An engineered microbe that contains a designed platform for the conversion of one-carbon substrates to chemical products is described. The designed platform embodies a new metabolic architecture that consolidates carbon fixation, central metabolism, and product synthesis into a single pathway. This is made possible by the key finding that 2-hydroxyacyl-CoA lyase, an enzyme in the co-oxidation pathway, is capable of catalyzing the C—C bond formation between formyl-CoA and aldehydes of different chain lengths, allowing for the elongation of the carbon backbone of said aldehyde by one-carbon units. These novel microbes present an opportunity for the production of chemicals from single-carbon feedstocks such as carbon dioxide, carbon monoxide, formate, formaldehyde, methanol or methane.

**Specification includes a Sequence Listing.**
Methane monooxygenase
NADH, H⁺, O₂ → NAD⁺, H₂O
CH₄ → methanol

Methanol dehydrogenase
NAD⁺ → NADH, H⁺
formaldehyde

Acylation aldehyde dehydrogenase
CoA, NAD⁺ → NADH, H⁺

Formate dehydrogenase; Electrocatalysis
CO₂ + H₂ → H⁺

Acetyl-CoA Synthetase
CoA, ATP + H₂O, AMP, PPI → formyl-CoA

Formyl-CoA

Formate kinase
ATP + ADP → P + CoA

Phosphate formyltransferase
formyl-phosphate

FIGURE 3
To elongation module

Acetyl-CoA

Acetyl-CoA reductase

Acyl-CoA

NADH, H^+

NAD^+, CoA

Enoyl-CoA hydratase

H_2O

3-hydroxyacyl-CoA dehydrogenase

3-hydroxyacyl-CoA thiolase

3-oxo-(C_4)-acyl-CoA

Fatty acids

Aldehydes

Alcohols

Alkanes

Etc.
FIGURE 6

An alcohol

An alcohol

A carboxylic acid

An aldehyde

An amine

An alkane

From elongation or termination modules

An acyl-CoA
octadecanoic acid

2-hydroxyoctadecanoic acid

2-hydroxyoctadecanoyl-CoA

2-hydroxyphytanoyl-CoA lyase

formyl-CoA

heptadecanol

aldehyde dehydrogenase

heptadecanoic acid

activation

β-oxidation

FIGURE 7
FIGURE 9

HACL1 ISOFORM 1 (Q9UJ83-1) SEQ ID NO. 1

MPDSNFAERS EEQVSGAKVI AQALKTDQDVE YIFGIVGIPV TEIAIAAQQL GIKYIGMRNE QAACAYAAASI GYLTSRPGVC LVVSGPGLIH ALGGMANANM NCWPLLVIGG SSERNQETMG AFQEPQPVEA CRLYKTFSAR PSSIEAPFV TEEAKVRSIIY GRPGACYVDDI PADFVNLQVN VNSIKYMER CVNPSSAAMET SAVCTAASVI RNAKQPLLII GKGAAAYAHAE ESIIKLVQY KLPLFLPTMPG KGVPDNHPY CVGAARSRAL QFADVIYLFQ ARNWNILHIFG LPPRYQPDVK FIQVDICAEEL LGNNVKKPAVT LLGNIHAVTK QLLEELDKTP WQYPPEKWW KTRLREKMKSN EAASKELASK KSLPMNYYTV FYHVQEQLPR DCFVVSEGAVN TMDIGRQTVLQ YNLPRHRIDQ GTFQGTDYGLQ GFAIAVA AAVKA DRRSPGQWII CVEGDSAFGQ SGMEVETICR YNLPIILLVY NNNGIYQGFQD TDWQVQTVLPD CLPPNSHYEQ VMFAGGKGGY FVQTPQEQPK SLRQSLADTT KPSLIMEIIE PQATRKAQDFH HW1 LTRSNM 578

HACL1 ISOFORM 2 (Q9UJ83-2) SEQ ID NO. 2

MPDSNFAERS EEQVSGAKVI AQALKTDQDVE YIFGIVGIPV TEIAI AQQQL GIKYIGMRNE QAACAYAAASI GYLTSRPLL VIGGSERNQEE TMGAQFQQSSP VEACRLYTKF SARPSSEIA PVIEKAVRVS SYYGRPGAC VDIPAFVNL QVNVNSIKYM ERCMSSPISI AETSVACTAA SVRNAKQPLL LIIGKAAAYA HAEESIKLVQ EQKLPQFLPT PMGKVVPDQ HPYCGLVAARS RALQFAADVIV LFGARLNWL HIQFGPQYQDP DKFQIVDIE AELGNVJNIK VTKQLEELID KTPWQYPPES KWWKTLREKMK SNEAASKEL ASKSLPMNY YTVFYHVQEQI LPRDCVFVSE GANTMDIGRT VLQNYLPRHR LDAGTFQTMG VGLGFAAIAA VAKDRPSQPG WICVGEQDS QDGSSMEVET ICRYNLPHI LQVNNIGYQ GFDTDWQVTK LKFOQDATAVQ PPMCLPNSH YEIQVMTAFGG KGYFQTPPEE LQKSLRQLSA DLTPKSLINI MEIPQATRKAQ DQFH HW1 LTRSNM 551

HACL1 ISOFORM 3 (Q9UJ83-3) SEQ ID NO. 3

MPDSNFAERS EEQVSGAKVI AQALKTDQDVE YIFGIVGIPV TEIAIAAQQL GIKYIGMRNE QAACAYAAASI GYLTSRPGVC LVVSGPGLIH ALGGMANANM NCWYMERCS PPISMAETS A VCTAASVIRN AKQPLLIIIG GAAAYAAEES IKKLVEQYYKL PFLPPTMGK VVDNHPLQ ARSARRALQF ADVIQEFLPQ LNWILHGGLP PRYPQDVKFI QVDCAEELG NNVKPAVIL GNIHAVTKQ LLEELDKTPQ WPPESKWWK LREIKMSN EAASKLE KSLPMNYYT VHYHVQEQLP DRC FVVSEGAN TM DIGRTVQNY IPRHRDAGT FQTMGVGLFG AIAAAVAKD RSPQQWIYCV EGDSAFCFEG MVEETICRYN LPIILLVNN NGIQYGFDTD WDQVQLFQD ATAVPQMLC LPNSHYEQVQ TADFGGKGYFQ QTEPEQLQSL RQLSLADTKP SLNIMIEPQ ATRKADDFHW LTRSNM 496

HACL ISOFORM 4 (Q9UJ83-4) SEQ ID NO. 4

MPDSNFAERS EEQVSGAKVI AQALKTDQDVE YIFGIVGIPV TEIAIAAQQL GIKYIGMRNE QAACAYAAASI GYLTSRPGVC LVVSGPGLIH ALGGMANANM NCWPLLLIGG SSERNQETMG AFQEPQPAVR SISYGRPGAC YVDIPADFYV LQNVNSIKY
FIGURE 9 CON'T

MERCMSPPIS MAETSAVCTA ASVIRNAKQP LLIGKGAAY AHAEEISKKL VEQYKLPFLP TPMGKGVVPD NHPYCVGAAR SRALQFADVVI VLGARLNWI LHFGLPFRYQ PDVKFIQVDI CAEELGNNVK PAVTLLGNIH AVTKQELASK KSLPMNYYTV FYHVQEQLPR DCFVVSEGANTMDIGRTVLQ NYLPRHRLDAGTFGTMGVGL GFAIAAVVAKDRSPGQWIICVEGDSAGF
SGMEVETICRYNLPIILLVV NNNGIYQGFD TDTWKEMLKF QDATAVVPMP CLLPNSHYEQ VMATFGGBKGY FVQTPEELQK
SLRQSLADTTKPSLINIMIE PQATRKAQDF HWLTRSNM 518
FIGURE 12

1: Ladder
2: Whole Cell
3: Crude Extract
4: Purified Lmo1179 Elution 1
5: Purified Lmo1179 Elution 2
FIGURE 13

Lmo1179 Assay - Excess CoA

Absorbance

Time (H:MM:SS)

△ No Enzyme  ■ No CoA  ● CoA

FIGURE 13
A peak corresponding to the expected size of formyl-CoA (796) was present in the enzyme assay containing Lmo1179.

FIGURE 14

No enzyme control

Lmo1179 sample
FIGURE 16
FIGURE 22A
FIGURE 22B

- pCDFDuet-1-P1-nll6-ecOxc - 5482nt
- 118...1813 ecOxc
- 1915...1984 LacO
- 1834...1934 LacO
- 2067...2111 S-tag
- 3,25 LacO
- 83...100 βHis
- T7 5466...3
- Lac 5219...4263
- Amp prom 3242...3214
Eadie-Hofstee plot for egTER

\[ y = -114.77x + 0.4155 \]
\[ R^2 = 0.86509 \]

FIGURE 25

MhpF Reduction of Butyryl-CoA

\[ y = -0.0007350x + 0.3521242 \]
\[ R^2 = 0.9566336 \]
\[ y = -0.0016880x + 0.3576305 \]
\[ R^2 = 0.9741849 \]
\[ y = -0.0023907x + 0.3550704 \]
\[ R^2 = 0.9933811 \]
\[ y = -0.0061428x + 0.3562672 \]
\[ R^2 = 0.9739360 \]
\[ y = -0.0026306x + 0.3555216 \]
\[ R^2 = 0.9851285 \]

FIGURE 26
BIOSYNTHESIS OF PRODUCTS FROM 1-CARBON COMPOUNDS

PRIOR RELATED APPLICATIONS

This application claims priority to U.S. Ser. No. 62/069,850, titled SYNTHE TESIS FROM 1-CARBON COMPOUNDS, filed Oct. 29, 2014 and incorporated by reference herein in its entirety for all purposes.

FEDERALLY SPONSORED RESEARCH STATEMENT

Not Applicable.

FIELD OF THE DISCLOSURE

The invention relates to engineered microorganisms that can make various chemicals for industrial use. In particular, the biological production of multi-carbon compounds directly from single carbon compounds without the need for central metabolism.

BACKGROUND OF THE DISCLOSURE

In the past, much of our energy and chemicals have derived from fossilized carbon resources, such as petroleum and coal. However, concerns about climate change, political instability, and depletion and cost of petroleum resources have recently ignited interest in the establishment of a bio-based industry and a number of strategies are being pursued to achieve the sustainable production of fuels and chemicals. One of the most promising approaches is the use of microorganisms to convert biomass into the desired product. The use of these feedstocks, however, is hindered by the difficulty of accessing the resources, their high price, and limited availability. One-carbon compounds, including CO₂, CO, formate, methanol, and methane have recently emerged as alternative renewable feedstocks. Conversion of these substrates to fuels and chemicals not only offers an attractive alternative in terms of process economics but also will further contribute to reduce the levels of greenhouse gases like CO₂ and methane.

