', HB101, JM83, JM101, JM103, JM105, JM107, JM109, JM110, MC1061, MC4100, MM294, NM522, NM554, TGI, χ 1776, XL1-Blue, and Y1089+, all of which are commercially available.

The term "hydrocarbon" refers to chemical compounds that contain the elements carbon (C) and hydrogen (H). All hydrocarbons consist of a carbon backbone and atoms of hydrogen attached to that backbone. Sometimes, the term is used as a shortened form of the term "aliphatic hydrocarbon." There are essentially three types of hydrocarbons: (1) aromatic hydrocarbons, which have at least about one aromatic ring; (2) saturated hydrocarbons, also known as alkanes, which lack double, triple or aromatic bonds; and (3) unsaturated hydrocarbons, which have one or more double or triple bonds between carbon atoms and include, for example, alkenes (e.g., dienes), and alkynes.

The term "identical" (or "identity"), in the context of two polynucleotide or polypeptide sequences, means that the residues in the two sequences are the same when aligned for maximum correspondence, as measured using a sequence comparison or analysis algorithm such as those described herein. For example, if when properly aligned, the corresponding segments of two sequences have identical residues at 5 positions out of 10, it is said that the two sequences have a 50% identity. Most bioinformatic programs report percent identity over aligned sequence regions, which are typically not the entire molecules. If an alignment is long enough and contains enough identical residues, an expectation value can be calculated, which indicates that the level of identity in the alignment is unlikely to occur by random chance.

The term "insertion," when used in the context of a polypeptide sequence, refers to an insertion in the amino acid sequence of a precursor polypeptide, resulting in a mutant polypeptide having an amino acid that is inserted between two existing contiguous amino acids, i.e., adjacent amino acids residues, which are present in the precursor polypeptide. The term "insertion," when used in the context of a polynucleotide sequence, refers to an insertion of one or more nucleotides in the precursor polynucleotide between two existing contiguous nucleotides, i.e., adjacent nucleotides, which are present in the precursor polynucleotides.

The term "introduced" refers to, in the context of introducing a polynucleotide sequence into a cell, any method suitable for transferring the polynucleotide sequence into the cell. Such methods for introduction include but are not limited to protoplast fusion, transfection, transformation, conjugation, and transduction (see, e.g., Ferrari et al., *Genetics*, in Hardwood et al, (eds.), *Bacillus*, Plenum Publishing Corp., pp. 57-72, 1989).

The term "isolated" or "purified" means a material that is removed from its original environment, for example, the natural environment if it is naturally occurring, or a culti-

vation broth if it is produced in a recombinant host cell cultivation medium. A material is said to be “purified” when it is present in a particular composition in a higher concentration than the concentration that exists prior to the purification step(s). For example, with respect to a composition normally found in a naturally-occurring or wild type organism, such a composition is “purified” when the final composition does not include some material from the original matrix. As another example, where a composition is found in combination with other components in a recombinant host cell cultivation medium, that composition is purified when the cultivation medium is treated in a way to remove some component of the cultivation, for example, cell debris or other cultivation products, through, for example, centrifugation or distillation. As another example, a naturally-occurring polynucleotide or polypeptide present in a living animal is not isolated, but the same polynucleotide or polypeptide, separated from some or all of the coexisting materials in the natural system, is isolated, whether such process is through genetic engineering or mechanical separation. Such polynucleotides can be parts of vectors. Alternatively, such polynucleotides or polypeptides can be parts of compositions. Such polynucleotides or polypeptides can be considered “isolated” because the vectors or compositions comprising thereof are not part of their natural environments. In another example, a polynucleotide or protein is said to be purified if it gives rise to essentially one band in an electrophoretic gel or a blot.

The term “mutant thioesterase” or “variant thioesterase” refers to a thioesterase that comprises a mutation with reference to a precursor thioesterase.

The term “mutation” refers to, in the context of a polynucleotide, a modification to the polynucleotide sequence resulting in a change in the sequence of a polynucleotide with reference to a precursor polynucleotide sequence. A mutant polynucleotide sequence can refer to an alteration that does not change the encoded amino acid sequence, for example, with regard to codon optimization for expression purposes, or that modifies a codon in such a way as to result in a modification of the encoded amino acid sequence. Mutations can be introduced into a polynucleotide through any number of methods known to those of ordinary skill in the art, including random mutagenesis, site-specific mutagenesis, oligonucleotide directed mutagenesis, gene shuffling, directed evolution techniques, combinatorial mutagenesis, site saturation mutagenesis among others.

“Mutation” or “mutated” means, in the context of a protein, a modification to the amino acid sequence resulting in a change in the sequence of a protein with reference to a precursor protein sequence. A mutation can refer to a substitution of one amino acid with another amino acid, an insertion or a deletion of one or more amino acid residues. Specifically, a mutation can also be the replacement of an amino acid with a non-natural amino acid, or with a chemically-modified amino acid or like residues. A mutation can also be a truncation (e.g., a deletion or interruption) in a sequence or a subsequence from the precursor sequence. A mutation may also be an addition of a subsequence (e.g., two or more amino acids in a stretch, which are inserted between two contiguous amino acids in a precursor protein sequence) within a protein, or at either terminal end of a protein, thereby increasing the length of (or elongating) the protein. A mutation can be made by modifying the DNA sequence corresponding to the precursor protein. Mutations can be introduced into a protein sequence by known methods in the art, for example, by creating synthetic DNA sequences that encode the mutation with reference to precursor proteins, or

chemically altering the protein itself. A “mutant” as used herein is a protein comprising a mutation. For example, it is also possible to make a mutant by replacing a portion of a thioesterase with a wild type sequence that corresponds to such portion but includes a desired variation at a specific position that is naturally-occurring in the wild type sequence.

A “naturally-occurring equivalent,” in the context of the present invention, refers to a naturally-occurring thioesterase, or a portion thereof that comprises a naturally-occurring residue.

The term “operably linked,” in the context of a polynucleotide sequence, refers to the placement of one polynucleotide sequence into a functional relationship with another polynucleotide sequence. For example, a DNA encoding a secretory leader (e.g., a signal peptide) is operably linked to a DNA encoding a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide. A promoter or an enhancer is operably linked to a coding sequence if it affects the transcription of the sequence. A ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, “operably linked” means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in the same reading frame.

The term “optimal alignment” refers to the alignment giving the highest overall alignment score.

“Overexpressed” or “overexpression” in a host cell occurs if the enzyme is expressed in the cell at a higher level than the level at which it is expressed in a corresponding wild-type cell.

The terms “percent sequence identity,” “percent amino acid sequence identity,” “percent gene sequence identity,” and/or “percent polynucleotide sequence identity,” with respect to two polypeptides, polynucleotides and/or gene sequences (as appropriate), refer to the percentage of residues that are identical in the two sequences when the sequences are optimally aligned. Thus, 80% amino acid sequence identity means that 80% of the amino acids in two optimally aligned polypeptide sequences are identical. The percent identities expressed herein with respect to a given named reference sequence are determined over the entire reference sequence, rather than only a portion thereof. Thus, an amino acid sequence at least about 80% identical to positions 28-317 of SEQ ID NO:4, for example, is at least about 80% identical to the entire sequence of positions 28-317 of SEQ ID NO:4, as opposed merely to subsequences thereof.

The term “plasmid” refers to a circular double-stranded (ds) DNA construct used as a cloning vector, and which forms an extrachromosomal self-replicating genetic element in some eukaryotes or prokaryotes, or integrates into the host chromosome.

The term “precursor thioesterase” refers a thioesterase protein from which the mutant thioesterase of the invention can be derived, through, for example, recombinant or chemical means. Examples of precursor thioesterases are naturally-occurring or wildtype thioesterases from plant, animal or microbial sources. A precursor thioesterase can also be a thioesterase that is non-naturally-occurring. An example of a non-naturally-occurring thioesterase is a thioesterase made through, for example, random mutation, chemical synthesis, molecular evolution, or site directed mutagenesis, which can serve as a useful starting point from which to design and/or make the mutant thioesterases of the invention.

A “production host” is a cell used to produce products. As disclosed herein, a production host is modified to express or

overexpress selected genes, or to have attenuated expression of selected genes. Non-limiting examples of production hosts include plant, animal, human, bacteria, yeast, cyanobacteria, algae, and/or filamentous fungi cells.

A “promoter” is a polynucleotide sequence that functions to direct transcription of a downstream gene. In preferred embodiments, the promoter is appropriate to the host cell in which the target gene is being expressed. The promoter, together with other transcriptional and translational regulatory polynucleotide sequences (also termed “control sequences”) is necessary to express a given gene. In general, the transcriptional and translational regulatory sequences include, but are not limited to, promoter sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences.

The terms “protein” and “polypeptide” are used interchangeably herein. The 3-letter code as well as the 1-letter code for amino acid residues as defined in conformity with the IUPAC-IUB Joint Commission on Biochemical Nomenclature (JCBN) is used throughout this disclosure. It is also understood that a polypeptide may be coded for by more than one polynucleotide sequence due to the degeneracy of the genetic code. An enzyme is a protein.

The term “recombinant,” when used to modify the term “cell” or “vector” herein, refers to a cell or a vector that has been modified by the introduction of a heterologous polynucleotide sequence, or that the cell is derived from a cell so modified. Thus, for example, recombinant cells express genes that are not found in identical form within the native (non-recombinant) form of the cells or express, as a result of deliberate human intervention, native genes that are otherwise abnormally expressed, underexpressed or not expressed at all. The terms “recombination,” “recombining,” and generating a “recombined” polynucleotide refer generally to the assembly of two or more polynucleotide fragments wherein the assembly gives rise to a chimeric polynucleotide made from the assembled parts.

The terms “regulatory segment,” “regulatory sequence,” or “expression control sequence” refer to a polynucleotide sequence that is operatively linked with another polynucleotide sequence that encodes the amino acid sequence of a polypeptide chain to effect the expression of that encoded amino acid sequence. The regulatory sequence can inhibit, repress, promote, or even drive the expression of the operably-linked polynucleotide sequence encoding the amino acid sequence.

The term “selectable marker” or “selective marker” refers to a polynucleotide (e.g., a gene) capable of expression in a host cell, which allows for ease of selection of those hosts containing the vector. Examples of selectable markers include but are not limited to antimicrobial markers. Thus, the term “selectable marker” refers to a gene that provides an indication when a host cell has taken up an incoming sequence of interest or when some other reaction has taken place. Typically, selectable markers are genes that confer antimicrobial resistance or a metabolic advantage on the host cells to allow the cells containing the exogenous sequences to be distinguished from the cells that have not received the exogenous sequences. A “residing selectable marker” is one that is located on the chromosome of the microorganism to be transformed. A residing selectable marker encodes a gene that is different from the selectable marker on the transforming construct. Selective markers are known to those of skill in the art. As indicated above, suitably the marker is an antimicrobial resistant marker, including, for example, *amp^R*; *phleo^R*; *spec^R*; *kan^R*; *ery^R*;

tet^R; *cmp^R*; and *neo^R*. See, e.g., Guerot-Fleury, *Gene*, 167: 335-337, 1995; Palmeros et al., *Gene*, 247:255-264, 2000; and Trieu-Cuot et al., *Gene*, 23:331-341, 1983. Other markers useful in accordance with the invention include, but are not limited to, auxotrophic markers, such as tryptophan; and detection markers, such as 6-galactosidase.

The term “selectable marker-encoding nucleotide sequence” refers to a polynucleotide sequence that is capable of expression in the host cells and where the expression of the selectable marker confers to the cells containing the expressed gene the ability to grow in the presence of a corresponding selective agent or in the absence of one or more essential nutrients.

The term “substantially identical,” in the context of two polynucleotides or two polypeptides refers to a polynucleotide or polypeptide that comprises at least 70% sequence identity, for example, at least 75%, at least 80%, at least 85%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity as compared to a reference sequence using the programs or algorithms (e.g., BLAST, ALIGN, CLUSTAL) using standard parameters.

“Substantially purified” means molecules that are at least about 60% free, preferably at least about 75% free, about 80% free, about 85% free, and more preferably at least about 90% free from other components with which they are naturally associated. As used herein, the term “purified” or “to purify” also refers to the removal of contaminants from a sample.

“Substitution” means replacing an amino acid in the sequence of a precursor protein with another amino acid at a particular position, resulting in a mutant of the precursor protein. The amino acid used as a substitute can be a naturally-occurring amino acid, or can be a synthetic or non-naturally-occurring amino acid.

The term “thioesterase” refers to an enzyme that has thioesterase activity. Thioesterases include thioester hydrolases, which are identified as members of Enzyme Classification E.C. 3.1.2.x and are obtainable from a variety of sources.

The term “thioesterase activity” refers to the capacity to catalyze a thioester cleavage reaction, which usually involves the hydrolysis of a thioester at a thiol group into an acid and a thiol, but can also include transesterification, wherein a thioester bond is cleaved and a new ester bond is formed. In general, an acyl-ACP thioesterase is capable of catalyzing the hydrolytic cleavage of fatty acyl-acyl carrier protein thioesters and/or fatty acyl-coenzyme A thioesters. Examples of enzymes having thioesterase activity include acetyl-CoA hydrolase, palmitoyl-CoA hydrolase, succinyl-CoA hydrolase, formyl-CoA hydrolase, acyl-CoA hydrolase, palmitoyl-protein thioesterase, and ubiquitin thioesterase. Thioesterase activity can be established by any of a number of assays described in U.S. Pat. No. 9,587,231, which is incorporated herein by reference.

The term “transformed” or “stably transformed” cell refers to a cell that has a non-native (heterologous) polynucleotide sequence integrated into its genome or as an episomal plasmid that is maintained for at least two generations.

“Vector” refers to a polynucleotide construct designed to introduce polynucleotides into one or more cell types. Vectors include cloning vectors, expression vectors, shuttle vectors, plasmids, cassettes and the like. In some embodiments, the polynucleotide construct comprises a polynucleotide sequence encoding a thioesterase (e.g., a precursor or a mature thioesterase) that is operably linked to a suitable

prosequence (e.g., a secretory pro-sequence) capable of effecting the expression of the polynucleotide or gene in a suitable host.

“Wild-type” means, in the context of gene or protein, a polynucleotide or protein sequence that occurs in nature. In some embodiments, the wild-type sequence refers to a sequence of interest that is a starting point for protein engineering.

The mutant thioesterases of the present invention herein can be used in place of the mutant thioesterases described in U.S. Pat. No. 9,587,231 for any embodiments described in U.S. Pat. No. 9,587,231.

The elements and method steps described herein can be used in any combination whether explicitly described or not.

All combinations of method steps as used herein can be performed in any order, unless otherwise specified or clearly implied to the contrary by the context in which the referenced combination is made.

As used herein, the singular forms “a,” “an,” and “the” include plural referents unless the content clearly dictates otherwise.

Numerical ranges as used herein are intended to include every number and subset of numbers contained within that range, whether specifically disclosed or not. Further, these numerical ranges should be construed as providing support for a claim directed to any number or subset of numbers in that range. For example, a disclosure of from 1 to 10 should be construed as supporting a range of from 2 to 8, from 3 to 7, from 5 to 6, from 1 to 9, from 3.6 to 4.6, from 3.5 to 9.9, and so forth.

All patents, patent publications, and peer-reviewed publications (i.e., “references”) cited herein are expressly incorporated by reference to the same extent as if each individual reference were specifically and individually indicated as being incorporated by reference. In case of conflict between the present disclosure and the incorporated references, the present disclosure controls.

It is understood that the invention is not confined to the particular construction and arrangement of parts herein illustrated and described, but embraces such modified forms thereof as come within the scope of the claims.

EXAMPLES

Summary

Medium-chain fatty acids (MCFA) are currently obtained from plants oils such as coconut and palm kernel oil or poorly selective chemical synthesis from fossil fuels. Consequently, strong demand for these molecules has contributed to the growth of oil-seed plantations and the deforestation of tropical habitats. Microbial conversion of renewable feedstocks to MCFA is one potential alternative to current practices. However, one of the challenges of microbial production of MCFA is the lack of enzymes that are both highly active and selective towards medium chain-length substrates. As a result, most microbial biocatalysts are either able to produce high titers of MCFA with mixed chain-lengths or low titers of products with a narrow chain-length distribution. One of the few enzymes involved in oleochemical metabolism possessing strong selectivity is the acyl-ACP thioesterase. This enzyme catalyzes the last step in microbial MCFA production strategies by hydrolyzing the thioester bond linking an acyl-chain to the acyl-carrier protein (ACP). In search of highly active and selective enzymes capable of producing octanoic acid, we developed a selection platform that relies on the lipolic acid requirement

of *E. coli*. This selection was used to find improved mutants in a library of randomly mutagenized gene variants derived from the C8 specific *Cuphea palustris* FatB1 thioesterase. Using this selection, we isolated a thioesterase that produced 1.7 g/L of octanoic acid with >90% specificity. In addition, we were able to show that a single chromosomal copy of this thioesterase was sufficient to achieve the titers mentioned above, a feat that is crucial when building industrially relevant strains. In vitro studies confirmed the mutant thioesterase possessed a large increase in kcat compared to its native counterpart.

Introduction

Oleochemicals are a large class of industrial chemicals used for making products in the bioenergy, plastics, surfactants, and personal care sectors. Oleochemicals include molecules such as free fatty acids (FFA), fatty acid methyl esters (FAME), fatty alcohols, and organosulfates (e.g. sodium dodecyl sulfate)^{1,2}. While the most desirable oleochemicals contain medium chains (C₆-C₁₂), most natural oleochemical sources are dominated by long acyl chains (>C₁₆). Of the major oil seed crops, only coconut and palm have large fractions of medium chain fatty acids (MCFA), with C₁₂ being the most abundant chain length (FIG. 1). Some plant oils contain high proportions of medium-chain acids, e.g. *Umbellularia californica* (California Bay laurel) contains high proportion of C₁₂ fatty acids³ and *Cuphea* species often contain high percentages of C₈ fatty acids⁴, but most are not cultivated in volumes capable of meeting oleochemical demand. The low availability of MCFA has motivated the development of microbial conversion strategies where renewable sugars are converted to specific oleochemicals in the medium chain category⁵. In this approach, metabolic engineering principles are used to redirect carbon flux through fatty acid biosynthesis to specific products^{6,7} leveraging heterologous enzymes capable of catalyzing desired biochemical transformations.

While many oleochemicals have been produced in microbes, there remains a dearth of enzymes capable of directing flux to products containing a specific chain length. The notable exception is the thioesterase⁸ which cleaves acyl-thioesters (CoA or acyl-carrier protein, ACP) to release FFA from biosynthetic pathways. Thioesterases are expressed in many organisms for various purposes. Microbial thioesterases often have proofreading roles in the cell and therefore act on a broad substrate range. In some cases, thioesterase selectivity can be tailored via protein engineering⁹, but complete selectivity remains an unmet challenge. In contrast, many plant thioesterases act on a narrow set of substrates. Plants synthesize fatty acids in the chloroplast and lipids in the cytosol¹². In order to transport acyl-chains across the chloroplast membrane, plants express thioesterases to release FFA in the chloroplast and reactivate them as acyl-CoA thioesters in the cytosol. Therefore, the substrate specificity of thioesterases often dictates the composition of plant oils. For this reason, plants have become a preferred source for isolating thioesterases with desired substrate preference. These enzymes can then be used in either transgenic crops or microbes to produce oleochemicals with desired chain lengths⁷. This approach is often made difficult in *Escherichia coli* by a loss of activity when plant thioesterases are heterologously expressed. Therefore, researchers remain motivated to isolate, evolve, and/or engineer improved thioesterases with desired selectivity and activity.

One challenge to thioesterase engineering is the lack of good screening methods to differentiate products with different chain lengths. The analysis of fatty acid chain length typically uses gas chromatography to separate fatty acid methyl esters (FAME) derived from biological samples. While accurate, this method requires considerable sample preparation time and instrument time that limits the number of samples that can be processed to less than a 200 per day per instrument. In protein engineering projects, a library size typically ranges from 10^3 to 10^8 samples. Therefore, gas chromatography is not an applicable method for screening large libraries for increased activity. Without high throughput screens, mutagenesis is limited to rational design. Although rational design of thioesterases has given some degree of success^{9,17}, there is still considerable room for improving these enzymes. One alternative to achieving this goal is the development of biosensors that use screening (change in observable phenotype) or selection (live/dead) as a way to differentiate improved enzymes from the rest. Biosensors for detecting fatty acids and other aliphatic molecules have been developed by others using transcriptional regulators linked to fluorescent proteins¹⁸ and G-protein coupled receptors¹⁹. However, these approaches have limited ability to tailor chain-length specificity. Here, we developed a genetic selection for acyl-chains containing exactly eight carbons using the lipoyl requirement of *E. coli* under aerobic conditions.

Lipoic acid is an essential vitamin in most of organisms. It is an important cofactor for function of several key enzymes involved in aerobic metabolism, such as pyruvate dehydrogenase, 2-oxoglutarate dehydrogenase, the glycine cleavage system, and the branched-chain 2-oxoacid dehydrogenase²⁰. Pyruvate dehydrogenase contains a lipoyl group in its E2 domain that translocates an activated acetyl moiety to the thiol of coenzyme A to form acetyl-CoA. This lipoyl group synthesis proceeds via one of two pathways in *E. coli*. The endogenous biosynthesis pathway branches from a central intermediate in fatty acid biosynthesis octanoyl-ACP (FIG. 2) which donates its octanoyl group to the E2-domain of pyruvate dehydrogenase via the enzyme LipB (octanoyl-ACP N-lipoyltransferase). Next, LipA, an iron-sulfur cluster enzyme, catalyzes a radical reaction to install two sulfur atoms at C₆ and C₈ positions of the octanoyl group to complete this biosynthesis. *E. coli* Δ lipB mutants are unable to grow on minimal media. However, *E. coli* has a salvage pathway to obtain and assimilate free lipoic acid from the environment. LplA, a lipoate-protein ligase, is an ATP-dependent enzyme that activates lipoic acid as well as octanoic acid and transfers it to the E2-domain of pyruvate dehydrogenase. As seen in FIG. 2, growth of *E. coli* Δ lipB mutants can be rescued by supplying lipoic acid as well as octanoic acid in the media²¹. The authors proposed that LplA could transfer octanoic acid to the E2 domain of pyruvate dehydrogenase and rejoin the native lipoic acid biosynthesis pathway where a LipA-catalyzed reaction completes the pathway. Conversely, other chain-length carboxylic acids were not able to rescue growth of the mutant (not shown). Based on these intriguing results, we hypothesized that an *E. coli* Δ lipB strain could be leveraged as a growth-based biosensor for octanoic acid presence.

In these examples, we describe how we used this novel screening approach to select for improved variants from a randomly mutagenized library of *Cuphea palustris* FatB1 thioesterase (CpFatB1) genes²². The best variants led to 5-7 fold improvements in octanoic acid titer while sustaining the enzyme's high selectivity towards 8-carbon chains. The best variant CpFatB1-M4 demonstrated a 15-fold increased k_{car}

The increased specific activity enabled us to place the gene on the chromosome in a single copy and achieve the same octanoic acid titer achieved by plasmid containing strains.

MATERIALS AND METHODS

Chemicals, Reagents, and Media

Chemicals were purchased from either Sigma Aldrich (St. Louis, Mo.) or Fisher Scientific (Waltham, Mass.). Oligonucleotides and gene fragments were purchased from Integrated DNA Technologies (Coralville, Iowa) or Thermo Fisher Scientific (Waltham, Mass.). Enzymes were purchased from New England Biolabs (Ipswich, Mass.). DNA purification kits were purchased from Qiagen (Venlo, Netherlands). All cultures were started from single colonies grown on LB agar isolated from freezer stocks stored in 15% glycerol. Overnight cultures of strains were grown in LB media at 30° C. in a rotary shaker at 250 r.p.m. When a selective pressure was necessary to for plasmid retention, media was supplemented with the appropriate antibiotics (carbenicillin, 100 μ g/mL; kanamycin, 50 μ g/mL; chloramphenicol, 34 μ g/mL).

DNA Synthesis and Cloning

Escherichia coli K12 MG1655 was used to create the Δ lipB selection strain. Here we used a CRISPR-Cas9 assisted homologous recombination protocol, modified from Li et al.,³² to delete the lipB coding sequence. Standard lambda red recombination³³ was used to introduce the deletion and Cas9 guided to lipB was used to destroy unmodified chromosomes. The repair template contained 30 bases upstream and downstream of the lagging strand of lipB.

NHL17 (*E. coli* K12 MG1655 Δ araBAD Δ fadD::trc-CpFatB1.2-M4-287) strain was created from *E. coli* K12 MG1655 Δ araBAD in the same manner by using a linear piece of dsDNA containing lacI-trc-CpFatB1.2-M4-287 between 500 base pairs of homology upstream and downstream of fadD coding sequence.

All plasmids made were constructed using Gibson Assembly of PCR products³⁴.

Random Mutagenesis of CpFatB1.2

The CpFatB1.2 library was constructed by error-prone PCR following the manufacturer's instruction (GeneMorph II, Agilent). The mutation frequency chosen was low (0 to 4.5 mutations/kb). The plasmid backbone was amplified by PCR using high fidelity polymerase Phusion. CpFatB1.2 library was assembled with designated backbones by Gibson assembly method. Primers used in for the creation of the library contained the start and stop codons in order to prevent mutations on them.

Lipoic Acid Selection

In order to find suitable conditions for the Δ lipB-based selection method, purified plasmid pBTRCK-CpFatB1.2 was transformed into *E. coli* Δ lipB strain and plated in MOPS minimal media agarose plates containing 0.2% glucose. In addition, the plates contained kanamycin to maintain the plasmid and different IPTG concentrations (0 μ M, 10 μ M, 20 μ M, 30 μ M and 50 μ M IPTG) to titrate the amount of CpFatB1.2 present in the cells. The 20 μ M induction condition was chosen because at this level of CpFatB1.2

expression, the cells needed an extra day (4 days) for rescuing growth compared to higher induction levels (3 days).

Gibson assembly reaction mixtures (2 μ L) containing a CpFatB1.2 library was transformed into 100 μ L of electro-competent *E. coli* Δ lipB. Following electroporation (1 mm cuvette, 2500 mV), 900 μ L of fresh LB was added and cells were allowed to recover for 1.5 hours. In order to remove any remaining lipoic acid from the rich media the cells were washed 3 times by spinning down at 10,000 rpm for 1 min followed by the 1 mL of M9 minimal media was added and the cells were resuspended. Finally, the washed cells were plated on MOPS minimal media-agarose plates containing 0.2% glucose and Kanamycin. The plates were incubated at 30° C. and after 3 days, the chosen 90 putative mutants growing colonies were streaked on LB plates with Kanamycin. The plates were incubated at 37° C. overnight to confirm the colony growth. Finally, the plasmids were purified and sequenced.

Fatty Acid Production

Plasmid based expression of thioesterases was performed in *E. coli* RL08ara³⁵ transformed with the appropriate plasmid. NHL17 strain contains a chromosomal copy of CpFatB1.2-M4-287 and therefore no plasmid was added.

For validating the 90 CpFatB1.2 variants (FIG. 7B, FIG. 8B, Table 2) as well as the truncations of CpFatB1 (FIG. 3B) and truncations in CpFatB1.2-M4 (FIG. 10B), isolated plasmids were transformed into *E. coli* RL08ara and single colonies were used to inoculate overnight cultures in LB media with kanamycin. Overnight cultures were used to inoculate 50 mL LB media with 0.4% glycerol and kanamycin at an initial optical density (OD₆₀₀) of 0.05. Cultures were incubated at 37° C. with shaking in 250 mL shake flasks. When cultures reached an OD₆₀₀ of 0.2-0.3 cells were induced with IPTG and moved to 30° C. for 24 hr. Mutants M1-M10 were tested using 20 μ M IPTG (FIG. 7B). Mutants M3, M4, and M11-M90 (FIG. 8B), truncations of CpFatB1 (FIG. 3B) and truncations in CpFatB1.2-M4 (FIG. 10B) where induced with 1 mM IPTG.

For experiments designed to test for high octanoic acid production (FIGS. 9, 11, 14A, and 14B) overnight cultures as described above were used to inoculate 50 mL of medium described in Kim, et al.²⁵ with the following changes: 1.39 mM Na₂HPO₄, no biotin, thiamine or sodium selenite added.

Minimal media experiments were carried out in MOPS minimal media²⁴ containing 1% glucose and 0.240 mM K₂HPO₄ in order to create phosphate limiting conditions³⁶.

Fatty Acid Extraction and Quantification

After 24 h post-induction, 2.5 mL of culture was transferred to 10 mL glass centrifuge tubes. 50 μ L of 12.5 mg/mL nonanoic acid, and 1.25 mg/ml pentadecanoic acid in ethanol solution was added as an internal standard. The nonanoic acid internal standard was used to quantify octanoic acid and the pentadecanoic acid internal standard was used to quantify C₁₀-C₁₈ chain lengths. Extraction and methylation process followed protocols described previously⁹.

Protein Expression and Purification of Apo-Acyl Carrier Protein (ACP)

E. coli K12 MG1655 acyl carrier protein (ACP) was cloned into the pET28t vector system fused to a N-terminal

polyhistidine tag coding the following peptide: MGSSHHHHHHSSENLYFQGGGG. The plasmid was transformed into BL21 (DE3) competent cells and grown LB media at 37° C. until OD₆₀₀ was 0.6-0.8. Cells were cooled to 18° C. in ice water, induced with 1 mM IPTG and incubated overnight at 18° C. with shaking. Cells were harvested by centrifugation at 8,000 \times g and pellets were stored at -80° C. for later use. Frozen pellets were resuspended in lysis buffer (50 mM Na₂HPO₄ pH8, 20 mM imidazole, 300 mM NaCl and 10% glycerol), sonicated, centrifuged at 12,000 RPM and filtered to clear the lysate. ACP was purified by Ni-NTA column following the manufacture's instruction (GE Healthcare Life Sciences). To the ACP protein solution, Tev protease was added at a molar ratio of 1:20 and dialysed against 50 mM Tris, pH 7.5 overnight. Cleaved ACP was then passed through the Ni-NTA column to remove Tev protease and the His tag peptide. The flow through was dialyzed against 50 mM Na₂HPO₄ pH8, 10% Glycerol for subsequent functionalization. The concentration of ACP was quantified via BCA assay (Thermo Fisher) using manufacturer's instructions.

Protein Expression and Purification of *Vibrio Harveyi* AasS and *Basillus sustilis* Sfp, CpFatB1.2 and CpFatB1.2-M4

Vibrio Harveyi AasS, *Basillus sustilis* Sfp, CpFatB1.2 and CpFatB1.2-M4 were cloned into pET28t vector system with an N-terminal poly-histidine tag as described for ACP. Proteins were purified as described for ACP with the exception that no Tev protease reaction was performed. Following purification in Ni-NTA column, proteins were concentrated and buffer-exchanged into 50 mM Na₂HPO₄ pH8, 30% Glycerol. Concentration of these proteins was quantified using the following extinction coefficients (280 nm): 67520 M⁻¹ cm⁻¹ for AasS, 30620 M⁻¹ cm⁻¹ Sfp, 56295 M⁻¹ cm⁻¹ for CpFatB1.2, and 50795 M⁻¹ cm⁻¹ for CpFatB1.2-M4.