Microbial catalysts have been designed and engineered to synthesize products of interest using feedstocks as diverse as sugars, glycerol, carbon dioxide, carbon monoxide, formate, methanol, and methane. For the case of 1-carbon feedstocks, such conversions are made possible by a general network of metabolic pathways that are organized as shown in FIG. 1 and include specialized pathways for carbon fixation, central metabolism, and product synthesis. This type of metabolic architecture has been exploited in all metabolic engineering efforts conducted to date to develop microbes for industrial applications.

This disclosure describes an alternative platform for the bioconversion of 1-carbon substrates, which consists of a single engineered metabolic pathway that allows for the direct synthesis of chemical products from one-carbon compounds using a ‘bottom up’ integrated design. The engineered pathway defines a new metabolic architecture that consolidates carbon fixation, central metabolism, and product synthesis into single pathway (FIG. 2). The pathway uses single carbon extension units, which bypasses the need for the production of common metabolic intermediates and allows for elongation of a carbon backbone iteratively in single carbon increments.

This is in contrast to current approaches, which require first the fixation of single carbon compounds to metabolic intermediates before elongation (FIG. 1) and which commonly elongate a carbon backbone by two or more carbons at a time. This synthetic platform was engineered from the ‘bottom-up’ and provides potential advantages over current ‘top-down’ methods, which suffer from difficulty of engineering and inefficiencies arising from the use of assimilating pathways to first produce metabolic intermediates before carbon chain elongation and product formation.

SUMMARY OF THE DISCLOSURE

Engineering biology from the ‘bottom-up’ has gained momentum recently. In particular, the field of synthetic biology has had particular focus on using this ‘bottom-up’ approach, with the end goal of designing systems with tunable control and well-defined response. Synthetically developed pathways can have advantages over natural pathways, for example by allowing for exogenous control over pathway enzyme expression. This potentially decouples the pathway performance with the organism’s complex and poorly understood regulatory mechanisms. Further decoupling can be achieved by synthetic pathways when designed to avoid the usage of central metabolic pathways, such as those generating acetyl-CoA, allowing the pathway performance to be independent of the organism’s central metabolism, which is also subject to regulation and interaction with other pathways. A bottom up design approach can thus allow for the creation of a pathway that is independent from the host system. Here, the bottom up design of a pathway capable of the conversion of C1 molecules to longer chain products is described.

The development of a synthetic pathway that consolidates fixation of single carbon compounds into the synthesis of products is described. The pathway allows for direct synthesis of products from single carbon compounds, avoiding the need to produce the cells typical intermediates and use of central metabolic pathways, as required by the existing approach. This synthetic pathway is inspired on the catalytic α-oxidation pathway (one carbon shortening of fatty acids) and based on the key recent finding in our laboratory that the enzyme hydroxyl-acyl-CoA lyase (HACl, part of the α-oxidation pathway) is reversible and able to catalyze C–C bond formation between formyl-CoA and an aldehyde. This mechanism for C–C bond formation had not been previously reported in the literature and allows for carbon-chain elongation by one-carbon units.

The pathway accomplishes four main functions:

First: single carbon compounds are converted to an active ‘extender’ form that can be incorporated into a carbon skeleton. This is done by enzymatically converting a one-carbon unit into an activated CoA derivative, and there are a variety of enzymes available to do this.

Second: the pathway allows for these single carbon extenders to iteratively elongate the carbon skeleton to desired length. An enzyme from the alpha oxidation pathway (hydroxyl-acyl-CoA lyases or HACLs) is used to catalyze the key condensation C–C bond formation reaction, being driven herein to run in the reverse direction. A series of dehydration and reduction reactions then converts the molecule to a form suitable for successive rounds of elongation.
[0013] Third: In addition, if the initial priming aldehyde has more than one carbon, a regeneration pathway to regenerate the starting aldehyde molecule is needed. However, if the starting aldehyde is formaldehyde or a single turn of the cycle is sufficient, then this functionality can be omitted.

[0014] Fourth: there is a method from which the desired products are synthesized from the elongated carbon skeleton. A large variety of termination enzymes can pull intermediates out of the cycle and if desired those can be further modified enzymatically.

[0015] The pathway described herein drew inspiration from the catastrophic α-oxidation pathway (one carbon shortening of fatty acids) and is based on the key recent finding in our laboratory that the enzyme 2-hydroxyacyl-CoA lyase (HACL, part of the α-oxidation pathway) is reversible and able to catalyze C—C bond formation between formyl-CoA and an aldehyde.

[0016] The synthetic pathway disclosed in this invention serves as a platform for the generation of valuable chemical products from less valuable and more abundant single carbon compounds, rather than from readily depleting fossil sources, such as petroleum. Methane, for example, can be converted into liquid “drop in” fuels for use in the current transportation infrastructure. Carbon dioxide can be sequestered and converted into useful products. The methods, materials and systems herein thus allow for single carbon molecules of varying reduction levels, from methane to carbon dioxide, to be assimilated directly into an elongating carbon backbone.

[0017] In the synthetic pathway herein disclosed, single carbon molecules are first activated to formyl-CoA, which can be directly ligated with an aldehyde by the enzyme 2-hydroxyacyl-CoA lyase. The resulting 2-hydroxyacyl-CoA is one carbon longer than the originating aldehyde and undergoes a series of reactions to return it to an aldehyde, allowing for the further elongation of the carbon backbone. Intermediates of the above reactions can be diverted from the cycle with a termination enzyme to generate valuable chemical products, such as (but not limited) to carboxylic acids, alcohols, or alkaines of varying chain lengths and functionalities.

[0018] The process involves performing traditional cultures using industrial organisms (such as E. coli or S. cerevisiae or Methylococcus capsulatus or Pichia pastoris) that convert single carbon compounds, such as methane, methanol, formaldehyde, formate, carbon monoxide, or carbon dioxide, into chemical products. These organisms are considered workhorses of modern biotechnology, and are easy to genetically engineer, and scale up for industrial production levels.

[0019] In another embodiment, however, one could provide any intermediate in the designed synthetic pathway as a starting compound, and need not make the entire molecule from scratch, extending the starting molecule once converted to aldehyde by one carbon, and pulling out any intermediate in the cycle using the termination module. Such embodiments may be useful where an intermediate is plentiful or a waste product from another process, and thus very inexpensive. Combined therewith, the regeneration module can be omitted, if the starting aldehyde is formaldehyde or a single turn of the cycle is sufficient.

[0020] It is also possible to use only a portion of the designed synthetic pathway, again if suitable given the starting molecule and final desired product. Thus, if just adding a single carbon will suffice, it may be possible to use just the HACL protein, as described herein. As yet another alternative, one or more downstream enzymes in the designed synthetic pathway are used in combination with the HACL. However, in most instances the most versatility comes from including all 4 modules.

[0021] The modules in a living system are generally made by transforming the microbe with an expression vector encoding one or more of the proteins, but the genes can also be added to the chromosome by recombining, homologous recombination, and similar techniques. Where the needed protein is endogenous, as is the case in some instances, it may suffice as is, but is usually overexpressed for better functionality and control over the level of active enzyme.

[0022] The use of the word “a” or “an” when used in conjunction with the term “comprising” in the claims or the specification means one or more than one, unless the context dictates otherwise.

[0023] The term “about” means the stated value plus or minus the margin of error of measurement or plus or minus 10% if no method of measurement is indicated.

[0024] The use of the term “or” in the claims is used to mean “and/or” unless explicitly indicated to refer to alternatives only or if the alternatives are mutually exclusive.

[0025] The terms “comprise”, “have”, “include” and “contain” (and their variants) are open-ended linking verbs and allow the addition of other elements when used in a claim.

[0026] As used herein, the expressions “microorganism,” “microbe,” “strain” and the like may be used interchangeably and all such designations include their progeny. It is also understood that all progeny may not be precisely identical in DNA content, due to deliberate or inadvertent mutations. Mutant progeny that have the same function or biological activity as screened for in the originally transformed cell are included. Where distinct designations are intended, it will be clear from the context.

[0027] Reference to proteins herein can be understood to include reference to the gene encoding such protein. Thus, a claimed “permease” protein can include the related gene encoding that permease. However, it is preferred herein to refer to the protein by standard name per ecoliwiki or HUGO since both enzymatic and gene names have varied widely, especially in the prokaryotic arts.

[0028] Once an exemplary protein is obtained, many additional examples of proteins of similar activity can be identified by BLAST search. Further, every protein record is linked to a gene record, making it easy to design overexpression vectors. Many of the needed enzymes are already available in vectors, and can often be obtained from cell repositories or from the researchers who cloned them. But, if necessary, new clones can be prepared based on available sequence information using RT-PCR techniques. Thus, it should be easily possible to obtain all of the needed enzymes for overexpression.

[0029] Another way of finding suitable enzymes/proteins for use in the invention is to consider other enzymes with the same EC number, since these numbers are assigned based on the reactions performed by a given enzyme. An enzyme that thus be obtained, e.g., from AddGene or from the author of the work describing that enzyme, and tested for functionality as described herein.

[0030] Understanding the inherent degeneracy of the genetic code allows one of ordinary skill in the art to design
multiple nucleotides that encode the same amino acid sequence. NCBI® provides codon usage databases for optimizing DNA sequences for protein expression in various species. Using such databases, a gene or cDNA may be "optimized" for expression in E. coli, yeast, algal or other species using the codon bias for the species in which the gene will be expressed.

**0031** Initial cloning experiments have proceeded in E. coli for convenience since most of the required genes are already available in plasmids suitable for bacterial expression, but the addition of genes to bacteria is of nearly universal applicability. Recombinant methods were invented in the 1970’s and are now so commonplace, even school children perform genetic engineering experiments using bacteria. Such species include e.g., Bacillus, Streptomyces, Azotobacter, Trichoderma, Rhizobium, Pseudomonas, Micrococcus, Nitrobra, Proteus, Lactobacillus, Pediococcus, Lactococcus, Salmonella, Streptococcus, Paracoccus, Methanosarcina, and Methylococcus, or any of the completely sequenced bacterial species. Indeed, hundreds of bacterial genomes have been completely sequenced, and this information greatly simplifies both the generation of vectors encoding the needed genes, as well as the planning of a recombinant engineering protocol. Such species are listed along with links at http://en.wikipedia.org/wiki/List_of_sequenced_bacterial_genomes.

**0032** Additionally, yeast are a common species used for microbial manufacturing, and many species can be successfully transformed. In fact, the alpha oxidation pathway is present in yeast and the alpha-oxidation enzyme hydroxylacyl-CoA lyases (HACL), which is a key part of this invention was successfully expressed in yeast Saccharomyces. Other species include but are not limited to Candida, Aspergillus, Arxula adeninivorans, Candida boidinii, Hansenula polymorpha (Pichia angusta), Kluyveromyces lactis, Pichia pastoris, and Yarrowia lipolytica, to name a few.

**0033** It is also possible to genetically modify many species of algae, including e.g., Spirulina, Apergillus, Chlamydomonas, Laminaria japonica, Undaria pinnatifida, Porphyra, Eucheuma, Kappaphycus, Gracilaria, Monostroma, Enteromorpha, Arthrospira, Chlorella, Dunaliella, Aphanizomenon, Isochrysis, Pavlova, Phaeodactylum, Ulkenia, Haematococcus, Chaetoceros, Nannochloropsis, Skelia, Valisneria, Thalassiosira, and Laminaria japonica, and the like. Indeed, the microalgae Pavlova lutheri is already being used as a source of economically valuable docosahexaenoic (DHA) and eicosapentaenoic acids (EPA), and Cryptophyllum calhounii is the heterotrophic algal species that is currently used to produce the DHA used in infant formulas.

**0034** Furthermore, a number of databases include vector information and/or a repository of vectors and can be used to choose vectors suitable for the chosen host species. See e.g., AddGene.org which provides both a repository and a searchable database allowing vectors to be easily located and obtained from colleagues. See also Plasmid Information Database (PlasmID) and DNASU having over 191,000 plasmids. A collection of cloning vectors of E. coli is also kept at the National Institute of Genetics as a resource for the biological research community. Furthermore, vectors (including particular ORFs therein) are usually available from colleagues.

**0035** The enzymes can be added to the genome or via expression vectors, as desired. Preferably, multiple enzymes are expressed in one vector or multiple enzymes can be combined into one operon by adding the needed signals between coding regions. Further improvements can be had by overexpressing one or more, or even all of the enzymes, e.g., by adding extra copies to the cell via plasmid or other vector. Initial experiments may employ expression plasmids hosting 3 or more ORFs for convenience, but it may be preferred to insert operons or individual genes into the genome for stability reasons.

**0036** Still further improvements in yield can be had by reducing competing pathways, such as those pathways for making e.g., acetate, formate, ethanol, and lactate, and it is already well known in the art how to reduce or knockout these pathways. See e.g., the William Marsh Rice University patent portfolio by Ka-Yiu San and George Bennett (U.S. Pat. No. 7,569,380, U.S. Pat. No. 7,262,046, U.S. Pat. No. 8,962,272, U.S. Pat. No. 8,795,991) and patents by these inventors (U.S. Pat. No. 8,129,157 and U.S. Pat. No. 8,691,552) (each incorporated by reference herein in its entirety for all purposes). Many others have worked in this area as well.

**0037** In calculating “% identity” the unaligned terminal portions of the query sequence are not included in the calculation. The identity is calculated over the entire length of the reference sequence, thus short local alignments with a query sequence are not relevant (e.g., % identity–number of aligned residues in the query sequence/length of reference sequence). Alignments are performed using BLAST homology alignment as described by Tatusova T A & Madden T L (1999) FEMS Microbiol. Lett. 174:247-250, and available through the NCBI website. The default parameters were used, except the filters were turned OFF.

**0038** “Operably associated” or “operably linked”, as used herein, refer to functionally coupled nucleic acid or amino acid sequences.

**0039** “Recombinant” is relating to, derived from, or containing genetically engineered material. In other words, the genetics of an organism was intentionally manipulated in some way.

**0040** “Reduced activity” or “inactivation” or “underexpressed” is defined herein to be at least a 75% reduction in protein activity, as compared with an appropriate control species (e.g., the wild type gene in the same host species). Preferably, at least 80, 85, 90, 95% reduction in activity is attained, and in the most preferred embodiment, the activity is eliminated (100%). Proteins can be inactivated with inhibitors, by mutation, or by suppression of expression or translation, by knock-out, by adding stop codons, by frame shift mutation, and the like.

**0041** By “null” or “knockout” what is meant is that the mutation produces undetectable active protein. A gene can be completely (100%) reduced by knockout or removal of part of all of the gene sequence. Use of a frame shift mutation, early stop codon, point mutations of critical residues, or deletions or insertions, and the like, can also completely inactivate (100%) gene product by completely preventing transcription and/or translation of active protein. All null mutants herein are signified by A.

**0042** “Overexpressed” or “overexpressed” is defined herein to be at least 150% of protein activity as compared with an appropriate control species, or any expression in a species that lacks the activity altogether. Preferably, the activity is increased 100-500%. Overexpression can be achieved by mutating the protein to produce a more active
form or a form that is resistant to inhibition, by removing inhibitors, or adding activators, and the like. Overexpression can also be achieved by removing repressors, adding multiple copies of the gene to the cell, or up-regulating the endogenous gene, and the like. All overexpressed genes or proteins are signified herein by “+”.

In certain species it is possible to genetically engineer the endogenous protein to be overexpressed by changing the regulatory sequences or removing repressors. However, overexpressing the gene by inclusion on selectable plasmids that exist in hundreds of copies in the cell may be preferred due to its simplicity and ease of exerting external controls, although permanent modifications to the genome may be preferred in the long term for stability reasons.

**BRIEF DESCRIPTION OF FIGURES**

**[0044]** FIG. 1. Current ‘top-down’ metabolic engineering approach based on editing existing architecture of natural metabolism.

**[0045]** FIG. 2. Overall structure of the designed pathway. “Bottom up” integrated design with a new metabolic architecture. Consolidation of F fixation, central metabolism, and product synthesis (shown in FIG. 1) into a single pathway that enables direct use of single carbon molecules for the synthesis of longer-chain (≥C2) products. See also FIG. 10 for more detail.

**[0046]** FIG. 3. Extender Generation Module for converting one carbon compound to an activated form that can be used for elongation. Refer to Table 1A for more detail.

**[0047]** FIG. 4A-B. Possible Elongation Modules for elongating a carbon backbone by one carbon per iteration. Refer to Table 1B for more detail.

**[0048]** FIG. 5. Some implementations include a Regeneration Module for allowing for the regeneration of an aldehyde to prime further elongation, thus allowing multiple cycles of the synthetic pathway and more than one carbon to be added to a growing chain. Refer to Table 1C for more detail.

**[0049]** FIG. 6. Termination module: Potential products derived from acyl-CoA intermediates of the pathway modules. Refer to Table 1D for more detail.

**[0050]** FIG. 7. Alpha Oxidation of straight chain fatty acids.

**[0051]** FIG. 8. Mechanism of the acylon condensation reaction by TPP dependent enzymes. Formyl-CoA is included, for comparison, as the donor molecule such that R1 is the S-CoA group. The acceptor molecule is any aldehyde. The ligation of these two molecules by this mechanism results in the formation of a 2-hydroxyacyl-CoA.

**[0052]** FIG. 9. Four isomers of human HACL1 (SEQ ID NOS 1-4).

**[0053]** FIG. 10A-B. Overview of some proposed implementations of the pathway. Numbers correspond to the reactions listed in Table 2.

**[0054]** FIG. 11. Vector construct containing the gene encoding N-terminal His-tagged acetylating aldehyde reductase Lmo1179 from *Lysteria monocytogenes* for expression in *E. coli*.

**[0055]** FIG. 12. SDS-PAGE showing expression and purification of *Lysteria monocytogenes* acetylating aldehyde reductase Lmo1179 from *E. coli*.

**[0056]** FIG. 13. Time course of absorbance at 340 nm corresponding to the production of NADH in the assay of *L. monocytogenes* acetylating aldehyde reductase Lmo1179.

**[0057]** FIG. 14. ESI-TOF MS data of the -CoA content of *L. monocytogenes* acetylating aldehyde reductase Lmo1179 reaction assay mixtures after solid phase extraction.

**[0058]** FIG. 15A-B. Expression vector constructs for *S. cerevisiae* (15A) and *E. coli* (15B) expression vector constructs containing the gene encoding an amino-terminal 6X His-tagged hydroxyl-acetyl-CoA lyase HACL1 from *Homo sapiens*.

**[0059]** FIG. 16. SDS-PAGE showing the purification of hydroxyl-acetyl-CoA lyase HACL1 from *Saccharomyces cerevisiae*. Lane 1 corresponds to the protein ladder. Lane 2 is the *S. cerevisiae* cell extract. Lane 3 corresponds to purified HACL1.

**[0060]** FIG. 17. GC-FID chromatograms of pentadecanal content in HACL1 degradative reaction mixtures after extraction with hexane. Top: pentadecanal standard; Middle: HACL1 assay sample; Bottom: no enzyme control. In samples containing HACL1, a pentadecanal peak is seen, while there is no peak in the sample in which enzyme was omitted.

**[0061]** FIG. 18. GC-FID chromatograms of HACL1 synthetic reaction mixtures after hydrolysis of acyl-CoAs and derivatization. From top to bottom: no enzyme control; HACL1 sample; 2-hydroxyhexadecanoyl-CoA standard; 2-hydroxyhexanoic acid standard. HACL1 was incubated with pentadecanal and formyl-CoA and was capable of ligating the molecules to 2-hydroxyhexadecanoyl-CoA as indicated by the peaks corresponding to the standards. FIG 18 and FIG 19 prove that the enzyme can run in reverse, as required for the abiotic pathway.

**[0062]** FIG. 19. HPLC chromatogram of HACL1 synthetic reaction samples incubated with acetaldehyde and formyl-CoA. Solid line: HACL1 sample; Dashed line: no enzyme control. The expected product of the ligations of acetaldehyde and formyl-CoA is lactoyl-CoA, which would be expected to be hydrolyzed to its acid form, lactate. A peak for lactate (26 min) was observed in samples containing purified HACL1 and was not observed in samples that did not contain enzyme.

**[0063]** FIG. 20. HPLC chromatogram of HACL1 synthetic reaction samples incubated with formaldehyde and formyl-CoA. Solid line: no enzyme control; Dashed line: HACL1 sample; Dotted line: glycolate standard. The expected product of the ligation of formaldehyde and formyl-CoA is glycosyl-CoA, which would be expected to be hydrolyzed to its acid form, glycolate. A peak matching a glycolate standard (25.2 min) was observed in samples containing purified HACL1 and was not observed in samples that did not contain enzyme.

**[0064]** FIG. 21A-B. NMR spectra of HACL1 assay samples incubated with acetaldehyde and formyl-CoA. (21A) HACL1 sample. (21B) No enzyme control. A peak corresponding to lactate was identified in the sample containing HACL1.

**[0065]** FIG. 22. Vector constructs encoding oxalyl-CoA decarboxylases from (A) *Oxalobacter formigenes* and (B) *E. coli*.

**[0066]** FIG. 23. Vector construct encoding benzaldehyde lyase from *Pseudomonas fluorescens*. 
FIG. 24. Vector construct containing the gene encoding LdABC from Clostridium propionicum for expression in E. coli. LdABC is an example of 2-hydroxycarbonyl-CoA dehydratases (enzyme 3 in FIG. 10 and Table 2).

FIG. 25. Eadie-Hofstee plot for the determination of egTER enzyme kinetics. egTER is an example of transenoyl-CoA reductase (enzyme 4 in FIG. 10 and Table 2).

FIG. 26. Time course of absorbance at 340 nm corresponding to the consumption of NADH in the assay of MnP.

FIG. 27. Time course of absorbance at 340 nm corresponding to the consumption of NADH in the assay of FueO.