Synthesis of Octanoyl-ACP

Octanoyl ACP synthesis was carried out by first functionalizing a 500 μ M mixture of apo-ACP and holo-ACP from *E. coli* into holo-ACP by incubating at 37° C. for 1 hr with 5 μ M purified Sfp, 10 mM MgCl₂, 5 mM Coenzyme A in 100 mM Na₂HPO₄ pH8 as has been described elsewhere²⁶. Next, 5 μ M of purified AssS, 10 mM ATP and 5 mM sodium octanoate are added to the reaction mixture and incubated overnight at 37° C. Samples were taken in between steps for characterization by HPLC. After incubation, octanoyl-ACP was passed through a Ni-NTA column to remove both AasS and Sfp followed by addition of an equimolar amount of 5,5'-dithiobis(2-nitrobenzoic acid) (DTNB) in order to react all the CoA remaining prior to the assays. DTNB and yellow TNB produced in this step were subsequently dialyzed out against 100 mM Na₂HPO₄, pH8, 10% glycerol before carrying out the enzymatic assays. Octanoyl-ACP concentration was quantified using BCA assay.

Liquid Chromatography of Octanoyl-ACP

To verify the functionalization and purity of Acyl carrier protein species, samples were separated via HPLC using a Harmony C4 column 2.1 \times 150 mm, 3.5 μ m (ES Industries). Mobile phases consisted of (1) aqueous solution of 0.05% (w/v) Trifluoroacetic acid and 0.05% (w/v) formic acid, and (2) 0.05% (w/v) Trifluoroacetic acid and 0.05% (w/v) formic

acid in acetonitrile. The samples were separated over 20 min by imposing a gradient of 20% aqueous mobile phase to 98% acetonitrile mobile phase. The oven temperature was kept at 30° C. and the flow rate was 0.2 ml/min with an injection volume of 10 μ L. Prior to injection, the samples were buffer exchanged into 50 mM ammonium acetate and treated with 0.1% (w/v) formic acid

In-vitro Analysis of CpFatB1.2-M4

Octanoyl-ACP thioesterase activity of CpFatB1.2, CpFatB1.2-M4 and TesA-R3.M4⁹ was analyzed in vitro by tracking the formation of holo-ACP using the thiol-dependant reduction of 5,5'-dithiobis(2-nitrobenzoic acid) (DTNB). TNB formation was monitored every 10 s for 2 min at Absorbance at 412 nm with a NanoDrop 2000c (Thermo Scientific) at a path length of 10 mm. Octanoyl-ACP was added to the assay in concentrations ranging 0-400 μ M (quantified via BCA assay). The conditions for the assay were as follows: 40 nM thioesterase, 8 μ g/mL BSA, 250 μ M DTNB, 100 mM phosphate buffer pH7.4, in 1 mL reaction volume. Assay was started with the addition of the thioesterase. All concentrations except 400 μ M were tested in triplicate.

Structural Modeling of CpFatB1.2-M3 and CpFatB1.2-M4-287

The CpFatB1.2 model was created using homology modeling of the CpFatB1.2 sequence and BTE (PDB: 5x04structure) as the template structure. Subsequently, the amino acid changes to create energy-minimized structures of the CpFatB1.2 mutants were made using Mutator.³⁷ The catalytic residues Asp220, Asn222, His224, Glu258, and Cys259 were identified using the *Umbellularia californica* thioesterase UaFatB1 (BTE) structure (PDB: 5x04) as a guide where the analogous residues have been reported to be Asp281, Asn283, His285, Glu319, and Cys320. 11 Cys320 mutants were seen to retain non-negligible catalytic activities, hence Cys259 was excluded from the list of catalytic residues in CpFatB1.2 model. The octanoyl-ACP (substrate) was docked such that the carbonyl carbon (C=O) of the thioester bond of the acyl-ACP molecule was close to the side-chain O atoms of Asp220 and Glu258. The catalytic distances corresponding to Asp220 and Glu258 were measured to be 3.5 and 3.7 Å, respectively. Subsequently, the BTE structure was used to identify the acyl-binding pocket residues which are important for controlling substrate specificity.²⁹ In order to understand the biophysical mechanism that underpins the catalytic activity in each of the enzyme variants, noncovalent contact maps were constructed similar

to Mendonça et al.³⁰ These contact networks (see FIGS. 17A and 18A) have been used to explain the possible path by which mutations away from the binding crevice or active site could affect enzyme activity. Python 2.7 scripts were written to identify heteroatoms of residues within 6 Å of an altered residue of the CpFatB1.2 mutant. Each of these residues form nodes of the contact map and edges are drawn to show that a noncovalent contact exists. The same procedure is repeated for the nodes identified in the previous step and a cascade of interactions is mapped. The process is terminated when one or more catalytic and acyl-binding pocket residues are identified within 6 Å of a residue identified in previous step. Subsequently, we parse the obtained information to classify the noncovalent contacts (edges) as hydrophobic or polar, depending on the nature of atoms involved in the interaction. Finally, we visualize the contacts using PyMOL visualizing software.

RESULTS

Establishment of a Baseline Thioesterase

Cupheapalustris FatB1 thioesterase (CpFatB1) is highly selective for C8:0-ACP when expressed in *E. coli*²³ albeit with lower activity relative to *E. coli* 'TesA and other commonly used thioesterases^{6,8}. The lower activity likely comes from a combination of poor expression and/or poor specific activity. Plant thioesterases are often associated with the chloroplast membrane and native genes contain membrane localization sequences. When heterologously expressed, these sequences can lead to insoluble or aggregated proteins. Therefore, one must construct an N-terminal truncation of a plant thioesterase to obtain high levels of soluble protein. We constructed three N-terminal truncations of CpFatB1 (SEQ ID NO:1 (nucleotide sequence) and SEQ ID NO:2 (protein sequence)) based on prior work¹⁵ and sequence alignment—CpFatB1.2 (SEQ ID NO:3 (nucleotide sequence) and SEQ ID NO:4 (protein sequence)), CpFatB1.3 (SEQ ID NO:5 (nucleotide sequence) and SEQ ID NO:6 (protein sequence)), and CpFatB1.4 (SEQ ID NO:7 (nucleotide sequence) and SEQ ID NO:8 (protein sequence)). Each was cloned into a high copy plasmid, pTRC99a (FIGS. 3A and 3B, Table 1) and transformed into *E. coli* RL08ara (Δ fadD) for testing. Each strain was grown in LB supplemented with 0.4% glycerol and 1 mM IPTG. Expression of CpFatB1.2 generated the highest titer of octanoic acid under these conditions. Therefore, we used this gene sequence as a starting point in our mutagenesis studies.

TABLE 1

Strains and plasmids used in the present examples.		
Strain/plasmid	Genotype	Source
<i>E. coli</i> K12 MG1655	F ⁻ λ ⁻ ilvG ⁻ rfb-50 rph-1	CGSG
Δ lipB	K12 MG1655 Δ lipB	This work
RL08ara	K-12 MG1655 Δ araBAD Δ fadD	13
araBAD	K-12 MG1655 Δ araBAD	38
NHL17	K-12 MG1655 Δ araBAD Δ fadD::trcCpFatB1.2-M4-287	This work
pBTRCK	ptrc promoter, pBBR1 origin, Kan ^R	39
ptrc99a	ptrc promoter, pBR322 origin, Amp ^R	40
pBad33	pBAD promoter, pACYC origin, Cm ^R	41
pBad33-CpFatB1.2	pBad33 with <i>Cuphea palustris</i> CpFatB1 gene truncated at MLLTAIT	This work
ptrc99a-CpFatB1	ptrc99a with <i>Cuphea palustris</i> CpFatB1 full gene sequence	This work

TABLE 1-continued

Strains and plasmids used in the present examples.		
Strain/plasmid	Genotype	Source
ptrc99a-CpFatB1.2	ptrc99a with <i>Cuphea palustris</i> CpFatB1 gene truncated at MLLTAIT	This work
ptrc99a-CpFatB1.3	ptrc99a with <i>Cuphea palustris</i> CpFatB1 gene truncated at MKSKRPN	This work
ptrc99a-CpFatB1.4	ptrc99a with <i>Cuphea palustris</i> CpFatB1 gene truncated at MGLVFRQ	This work
pBTRCK-CpFatB1.2	pBTRCK with <i>Cuphea palustris</i> CpFatB1.2 under trc promoter	This work
pBTRCK-CpFatB1.2-M4	pBTRCK with CpFatB1.2-M4 sequence	
pBTRCK-CpFatB1.2-M4-287	pBTRCK with CpFatB1.2-M4 sequence truncated	
pBTRCK-CpFatB1.2-M4-288	pBTRCK	
pBTRCK-CpFatB1.2-M4-289	pBTRCK	
pBTRCK-CpFatB1.2-M4-290	pBTRCK	
pBTRCK-CpFatB1.2-M4-291	pBTRCK	
ptrc99a-CpFatB1.2-M3		
ptrc99a-CpFatB1.2-M4		
ptrc99a-CpFatB1.2-M9		
ptrc99a-CpFatB1.2-M4-287		

Development of a Lipic Acid-Based Selection

As discussed above, *E. coli* requires small amounts of lipic acid to enable pyruvate decarboxylase activity under aerobic conditions. As little as 50 μM (7.2 mg/L) octanoic acid can restore growth of an *E. coli* ΔlipB strain²¹. This amount is less than the ~ 200 mg/L of octanoic acid produced from the plasmid-based CpFatB1.2 described above. Therefore, to use the lipic acid requirement as a selection, the overall activity of the thioesterase must be reduced, such that the baseline enzyme cannot complement a ΔlipB mutation. To do this we reduced the expression of CpFatB1.2 by swapping the promoter for a weaker P_{araBAD} and moved the expression cassette to a plasmid maintained at a lower copy number (pACYC origin). Unfortunately, *E. coli* ΔlipB pBAD33-CpFatB1.2 grew on MOPS-minimal media-agar²⁴ containing 0.2% arabinose to induce expression (FIG. 4). In a second attempt to reduce activity, we cloned the original P_{TRC} -CpFatB1.2 expression cassette onto a low-copy plasmid, pBTRCK (pBBR1 origin), and used a series of low IPTG concentrations to vary expression (FIG. 5). Interestingly, cells grown on plates with 20 μM IPTG took an extra day to grow (4 days), compared to cells grown (3 days) on the same media with 30 μM IPTG (FIG. 5). Given this difference in growth rates, we hypothesized that we had found a window in which cells expressing thioesterase variants with improved specific activity or mutations that increased protein production would be identified before the cells carrying the parent gene became visible colonies. Therefore, we used the low-copy pBTRCK plasmid and 20 μM IPTG in our mutagenesis study.

Library of CpFatB1.2 Mutants

To introduce mutations, we generated a library of CpFatB1.2 variants by error-prone PCR covering the full coding sequence. PCR products were cloned into pBTRCK, plasmids were transformed into *E. coli* ΔlipB , and cells were plated on MOPS minimal media containing 20 μM IPTG. Ninety colonies appeared after three days (FIG. 6). To validate each hit, plasmids were isolated from each colony, retransformed into fresh *E. coli* ΔlipB cells, and cells were grown under selecting conditions. All of the variants rescued growth within three days (not shown), indicating that

growth-conferring mutations were plasmid-based. When cultured in liquid media, cells harboring the variant thioesterases produced more octanoic acid than cultures expressing the original CpFatB1.2 (FIG. 7B). In particular, variants CpFatB1.2-M3 (SEQ ID NO:9 (nucleotide sequence) and SEQ ID NO:10 (protein sequence)), CpFatB1.2-M4 (SEQ ID NO:11 (nucleotide sequence) and SEQ ID NO:12 (protein sequence)), and CpFatB1.2-M9 exhibited 4-fold, 5.3-fold and 2.3-fold improvements, respectively, when expressed from the low-copy plasmid. After analyzing the first 10 mutants, we repeated the protocol on the remainder of the 90 putative mutants, finding several additional improved variants, but none superior to M3 or M4 (FIG. 8B).

Plasmids isolated from each of the hits that generated more than a 2-fold increase in octanoic acid were sequenced. A small family of mutations was observed in these hits. Interestingly, one mutation, D293V appeared independently in five of the sequenced mutants, CpFatB1.2-M20, CpFatB1.2-M40, CpFatB1.2-M47, CpFatB1.2-M66, and CpFatB1.2-M73. Mutant CpFatB1.2-M40 and CpFatB1.2-M66 contained only the D293V mutation, indicating that it provided on average a 2.3-fold increase in activity over CpFatB1.2. In addition to two point mutations (N28S, I65M), CpFatB1.2-M4, the best variant, contained a frameshifting deletion which introduced a premature stop codon. CpFatB1.2-M3 contained two mutations (A59S and K296R) that were also found in other mutants. Given the superior performance of CpFatB1.2-M3, CpFatB1.2-M4, and CpFatB1.2-M9, we focused the remainder of the study on these variants.

Mutants CpFatB1.2-M3, CpFatB1.2-M4, and CpFatB1.2-M9 were subcloned into high copy plasmid pTRC99a to determine if the improvements found under screening conditions would be maintained under optimal production conditions. Plasmids were transformed into *E. coli* RL08ara (ΔfadD) and cells were grown in MOPS media enriched²⁵ with tryptone, yeast extract, and 1 mM IPTG to maximize induction. Cells expressing the M3 and M4 variants produced 1751 mg/L and 1263 mg/L of octanoic acid respectively. These titers represent a 3-4 fold-increase relative to cells expressing CpFatB1.2 which produced 375 mg/L (FIG. 9). Cells expressing the M9 variant produced approximately 500 mg/L, a smaller relative value to CpFatB1.2 than seen

under screening conditions. Conveniently, the increased activity did not come at the expense of octanoic acid selectivity. Variants M3 (92 mol %) and M4 (94 mol %) generated equivalent if not larger percentages of octanoic acid compared to CpFatB1.2 (89% mol). These data show that the lipoic acid selection was capable of identifying useful mutations.

Characterization of M4 Variant In Vivo

The CpFatB1.2-M4 (ΔA_{54} , N28S, I65M) variant contained two point mutations and an early nucleotide deletion (ΔA_{54}) that led the original open reading frame to an early stop codon (FIG. 10A). Since CpFatB1.2-M4 is active, we suspected that translation was restarting at a different in-frame methionine using a suboptimal ribosome binding site (RBS). If true, we hypothesized that the M4 variant could generate more activity, if expressed with an optimal RBS. To determine which protein was being made, we cloned five in-frame CpFatB1.2-M4 variants based on the next five in-frame methionines as start codons, creating CpFatB1.2-M4-287 (SEQ ID NO:13 (nucleotide sequence) and SEQ ID NO:14 (protein sequence)), -288, -289, -290 and -291. These genes were cloned into a pBTRCk vector to position the new start adjacent to the original, strong RBS. Plasmids harboring each variant were individually transformed into RL08ara (Δ fadD) strain and cells were grown in LB supplemented with 0.4% glycerol and 1 mM IPTG. Only variant CpFatB1.2-M4-287 showed a significant level of octanoic acid production (FIG. 10B). Moreover, variant CpFatB1.2-M4-287 generated more octanoic acid than the isolated variant CpFatB1.2-M4. This suggests that the hypothesis of CpFatB1.2-M4 being translated from a non-specific RBS was correct.

Further, the pFatB1.2-M4-287 demonstrated a ~20-fold increase in octanoic acid production under the low expression conditions tested (FIG. 10B), suggesting that more activity could be obtained if overexpressed. Therefore, we cloned CpFatB1.2-M4-287 onto a high copy plasmid, pTRC99a, transformed the plasmid into *E. coli* RL08ara (Δ fadD), and cultured cells in MOPS media enriched²⁵ with tryptone and yeast extract (see methods) and different concentrations of IPTG to optimize expression (FIG. 11). The resulting fatty acid profiles extracted from these cultures showed that a subsaturating concentration of 50 μ M IPTG gave maximum activity. At saturating concentrations of 1 mM IPTG, we observed a growth defect and a drastic decrease in C_{16} fatty acid species (FIG. 11). This data strongly suggested that we had dramatically increased the specific activity of CpFatB1, because it could no longer be maximally expressed.