FIG. 28. HPLC chromatogram of the in vitro assembly of the trans-2-enoyl-CoA reductase and acyl-CoA reductase steps of the elongation module. A peak corresponding to butyraldehyde (butyraldehyde standard in bold) was present in samples containing Treponema denticola TER and E. coli MnP (solid, thin line), but not the control containing Treponema denticola TER only (dashed line).

FIG. 29. Eadie-Hofstee plot for the determination of FadB enzyme kinetics.

FIG. 30. Eadie-Hofstee plot for the determination of AtoB enzyme kinetics.

FIG. 31. Time course of absorbance at 412 nm corresponding to the production of TNB in the assay of the thioesterases: yciA, fadM, ydhB, and psd.

FIG. 32. HPLC chromatogram of assays of the combined regeneration and termination modules.

FIG. 33. HPLC chromatogram of an in vitro implementation of the pathway consisting of a one unit elongation catalyzed by HACL1, followed by direct termination to product. In this case, HACL1 catalyzed the ligation of formylglycine and formyl-CoA to produce glycocoly-CoA. E. coli YciA included in the reaction mixture converted glycocoly-CoA to glycocolic acid.

FIG. 34. Comparison of the ability of the engineered pathway to support growth of E. coli on different single carbon feedstocks as evaluated by a genome scale model.

Table 1A-D. Enzyme list for each of the four modules. These are only a few of the enzymes that can be used, but provides a representative listing.

Detailed Description of the Disclosure

Herein, our focus will be a mammalian version of the alpha oxidation pathway known to operate on straight chain fatty acids. See FIG. 7. In this pathway, fatty acids are first hydroxylated at the α-position, and the resulting 2-hydroxy fatty acid is activated to the acyl-CoA derivative. The key enzyme, 2-hydroxyacyl-CoA lyase (“HACL”), cleaves the 2-hydroxyacyl-CoA into formyl-CoA and an aldehyde one carbon shorter than the starting fatty acid. This forms the foundation of the second part of the hypothesis—that HACL can catalyze the reverse reaction, elongating a fatty aldehyde by one carbon using formyl-CoA as an extender.

Just as the thiolase reaction in the β-oxidation pathway was shown to be reversible (see e.g., US201303166413, US2040273110, incorporated by reference herein in its entirety for all purposes), it was anticipated that the HACL reaction of α-oxidation could be reversed. Thermodynamically and mechanistically, the hypothesis seemed plausible. Our preliminary experiments were expected to confirm this, based on the Gibbs free energy of the reaction, and our subsequent results show that we were correct. The overall design of the new synthetic anabolic pathway is shown in FIG. 2 and additional detail is in FIG. 10.

The gene encoding a 2-hydroxyacyl-CoA lyase from Homo sapiens, HACL1 (FIG. 9), was cloned into E. coli. Cloning of Homo sapiens HACL1 into a yeast expression vector, pYES2, has also been completed (see FIG. 15A-B), and expression experiments in yeast have been performed. Using an overexpressed HACL enzyme, we were able to drive the reaction in reverse (see EQ 1), thus providing an elongation module for use. See FIG. 18/19.

Although we used the human HACL gene herein (see UniProt Q9U383), that was for convenience, and any mammalian HACL gene could be used, provided it catalyzes the same reaction. The HACL gene is EC number 4.1 (incompletely characterized to date), and other proteins that could suffice include gorilla (XP_004033753.1 = 577/578 (99%) identity), chimp (NP_001267049.1 = 574/578 (99%)); yak (XP_005910699.1 = 515/581 (89%)); dog (XP_013961990.1 = 514/581 (88%)); cat (XP_003992184.1 = 520/581 (90%)); mouse (NP_064359.2 = 508/581 (87%), and the like. In addition, non-mammalian species may also suffice, as homologs are known to exist in the frog (GenBank NP_001017332) and zebrafish (GenBank NP_958250). Recombinant protein is even commercially available from Arabidopsis (MyBioSource).

The HACL reaction will of course normally run in the physiologically forward direction, making 2-hydroxyacyl-CoA molecules shorter by one carbon (the catabolic direction). However, it can be forced to run in the reverse elongation direction (see EQ 1) by overexpressing it at sufficient levels, and by increasing substrate levels or by removing products, thus pushing the reaction in the opposite direction. We were able to make it run in the reverse anabolic or extension direction by providing extra substrate. Additionally, the temperature of the reaction was lowered from 37°C to room temperature and the pH of the reaction was lowered from 7.4 to 5.4.

In total, four modules comprise the proposed invention, and these are generally described in turn.

Extender Generation Module:

The extender generation module, illustrated in FIG. 3 and summarized in Table 1A, is intended to activate a single carbon molecule into the form in which it can be used for ligation. In the proposed scheme using 2-hydroxyacyl-CoA lyase (“HACL”) as the extender enzyme, this activated form is the molecule formyl-CoA. Methods for the conversion of formate to formyl-CoA will be described in detail here.

Formate serves as a good starting point for single carbon utilization because it can be reduced from CO2, enzymatically or electrocatalytically, and oxidized from
more reduced molecules such as methanol, methylal, or formaldehyde. There are at least 3 potential routes for formate activation to formyl-CoA.

[0088] 1) The first route uses one enzyme, an acyl-CoA synthase (ACS), to directly produce the thioester. In the process, this hydrolyzes one ATP to AMP, effectively consuming two ATP equivalents. One such AMP forming acyl-CoA synthase with activity on formate has been found in the organism \textit{Pyrococcus aerophilus}. The ACS, encoded by the open reading frame (ORF) PAE2867, showed an unusually broad range of substrate specificity from formate to butyrate.

[0089] 2) The alternative route uses two enzymes, but produces ADP, utilizing one ATP equivalent. The enzymes capable of catalyzing these reactions were first identified in \textit{Clostridium cylindrosporum}. This organism contains two formate kinases, which catalyze the formation of formyl-phosphate from formate and ATP, and a phosphate formyl-transferase, which is capable of synthesizing formyl-CoA from formyl-phosphate. Since then, other enzymes have been discovered with activity on formate, such as acetate kinase from \textit{Salmonella typhimurium}. These enzymes warrant investigation for the generation of formyl-CoA.

[0090] 3) An additional alternative allows for the direct conversion of formaldehyde to formyl-CoA using the acetylating dehydrogenase enzyme. Variants of this enzyme have been discovered in species such as \textit{Giardia intestinalis}, \textit{Pseudomonas} sp. CF600, and \textit{Acinetobacter} sp. H1S-2 with experimental evidence of activity on formaldehyde. These enzymes catalyze the conversion of formaldehyde to formyl-CoA with the generation of one molecule of NADH. Much like the methylotrophic pathways of single carbon metabolism, this enzyme enables the more efficient utilization of single carbon molecules at the same oxidation state or more reduced than formaldehyde, including methanol and methane.

[0091] Elongation Module:

[0092] The elongation module, shown in FIG. 4A-B and summarized in Table 1B, allows for the iterative elongation of the carbon backbone. Using HACL, an exemplary one-carbon carbon-carbon bond-forming enzyme, an aldehyde is ligated with formyl-CoA to produce a 2-hydroxyacetyl-CoA that is one carbon longer than the originating aldehyde.

[0093] To allow for additional rounds of elongation, the 2-hydroxyacetyl-CoA must be converted back to the aldehyde form. In one implementation, FIG. 4A, from the 2-hydroxyacetyl-CoA, the next step is dehydration of the molecule to the trans-2-enoyl-CoA. An enzyme with this activity, lactal-CoA dehydratase, has been discovered in \textit{Clostridium propionicum} encoded by gene \textit{ldA}. This enzyme was found to catalyze the conversion of 2-hydroxy substrates such as 2-hydroxypropionyl-CoA, also known as lactal-CoA, and 2-hydroxybutyryl-CoA to prop-2-enoyl-CoA (acryloyl-CoA) and but-2-enoyl-CoA (crotonyl-CoA), respectively.

[0094] In the next step, the resulting trans-2-enoyl-CoA is converted to a saturated acetyl-CoA. A number of enzymes have been identified with this desired activity. To follow the previous example, acryloyl-CoA has been shown to be reduced to propionyl-CoA by an acryloyl-CoA reductase encoded by \textit{Rhodobacter sphaeroides} acr.

[0095] Other trans-2-enoyl-CoA reductases with broad chain specificity have been used in engineered pathways such as the reverse β-oxidation pathway, which, similar to this pathway, dehydrates a 3-hydroxyacyl-CoA to a trans-2-enoyl-CoA before reducing it to the saturated acyl-CoA. In this application, the trans-2-enoyl-CoA reductase from \textit{Englera gracilis} has already been successfully used. Increasing evidence also indicates that the native \textit{E. coli} gene \textit{fbc}, which encodes an enoyl-ACP reductase used in the fatty acid biosynthesis pathway, also has activity on enoyl-CoA substrates (Vick et al., 2014).

[0096] From the acetyl-CoA, the aldehyde can be directly generated. \textit{Salmonella enterica} \textit{EutI} encodes an acyl-CoA reductase that is capable of catalyzing the conversion of propionyl-CoA to propionaldehyde. (Zhu et al., 2011) Similarly, the acetyl-CoA reductase from \textit{Clostridium beijerinckii} gene \textit{ald} has been shown to have activity on butyryl-CoA in its conversion to butanal. This step completes the cycle, allowing the carbon chain to undergo another round of elongation.

[0097] In an alternative implementation of the elongation module, 2-hydroxyacyl-CoA can be converted to its corresponding aldehyde via an alternative arrangement of reductions and dehydrations to those described above (FIG. 4B). Lyase is first reduced to a 2-hydroxyaldehyde by an acyl-CoA reductase, examples of which are provided above. The 2-hydroxyaldehyde is then reduced further to a 1,2-diol by a 1,2-diol oxidoreductase. \textit{E. coli} \textit{fucO} has been shown to encode one such 1,2-diol oxidoreductase. Finally, the 1,2-diol can be converted to the aldehyde by a diol dehydratase. \textit{Klebsiella oxytoca} \textit{pdhABC} has been shown to encode an enzyme capable of performing the dehydration of C2-C4 1,2-diols to their corresponding aldehydes. This aldehyde can be used for further elongation of the carbon backbone.

[0098] Regeneration Module:

[0099] A pathway that allows for the regeneration of a priming aldehyde is described herein and illustrated in FIG. 5. This would allow for the generation of products without necessitating the diversion of cellular resources to produce a starting aldehyde.

[0100] Starting from the trans-2-enoyl-CoA, an enoyl-CoA hydratase, such as native \textit{E. coli} \textit{FadB}, catalyzed the conversion to a 3-hydroxyacyl-CoA. \textit{FadB} has also been shown to have 3-hydroxyacyl-CoA dehydrogenase activity, which catalyzes the conversion of the 3-hydroxyacyl-CoA to a 3-ketoacyl-CoA. A thiolase, such as the \textit{E. coli} enzymes \textit{fadA} or \textit{atoB}, can then cleave an acetyl-CoA from the 3-ketoacyl-CoA, leaving an acetyl-CoA. The acetyl-CoA can be reduced, for example by the aforementioned \textit{Salmonella enterica} \textit{EutE}, to acetaldehyde and used to prime a further elongation cycles, while the acetyl-CoA can be converted to products.