Finally, we made combinations of the constitutive mutations found in CpFatB1.2-M4-287 (N28S, I65M, 287-truncation) to determine which contributed to enhanced activity. CpFatB1.2 variants containing 1, 2 or all three mutations were cloned into pTRC99A and cultured in *E. coli* RL08ara as described for FIG. 11. As a negative control, we cloned a variant with a H224A mutation that renders the enzyme catalytically inactive. Interestingly, only I65M was observed to increase activity above the baseline variant, CpFatB1.2 (FIG. 12). Only the triple mutant was able to drastically outperform the CpFatB1.2. This also suggests a possible

explanation for the CpFatB1.2-M4-288 truncation (which excludes the N28S mutation) being less active than the CpFatB1.2-M4 (FIG. 10B).

Characterization of M4 Variant In Vitro

Our in vivo data suggested that the M4 variant had increased specific activity towards C_8 -acyl ACPs. To prove this hypothesis, we measured the reaction rate in vitro using Ellman's reagent (DTNB) to monitor release of free thiols in holo-ACP (FIG. 13A). As hydrolysis of octanoyl-ACP occurs, holo-ACP formed reacts with DTNB forming the colored compound TNB that absorbs light at 412 nm. For substrate, we synthesized octanoyl-ACP in vitro from apo-ACP (purified from *E. coli*), coenzyme A (which donated the 4'-phosphopantetheine prosthetic group to convert apo-ACP to holo-ACP), and octanoate, using methods described elsewhere²⁶. Complete synthesis of octanoyl-ACP was confirmed by HPLC (FIG. 13B). Using the DTNB assay, we measured the initial rate of the thioesterase reaction for a range of substrate concentrations. We found that mutant CpFatB1.2-M4 has dramatically improved activity towards octanoyl-ACP compared to both CpFatB1.2 as well as a TesA variant (TesA-R3.M4) that we designed computationally in previous work⁹ to produce octanoic acid. The major contribution to CpFatB1.2-M4 improvement observed was due to a 15.7 fold increase in V_{max} over CpFatB1.2 while the K_m remained relatively low (FIGS. 13C and 13D). Interestingly, we found that our TesA-R3.M4 mutant had a high V_{max} as well but very low affinity for octanoyl-ACP with K_m 17-fold higher than CpFatB1.2-M4. These data confirm that our lipoic acid selection isolated a variant with improved specific activity.

Optimizing Expression in *E. Coli*

When building stable, industrially-relevant strains, it is beneficial to remove any requirement for antibiotics for maintaining plasmids and to reduce the cellular burden associated with protein overexpression. In other words, it is preferable to achieve a desired activity by increasing specific activity of essential enzymes such that each enzyme can be expressed at a modest level. Here, we wanted to test the ability of CpFatB1.2M4 to provide thioesterase activity when expressed from low copy plasmids or the chromosome. Therefore, we created the low copy plasmid, pBTRCK-CpFatB1.2-M4, with a n optimized RBS and tested FFA production under various induction levels (FIG. 14A). CpFatB1.2-M4 was optimally expressed from this construct when no IPTG was added, suggesting that the copy number could be decreased further. Moreover, at high induction we observed a growth defect, which reduced the final octanoic acid titers, a phenomenon that has been seen before¹³. Next, we took the same construct and inserted it into the *E. coli* chromosome in the fadD locus using CRISPR-Cas9 mediated homologous recombination. This yielded *E. coli* strain NHL17 (FIG. 14B, Table 1). As can be seen from FIG. 14B, a single copy CpFatB1.2-M4 in the chromosome when fully induced was sufficient to yield the levels of production obtained from the plasmid.

TABLE 2

Fatty acid profile of 90 putative mutants. Each of the colonies selected was grown to isolate its plasmid. Isolate plasmids were transformed into the RL08ara (Δ fadD) strain and grown in LB 0.4% glycerol and 20 μ M IPTG (mutants M1-M10) and 1 mM IPTG (mutants M3, M4, and M11-M90).

Mutant	# Residue Changes	Mutations	C8:0 (mg/L)	C10:0 (mg/L)	C10:1 (mg/L)	C12:0 (mg/L)	C12:1 (mg/L)	C14:0 (mg/L)	C14:1 (mg/L)	C16:0 (mg/L)	C16:1 (mg/L)	C18:0 (mg/L)	C18:1 (mg/L)	Total FFA (mg/L)	Induction Lev Tested (μ M)	C8:0 Fold Increase Relative to CpFatB1.2 at Same Induction Level
CpFatB1.2	0	n/a	12.4	5.6	0.9	0.0	1.5	0.6	9.4	10.1	52.6	0.0	0.0	93.2	20	1.0
			20.7 \pm 1.75	1.7 \pm 0.48	1.5 \pm 0.07	0.0	1.7 \pm 0.12	0.7 \pm 0.1	8.4 \pm 1.1	13.2 \pm 1.2	56.0 \pm 2.9	0.0	1.3 \pm 0.04	107.1	1000	1.0
CpFatB1.2-M1	0	n/a	20.7	4.7	1.6	0.0	1.8	0.8	9.8	12.8	58.2	0.0	0.0	110.2	20	1.7
CpFatB1.2-M2	3	M29T, T117S, Q163L	15.0	2.8	1.3	0.0	1.6	0.8	10.5	17.5	69.7	0.0	1.5	120.7	20	1.2
CpFatB1.2-M3	2	A59S, K296R	49.5	2.7	4.8	0.6	3.6	1.1	9.4	11.7	51.8	0.0	1.4	136.7	20	4.0
			82.9 \pm 5.86	1.8 \pm 0.34	7.0 \pm 0.26	1.0 \pm 0.04	4.9 \pm 0.09	1.4 \pm 0.04	8.6 \pm 0.4	10.1 \pm 0.3	51.1 \pm 0.8	0	1.1 \pm 0.9	169.9	1000	4.0
CpFatB1.2-M4	2	Δ A54 (new start codon at M19), N28S, I65M	66.8	2.9	4.8	0.6	3.8	0.9	8.3	12.6	46.9	0.0	1.4	149.0	20	5.4
			155.3 \pm 16.2	2.1 \pm 0.18	10.8 \pm 0.96	1.3 \pm 0.12	6.1 \pm 0.26	1.6 \pm 0.1	8.8 \pm 0.4	9.1 \pm 0.8	47.8 \pm 1.6	0	1.2 \pm 1.0	243.9	1000	7.5
CpFatB1.2-M5	2	W17R, T204S	24.9	4.3	1.9	0.0	2.0	0.7	8.4	12.1	52.8	0.0	0.0	107.0	20	2.01
CpFatB1.2-M6	2	L251M, L265I	17.4	4.2	1.5	0.0	1.6	0.7	8.5	12.8	58.4	0.0	0.0	105.0	20	1.40
CpFatB1.2-M7	0	n/a	20.5	5.9	1.3	0.0	1.8	0.6	7.7	12.6	56.4	0.0	1.3	108.0	20	1.65
CpFatB1.2-M8	2	K15E, S207T	16.4	5.5	1.1	0.0	1.6	0.7	8.6	10.8	54.3	0.0	0.0	99.0	20	1.32
CpFatB1.2-M9	1	R261S	28.6	4.4	2.4	0.0	2.5	0.8	8.4	11.4	53.3	0.0	0.0	111.8	20	2.31
CpFatB1.2-M10	0	n/a	12.4	5.6	0.9	0.0	1.5	0.6	9.4	10.1	52.6	0.0	0.0	93.2	20	1.27
CpFatB1.2-M11	ND	ND	26.4	0.0	1.0	0.0	1.7	0.0	7.6	12.1	69.0	0.0	14.5	132.2	1000	1.3
CpFatB1.2-M12	1	M136V	61.0	1.3	2.8	0.0	3.8	0.0	7.9	11.8	60.2	0.0	21.4	170.2	1000	2.9
CpFatB1.2-M13	ND	ND	13.9	0.0	0.0	0.0	0.8	0.0	3.3	3.7	47.1	0.0	11.9	80.7	1000	0.7
CpFatB1.2-M14	ND	ND	16.5	0.0	0.9	0.0	1.4	0.0	5.3	8.9	68.5	0.0	15.3	116.8	1000	0.8
CpFatB1.2-M15	ND	ND	15.7	0.0	0.7	0.0	1.1	0.0	3.1	3.7	45.5	0.0	12.2	81.9	1000	0.8
CpFatB1.2-M16	1	K296R	43.2	0.8	2.3	0.0	3.0	0.0	6.9	10.7	61.8	0.0	19.2	147.8	1000	2.1
CpFatB1.2-M17	ND	ND	26.9	0.6	1.5	0.0	1.8	0.0	9.4	14.2	68.0	0.0	18.0	140.3	1000	1.3

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TABLE 2-continued

Fatty acid profile of 90 putative mutants. Each of the colonies selected was grown to isolate its plasmid. Isolate plasmids were transformed into the RL08ara (AfadD) strain and grown in LB 0.4% glycerol and 20 μ M IPTG (mutants M1-M10) and 1 mM IPTG (mutants M3, M4, and M11-M90).

Mutant	# Residue Changes	Mutations	C8:0 (mg/L)	C10:0 (mg/L)	C10:1 (mg/L)	C12:0 (mg/L)	C12:1 (mg/L)	C14:0 (mg/L)	C14:1 (mg/L)	C16:0 (mg/L)	C16:1 (mg/L)	C18:0 (mg/L)	C18:1 (mg/L)	Total FFA (mg/L)	Induction Lev Tested (μ M)	C8:0 Fold Increase Relative to CpFatB1.2 at Same Induction Level
CpFatB1.2-M18	ND	ND	27.2	1.1	1.5	0.0	2.2	0.0	10.0	17.4	86.7	0.0	24.2	170.2	1000	1.3
CpFatB1.2-M19	ND	ND	18.5	1.1	0.9	0.0	1.5	0.0	6.9	12.2	65.5	0.0	16.6	123.1	1000	0.9
CpFatB1.2-M20	2	W17ST OP D293V	61.5	0.7	3.4	0.0	3.6	0.0	7.0	8.7	57.2	0.0	18.3	160.3	1000	3.0
CpFatB1.2-M21	ND	ND	17.4	0.0	0.9	0.0	1.9	0.0	10.1	13.2	63.5	1.1	14.5	122.7	1000	0.8
CpFatB1.2-M22	ND	ND	38.9	0.0	2.0	0.0	3.2	0.0	9.1	13.3	60.0	1.0	19.1	146.6	1000	1.9
CpFatB1.2-M23	ND	ND	21.5	0.0	1.2	0.0	2.3	0.0	9.2	12.7	60.4	0.9	15.5	123.7	1000	1.0
CpFatB1.2-M24	ND	ND	17.4	0.0	1.0	0.0	2.0	0.0	10.6	13.8	69.2	1.5	13.6	129.1	1000	0.8
CpFatB1.2-M25	ND	ND	17.5	0.0	1.0	0.0	2.0	0.0	11.9	14.3	63.3	1.0	14.9	125.8	1000	0.8
CpFatB1.2-M26	ND	ND	20.3	0.0	1.1	0.0	1.9	0.0	7.6	12.6	68.1	1.2	14.4	127.3	1000	1.0
CpFatB1.2-M27	ND	ND	15.8	0.0	0.9	0.0	2.2	0.0	11.7	13.4	64.4	1.2	13.2	122.9	1000	0.8
CpFatB1.2-M28	ND	ND	16.9	0.0	0.9	0.0	2.0	0.0	10.8	14.1	68.4	1.4	13.5	128.1	1000	0.8
CpFatB1.2-M29	ND	ND	21.9	0.0	1.2	0.0	2.0	0.0	5.7	9.4	65.5	1.4	13.1	120.1	1000	1.1
CpFatB1.2-M30	5	M19T, G35D, T117S, T121A, M138I	28.4	0.0	1.8	0.0	4.0	0.0	9.7	12.7	66.2	1.1	18.8	142.7	1000	1.4
CpFatB1.2-M31	ND	ND	15.2	0.0	1.0	0.0	2.0	0.0	10.1	13.5	64.0	3.5	11.7	121.0	1000	0.7
CpFatB1.2-M32	1	R22H	31.4	0.0	1.8	0.0	2.7	0.0	9.2	13.8	64.3	3.1	19.1	145.3	1000	1.5
CpFatB1.2-M33	ND	ND	12.8	0.0	0.8	0.0	1.6	0.0	6.1	10.8	63.5	3.5	8.7	107.9	1000	0.6
CpFatB1.2-M34	ND	ND	8.0	0.0	0.5	0.0	0.9	0.0	3.1	1.1	36.9	2.5	3.1	56.2	1000	0.4
CpFatB1.2-M35	ND	ND	14.2	0.0	0.9	0.0	1.7	0.0	7.9	12.8	65.6	3.8	10.6	117.6	1000	0.7
CpFatB1.2-M36	ND	ND	11.8	0.0	0.8	0.0	2.2	0.0	8.7	11.5	64.0	3.4	10.2	112.7	1000	0.6
CpFatB1.2-M37	ND	ND	11.4	0.0	0.7	0.0	1.3	0.0	5.4	6.9	63.9	3.6	6.8	100.0	1000	0.6

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TABLE 2-continued

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Mutant	# Residue Changes	Mutations	C8:0 (mg/L)	C10:0 (mg/L)	C10:1 (mg/L)	C12:0 (mg/L)	C12:1 (mg/L)	C14:0 (mg/L)	C14:1 (mg/L)	C16:0 (mg/L)	C16:1 (mg/L)	C18:0 (mg/L)	C18:1 (mg/L)	Total FFA (mg/L)	Induction Lev Tested (μ M)	C8:0 Fold Increase Relative to CpFatB1.2 at Same Induction Level
CpFatB1.2-M38	ND	ND	12.1	0.0	0.7	0.0	1.2	0.0	4.6	6.1	61.1	3.3	9.2	98.5	1000	0.6
CpFatB1.2-M39	ND	ND	17.3	0.0	1.1	0.0	1.9	0.0	6.5	12.3	76.2	4.6	11.2	131.0	1000	0.8
CpFatB1.2-M40	1	D293V	43.1	0.0	2.9	0.0	3.7	0.0	7.4	10.0	56.0	2.8	17.1	143.1	1000	2.1
CpFatB1.2-M41	ND	ND	8.6	0.0	0.0	0.0	1.0	0.0	3.3	2.4	39.8	0.7	6.8	62.6	1000	0.4
CpFatB1.2-M42	ND	ND	8.3	0.0	0.5	0.0	1.0	0.0	3.6	3.8	48.0	0.8	9.6	75.7	1000	0.4
CpFatB1.2-M43	ND	ND	13.8	0.0	0.8	0.0	1.6	0.0	5.5	8.9	71.5	1.2	15.5	118.8	1000	0.7
CpFatB1.2-M44	ND	ND	16.0	0.0	0.9	0.0	2.0	0.0	6.6	12.8	62.2	1.2	13.3	115.0	1000	0.8
CpFatB1.2-M45	ND	ND	8.4	0.0	0.5	0.0	1.0	0.0	3.2	3.5	42.6	0.7	9.0	69.0	1000	0.4
CpFatB1.2-M46	ND	ND	14.2	0.0	0.8	0.0	1.6	0.0	5.1	9.2	63.5	1.2	11.2	106.7	1000	0.7
CpFatB1.2-M47	3	N146K, D293V, N309D	69.2	0.0	4.3	0.0	4.6	0.0	6.1	8.9	62.7	1.2	21.0	178.0	1000	3.3
CpFatB1.2-M48	ND	ND	24.7	0.0	1.3	0.0	2.2	0.0	7.4	11.1	58.8	1.0	12.7	119.3	1000	1.2
CpFatB1.2-M49	ND	ND	8.3	0.0	0.5	0.0	1.0	0.0	3.4	3.6	45.0	0.8	9.9	72.5	1000	0.4
CpFatB1.2-M50	ND	ND	18.1	0.0	1.0	0.0	1.3	0.0	4.7	7.6	62.7	1.2	9.1	105.7	1000	0.9
CpFatB1.2-M51	ND	ND	13.2	0.0	0.0	0.0	1.5	0.0	6.1	0.0	61.9	0.0	11.8	94.5	1000	0.6
CpFatB1.2-M52	ND	ND	7.9	0.0	0.0	0.0	0.9	0.0	3.0	0.0	39.5	0.0	5.8	57.2	1000	0.4
CpFatB1.2-M53	ND	ND	14.7	0.0	0.0	0.0	1.5	0.0	4.7	0.0	60.0	0.0	9.3	90.2	1000	0.7
CpFatB1.2-M54	ND	ND	9.9	0.0	0.0	0.0	1.2	0.0	4.6	0.0	60.5	0.0	12.6	88.8	1000	0.5
CpFatB1.2-M55	ND	ND	23.5	0.0	0.0	0.0	2.0	0.0	5.9	0.0	55.3	0.0	13.1	99.9	1000	1.1
CpFatB1.2-M56	ND	ND	16.0	0.0	0.0	0.0	2.0	0.0	7.7	0.0	56.9	0.0	12.6	95.2	1000	0.8
CpFatB1.2-M57	ND	ND	19.8	0.0	0.0	0.0	2.0	0.0	7.1	0.0	60.5	0.0	14.4	103.8	1000	1.0
CpFatB1.2-M58	ND	ND	20.2	0.0	0.0	0.0	2.3	0.0	9.3	0.0	57.5	0.0	15.1	104.3	1000	1.0
CpFatB1.2-M59	ND	ND	18.6	0.0	0.0	0.0	1.7	0.0	6.0	0.0	64.6	0.0	13.4	104.4	1000	0.9