[0101] Termination Module:

[0102] From the intermediates of the above reactions, there are a number of possible ways by which products can be generated by pulling intermediates out of the pathway with one or more enzyme reactions. Termination pathways are highly inspired by our work on the reverse beta oxidation, which has provided a great number of termination enzymes and pathways that can be employed herein. Thus, the reader is referred to US20130316413, US20140273110, et seq, and the associated publications for additional termination enzymes, including the selection thereof to control chain length, and at which point in the cycle an intermediate is removed.

[0103] Resulting acyl-CoAs from the pathway can be converted to products such as fatty acids by expression of a thioesterase, alcohols by expression of an acyl-CoA reduc-
tase and an aldehyde reductase, or alkanes by expression of an acyl-CoA reductase and aldehyde decarboxylase. See Figs. 6.

[0104] Tables 1A-D present a summary of the enzymes corresponding to the proposed pathway design and Fig. 10A-B provide a representation of the overall pathway in the forms proposed above. As written, starting from formate, the pathway uses one ATP equivalent for every one carbon elongation or, for ease of comparison to alternatives, two ATP equivalents for two carbon elongation.

[0105] It is important to note, though, that starting from formaldehyde, which can be readily oxidized from more reduced compounds, the pathway requires no ATP equivalents for elongation. This ATP efficiency is comparable to even the more efficient alternatives, for example the combination of the reductive acetyl-CoA or Ribulose Monophosphate Pathway (RuMP) pathways with the reverse \( \beta \)-oxidation pathway, which consume approximately one ATP equivalent from \( \text{CO}_2 \) or no ATP from formaldehyde, respectively, for two carbon elongation.

[0106] Following the construction of a suitable strain containing the engineered pathway, culturing of the developed strains can be performed to evaluate the effectiveness of the pathway at its intended goal, the production of products from single carbon compounds. The organism can be evaluated for growth on a variety of single carbon substrates, from methane to \( \text{CO}_2 \) and \( \text{H}_2 \), either autotrophically or mixotrophically, with the inclusion of an additional carbon source. The products produced by the organism can be measured by HPLC or GC, and indicators of performance such as growth rate, productivity, titr, yield, and carbon efficiency can be determined.

[0107] Further, evaluation of the interaction of the heterologously expressed pathway enzymes with each other and with the host system can allow for the optimization of pathway performance and minimization deleterious effects. Because the pathway is under synthetic control, rather than under the organism’s native evolved regulatory mechanisms, the expression of the pathway must be manually tuned to avoid potential issues that slow cell growth or production and to optimize production of desired compounds.

[0108] Additionally, an imbalance in relative enzyme activities might restrict overall carbon flux throughout the pathway, leading to suboptimal production rates and the buildup of pathway intermediates, which can inhibit pathway enzymes or be cytotoxic. Analysis of the cell cultures by HPLC or GC can reveal the metabolic intermediates produced by the constructed strains. This information can point to potential pathway issues.

[0109] As an alternative to the in vivo expression of the pathway, a cell free, in vitro, version of the pathway can be constructed. By purifying the relevant enzyme for each reaction step, the overall pathway can be assembled by combining the necessary enzymes. With the addition of the relevant cofactors and single carbon compounds, the pathway can be assessed for its performance independently of a host. As yet another alternative, whole wet or dried cells can be used as bioreactors. However, a living, growing culture system is preferred.

[0110] The primary focus of the proposed work was the development of a pathway for the direct synthesis of products from single carbon compounds. An additional area of interest, however, involves the use of the enzymes related to \( \alpha \)-oxidation for the production of \( \alpha \)-functionalized products. In particular, fatty acid 2-hydroxylases allow for the production of \( \alpha \)-hydroxy acids from fatty acids, which can be further converted to \( \alpha \)-hydroxy aldehydes, alcohols, ketones, or esters, or further into \( \alpha \)-keto compounds. Such \( \alpha \)-functionalized products have been found to be particularly useful bioactive molecules and have found applications in cosmetics as well as for drug delivery. The enzymes required for these transformations can be expressed in \( E. \ coli \), yeast or algae and tested in vitro, as described above. When used in conjunction with a product-synthesizing platform, such as the \( \beta \)-oxidation reversal pathway, the microbial production of \( \alpha \)-functionalized products can be achieved.

[0111] The following description of experiments provides additional details, any one of which can be subject to patenting in combination with any other. The specification in its entirety is to be treated as providing a variety of details that can be used interchangeably with other details, as it would be of inordinate length if one were to list every possible combination of genes/vectors/enzymes/hosts that can be made to run a reverse alpha oxidation pathway.

Materials & Methods

[0112] Enzymes of interest can be expressed from vectors such as pCDFDuet-1 (MERCK, Germany), which makes use of the DE3 expression system. Genes can be codon optimized according to the codon usage frequencies of the host organism and synthesized by a commercial vendor or in-house. However, thousands of expression vectors and hosts are available, and this is a matter of convenience.

[0113] The genes can be amplified by PCR using primers designed with 15-22 base pairs of homology for the appropriate vector cut site. For enzymes that will not require a 6X-histadine tag fusion for purification, pCDFDuet-1 can be linearized with NcoI and EcoRI. Enzymes that will be purified by Ni-NTA column will make use of the 6X-HIS tag in pCDFDuet-1. The vector can be linearized using only EcoRI in this case.

[0114] The PCR product can be inserted into the vector using e.g., the In-Fusion HD EcoDry Cloning System and the vector transformed by heat shock into competent \( E. \ coli \) cells. Transformants can be selected on solid media containing the appropriate antibiotic. Plasmid DNA can be isolated using any suitable method, including QIAprep Spin Miniprep Kit (QIAGEN, Limburg), and the construct confirmed by PCR and sequencing. Confirmed constructs can be transformed by e.g., electroporation into a host strain such as \( E. \ coli \) for expression, but other host species can be used with suitable expression vectors and possible codon optimization for that host species.

[0115] Expression of the desired enzymes from the constructed strain can be conducted in liquid culture, e.g., shaking flasks, bioreactors, chemostats, fermentation tanks and the like. Gene expression is typically induced by the addition of a suitable inducer, when the culture reaches an OD550 of approximately 0.5-0.8. Induced cells can be grown for about 4-8 hours, at which point the cells can be pelleted and saved to ~20°C. Expression of the desired protein can be confirmed by running cell pellet samples on SDS-PAGE.

[0116] The expressed enzyme can be directly assayed in crude cell lysates, simply by breaking the cells by chemical, enzymatic, heat or mechanical means. Depending on the expression level and activity of the enzyme, however, pur-
fication may be required to be able to measure enzyme activity over background levels. Purified enzymes can also allow for the in vitro assembly of the pathway, allowing for its controlled characterization.

[0117] N- or C-terminal HIS-tagged proteins can be purified using e.g., a Ni-NTA Spin Kit (Qugen, Venlo, Limburg) following the manufacturer’s protocol, or other methods could be used. The HIS-tag system was chosen for convenience only, and other tags are available for purification uses. Further, the proteins in the final assembled pathway need not be tagged if they are for in vivo use. Tagging was convenient, however, for the enzyme characterization work performed herein.

[0118] Reaction conditions for enzyme assays can vary greatly with the type of enzyme to be tested. In general, however, enzyme assays follow a similar general protocol. Purified enzyme or crude lysate is added to suitable reaction buffer. Reaction buffers typically contain salts, necessary enzyme cofactors, and are at the proper pH. Buffer compositions often change depending on the enzyme or reaction type. The reaction is initiated by the addition of substrate, and some aspect of the reaction related either to the consumption of a substrate or the production of a product is monitored.

[0119] Choice of the appropriate monitoring method depends on the compound to be measured. Spectrophotometric assays are convenient because they allow for the real time determination of enzyme activity by measuring the concentration dependent absorbance of a compound at a certain wavelength. There are not always compounds with a measurable absorbance at convenient wavelengths in the reaction, unfortunately. In these situations, other methods of chemical analysis may be necessary to determine the concentration of the involved compounds.

[0120] Gas chromatography (GC) is convenient for the quantification of volatile substances, of which fatty acids and aldehydes are of particular relevance. Internal standards, typically one or more molecules of similar type not involved in the reaction, are added to the reaction mixture, and the reaction mixture is extracted with an organic solvent, such as hexane. Fatty acid samples, for example, can be dried under a stream of nitrogen and converted to their trimethylsilyl derivatives using BSTFA and pyridine in a 1:1 ratio. After 30 minutes incubation, the samples are once again dried and redissolved in hexane to be applied to the GC. Aldehyde samples do not need to be derivatized. Samples can be run e.g., on a Varian CP-3800 gas chromatograph (VARIAN ASSOCIATES, Palo Alto, Calif.) equipped with a flame ionization detector and HP-5 capillary column (AGILENT Tech., Santa Clara, Calif.).

[0121] Once the pathway has been fully studied in vitro, the pathway can be constructed in vivo with greater confidence. The strain construction for the in vivo pathway operation should allow for the well-defined, controlled expression of the enzymes of the pathway. As before, E. coli or yeast will be a host of choice for the in vivo pathway, but other hosts could be used. The Duet system, for example, allows for the simultaneous expression of up to eight proteins by induction with IPTG in E. coli; and initial experiments will use this host.

[0122] Pathway enzymes can also be inserted into the host chromosome, allowing for the maintenance of the pathway without requiring antibiotics to ensure the continued upkeep of plasmids. There are also, theoretically, an infinite number of genes that can be placed on the chromosome, as chromosomal expression does not require separate origins of replication as is the case with plasmid expression.

[0123] DNA constructs for chromosomal integration usually include an antibiotic resistance marker with flanking FRT sites for removal, as described by Datsenko and Wanner, a well characterized promoter, a ribosome binding site, the gene of interest, and a transcriptional terminator. The overall product is a linear DNA fragment with 50 base pairs of homology for the target site on the chromosome flanking each side of the construct.

[0124] However, the Flp-FRT recombination method is only one system for adding genes to a chromosome, and other systems are available, such as the RecICD pathway, the RecA pathway, RecA recombinase, non-homologous end joining (NHEJ), Cre-Lox combination, TVR recombinases and integrases, SIR resolvases/invertases, SIR integrases, PhiC31 Integrase, and the like. Chromosomal modifications in E. coli can also achieved by the method of recombineering, as originally described by Datsenko and Wanner.