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TABLE 2-continued

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Mutant	# Residue Changes	Mutations	C8:0 (mg/L)	C10:0 (mg/L)	C10:1 (mg/L)	C12:0 (mg/L)	C12:1 (mg/L)	C14:0 (mg/L)	C14:1 (mg/L)	C16:0 (mg/L)	C16:1 (mg/L)	C18:0 (mg/L)	C18:1 (mg/L)	Total FFA (mg/L)	Induction Lev Tested (μ M)	C8:0 Fold Increase Relative to CpFatB1.2 at Same Induction Level
CpFatB1.2-M60	ND	ND	19.1	0.0	0.0	0.0	1.8	0.0	6.4	0.0	55.8	0.0	12.7	95.7	1000	0.9
CpFatB1.2-M61	ND	ND	13.6	0.0	0.0	0.0	1.0	0.0	3.2	1.4	39.6	0.7	4.3	63.8	1000	0.7
CpFatB1.2-M62	ND	ND	12.6	0.0	0.0	0.0	1.0	0.0	3.2	1.7	40.4	0.8	4.9	64.6	1000	0.6
CpFatB1.2-M63	ND	ND	19.7	0.0	0.0	0.0	1.6	0.0	6.7	12.7	66.2	1.3	10.4	118.6	1000	1.0
CpFatB1.2-M64	ND	ND	22.1	0.0	0.0	0.0	2.1	0.0	12.3	16.5	69.2	1.5	13.9	137.5	1000	1.1
CpFatB1.2-M65	ND	ND	20.5	0.0	0.0	0.0	1.7	0.0	6.6	10.3	73.9	1.3	10.0	124.3	1000	1.0
CpFatB1.2-M66	1	D293V	54.3	0.0	1.3	0.0	3.7	0.0	7.8	10.9	58.6	1.1	15.2	152.9	1000	2.6
CpFatB1.2-M67	ND	ND	12.7	0.0	0.0	0.0	0.9	0.0	3.2	1.0	39.4	0.8	3.6	61.6	1000	0.6
CpFatB1.2-M68	1	T245N	28.4	0.0	0.0	0.0	2.0	0.0	8.1	15.4	64.8	1.2	14.3	134.3	1000	1.4
CpFatB1.2-M69	ND	ND	21.2	0.0	0.0	0.0	1.7	0.0	7.9	14.8	65.2	1.1	12.0	123.7	1000	1.0
CpFatB1.2-M70	ND	ND	21.8	0.0	0.0	0.0	1.6	0.0	7.3	14.2	67.9	1.2	12.3	126.4	1000	1.1
CpFatB1.2-M71	1	M136I	46.7	0.0	1.0	0.0	2.8	0.0	5.2	3.9	85.2	1.8	18.0	164.6	1000	2.3
CpFatB1.2-M72	ND	ND	22.9	0.0	0.0	0.0	1.8	0.0	5.2	8.5	61.5	1.3	11.1	112.3	1000	1.1
CpFatB1.2-M73	3	V268I, R279H, D293V	61.2	0.0	1.4	0.0	4.2	0.0	7.8	8.5	56.0	0.9	16.4	156.5	1000	3.0
CpFatB1.2-M74	ND	ND	21.6	0.0	0.0	0.0	2.1	0.0	9.1	12.8	60.1	0.9	14.1	120.6	1000	1.0
CpFatB1.2-M75	ND	ND	18.0	0.0	0.0	0.0	1.4	0.0	4.3	7.3	50.2	1.0	9.2	91.2	1000	0.9
CpFatB1.2-M76	2	K23E, L86Q	46.6	0.0	0.6	0.0	2.5	0.0	6.6	11.9	58.6	0.9	16.9	144.6	1000	2.2
CpFatB1.2-M77	ND	ND	21.4	0.0	0.0	0.0	1.9	0.0	8.2	14.4	62.9	1.1	15.5	125.4	1000	1.0
CpFatB1.2-M78	ND	ND	28.2	0.0	0.0	0.0	2.4	0.0	8.7	12.6	66.9	1.2	15.6	135.7	1000	1.4
CpFatB1.2-M79	3	V9M, K15E, E236A	42.0	0.0	0.6	0.0	3.3	0.0	8.5	12.6	59.1	1.0	17.9	144.9	1000	2.0
CpFatB1.2-M80	ND	ND	19.9	0.0	0.0	0.0	1.3	0.0	6.3	10.4	59.8	1.1	12.4	111.2	1000	1.0

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TABLE 2-continued

Fatty acid profile of 90 putative mutants. Each of the colonies selected was grown to isolate its plasmid. Isolate plasmids were transformed into the RL08ara (*AfadD*) strain and grown in LB 0.4% glycerol and 20 μ M IPTG (mutants M1-M10) and 1 mM IPTG (mutants M3, M4, and M11-M90).

Mutant	# Residue Changes	Mutations	C8:0 (mg/L)	C10:0 (mg/L)	C10:1 (mg/L)	C12:0 (mg/L)	C12:1 (mg/L)	C14:0 (mg/L)	C14:1 (mg/L)	C16:0 (mg/L)	C16:1 (mg/L)	C18:0 (mg/L)	C18:1 (mg/L)	Total FFA (mg/L)	Induction Lev Tested (μ M)	C8:0 Fold Increase Relative to CpFatB1.2 at Same Induction Level
CpFatB1.2-M81	ND	ND	8.5	0.0	0.4	0.0	1.2	0.0	5.0	3.2	72.0	1.6	8.3	100.1	1000	0.4
CpFatB1.2-M82	ND	ND	11.3	0.0	0.6	0.0	1.8	0.0	4.9	6.4	71.8	1.6	15.2	113.6	1000	0.5
CpFatB1.2-M83	ND	ND	18.8	0.0	1.2	0.0	2.6	0.0	5.6	4.4	69.2	1.5	14.6	117.8	1000	0.9
CpFatB1.2-M84	ND	ND	9.2	0.0	0.4	0.0	1.2	0.0	4.6	2.4	66.2	1.5	7.4	93.0	1000	0.4
CpFatB1.2-M85	ND	ND	9.4	0.0	0.5	0.0	1.4	0.0	4.6	5.0	69.1	1.6	10.3	101.8	1000	0.5
CpFatB1.2-M86	ND	ND	16.2	0.0	1.0	0.0	2.1	0.0	5.3	6.6	71.7	1.4	16.1	120.3	1000	0.8
CpFatB1.2-M87	ND	ND	27.2	0.0	1.6	0.0	1.9	0.0	4.9	2.7	65.6	1.6	9.4	114.9	1000	1.3
CpFatB1.2-M88	ND	ND	12.0	0.0	0.7	0.0	1.8	0.0	5.1	6.7	74.5	1.6	16.4	118.8	1000	0.6
CpFatB1.2-M89	ND	ND	8.6	0.0	0.4	0.0	1.2	0.0	4.3	2.4	63.9	1.5	7.7	90.0	1000	0.4
CpFatB1.2-M90	ND	ND	11.1	0.0	0.6	0.0	1.5	0.0	5.1	5.2	72.8	1.5	12.7	110.6	1000	0.5

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Production of Octanol Via Fatty Acid Biosynthesis

Expression of the CpFatB1.2 variants enables high flux to octanoic acid. Analogous to work with the BTE and conversion of dodecanoic acid to dodecanol (U.S. Pat. No. 9,708,630), we can co-express the CpFatB1.2 variants, an acyl-CoA synthetase, and a hybrid acyl-CoA reductase/aldehyde reductase to produce octanol. FadD, the native acyl-CoA synthetase used in the prior dodecanol work (U.S. Pat. No. 9,708,630) has poor activity against octanoic acid, so we replaced it with variants from other organisms. The best variant was FadD6 from *Mycobacterium tuberculosis* (SEQ ID NO:15 (nucleotide coding sequence) and SEQ ID NO:16 (protein sequence)). When these genes were co-expressed from plasmids in *E. coli* NHL13 (Δ fadD::fadD6 Δ pta Δ poxB Δ ldhA), we observed up to 1.1 g/L titers from cultures grown in Clomberg media. The CpFatB1.2-M4 variant produced the most octanol (FIG. 16). Another suitable acyl-CoA synthetase with high activity on C8:0 is *Pseudomonas putida* PP0763 (SEQ ID NO:17 (nucleotide coding sequence) and SEQ ID NO:18 (protein sequence)).

Structural Modeling Insights

It has been experimentally shown that the CpFatB1.2-M4-287 mutant exhibits 15-fold higher specific activity in vivo (FIG. 11) relative to the natural parent enzyme. The CpFatB1.2-M4-287 enzyme variant the N28S, I65M double mutation and an 18-residue N-terminal truncation. To evaluate how these mutations impacted activity, we constructed a computational model of the mutant and parent enzymes. These were prepared by homology modeling using the published structure of *U. californica* UcFatB1 (PDB:5x04)²⁹ as a template. Unfortunately, we could not predict the structure of the N-terminus of CpFatB1.2-M4-287 as the 12 N-terminal residues of M4-287 were not conserved with the crystallized protein and no empirical 3D template structure could be identified with >30% sequence identity to this region. This means that our CpFatB1.2 model structure has a 30-residue N-terminus truncation relative to CpFatB1.2 and thus excluding the important N28S mutation and the truncation introduced in variant M4-287. On the other hand, the model was able to project where the I65M mutation resided relative to the active site (FIGS. 17A and 17B). Ile65 was positioned at one end of the acyl-binding crevice farthest from the crevice opening, where the catalytic residues line the periphery of the opening. We hypothesize that the bulkier side chain of the methionine is occluding the crevice end introducing steric clashes with the omega-1 acyl carbon of C₁₂-ACP, the preferred substrate of the template (PDB: 5x04) thioesterase. This residue is thus seemingly important for altering C₈-specificity, not activity. However, Mendonça et al.³⁰ show that mutations to amino acids connected to an active site residue by three or less noncovalent interactions can have a high impact on enzyme activity, a feat consistent with I65M according to the contact map for this residue (FIG. 17C).

The CpFatB1.2-M3 also exhibited elevated enzymatic activities (~4 folds higher than wild-type in vivo). Our CpFatB1.2 computational model with the 30 amino acid truncation at the N-terminus was used as the starting point to generate the variant model. Unlike CpFatB1.2-M4-287, we could capture the effects of both A59S and K296R mutations in this model. We hypothesize K296R has indirect and A59S has direct effects on enzyme activity based on the number of noncovalent bonds that connect these residues to one or more catalytic residues. Lys296 is a surface residue

that has its side chain facing away from the acyl-binding pocket. However, a K296R mutation introduces a stable salt bridge of interaction (~3.2 Å) between the positively charged N atom of Arg296 side chain and side chain O atom of Glu254 (FIGS. 18A-18C). Prior to mutation the orientation of the Lys296 side chain did not result in a salt-bridge (~5.3 Å apart). Contact map (FIG. 18C) reveals Lys296 needs at least five noncovalent interactions to reach an active site residue which is shorter than that before the K296R mutation, but is less likely to have as much effect as I65M from CpFatB1.2-M4-287 (separated by 3 noncovalent interactions). However, CpFatB1.2-M3 accounts for its increase in activity by the A59S mutation which is adjacent to a Asp220 (a key catalytic residue). The side chain OH of Ser59 is linked to the backbone O atom of Asp220 (~4.5 Å) (FIG. 18C). The aliphatic methyl-side chain of Ala59 before mutation prevented any such polar contacts.

The CpFatB1.2-M4-287 and CpFatB1.2-M3 contact maps reveal the importance of Met69 (see FIGS. 17C and 18C) in connecting the mutated residues to the active site residues as it is the most connected node in the map. It is noteworthy that Met69 maintains these contact networks by mostly hydrophobic interactions higher up the cascade and polar interactions lower down. The polar interactions are controlled by the backbone O atom whereas the hydrophobic ones require the side chain C atoms. We hypothesize that Met69 is important for ensuring a connected contact map for both these mutants and altering M69 to a charged amino acid or a small-side chain hydrophobic residue can significantly reduce the enhanced activity of the CpFatB1 mutants.

Conclusions

Using a lipoic acid selection, we isolated a mutated octanoyl-ACP thioesterase capable of high rates of hydrolysis while maintaining >90% specificity towards C₈ acyl chains. A cell harboring a single chromosomal copy of this thioesterase gene is capable of achieving the same high level of production observed from plasmids expressing the parent enzyme. Under the conditions tested we demonstrated a more than 3-fold improvement over the highest reported octanoic acid titers in the literature. In light of the improved activity, we conclude that this work removes the thioesterase bottleneck for producing C₈ compounds in *E. coli*. Additional examples and discussion can be found in Hernandez Lozada et al.⁴², which is incorporated herein by reference in its entirety. Future work can now focus on optimizing the flux from octanoic acid to desired 8-carbon products with other chemical functionalities.

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Gly Leu Thr Pro Gly Trp Tyr Asp Leu Asp Val Asn Gln His Val Ser
 305 310 315 320

Asn Val Lys Tyr Ile Gly Trp Ile Leu Glu Ser Met Pro Thr Glu Val
 325 330 335

Leu Glu Thr Gln Glu Leu Cys Ser Leu Thr Leu Glu Tyr Arg Arg Glu
 340 345 350

Cys Gly Arg Asp Ser Val Leu Glu Ser Val Thr Ser Met Asp Pro Ser
 355 360 365

Lys Val Gly Asp Arg Phe Gln Tyr Arg His Leu Leu Arg Leu Glu Asp
 370 375 380

Gly Ala Asp Ile Met Lys Gly Arg Thr Glu Trp Arg Pro Lys Asn Ala
 385 390 395 400

Gly Thr Asn Gly Ala Ile Ser Thr Gly Lys Thr
 405 410

<210> SEQ ID NO 3
 <211> LENGTH: 954
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB1 1.2 Trucation

<400> SEQUENCE: 3

atgctgctca ccgcgattac caccgtgttc gttgcgccgg aaaaacgttg gactatgttc	60
gaccgcaaat ctaaacgtcc gaacatgctg atggatagct tcggtctgga acgtgttgtc	120
caggatggtc tggttttocg ccaatccttc tccattcgta gctacgaaat ctgtgcggac	180
cgcacggcga gcatcgagac cgtgatgaac cacgttcagg aaaccagcct gaaccagtgt	240
aagtccatcg gcctgctgga cgacggtttc ggtcgtcac cggaatgtg taaacgcgac	300

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ctgatttggg ttgtaaccog tatgaaaatt atggtaaacc gttatccgac ctggggcgac   360
accattgaag tctccacctg gctgtcccag tccggtaaaa ttggcatggg tccgactgctg   420
ctgattagcg attgtaatac cggtgaaata ctggttcgtg caacctcagt ttacgccatg   480
atgaaccaga aaacgcgtcg cttctctaaa ctgccccacg aagtacgcca ggagtgcgctg   540
ccgcacttct tagactcccc gccggccatt gaagacaacg acggtaaact gcagaaattc   600
gatgtgaaaa ccggtgatcc catccgcaaa ggtctgaccc cgggctggta cgacctggac   660
gtaaaccagc acgtttccaa tgtcaaatat atcggttgga tcctggaatc catgcccacc   720
gaagtcttgg aaaccaaga actttgcagc ctgactctcg aataccgtcg cgaatgcggt   780
cgcgactcag tacttgagag cgtaacctcc atggaccctg ctaaagttgg tgatcgtttt   840
cagtatcgtc atctgctgcg tctggaagat ggtgcggaca ttatgaaggg ccgtaccgaa   900
tggcgtccga aaaacgcggg cactaatggt gctatctcca ctgggaaaac ataa       954

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<210> SEQ ID NO 4

<211> LENGTH: 317

<212> TYPE: PRT

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: CpFatB 1.2 Truncation

<400> SEQUENCE: 4

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Met Leu Leu Thr Ala Ile Thr Thr Val Phe Val Ala Pro Glu Lys Arg
1           5           10          15
Trp Thr Met Phe Asp Arg Lys Ser Lys Arg Pro Asn Met Leu Met Asp
          20          25          30
Ser Phe Gly Leu Glu Arg Val Val Gln Asp Gly Leu Val Phe Arg Gln
          35          40          45
Ser Phe Ser Ile Arg Ser Tyr Glu Ile Cys Ala Asp Arg Thr Ala Ser
          50          55          60
Ile Glu Thr Val Met Asn His Val Gln Glu Thr Ser Leu Asn Gln Cys
          65          70          75          80
Lys Ser Ile Gly Leu Leu Asp Asp Gly Phe Gly Arg Ser Pro Glu Met
          85          90          95
Cys Lys Arg Asp Leu Ile Trp Val Val Thr Arg Met Lys Ile Met Val
          100         105         110
Asn Arg Tyr Pro Thr Trp Gly Asp Thr Ile Glu Val Ser Thr Trp Leu
          115         120         125
Ser Gln Ser Gly Lys Ile Gly Met Gly Arg Asp Trp Leu Ile Ser Asp
          130         135         140
Cys Asn Thr Gly Glu Ile Leu Val Arg Ala Thr Ser Val Tyr Ala Met
          145         150         155         160
Met Asn Gln Lys Thr Arg Arg Phe Ser Lys Leu Pro His Glu Val Arg
          165         170         175
Gln Glu Phe Ala Pro His Phe Leu Asp Ser Pro Pro Ala Ile Glu Asp
          180         185         190
Asn Asp Gly Lys Leu Gln Lys Phe Asp Val Lys Thr Gly Asp Ser Ile
          195         200         205
Arg Lys Gly Leu Thr Pro Gly Trp Tyr Asp Leu Asp Val Asn Gln His
          210         215         220
Val Ser Asn Val Lys Tyr Ile Gly Trp Ile Leu Glu Ser Met Pro Thr
          225         230         235         240
Glu Val Leu Glu Thr Gln Glu Leu Cys Ser Leu Thr Leu Glu Tyr Arg
          245         250         255

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Arg Glu Cys Gly Arg Asp Ser Val Leu Glu Ser Val Thr Ser Met Asp
 260 265 270

Pro Ser Lys Val Gly Asp Arg Phe Gln Tyr Arg His Leu Leu Arg Leu
 275 280 285

Glu Asp Gly Ala Asp Ile Met Lys Gly Arg Thr Glu Trp Arg Pro Lys
 290 295 300

Asn Ala Gly Thr Asn Gly Ala Ile Ser Thr Gly Lys Thr
 305 310 315

<210> SEQ ID NO 5
 <211> LENGTH: 891
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB 1.3 Truncation

<400> SEQUENCE: 5

atgaaatcta aacgtccgaa catgctgatg gatagcttcg gtctggaacg tgttgtccag 60
 gatggtctgg ttttccgcc atccttctcc attcgtagct acgaaatctg tgcggaccgc 120
 acggcgagca tcgagaccgt gatgaaccac gttcaggaaa ccagcctgaa ccagtgtaa 180
 tccatcggcc tgctggacga cggtttcggt cgctcaccgg aaatgtgtaa acgcgacctg 240
 atttgggttg taaccctgat gaaaattatg gtaaacctt atccgacctg gggcgacacc 300
 attgaagtct ccacctggct gtcccagtcc ggtaaaattg gcatgggtcg cgactggctg 360
 attagcgatt gtaataccgg tgaatactg gttcgtgcaa cctcagttta cgccatgatg 420
 aaccagaaaa cgcgtcgtct ctctaaactg ccccacgaag tacgccagga gttcgcgccg 480
 cacttcttag actccccgcc ggccattgaa gacaacgacg gtaaaactgca gaaattcgat 540
 gtgaaaacgg gtgattccat ccgcaaaggc ctgaccccggt gctggtaacga cctggacgta 600
 aaccagcacg tttccaatgt caaatatctc ggttggatcc tggaatccat gccgaccgaa 660
 gttctgaaaa cccaagaact ttgcagcctg actctcgaat accgtcgcga atgcggctcg 720
 gactcagtac ttgagagcgt aaacctcatg gaccctgcta aagttggtga tcgttttcag 780
 tatcgtcctc tgctgcgtct ggaagatggt gcggacatta tgaagggccg taccgaatgg 840
 cgccgaaaa acgcgggcac taatggtgct atctccactg ggaaaacata a 891

<210> SEQ ID NO 6
 <211> LENGTH: 296
 <212> TYPE: PRT
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB 1.3 Truncation

<400> SEQUENCE: 6

Met Lys Ser Lys Arg Pro Asn Met Leu Met Asp Ser Phe Gly Leu Glu
 1 5 10 15

Arg Val Val Gln Asp Gly Leu Val Phe Arg Gln Ser Phe Ser Ile Arg
 20 25 30

Ser Tyr Glu Ile Cys Ala Asp Arg Thr Ala Ser Ile Glu Thr Val Met
 35 40 45

Asn His Val Gln Glu Thr Ser Leu Asn Gln Cys Lys Ser Ile Gly Leu
 50 55 60

Leu Asp Asp Gly Phe Gly Arg Ser Pro Glu Met Cys Lys Arg Asp Leu
 65 70 75 80

Ile Trp Val Val Thr Arg Met Lys Ile Met Val Asn Arg Tyr Pro Thr
 85 90 95

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Trp Gly Asp Thr Ile Glu Val Ser Thr Trp Leu Ser Gln Ser Gly Lys
 100 105 110

Ile Gly Met Gly Arg Asp Trp Leu Ile Ser Asp Cys Asn Thr Gly Glu
 115 120 125

Ile Leu Val Arg Ala Thr Ser Val Tyr Ala Met Met Asn Gln Lys Thr
 130 135 140

Arg Arg Phe Ser Lys Leu Pro His Glu Val Arg Gln Glu Phe Ala Pro
 145 150 155 160

His Phe Leu Asp Ser Pro Pro Ala Ile Glu Asp Asn Asp Gly Lys Leu
 165 170 175

Gln Lys Phe Asp Val Lys Thr Gly Asp Ser Ile Arg Lys Gly Leu Thr
 180 185 190

Pro Gly Trp Tyr Asp Leu Asp Val Asn Gln His Val Ser Asn Val Lys
 195 200 205

Tyr Ile Gly Trp Ile Leu Glu Ser Met Pro Thr Glu Val Leu Glu Thr
 210 215 220

Gln Glu Leu Cys Ser Leu Thr Leu Glu Tyr Arg Arg Glu Cys Gly Arg
 225 230 235 240

Asp Ser Val Leu Glu Ser Val Thr Ser Met Asp Pro Ser Lys Val Gly
 245 250 255

Asp Arg Phe Gln Tyr Arg His Leu Leu Arg Leu Glu Asp Gly Ala Asp
 260 265 270

Ile Met Lys Gly Arg Thr Glu Trp Arg Pro Lys Asn Ala Gly Thr Asn
 275 280 285

Gly Ala Ile Ser Thr Gly Lys Thr
 290 295

<210> SEQ ID NO 7
 <211> LENGTH: 831
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB 1.4 Truncation

<400> SEQUENCE: 7

atgggtctgg ttttccgcc atccttctcc attcgtagct acgaaatctg tgcggaccgc 60
 acggcgagca tcgagaccgt gatgaaccac gttcaggaaa ccagcctgaa ccagtgtaa 120
 tccatcgccc tgctggacga cggtttcggc cgctcaccgg aaatgtgtaa acgogacctg 180
 atttgggttg taaccgtat gaaaattatg gtaaaccgtt atccgacctg gggcgacacc 240
 attgaagtct ccacctggct gtcccagtcc ggtaaaattg gcatgggtcg cgactggctg 300
 attagcgatt gtaataccgg tgaataactg gttcgtgcaa cctcagttta cgccatgatg 360
 aaccagaaaa cgcgtcgctt ctctaactg cccaacgaag tacgccagga gttcgcgccc 420
 cacttcttag actccccgcc ggccattgaa gacaacgacg gtaaactgca gaaattcgat 480
 gtgaaaaccg gtgattccat ccgcaaggt ctgacccccg gctggtaacga cctggacgta 540
 aaccagcacg tttccaatgt caaatatc ggttggatcc tggaatccat gccgaccgaa 600
 gttctggaaa cccaagaact ttgcagcctg actctcgaat accgtcgcga atgcggtcgc 660
 gactcagtac ttgagagcgt aacctccatg gaccctgcta aagttgggta tcgttttcag 720
 tatcgtcatc tgctgcgtct ggaagatggt gcggacatta tgaagggccg taccgaatgg 780
 cgtccgaaaa acgcggggc ac taatggtgct atctccactg ggaaaacata a 831

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<210> SEQ ID NO 8
 <211> LENGTH: 276
 <212> TYPE: PRT
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB 1.4 Truncation

<400> SEQUENCE: 8

Met Gly Leu Val Phe Arg Gln Ser Phe Ser Ile Arg Ser Tyr Glu Ile
 1 5 10 15
 Cys Ala Asp Arg Thr Ala Ser Ile Glu Thr Val Met Asn His Val Gln
 20 25 30
 Glu Thr Ser Leu Asn Gln Cys Lys Ser Ile Gly Leu Leu Asp Asp Gly
 35 40 45
 Phe Gly Arg Ser Pro Glu Met Cys Lys Arg Asp Leu Ile Trp Val Val
 50 55 60
 Thr Arg Met Lys Ile Met Val Asn Arg Tyr Pro Thr Trp Gly Asp Thr
 65 70 75 80
 Ile Glu Val Ser Thr Trp Leu Ser Gln Ser Gly Lys Ile Gly Met Gly
 85 90 95
 Arg Asp Trp Leu Ile Ser Asp Cys Asn Thr Gly Glu Ile Leu Val Arg
 100 105 110
 Ala Thr Ser Val Tyr Ala Met Met Asn Gln Lys Thr Arg Arg Phe Ser
 115 120 125
 Lys Leu Pro His Glu Val Arg Gln Glu Phe Ala Pro His Phe Leu Asp
 130 135 140
 Ser Pro Pro Ala Ile Glu Asp Asn Asp Gly Lys Leu Gln Lys Phe Asp
 145 150 155 160
 Val Lys Thr Gly Asp Ser Ile Arg Lys Gly Leu Thr Pro Gly Trp Tyr
 165 170 175
 Asp Leu Asp Val Asn Gln His Val Ser Asn Val Lys Tyr Ile Gly Trp
 180 185 190
 Ile Leu Glu Ser Met Pro Thr Glu Val Leu Glu Thr Gln Glu Leu Cys
 195 200 205
 Ser Leu Thr Leu Glu Tyr Arg Arg Glu Cys Gly Arg Asp Ser Val Leu
 210 215 220
 Glu Ser Val Thr Ser Met Asp Pro Ser Lys Val Gly Asp Arg Phe Gln
 225 230 235 240
 Tyr Arg His Leu Leu Arg Leu Glu Asp Gly Ala Asp Ile Met Lys Gly
 245 250 255
 Arg Thr Glu Trp Arg Pro Lys Asn Ala Gly Thr Asn Gly Ala Ile Ser
 260 265 270
 Thr Gly Lys Thr
 275

<210> SEQ ID NO 9
 <211> LENGTH: 954
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB 1.2 M3 Mutant

<400> SEQUENCE: 9

atgctgctca ccgcatctac caccgtgttc gttgcgccgg aaaaacgttg gactatgttc 60
 gaccgcaaat ctaaactgtcc gaacatgctg atggatagct tcggtctgga acgtgttgtc 120
 caggatggtc tggttttccg ccaatccttc tccattcgta gctacgaaat ctgttcggac 180

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cgcacggcga gcatcgagac cgtgatgaac cacgttcagg aaaccagcct gaaccagtgt 240
aagtccatcg gcctgctgga cgacggtttc ggtcgctcac cggaatgtg taaacgcgac 300
ctgatttggg ttgtaaccg tatgaaaatt atggtaaacc gttatccgac ctggggcgac 360
accattgaag tctccacctg gctgtcccag tccggtaaaa ttggcatggg tcgogactgg 420
ctgattagcg attgtaatac cggtgaaata ctggttcgtg caacctcagt ttacgccatg 480
atgaaccaga aaacgcgctg cttctctaaa ctgccccacg aagtacgcca ggagttcgcg 540
ccgcacttct tagactcccc gccggccatt gaagacaacg acggtaaaact gcagaaattc 600
gatgtgaaaa ccggtgatcc catccgcaaa ggtctgacct cgggctggta cgacctggac 660
gtaaaccagc acgtttccaa tgtcaaatat atcggttgga tcctggaate catgccgacc 720
gaagttctgg aaaccaaga actttgcagc ctgactctcg aataccgtcg cgaatgcggt 780
cgcgactcag tacttgagag cgtaacctcc atggacctgt ctaaagtgg tgatcgtttt 840
cagtatcgtc atctgctgcg tctggaagat ggtgctggaca ttatgagggg ccgtaccgaa 900
tggcgtccga aaaacgcggg cactaatggt gctatctcca ctgggaaaac ataa 954
    
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<210> SEQ ID NO 10
<211> LENGTH: 317
<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: CpFatB1.2 M3 Mutant
    