[0125] In a recombineering method, for example, the cells are prepared for electroporation following standard techniques, and the cells transformed with linear DNA that contains flanking 50 base pair targeting homology for the desired modification site. For seamless integration of a DNA construct, a two-step approach can be taken using a cassette that contains both positive and negative selection markers, such as the combination of cat and sacB. In the first round of recombineering, the cat-sacB cassette with targeting homology for the desired modification site is introduced to the cells. The cat gene provides resistance to chloramphenicol, which allows for positive recombaints to be selected for on solid medium containing chloramphenicol. A positive isolate can be subjected to a second round of recombineering introducing the desired DNA construct with targeting homology for sites that correspond to the removal of the cat-sacB cassette. The sacB gene encodes for an enzyme that provides sensitivity to sucrose. Thus, growth on media containing sucrose allows for the selection of recombaints in which the cat-sacB construct was removed. Plague lysates can be made from isolates confirmed by PCR and sequencing. The lysates can be used to transduce the modification into desired strains, as described previously.

[0126] Engineered strains expressing the designed pathway can be cultured under the following or similar conditions. Overnight cultures started from a single colony can be used to inoculate flasks containing appropriate media. Cultures are grown for a set period of time, and the culture media analyzed. The conditions will be highly dependent on the specifications of the actual pathway and what exactly is to be tested. For example, the ability for the pathway to be used for autotrophic growth can be tested by the use of formate or formaldehyde as a substrate in MOPS minimal media, as described by Neidhardt, supplemented with appropriate antibiotics, and inducers. Mixotrophic growth can be characterized by the addition of both single carbon compounds and glucose or glycerol.

[0127] Analysis of culture media after fermentation provides insight into the performance of the engineered pathway. Quantification of longer-chain fatty acid products can be analyzed by GC. Other metabolites, such as short chain organic acids and substrates such as glucose or glycerol can be analyzed by HPLC. Once the pathway is fully functional,
the cultures can be grown in chemostat, providing continuous uninterrupted production of product if desired.

Various -omics techniques, such as microarray or 2D-PAGE can give information about gene expression or protein expression, respectively. Genome scale modeling allows for the identification of additional modifications to the host strain that might lead to improved performance. Deletion of competing pathways, for example, might increase carbon flux through the engineered pathway for product production.

Standard molecular biology techniques were used for gene cloning, plasmid isolation, and E. coli transformation. Native E. coli genes were amplified from E. coli MG1655 genomic DNA using primers to append 15 bp of homology on each end of the gene insert for recombination into the vector backbone. Genes from other organisms were codon optimized and synthesized by either GeneArt (LIFE TECH, CA or GENSCRIPT, NJ). Plasmids were linearized by the appropriate restriction enzymes and recombined with the gene inserts using the In-Fusion HD Eco-Dry Cloning system (CLONTECH Lab., CA). The mixture was subsequently transformed into Stellar competent cells (CLONTECH Lab).

Transformants that grew on solid media (LB+Agar) supplemented with the appropriate antibiotic were isolated and screened for the gene insert by PCR. Plasmid was isolated from the verified transformants and the sequence of the gene insert was further confirmed by DNA sequencing (LONE STAR LABS, TX). Plasmids (also referred to as vectors) in each case contain at least one promoter, a ribosome binding site for each gene, the gene(s) of interest, at least one terminator, an origin of replication, and an antibiotic resistance marker. Exemplary plasmids are shown in FIGS. 11, 15, and 22-24.

2-hydroxyhexadecanoyl-CoA was prepared by the n-hydroxysuccinimide method (Blecher, 1981). In summary, the n-hydroxysuccinimide ester of 2-hydroxyhexadecanoic acid was prepared by reacting n-hydroxysuccinimide with the acid in the presence of dicyclohexylcarbodiimide. The product was filtered and purified by recrystallization from methanol to give pure n-hydroxysuccinimide ester of 2-hydroxyhexadecanoic acid. The ester was reacted with CoASH in presence of thio-erythritol to give 2-hydroxyhexadecanoyl-CoA. The 2-hydroxyhexadecanoyl-CoA was purified precipitation using perchloric acid, filtration, and washing the filtrate with perchloric acid, diethyl ether, and acetone.

Formyl-CoA was prepared by first forming formic ethylcarboxy anhydride as previously described (Parasaran & Tarbell, 1964). Briefly, formic acid (0.4 mmol) and ethyl chloroformate (0.4 mmol) were combined in 4 mL anhydrous diethyl ether and cooled to −20°C. 0.4 mmol triethylamine was added to the mixture and the reaction was allowed to proceed at −20°C for 30 minutes. The reaction mixture was filtered over glass wool to a solution containing formic ethylcarboxy anhydride in diethyl ether. To obtain formyl-CoA, 7 μmol CoASH was dissolved in 5 mL 3.2 water:triethylammonium, to which 10 mg of sodium bicarbonate were added. The solution of formic ethylcarboxy anhydride was added dropwise to the CoASH solution with vigorous agitation, after which the organic phase was evaporated under a stream of nitrogen. The mixture was kept at 4°C for two hours, after which any remaining diethyl ether was evaporated under nitrogen. Solid phase extraction using a C18 column was used to purify formyl-CoA from the reaction mixture. Formyl-CoA was eluted from the C18 column in methanol and stored in 2:1 methanol:ammonium acetate pH 5.5.

A plasmid containing the codon optimized gene encoding 6X HS-tagged Lmo1179 from Lysteria monocytogenes was constructed as described above. The resulting construct, FIG. 11, was transformed into E. coli BL21(DE3) for expression. The resulting strain was cultured in 50 mL of TB media containing 50 μg/mL spectinomycin in a 250 mL flask. When the culture reached an OD550 of approximately 0.6, expression was induced by the addition of 0.1 mM IPTG, and the cells were harvested by centrifugation after overnight incubation at room temperature.

The resulting cell pellet was resuspended in Bacterial Protein Extraction Reagent (B-PER) (THERMO SCI, MA) to an OD550 of approximately 40, to which approximately 5000 U of lysozyme and approximately 250 U of Benzonase nuclease (SIGMA-ALDRICH Co., MO) were added. The cell mixture was left at room temperature until completely clarified to give the cell extract. 1 M stock solution of imidazole was added to provide a final concentration of 10 mM imidazole in the cell extract.

The HS-tagged Lmo1179 protein was purified from the cell extract using Talon Metal Affinity Resin (CLONTECH Lab.). In short, a 250 μL resin bed was equilibrated twice using 2.5 mL of a buffer containing 50 mM sodium phosphate, 300 mM NaCl, and 10 mM imidazole at pH 7.5 (NPI-10). The cell extract was added to the resin and the mixture shaken gently for 20 minutes on ice. The resin was then washed twice with 2.5 mL buffer NPI-20 (same as NPI-10 but with 20 mM imidazole), shaking gently on ice for 15 minutes each wash. The resin was then transferred to a gravity column and washed once with 1.25 mL NPI-20. Finally, the desired protein was eluted using 1.25 mL of buffer NPI-250 (same as buffer NPI-10 but with 250 mM imidazole), and the eluate collected in 500 μL fractions. An sample purification of Lmo1179 is shown in FIG. 12.

A plasmid containing the codon optimized gene encoding human HS-tagged HACL1 was constructed as described. The resulting construct, FIG. 15, was transformed into S. cerevisiae InxSCL (Life Tech.). The resulting strain was cultured in 50 mL of SC-URA media containing 2% glucose at 30°C for 24 hours. The cells were pelleted and the required amount of cells were used to inoculate a 250 mL culture volume of SC-URA media containing 0.2% galactose, 1 mM MgCl2, and 0.1 mM thiamine to 0.4 OD600. After 20 hours incubation with shaking at 30°C, the cells were pelleted and saved.

When needed, the cell pellets were resuspended to an OD600 of approximately 100 in a buffer containing 50 mM potassium phosphate pH 7.4, 0.1 mM thiamine pyrophosphate, 1 mM MgCl2, 0.5 mM AEBSF, 10 mM imidazole, and 250 units of Benzonase nuclease. To the cell suspension, approximately equal volumes of 425-600 μm glass beads were added. Cells were broken in four cycles of 30 seconds of vortexing at 3000 rpm followed by 30 seconds on ice. The glass beads and cell debris were pelleted by centrifugation and supernatant containing the cell extract was collected. The HACL1 was purified from the cell extract using Talon Metal Affinity Resin as described above, with the only modification being the resin bed volume and all subsequent washes were halved. The eluate
was collected in two 500 μL fractions. A sample purification of HAC11 is shown in Fig. 16.

**E. coli** fadA and fadB genes were cloned from MG1655 genomic DNA into the pUCB-Ntl6 vector to yield a constitutively expressed gene with an aminoterminal HIS-tag that can be cleaved by Thrombin. For AtoB and thioesterase characterization assays, the pCA24N-gene(-gfp) plasmids from the ASKA collection were used (Kitagawa et al., 2005). For expression of *E. gracilis* TER for kinetic characterization, the egTER gene was cloned into pTrcHis2A using the In-Fusion protocol (CLONTECH LAB.) following PCR amplification from the aforementioned codon-optimized egTER containing plasmid.

**Cultures for enzymatic assays** were grown overnight in 100 mL of LB media at 37°C in 250 mL baffled flasks (WHEATON IND., NJ) in *E. coli* BL21 (DE3) cells induced with either 1 mM IPTG (pCA24N, pTrcHis2A) or 1 mM Arabinose (pUCB-pBAD) at an OD600 of about 0.6, or expressed constitutively (pUCB-HIS6). Reactions were monitored on either a Synergy HT plate reader (BIOTEK INSTR., VT) at 25°C. (for reactions monitored at 300 nm or higher) or in a Biomek 5 Spectrophotometer (THERMO SCL,) for reactions at 263 nm.

**Cells were lysed** using Bacterial Protein Extraction Reagent (B-PEER) (Thermo Sci.) as per the prescribed protocol in order to obtain the supernatant containing the active enzymes. FadE and YidO were purified following previously established methods using the same growth conditions mentioned above. Cell pellets were resuspended in 40 mL of 50 mM potassium phosphate buffer pH 7.2 and broken by disruption with a homogenizer (AVESTIN, ON). Disrupted cells were then spun for 90 min at 4°C at 120,000g in an Optima L-80XP Ultracentrifuge (BECKMAN-COULTER, IL) to produce the supernatant used for assays.

For specific activity assays (reported in μmol substrate/mg protein/min) these supernatant fractions were utilized and protein concentration was established using the Bradford Reagent (Thermo Sci.) using BSA as the protein standard. Linearity was established for each reaction and the background non-enzymatic rate was subtracted to establish the activity. HIS-tagged proteins of AtoB, FadB, and egTER were purified from the B-PEER supernatant fractions using Talon Metal Affinity Resin (CLONTECH LAB.) using gravity purification.

In short, the supernatant was mixed for 1 hr at room temperature on a LabQuake rotator (FISHER SCL., PA) with approximately 2 mL of Talon Resin (1 mL resin/0.3 mg supernatant protein) that was pre-washed twice with Buffer A (50 mM Tris pH 7.9, 5 mM MgCl2, 100 mM NaCl, 5 mM imidazole). Resin was then spun at 700g for 5 min to remove the non-bound proteins, washed with 20x (40 mL) Buffer A, resuspended in 20x (40 mL) Buffer B (Buffer A with 20 mM imidazole) and loading onto a gravity column. Buffer B was then drained off and the protein was eluted with 20 mL of Buffer C (Buffer A with 250 mM imidazole). The eluted fraction was then concentrated and used for kinetic characterizations. Enzyme concentration was established by measuring the absorbance at 260 nm and extinction coefficients predicted for each enzyme by the ProtParam program (web.expasy.org/protparam/).

**For crude extract assays,** cells (JST07-atoB<sup>C75</sup>-fadB<sup>C75</sup>, JST07-atoB<sup>C75</sup>-fadB<sup>C75</sup>/pCDF-yciA) were grown in 25 mL LB medium in 125 mL Erlenmeyer flasks at 37°C and 200 rpm. Cumarate 0.2 mM and IPTG 0.1 mM were added at OD550 0.3-0.5 for the overexpression of fadB, atoB under control of cymA promoter (CT5) and yciA under T7 promoter respectively followed by additional cultivation for 4 hr under the same conditions before harvested by centrifugation at 5000g for 10 min. Harvested cells were washed once with a 9 g/L NaCl solution, and stored as cell pellets at ~80°C until used. Frozen cells were resuspended in Tris-HCl (50 mM, pH 7.5) buffer supplemented with 1 mM diethiothreitol (DTT) for the cell disruption carried out by using glass (0.1 mm diameter) beads.

In Vitro Pathway Characterization

**Genes that encode the enzymes of the engineered pathway were cloned and expressed as described above. The purified enzymes were assessed for their ability to catalyze the proposed reactions as summarized in Table 2 and Fig. 10A-B. The engineered pathway consists of at least an extender generation module, which results in the production of formyl-CoA; an elongation module, which allows for the iterative elongation of a carbon backbone; and a termination module, which allows for the production of products from intermediates of the other modules. A regeneration module is also described, but is optional, and can be omitted if the cycle runs only once. Here, we described testing in vitro the steps that make up an implementation that uses formaldehyde to produce saturated fatty acids.**

### TABLE 2

<table>
<thead>
<tr>
<th>Extender Generation</th>
<th>Enzyme class</th>
<th>Enzyme</th>
<th>Measured specific activity (μmol/mg protein/min)</th>
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<td>Acylating aldehyde reductase</td>
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<td>Elongation</td>
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<td></td>
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<tr>
<td>2</td>
<td>2-hydroxyacyl-CoA lyase</td>
<td><em>H. sapiens</em> HACL1</td>
<td>Activity observed</td>
<td>This work (HOFMEISTER &amp; BUCKEL, 1992)</td>
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<td>Enzyme</td>
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</tr>
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<td>E. coli YclA</td>
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[0145] Extender Generation Module:

L. monocytogenes Lmo1179 was cloned (FIG. 11), expressed, and purified (FIG. 12) in E. coli as described above. The purified enzyme was evaluated for its ability to convert formyldehydrogenate into the extender unit formyl-CoA. Enzyme assays were performed in 25 mM potassium phosphate buffer pH 7.0, 1 mM CoASH, 0.5 mM NAD⁺, 20 mM 2-mercaptoethanol, and 50 mM formaldehyde. The reaction was monitored by measuring absorbance at 340 nm, corresponding to the production of NADH. CoA compounds were extracted from the reaction mixture by solid phase extraction (SPE) using a C18 column, and the masses of the extracted CoAs were determined by ESI-TOF MS. The inclusion of Lmo1179 in the reaction mixture resulted in the conversion of formyldehydrogenate to formyl-CoA as indicated by the coproduction of NADH (FIG. 13). Mass spectrometry analysis confirmed the production of formyl-CoA by Lmo1179. A peak at the expected mass of formyl-CoA (796) was identified in the sample incubated with enzyme, as shown in FIG. 14. The peak was not present in the no enzyme control (FIG. 14), indicating that Lmo1179 produced formyl-CoA.

[0147] Elongation Module:

The elongation module consists of the ligation of an aldehyde and formyl-CoA, followed by steps that convert the 2-hydroxyacyl-CoA back to an aldehyde 1 carbon longer than the starting aldehyde.

[0149] To catalyze the ligation step, human HACL1 was cloned (FIG. 15A-B), expressed, and purified (FIG. 16) in S. cerevisiae as described above. Purified HACL1 was tested for its native catalytic activity by assessing its ability to cleave 2-hydroxyhexadecanoyl-CoA to pentadecanal and formyl-CoA. Enzyme assays were performed in 50 mM tris-HCl pH 7.5, 0.8 mM MgCl₂, 0.02 mM TPP, 6.6 µM BSA, and 0.3 mM 2-hydroxyhexadecanoyl-CoA. The assay mixtures were incubated for one hour at 37° C, after which the presence of pentadecanal was assessed by extraction with hexane and analysis by GC-FID. As shown in FIG. 17, pentadecanal was produced in the sample containing HACL1, but not in the control sample, which did not contain HACL1, indicating that the protein was expressed and purified in an active form.

[0150] The ability of purified HACL1 to run in the anaerobic direction (reverse from the physiological direction) was also determined. An aldehyde and formyl-CoA were tested for ligation in a buffer comprised of 60 mM potassium phosphate pH 5.4, 2.5 mM MgCl₂, 0.1 mM TPP, 6.6 µM BSA, 5 mM aldehyde, 20% DMSO, approximately 1 mM freshly prepared formyl-CoA, and approximately 0.5 mg/mL purified HACL1. The reaction was allowed to take place at room temperature for 16 hours, after which acyl-CoAs were hydrolyzed to their corresponding acids by adjusting to pH>12.0. For instances in which a short carbon chain product was expected, for example lactate production from acetaldheyde, samples were analyzed by HPLC. In the case of longer products, for example the production of 2-hydroxyhexadecanoic acid from pentadecanal, samples were acidified with HCl and extracted with diethyl ether. The extracted diethyl ether was evaporated to dryness under a stream of nitrogen and derivatized by the addition of 1:1 BSTFA:pyridine. After incubation at 70° C for 30 min, these samples were analyzed by GC-FID.

[0151] When the purified enzyme was supplied with pentadecanal and formyl-CoA, as in FIG. 18, HACL1 was shown to catalyze the ligation of these molecules to 2-hydroxyhexadecanoyl-CoA as hypothesized. After hydrolysis of acyl-CoAs, the chromatogram of the sample containing enzyme shows the peaks to the 2-hydroxyhexadecanoyl-CoA spiked standard, which are absent from the sample containing no enzyme.

[0152] The purified HACL1 was further tested for activity on shorter aldehydes, such as the ligation of acetaldehyde or formaldehyde with formyl-CoA to produce lactyl-CoA or glycolyl-CoA, respectively. After hydrolysis of acyl-CoAs to their acid forms, these samples were analyzed by HPLC. The presence of lactate from elongation of acetaldehyde and formyl-CoA was identified in the sample containing HACL1, but not in the no enzyme control as shown in FIG. 19.

[0153] Similar results were observed for glycolate from formaldehyde and formyl-CoA as shown in FIG. 20. The presence of lactate in the relevant samples was confirmed by NMR, as shown in FIG. 21A-B. This demonstrates that HACL1 is capable of catalyzing the ligation of aldehydes with chain lengths ranging at least from C1-C15 with formyl-CoA, making it suitable for the engineered iterative pathway.

[0154] The 2-hydroxyacyl-CoA then undergoes dehydration to its corresponding trans-2-enoyl-CoA. A 2-hydroxyacyl-CoA dehydratase was identified as LcdABC from
*Clostridium propionicium*. LdcABC has been characterized by Hofmeister and Buckel for the dehydration of 2-hydroxybutyryl-CoA to crotonyl-CoA, with specific activity corresponding to 1.21±0.08 μmol/min/mg protein (Hofmeister & Buckel, 1992). The genes encoding LcdABC were cloned as described above (FIG. 24).

**0155** The trans-2-enoyl-CoA is then reduced to the saturated acyl-CoA by a trans-2-enoyl-CoA reductase. A variant from *Euglena gracilis*, EgTER, was identified and cloned as described above. After expression and purification of EgTER, in vitro assays were performed by monitoring the loss of NADH absorbance in the presence of 100 mM Tris HCl pH 7.5 and 0.2 mM NADH in a final volume of 200 μL at 25°C. This revealed that the enzyme is capable of catalyzing the conversion of crotonyl-CoA to butyryl-CoA with specific activity corresponding to 1.21±0.08 μmol/min/mg protein (FIG. 25).

**0156** The saturated acyl-CoA is finally converted to its corresponding aldehyde by an acyl-CoA reductase. Native *E. coli* acyl-CoA reductase MhpF was tested for its ability to convert butyryl-CoA to butyraldehyde (FIG. 26). *E. coli* MhpF was expressed from an ASKA collection strain (Kitagawa et al., 2005). Strains were grown anaerobically in 10 mL with 100 mM Tris pH 8.0, 10 g/L glucose, 50 μM FeSO₄, 5 μM NaHSO₃, and 5 μM (NH₄)₆Mo₇O₂⁴. The cells were grown to 0.4 OD550 at which point protein expression was induced by the addition of 0.1 mM IPTG.

**0157** After 3 hours growth, cells were pelleted and the pellet was resuspended to 40 OD in 100 mM MOPS pH 7.5. 1 mL of the cell suspension was added to 0.75 g of glass beads and the cells were disrupted for 3 minutes using a cell disruptor (Scientific Industries, Bohemia, N.Y., USA). The cell debris and glass beads were pelleted by centrifugation and the supernatant comprising the cell extract was used for assays.

**0158** The assay mixture consisted of 100 mM MOPS pH 7.5, 6 mM DTT, 5 mM MgSO₄, 0.3 mM (NH₄)₂Fe(SO₄)₂, 0.3 mM NADH, and 0.2 mM butyryl-CoA. The reaction was monitored by loss of absorbance at 340 nm corresponding to the consumption of NADH, as shown in FIG. 24. MhpF was capable of catalyzing the conversion with a specific activity of 0.009±0.003 μmol/min/mg protein (FIG. 26).

**0159** *E. coli* FucO was assayed for its ability to perform an analogous reaction, the reduction of butyraldehyde to butanol. FucO was expressed from an ASKA collection strain and purified as described above. Purified FucO was assayed in a buffer containing 100 mM Tris-HCl pH 7.5, 0.3 mM NADH, and 10 mM butyraldehyde. The reaction was monitored by loss of absorbance at 340 nm corresponding to the consumption of NADH, as shown in FIG. 27. FucO was capable of catalyzing the reduction of butyraldehyde to butanol with a specific activity of 5.08±0.08 μmol/min/mg protein.