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<400> SEQUENCE: 10

```

Met Leu Leu Thr Ala Ile Thr Thr Val Phe Val Ala Pro Glu Lys Arg
1          5          10          15

Trp Thr Met Phe Asp Arg Lys Ser Lys Arg Pro Asn Met Leu Met Asp
          20          25          30

Ser Phe Gly Leu Glu Arg Val Val Gln Asp Gly Leu Val Phe Arg Gln
          35          40          45

Ser Phe Ser Ile Arg Ser Tyr Glu Ile Cys Ser Asp Arg Thr Ala Ser
          50          55          60

Ile Glu Thr Val Met Asn His Val Gln Glu Thr Ser Leu Asn Gln Cys
65          70          75          80

Lys Ser Ile Gly Leu Leu Asp Asp Gly Phe Gly Arg Ser Pro Glu Met
          85          90          95

Cys Lys Arg Asp Leu Ile Trp Val Val Thr Arg Met Lys Ile Met Val
          100          105          110

Asn Arg Tyr Pro Thr Trp Gly Asp Thr Ile Glu Val Ser Thr Trp Leu
          115          120          125

Ser Gln Ser Gly Lys Ile Gly Met Gly Arg Asp Trp Leu Ile Ser Asp
          130          135          140

Cys Asn Thr Gly Glu Ile Leu Val Arg Ala Thr Ser Val Tyr Ala Met
          145          150          155          160

Met Asn Gln Lys Thr Arg Arg Phe Ser Lys Leu Pro His Glu Val Arg
          165          170          175

Gln Glu Phe Ala Pro His Phe Leu Asp Ser Pro Pro Ala Ile Glu Asp
          180          185          190

Asn Asp Gly Lys Leu Gln Lys Phe Asp Val Lys Thr Gly Asp Ser Ile
          195          200          205

Arg Lys Gly Leu Thr Pro Gly Trp Tyr Asp Leu Asp Val Asn Gln His
          210          215          220
    
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Val Ser Asn Val Lys Tyr Ile Gly Trp Ile Leu Glu Ser Met Pro Thr
 225 230 235 240

Glu Val Leu Glu Thr Gln Glu Leu Cys Ser Leu Thr Leu Glu Tyr Arg
 245 250 255

Arg Glu Cys Gly Arg Asp Ser Val Leu Glu Ser Val Thr Ser Met Asp
 260 265 270

Pro Ser Lys Val Gly Asp Arg Phe Gln Tyr Arg His Leu Leu Arg Leu
 275 280 285

Glu Asp Gly Ala Asp Ile Met Arg Gly Arg Thr Glu Trp Arg Pro Lys
 290 295 300

Asn Ala Gly Thr Asn Gly Ala Ile Ser Thr Gly Lys Thr
 305 310 315

<210> SEQ ID NO 11
 <211> LENGTH: 953
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB1.2 M4 Mutant

<400> SEQUENCE: 11

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atgctgctca ccgcgattac caccgtgttc gttgcgccgg aaaacgttgg actatgtttg    60
accgcaaadc taaacgtccg agcatgctga tggatagctt cggctctggaa cgtgttgctc    120
aggatggtct gggtttccgc caatccttct ccattcgtag ctacgaaadc tgtgcggacc    180
gcacggcgag catggagacc gtgatgaacc acgttcagga aaccagcctg aaccagtgta    240
agtccatcgg cctgctggac gacggtttcg gtcgctcacc ggaaatgtgt aaacgcgacc    300
tgatttgggt tgtaaccctg atgaaaatta tggtaaaccg ttatccgacc tggggcgaca    360
ccattgaagt ctcccactgg ctgtcccagt cgggtaaaat tggcatgggt cgcgactggc    420
tgattagcga ttgtaatacc ggtgaaatac tggttcgtgc aacctcagtt tacgccatga    480
tgaaccagaa aacgcgtcgc ttctctaaac tgccccacga agtacgccag gagttcgcgc    540
cgcacttctt agactccccg ccggccattg aagacaacga cggtaaaactg cagaaattcg    600
atgtgaaaac cgggtattcc atccgcaaag gtctgacccc gggctggtac gacctggacg    660
taaaccagca cgtttccaat gtcaaatata tcggttggat cctggaatcc atgccgaccg    720
aagttctgga aacccaagaa ctttgcagcc tgactctcga ataccgtcgc gaatgcggtc    780
gcgactcagt acttgagagc gtaacctcca tggaccctgc taaagttggt gatcgttttc    840
agtatcgtca tctgctcgct ctggaagatg gtgcggacat tatgaagggc cgtaccgaa    900
ggcgtccgaa aaacgcgggc actaatggtg ctatctccac tgggaaaaca taa          953
    
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<210> SEQ ID NO 12
 <211> LENGTH: 317
 <212> TYPE: PRT
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB1.2 M4 Mutant

<400> SEQUENCE: 12

Tyr Ala Ala His Arg Asp Tyr His Arg Val Arg Cys Ala Gly Lys Arg
 1 5 10 15

Trp Thr Met Phe Asp Arg Lys Ser Lys Arg Pro Ser Met Leu Met Asp
 20 25 30

Ser Phe Gly Leu Glu Arg Val Val Gln Asp Gly Leu Val Phe Arg Gln
 35 40 45

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Ser Phe Ser Ile Arg Ser Tyr Glu Ile Cys Ala Asp Arg Thr Ala Ser
 50 55 60

Met Glu Thr Val Met Asn His Val Gln Glu Thr Ser Leu Asn Gln Cys
 65 70 75 80

Lys Ser Ile Gly Leu Leu Asp Asp Gly Phe Gly Arg Ser Pro Glu Met
 85 90 95

Cys Lys Arg Asp Leu Ile Trp Val Val Thr Arg Met Lys Ile Met Val
 100 105 110

Asn Arg Tyr Pro Thr Trp Gly Asp Thr Ile Glu Val Ser Thr Trp Leu
 115 120 125

Ser Gln Ser Gly Lys Ile Gly Met Gly Arg Asp Trp Leu Ile Ser Asp
 130 135 140

Cys Asn Thr Gly Glu Ile Leu Val Arg Ala Thr Ser Val Tyr Ala Met
 145 150 155 160

Met Asn Gln Lys Thr Arg Arg Phe Ser Lys Leu Pro His Glu Val Arg
 165 170 175

Gln Glu Phe Ala Pro His Phe Leu Asp Ser Pro Pro Ala Ile Glu Asp
 180 185 190

Asn Asp Gly Lys Leu Gln Lys Phe Asp Val Lys Thr Gly Asp Ser Ile
 195 200 205

Arg Lys Gly Leu Thr Pro Gly Trp Tyr Asp Leu Asp Val Asn Gln His
 210 215 220

Val Ser Asn Val Lys Tyr Ile Gly Trp Ile Leu Glu Ser Met Pro Thr
 225 230 235 240

Glu Val Leu Glu Thr Gln Glu Leu Cys Ser Leu Thr Leu Glu Tyr Arg
 245 250 255

Arg Glu Cys Gly Arg Asp Ser Val Leu Glu Ser Val Thr Ser Met Asp
 260 265 270

Pro Ser Lys Val Gly Asp Arg Phe Gln Tyr Arg His Leu Leu Arg Leu
 275 280 285

Glu Asp Gly Ala Asp Ile Met Lys Gly Arg Thr Glu Trp Arg Pro Lys
 290 295 300

Asn Ala Gly Thr Asn Gly Ala Ile Ser Thr Gly Lys Thr
 305 310 315

<210> SEQ ID NO 13
 <211> LENGTH: 900
 <212> TYPE: DNA
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: CpFatB1.2 M4-287 Mutant

<400> SEQUENCE: 13

atggttgacc gcaaatctaa acgtccgagc atgctgatgg atagcttcgg tctggaacgt 60
 gttgtccagg atggtctggt tttccgccaa tccttctcca ttcgtagcta cgaaatctgt 120
 gcggaccgca cggcgagcat ggagaccgtg atgaaccacg ttcaggaaac cagcctgaac 180
 cagtgttaagt ccctcggcct gctggacgac ggtttcggtc gctcaccgga aatgtgtaaa 240
 cgcgacctga tttgggttgt aaccggtatg aaaattatgg taaaccgtta tccgacctgg 300
 ggcgacacca ttgaagtctc cacctggctg tcccagtcgg gtaaaaattgg catgggtcgc 360
 gactggctga ttagcgattg taataccggt gaaatactgg ttcgtgcaac ctcagtttac 420
 gccatgatga accagaaaac gcgtcgcttc tctaaactgc cccacgaagt acgccaggag 480
 ttcgcgccgc acttcttaga ctccccgcgg gccattgaag acaacgacgg taaactgcag 540

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aaattcgatg tgaaaaccgg tgattccatc cgcaaaggtc tgaccccggg ctggtacgac   600
ctggacgtaa accagcacgt ttccaatgtc aaatatatcg gttggatcct ggaatccatg   660
cggaccgaag ttctggaaac ccaagaactt tgcagcctga ctctogaata cegtcgcgaa   720
tgcggtcgcg actcagtact tgagagcgta acctccatgg acccgtctaa agttggtgat   780
cgttttcagt atcgatcatc gctgcgctctg gaagatggtg cggacattat gaagggccgt   840
accgaatggc gtccgaaaaa cgcgggcact aatggtgcta tctccactgg gaaaacataa   900

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<210> SEQ ID NO 14
<211> LENGTH: 299
<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: CpFatB1.2 M4-287 Mutant

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<400> SEQUENCE: 14

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Met Phe Asp Arg Lys Ser Lys Arg Pro Ser Met Leu Met Asp Ser Phe
 1           5           10           15
Gly Leu Glu Arg Val Val Gln Asp Gly Leu Val Phe Arg Gln Ser Phe
          20           25           30
Ser Ile Arg Ser Tyr Glu Ile Cys Ala Asp Arg Thr Ala Ser Met Glu
          35           40           45
Thr Val Met Asn His Val Gln Glu Thr Ser Leu Asn Gln Cys Lys Ser
          50           55           60
Ile Gly Leu Leu Asp Asp Gly Phe Gly Arg Ser Pro Glu Met Cys Lys
 65           70           75           80
Arg Asp Leu Ile Trp Val Val Thr Arg Met Lys Ile Met Val Asn Arg
          85           90           95
Tyr Pro Thr Trp Gly Asp Thr Ile Glu Val Ser Thr Trp Leu Ser Gln
          100          105          110
Ser Gly Lys Ile Gly Met Gly Arg Asp Trp Leu Ile Ser Asp Cys Asn
          115          120          125
Thr Gly Glu Ile Leu Val Arg Ala Thr Ser Val Tyr Ala Met Met Asn
          130          135          140
Gln Lys Thr Arg Arg Phe Ser Lys Leu Pro His Glu Val Arg Gln Glu
          145          150          155          160
Phe Ala Pro His Phe Leu Asp Ser Pro Pro Ala Ile Glu Asp Asn Asp
          165          170          175
Gly Lys Leu Gln Lys Phe Asp Val Lys Thr Gly Asp Ser Ile Arg Lys
          180          185          190
Gly Leu Thr Pro Gly Trp Tyr Asp Leu Asp Val Asn Gln His Val Ser
          195          200          205
Asn Val Lys Tyr Ile Gly Trp Ile Leu Glu Ser Met Pro Thr Glu Val
          210          215          220
Leu Glu Thr Gln Glu Leu Cys Ser Leu Thr Leu Glu Tyr Arg Arg Glu
          225          230          235          240
Cys Gly Arg Asp Ser Val Leu Glu Ser Val Thr Ser Met Asp Pro Ser
          245          250          255
Lys Val Gly Asp Arg Phe Gln Tyr Arg His Leu Leu Arg Leu Glu Asp
          260          265          270
Gly Ala Asp Ile Met Lys Gly Arg Thr Glu Trp Arg Pro Lys Asn Ala
          275          280          285
Gly Thr Asn Gly Ala Ile Ser Thr Gly Lys Thr
          290          295

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<210> SEQ ID NO 15
<211> LENGTH: 1812
<212> TYPE: DNA
<213> ORGANISM: Mycobacterium tuberculosis

<400> SEQUENCE: 15

atgagcgatt attatggtgg tgcacatacc accgttcgtc tgattgatct ggcaaccctg      60
atgcctcgtg ttctggcaga tacaccggtt attgttcggt gtgcaatgac cggctctgctg     120
gcacgtccga atagcaaagc aagcattggc accgtttttc aggatcgtgc agcacgttat     180
ggtgatcgtg tttttctgaa atttggatgat cagcaactga cctatcgtga tgcaaatgca     240
accgcaaate gttatgcagc agtgcctggc gcacgtgggt ttggtccggg tgatggtggt     300
ggtattatgc tgcgtaatag cccgagcacc gttctggcaa tgctggcaac cgttaaatgt     360
ggtgccattg caggtatgct gaattatcat cagcgtgggt aagttctggc acatagcctg     420
ggcctgctgg atgcaaaagt tctgattgca gaaagcgatc tggttagcgc agttgcagaa     480
tgcggtgcaa gccgtggctg tgttgccggt gatgttctga ccgttgaaga tgttgaacct     540
tttgcaacca ccgctccggc aaccaatccg gcaagcga ggcgagttca ggcaaaagat     600
accgcctttt atatctttac cagcggcacc accggttttc cgaaagcaag cgttatgacc     660
catcatcggt ggctgcgtgc actggcagtt tttggtggta tgggtctgcg tctgaaaggt     720
agcgatcccc tgtatagctg tctgccgctg taccataata atgcaactgac cgtggcagtt     780
tccagcgtga ttaatagcgg tgcaaacctg gcaactggta aaagttttag cgcaagccgt     840
ttttgggatg aagttattgc aaatcgtgca accgcctttg tgtatattgg tgaattttgt     900
cgctatctgc tgaatcagcc tgcaaaaaccg accgatcgtg cacatcaggt tcgtgttatt     960
tgtggtaatg gcctgcgtcc ggaaactctg gatgaattta ccaccggtt ttggtgttgc     1020
cgtgtttgtg agttttatgc agccagcga ggtaatagcg cctttattaa catttttaac     1080
gttccgcgta ccgcaaggct tagcccgatg ccgctggcat ttgttgaata tgatctggat     1140
accggtgatc cgctgcgtga tgcgagcggc cgtgttcgct gtgtgccgga tggatgaaccg     1200
ggtctgctgc tgagccgctg taatcgtctg caaccgtttg atggttatac cgatccggtt     1260
gcaagcgaaa aaaaactggt tcgtaatgca ttctgctgat gcgattgttg gtttaataca     1320
ggtgatgta tgagtcgca gggatggtgc catgcagcct ttgttgatcg tctgggtgat     1380
acctttcgtt ggaaaggtga aaatgttgc accaccagc ttgaagcagc actggcaagc     1440
gatcagaccg tggagaatg taccgtttat ggtgtgcaga ttctctgtac cgggtggtcgt     1500
gccggtatgg cagcaattac cctgcgtgcc ggtgcagaat ttgatggtca ggcactggca     1560
cgcaccgtgt atggtcatct gcttggttat gcaactgctc tgtttgttcg tgtgtgggt     1620
agcctggcgc ataccacaac ctttaaaagc cgtaaagttg aactgcgtaa tcaggcctat     1680
ggtgcagata ttgaagatcc gctgatgta ctggcaggtc cggatgaagg ttatgttccg     1740
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caccatcatt aa                                                                1812

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<210> SEQ ID NO 16
<211> LENGTH: 603
<212> TYPE: PRT
<213> ORGANISM: Mycobacterium tuberculosis

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<400> SEQUENCE: 16

Met Ser Asp Tyr Tyr Gly Gly Ala His Thr Thr Val Arg Leu Ile Asp
 1 5 10 15
 Leu Ala Thr Arg Met Pro Arg Val Leu Ala Asp Thr Pro Val Ile Val
 20 25 30
 Arg Gly Ala Met Thr Gly Leu Leu Ala Arg Pro Asn Ser Lys Ala Ser
 35 40 45
 Ile Gly Thr Val Phe Gln Asp Arg Ala Ala Arg Tyr Gly Asp Arg Val
 50 55 60
 Phe Leu Lys Phe Gly Asp Gln Gln Leu Thr Tyr Arg Asp Ala Asn Ala
 65 70 75 80
 Thr Ala Asn Arg Tyr Ala Ala Val Leu Ala Ala Arg Gly Val Gly Pro
 85 90 95
 Gly Asp Val Val Gly Ile Met Leu Arg Asn Ser Pro Ser Thr Val Leu
 100 105 110
 Ala Met Leu Ala Thr Val Lys Cys Gly Ala Ile Ala Gly Met Leu Asn
 115 120 125
 Tyr His Gln Arg Gly Glu Val Leu Ala His Ser Leu Gly Leu Leu Asp
 130 135 140
 Ala Lys Val Leu Ile Ala Glu Ser Asp Leu Val Ser Ala Val Ala Glu
 145 150 155 160
 Cys Gly Ala Ser Arg Gly Arg Val Ala Gly Asp Val Leu Thr Val Glu
 165 170 175
 Asp Val Glu Arg Phe Ala Thr Thr Ala Pro Ala Thr Asn Pro Ala Ser
 180 185 190
 Ala Ser Ala Val Gln Ala Lys Asp Thr Ala Phe Tyr Ile Phe Thr Ser
 195 200 205
 Gly Thr Thr Gly Phe Pro Lys Ala Ser Val Met Thr His His Arg Trp
 210 215 220
 Leu Arg Ala Leu Ala Val Phe Gly Gly Met Gly Leu Arg Leu Lys Gly
 225 230 235 240
 Ser Asp Thr Leu Tyr Ser Cys Leu Pro Leu Tyr His Asn Asn Ala Leu
 245 250 255
 Thr Val Ala Val Ser Ser Val Ile Asn Ser Gly Ala Thr Leu Ala Leu
 260 265 270
 Gly Lys Ser Phe Ser Ala Ser Arg Phe Trp Asp Glu Val Ile Ala Asn
 275 280 285
 Arg Ala Thr Ala Phe Val Tyr Ile Gly Glu Ile Cys Arg Tyr Leu Leu
 290 295 300
 Asn Gln Pro Ala Lys Pro Thr Asp Arg Ala His Gln Val Arg Val Ile
 305 310 315 320
 Cys Gly Asn Gly Leu Arg Pro Glu Ile Trp Asp Glu Phe Thr Thr Arg
 325 330 335
 Phe Gly Val Ala Arg Val Cys Glu Phe Tyr Ala Ala Ser Glu Gly Asn
 340 345 350
 Ser Ala Phe Ile Asn Ile Phe Asn Val Pro Arg Thr Ala Gly Val Ser
 355 360 365
 Pro Met Pro Leu Ala Phe Val Glu Tyr Asp Leu Asp Thr Gly Asp Pro
 370 375 380
 Leu Arg Asp Ala Ser Gly Arg Val Arg Arg Val Pro Asp Gly Glu Pro
 385 390 395 400
 Gly Leu Leu Leu Ser Arg Val Asn Arg Leu Gln Pro Phe Asp Gly Tyr
 405 410 415

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Thr Asp Pro Val Ala Ser Glu Lys Lys Leu Val Arg Asn Ala Phe Arg
 420 425 430

Asp Gly Asp Cys Trp Phe Asn Thr Gly Asp Val Met Ser Pro Gln Gly
 435 440 445

Met Gly His Ala Ala Phe Val Asp Arg Leu Gly Asp Thr Phe Arg Trp
 450 455 460

Lys Gly Glu Asn Val Ala Thr Thr Gln Val Glu Ala Ala Leu Ala Ser
 465 470 475 480

Asp Gln Thr Val Glu Glu Cys Thr Val Tyr Gly Val Gln Ile Pro Arg
 485 490 495

Thr Gly Gly Arg Ala Gly Met Ala Ala Ile Thr Leu Arg Ala Gly Ala
 500 505 510

Glu Phe Asp Gly Gln Ala Leu Ala Arg Thr Val Tyr Gly His Leu Pro
 515 520 525

Gly Tyr Ala Leu Pro Leu Phe Val Arg Val Val Gly Ser Leu Ala His
 530 535 540

Thr Thr Thr Phe Lys Ser Arg Lys Val Glu Leu Arg Asn Gln Ala Tyr
 545 550 555 560

Gly Ala Asp Ile Glu Asp Pro Leu Tyr Val Leu Ala Gly Pro Asp Glu
 565 570 575

Gly Tyr Val Pro Tyr Tyr Ala Glu Tyr Pro Glu Glu Val Ser Leu Gly
 580 585 590

Arg Arg Pro Gln Gly His His His His His His
 595 600

<210> SEQ ID NO 17
 <211> LENGTH: 1701
 <212> TYPE: DNA
 <213> ORGANISM: Pseudomonas putida

<400> SEQUENCE: 17

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cagatgcggc tgacgtatcc acagctcaac gagcgcattg cccgcctggc caacgtgctg 180
accgagggcc gggtaaacgc cggtgacacc gtggcgggta tggactggga cagccatcgc 240
tacctggaat gcattgttcg catcccgatg atcggcgctg tgggtgcacac catcaacgtg 300
cgctgtcgc cggagcagat cctctacacc atgaaccatg ccgaagaccg cgtggtgctg 360
gtcaacagcg acttcgctcg cctgtaccag gccatcgccg ggcagctgac cactgtcgac 420
aagaccctgc tactgaccga tggcccggac aagactgccg aactgcccgg tctggtcggc 480
gagtatgagc agctgctggc tgctgccagc ccgcgctacg acttcccgga tttcgacgag 540
aattcggtgg ccaactacct ctacaccact ggcaccaccg gtaaccccaa gggcgtgat 600
ttcagtcacc gccagctggt gctgcacacc ctggccgagg cctcggtcac cggcagtatc 660
gacagcgtgc gcctgctggg cagcaacgat gtgtacatgc ccatcaccce gatgttccac 720
gtgcatgcct ggggcatccc ctacgctgcc accatgctcg gcatgaagca ggtgtaacca 780
gggcgctaag agccggacat gctggtaaac ctttggcgtg aagagaaggt cactttctcc 840
cactgcgtgc cgaccatcct gcagatgctg ctcaactgcc cgaacgccca ggggcaggac 900
ttcggcggct ggaagatcat catcggcggc agctcgtcga accgttcgct gtaccaggcc 960
gccctggcgc gggcatcca gctgaccgcc gcgtatggca tgcggaaac ctgcccgctg 1020
atctccggcg cacacctgaa cgatgaactg caggccggca gcgaggatga gcgcgtcact 1080
    
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taccgtatca aggccgggtgt gccgggtgccg ttggtcgaag cggccatcgt cgacggcgaa 1140
ggcaacttcc tgccccgcga tgggtgaacc cagggcgagc tggtaactgcg tgcgcccgtgg 1200
ctgaccatgg gctacttcaa ggagccggag aagagcgagg agctgtggca gggcggtgg 1260
ctgcacacog gtgacgtgc caccctogac ggcattgggt acatcgacat ccgogaccgc 1320
atcaaggatg tgatcaagac cgggtggcgag tgggtttcct cgctcgacct ggaagacctg 1380
atcagccgcc acccgccgt ggcggaagtg gcggtggtgg ggggtggcga cccgcagtgg 1440
ggtgagcgcc cgtttgccct gctggtggca cgtgacggcc acgatatcga cgccaaggcg 1500
ctgaaggaac acctcaagcc attcgtcgag caaggtcata tcaacaagtg ggcgattcca 1560
agccagatcg cccttgttac tgaattccc aagaccagtg tgggcaagct cgacaagaaa 1620
cgcattcgcc aggacatcgt ccagtggcag gccagcaaca gcgcttctct ttccacgttg 1680
catcaccatc accatcacta a 1701

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<210> SEQ ID NO 18

<211> LENGTH: 566

<212> TYPE: PRT

<213> ORGANISM: Pseudomonas putida

<400> SEQUENCE: 18

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Met Leu Gln Thr Arg Ile Ile Lys Pro Ala Glu Gly Ala Tyr Ala Tyr
1           5           10          15
Pro Leu Leu Ile Lys Arg Leu Leu Met Ser Gly Ser Arg Tyr Glu Lys
20          25          30
Thr Arg Glu Ile Val Tyr Arg Asp Gln Met Arg Leu Thr Tyr Pro Gln
35          40          45
Leu Asn Glu Arg Ile Ala Arg Leu Ala Asn Val Leu Thr Glu Ala Gly
50          55          60
Val Lys Ala Gly Asp Thr Val Ala Val Met Asp Trp Asp Ser His Arg
65          70          75          80
Tyr Leu Glu Cys Met Phe Ala Ile Pro Met Ile Gly Ala Val Val His
85          90          95
Thr Ile Asn Val Arg Leu Ser Pro Glu Gln Ile Leu Tyr Thr Met Asn
100         105         110
His Ala Glu Asp Arg Val Val Leu Val Asn Ser Asp Phe Val Gly Leu
115         120         125
Tyr Gln Ala Ile Ala Gly Gln Leu Thr Thr Val Asp Lys Thr Leu Leu
130         135         140
Leu Thr Asp Gly Pro Asp Lys Thr Ala Glu Leu Pro Gly Leu Val Gly
145         150         155         160
Glu Tyr Glu Gln Leu Leu Ala Ala Ala Ser Pro Arg Tyr Asp Phe Pro
165         170         175
Asp Phe Asp Glu Asn Ser Val Ala Thr Thr Phe Tyr Thr Thr Gly Thr
180         185         190
Thr Gly Asn Pro Lys Gly Val Tyr Phe Ser His Arg Gln Leu Val Leu
195         200         205
His Thr Leu Ala Glu Ala Ser Val Thr Gly Ser Ile Asp Ser Val Arg
210         215         220
Leu Leu Gly Ser Asn Asp Val Tyr Met Pro Ile Thr Pro Met Phe His
225         230         235         240
Val His Ala Trp Gly Ile Pro Tyr Ala Ala Thr Met Leu Gly Met Lys
245         250         255

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Gln Val Tyr Pro Gly Arg Tyr Glu Pro Asp Met Leu Val Lys Leu Trp
260 265 270

Arg Glu Glu Lys Val Thr Phe Ser His Cys Val Pro Thr Ile Leu Gln
275 280 285

Met Leu Leu Asn Cys Pro Asn Ala Gln Gly Gln Asp Phe Gly Gly Trp
290 295 300

Lys Ile Ile Ile Gly Gly Ser Ser Leu Asn Arg Ser Leu Tyr Gln Ala
305 310 315 320

Ala Leu Ala Arg Gly Ile Gln Leu Thr Ala Ala Tyr Gly Met Ser Glu
325 330 335

Thr Cys Pro Leu Ile Ser Ala Ala His Leu Asn Asp Glu Leu Gln Ala
340 345 350

Gly Ser Glu Asp Glu Arg Val Thr Tyr Arg Ile Lys Ala Gly Val Pro
355 360 365

Val Pro Leu Val Glu Ala Ala Ile Val Asp Gly Glu Gly Asn Phe Leu
370 375 380

Pro Ala Asp Gly Glu Thr Gln Gly Glu Leu Val Leu Arg Ala Pro Trp
385 390 395 400

Leu Thr Met Gly Tyr Phe Lys Glu Pro Glu Lys Ser Glu Glu Leu Trp
405 410 415

Gln Gly Gly Trp Leu His Thr Gly Asp Val Ala Thr Leu Asp Gly Met
420 425 430

Gly Tyr Ile Asp Ile Arg Asp Arg Ile Lys Asp Val Ile Lys Thr Gly
435 440 445

Gly Glu Trp Val Ser Ser Leu Asp Leu Glu Asp Leu Ile Ser Arg His
450 455 460

Pro Ala Val Arg Glu Val Ala Val Val Gly Val Ala Asp Pro Gln Trp
465 470 475 480

Gly Glu Arg Pro Phe Ala Leu Leu Val Ala Arg Asp Gly His Asp Ile
485 490 495

Asp Ala Lys Ala Leu Lys Glu His Leu Lys Pro Phe Val Glu Gln Gly
500 505 510

His Ile Asn Lys Trp Ala Ile Pro Ser Gln Ile Ala Leu Val Thr Glu
515 520 525

Ile Pro Lys Thr Ser Val Gly Lys Leu Asp Lys Lys Arg Ile Arg Gln
530 535 540

Asp Ile Val Gln Trp Gln Ala Ser Asn Ser Ala Phe Leu Ser Thr Leu
545 550 555 560

His His His His His His
565

What is claimed is:

1. An unnatural, mutated protein comprising an amino acid sequence at least about 80% identical to positions 19-317 of SEQ ID NO:4, wherein:
 - the amino acid sequence comprises:
 - a residue other than asparagine at a position corresponding to position 28 of SEQ ID NO:4; and
 - a residue other than isoleucine at a position corresponding to position 65 of SEQ ID NO:4;
 - the protein lacks an N-terminal portion having an amino acid sequence identical to positions 1-18 of SEQ ID NO:4; and
 - the protein exhibits thioesterase activity.
2. The protein of claim 1, wherein the amino acid sequence at least about 80% identical to positions 19-317 of SEQ ID NO:4 comprises:
 - a serine, cysteine, threonine, tyrosine, or glutamine at the position corresponding to position 28 of SEQ ID NO:4; and
 - a methionine, alanine, leucine, phenylalanine, valine, proline, or glycine at the position corresponding to position 65 of SEQ ID NO:4.
3. The protein of claim 2, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.
4. The protein of claim 1, wherein the amino acid sequence at least about 80% identical to positions 19-317 of SEQ ID NO:4 comprises:
 - a serine at the position corresponding to position 28 of SEQ ID NO:4; and

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a methionine at the position corresponding to position 65 of SEQ ID NO:4.

5. The protein of claim 4, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

6. The protein of claim 1, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

7. The protein of claim 1, wherein the amino acid sequence at least about 80% identical to positions 19-317 of SEQ ID NO:4 is an amino acid sequence at least about 85% identical to positions 19-317 of SEQ ID NO:4.

8. The protein of claim 7, wherein the amino acid sequence at least about 85% identical to positions 19-317 of SEQ ID NO:4 comprises:

a serine, cysteine, threonine, tyrosine, or glutamine at the position corresponding to position 28 of SEQ ID NO:4; and

a methionine, alanine, leucine, phenylalanine, valine, proline, or glycine at the position corresponding to position 65 of SEQ ID NO:4.

9. The protein of claim 8, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

10. The protein of claim 7, wherein the amino acid sequence at least about 85% identical to positions 19-317 of SEQ ID NO:4 comprises:

a serine at the position corresponding to position 28 of SEQ ID NO:4; and

a methionine at the position corresponding to position 65 of SEQ ID NO:4.

11. The protein of claim 10, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

12. The protein of claim 7, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

13. The protein of claim 1, wherein the amino acid sequence at least about 80% identical to positions 19-317 of SEQ ID NO:4 is an amino acid sequence at least about 90% identical to positions 19-317 of SEQ ID NO:4.

14. The protein of claim 13, wherein the amino acid sequence at least about 90% identical to positions 19-317 of SEQ ID NO:4 comprises:

a serine, cysteine, threonine, tyrosine, or glutamine at the position corresponding to position 28 of SEQ ID NO:4; and

a methionine, alanine, leucine, phenylalanine, valine, proline, or glycine at the position corresponding to position 65 of SEQ ID NO:4.

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15. The protein of claim 14, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

16. The protein of claim 13, wherein the amino acid sequence at least about 90% identical to positions 19-317 of SEQ ID NO:4 comprises:

a serine at the position corresponding to position 28 of SEQ ID NO:4; and

a methionine at the position corresponding to position 65 of SEQ ID NO:4.

17. The protein of claim 16, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

18. The protein of claim 13, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

19. The protein of claim 1, wherein the amino acid sequence at least about 80% identical to positions 19-317 of SEQ ID NO:4 is an amino acid sequence at least about 95% identical to positions 19-317 of SEQ ID NO:4.

20. The protein of claim 19, wherein the amino acid sequence at least about 95% identical to positions 19-317 of SEQ ID NO:4 comprises:

a serine, cysteine, threonine, tyrosine, or glutamine at the position corresponding to position 28 of SEQ ID NO:4; and

a methionine, alanine, leucine, phenylalanine, valine, proline, or glycine at the position corresponding to position 65 of SEQ ID NO:4.

21. The protein of claim 20, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

22. The protein of claim 19, wherein the amino acid sequence at least about 95% identical to positions 19-317 of SEQ ID NO:4 comprises:

a serine at the position corresponding to position 28 of SEQ ID NO:4; and

a methionine at the position corresponding to position 65 of SEQ ID NO:4.

23. The protein of claim 22, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

24. The protein of claim 19, wherein the protein comprises, relative to a protein consisting of an amino acid sequence of SEQ ID NO:4, an N-terminal truncation of positions 1-18 of SEQ ID NO:4.

* * * * *