**0160** A portion of the elongation module was assembled in vitro to demonstrate the functionality of the combined pathway steps. The trans-2-enoyl-CoA reductase and acyl-CoA reductase steps were combined to assess the overall conversion of crotonyl-CoA to butyraldehyde. Experiments were carried out carried out in 50 mM Tris buffer, pH 7.5 containing 1 mM DTT at 37°C. The reaction with MhpF contained 0.15 g/L TdTer, 0.1 g/L of MhpF, 7.5 mM NADH and 1.7 mM crotonyl-CoA. Reaction with Lmo1179 contained 0.15 g/L TdTer, 0.69 g/L Lmo1179, 7.5 mM NADH and 1.7 mM crotonyl-CoA. The assay was monitored by measuring production of butyraldehyde with HPLC. When assayed under the conditions described above, the combination of trans-2-enoyl-CoA reductase from *Treponema denticola* and *E. coli* enzyme MhpF was shown to produce butyraldehyde from crotonyl-CoA to a final concentration of 0.55 mM (FIG. 28).

**0161** Regeneration Module:

**0162** Starting from the enoyl-CoA of the elongation module, an enoyl-CoA hydratase converts the enoyl-CoA to the corresponding 3-hydroxyacetyl-CoA. The 3-hydroxyacetyl-CoA is then converted to a 3-ketoacyl-CoA by a 3-hydroxyacyl-CoA dehydratase. A 3-ketoacyl-CoA thiolase cleaves the 3-ketoacyl-CoA into an acetyl-CoA and a two carbon shortened acyl-CoA. In this implementation, the acetyl-CoA is converted to an aldehyde by one of the aforementioned acyl-CoA reductases and is used to prime the elongation cycle in another round of the cycle. The remaining acyl-CoA is converted to a carboxylic acid product by a thioesterase or otherwise handled by the termination module.

**0163** Native *E. coli* FadB has been shown to have both enoyl-CoA hydratase and 3-hydroxyacyl-CoA dehydratase activity. *E. coli* FadB was purified and assayed in vitro in the opposite direction (FIG. 29), though its physiological activity in the desired direction has been well characterized (Binstock & Schulte, 1981). The β-hydroxybutyryl-CoA dehydratase assays were performed in the presence of 1.5 mM DTT, 4.5 mM MgCl₂, 100 mM Tris HCl pH 7.5 and 0.2 mM NADH in a total volume of 200 μL at 25°C. β-hydroxybutyryl-CoA dehydratase activity was monitored by measuring the oxidation of NADH at 340 nm. Enoyl-CoA hydratase activity was monitored by following the loss of crotonyl-CoA at 263 nm (ε=6.7 mM⁻¹ cm⁻¹) in the presence of 100 mM Tris HCl pH 7.5 in 200 μL total volume. In these conditions, enoyl-CoA hydratase specific activity was measured to be 0.051±0.004 μmol/min/mg protein, while 3-hydroxyacyl-CoA dehydratase activity was measured to be 0.185±0.001 μmol/min/mg protein (FIG. 29).

**0164** Native *E. coli* thiolases AtoB and FadA were assayed for their activity on cleaving 3-ketobutyryl-CoA to two molecules of acetyl-CoA (FIG. 30). Thioleolate activity was determined in the presence of 0.5 mM DTT, 4.5 mM MgCl₂, 100 mM Tris HCl pH 7.5 and 2 mM CoA in a total volume of 200 μL at 25°C. The specific activities were measured as 0.36±0.05 μmol/min/mg protein for AtoB and 0.013±0.002 μmol/min/mg protein for FadA.

**0165** Termination Module:

**0166** Native *E. coli* thioesterases FadM, TesA, TesB, YbgC, YciA, and Ydil were characterized in vitro by following the production of TBN at 412 nm (ε=4.3 mM⁻¹ cm⁻¹). Reactions were carried out in the presence of 100 mM Tris pH 7.5, 200 mM KCl, 25 mM DTNB and 200 μM of the ‘CoA’ substrate in a volume of 200 μL at 25°C. Due to their varying substrate chain length specificities, selection
of the proper thioesterase for the desired product can be important to the optimal functionality of the pathway. The results of the in vitro characterization of the thioesterases on acetyl-CoA are summarized in Table 2 and in FIG. 31.

[0167] The regeneration and termination modules were assembled in vivo and crude extracts of cells expressing the module enzymes were assayed in vitro to demonstrate the functionality of the module. JST07, an E. coli MG1655 strain with native thioesterases knocked out, was used as a background from which AtoB and FadB, comprising the regeneration module, and YciA, comprising the termination module were overexpressed. With crotonyl-CoA as a substrate, this combination of regeneration and termination modules would be expected to produce acetate as a product. The reaction mixture contained 100 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, 200 mM KCl, 1.7 mM crotonyl-CoA, 1 mM DTT, 3.5 mM NAD⁺, 5 mM CoA and crude extract in a final volume of 300 µL. Reactions were carried out at 30°C and generation of acetic acid was measured by HPLC. Assay mixture for acetic acid analysis was filtered through Amicon spin filters (molecular cutoff of 10 kDa) and stored at 4°C prior to HPLC analysis. As shown in FIG. 32, acetate was observed in samples containing the entire regeneration and termination modules, but not in samples in which the termination module, YciA, was omitted. These results indicate the functional construction and expression of the combined regeneration and termination modules.

[0168] A simple implementation of the proposed pathway was implemented in vitro for the production of glycolic acid. In this implementation, formaldehyde was elongated by the one carbon unit formyl-CoA and the resulting 2-hydroxyacetyl-CoA, glycolyl-CoA, was diverted to the termination module consisting of a thioesterase to give glycolic acid. In the in vitro implementation, HACL1 and E. coli YciA were combined in a reaction mixture comprised of 60 mM potassium phosphate pH 5.4, 2.5 mM MgCl₂, 0.1 mM TPP, 6.6 µM BSA, 5 mM aldehyde, and approximately 1 mM formyl-CoA. The mixture was incubated at room temperature for 30 minutes and analyzed directly by HPLC, which revealed the glycolic acid was produced by the pathway FIG. 33.

In Silico Pathway Modeling

[0169] A mathematical model of the pathway implemented in E. coli as a host was developed and evaluated to assess the feasibility of the pathway for supporting product formation and organism growth on various one-carbon feedstocks. The genome-scale model (GEM) iJO1366 for E. coli was used as starting point (Orth et al., 2011). This model includes 1366 genes, 2251 reactions, and 1136 unique metabolites. While adapting GEM iJO1366 to simulate the fermentative metabolism of glycerol by wild-type E. coli in minimal medium, this model predicted operation of a number of pathways that are not functional under the simulated conditions but that artificially enabled redox balance. Consequently, we restricted the reactions involved exclusively in amino acid degradation, nucleotide degradation and export of amino acids, spermidine, 5-methylthio-d-ribose production, ferrous and hydrogen sulfide to zero flux (Cintolesi, et al., 2014). The reactions comprising the proposed pathway were then added to the GEM.

[0170] For simplification, the net reaction for a two-turn elongation cycle was used to give production of acetyl-CoA, which in turn allows for cell growth. This net reaction can be written:

2formyl-CoA+2NADH+2H⁺→1acetetyl-CoA+
1CoASH+2NAD++2H₂O.

[0171] In addition, reactions for the conversion of different single carbon substrates to formyl-CoA were added to the model. For formaldehyde the reaction took the form:

1formaldehyde+C₃H₅O→1formyl-CoA+
1NADH+1H⁺.

[0172] For formic acid, the reaction was written:

1HCOOH+1ATP+1CoA→1formyl-CoA+1ADP+1P₁.

[0173] In addition, the conversion of CO₂ and hydrogen to formic acid by CO₂ reductase (Schuchmann & Müller, 2013) was written:

1CO₂+1H₂→1HCOOH.

[0174] Conversion of methanol to formaldehyde was written:

1methanol+1NAD⁺→1formaldehyde+1NADH+1H⁺.

[0175] Conversion of methane to methanol by methane monooxygenases was written:

1methane+1O₂+1NADH+1H⁺→1methanol+1NAD⁺+
H₂O.

[0176] Flux balance analysis was used to evaluate the model. The evaluation was done using COBRA toolbox v2.0 (Schellenberger et al., 2011) in Matlab (THE MATHWORKS, MA). For comparison, the uptake of the one carbon substrates were set equal on a carbon mole basis (arbitrarily, 10 mmol C/g DCW/hr).

[0177] The ability for the pathway to support cell growth by the production of acetyl-CoA was then modeled. Because E. coli cannot natively grow on single carbon substrates, cell growth as a result of the inclusion of the pathway in the E. coli model indicates a functional pathway. Indeed, the results indicated that E. coli containing the designed pathway was able to grow on single carbon substrates ranging from CO₂ to methane (FIG. 34). Methanol appeared to allow for the highest growth rate, most likely due to the additional reducing equivalent obtained from methanol oxidation, which can be used to produce ATP for growth. Growth on formate and CO₂ was more challenging compared to the other substrates due to the ATP requirement of their conversion to formyl-CoA and because of their high oxidation state. This is a proof of principle demonstration that the pathway is capable of functioning as designed in the context of E. coli metabolism, as well as the metabolism of other organisms.
<table>
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<tr>
<th>Reaction</th>
<th>Illustration</th>
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<td>Methanol → Formaldehyde</td>
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<td>1.1.1.244 1.1.2.7 1.1.99.37</td>
<td>Methanol dehydrogenase</td>
<td><em>Bacillus methanocorpus</em> muh <em>Mycobacterium sp. DSM 3805</em> tulo <em>Methylobacterium cattouaens</em> mnoX, mnoY</td>
<td>P31005, CSMR78, P14775, P16027</td>
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<td>Formaldehyde → Formyl-CoA</td>
<td><img src="image7" alt="Diagram" /></td>
<td>1.2.1.10</td>
<td>Acylating aldehyde dehydrogenase</td>
<td><em>E. coli</em> mplF <em>Pseudomonas sp. CF600</em> dirpF</td>
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### TABLE 1B

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<td>Formyl-CoA + C(n+1)-acyl-CoA → 2-hydroxy-C(n+1)-acyl-CoA</td>
<td><img src="image1" alt="Illustration" /></td>
<td>4.1.--</td>
<td>2-hydroxyacyl-CoA lyase, 2-hydroxyacyl-CoA decarboxylase</td>
<td><em>Homo sapiens</em> lacI</td>
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<td>2-hydroxy-C(n+1)-acyl-CoA → &amp;trans-2-(C(n+1)-enoyl-CoA</td>
<td><img src="image2" alt="Illustration" /></td>
<td>4.2.1.54</td>
<td>2-hydroxyacyl-CoA dehydratase</td>
<td><em>Clostridium propionicum</em> IcdABC, <em>Clostridium difficile</em> lalBCI</td>
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### TABLE 1B-continued

**Summary of pathway reactions and enzymes**

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<td>1,2-C(n + 1)-dial → C(n + 1)-aldehyde</td>
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<td>4.2.1.17</td>
<td>Diol dehydratase</td>
<td><em>Klebsiella oxytoca</em></td>
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<td>1,2-C(n + 1)-dial</td>
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<td>4.2.1.13, 4.2.1.119</td>
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<td><em>E. coli</em> phdCDE</td>
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<td>C(n + 1)-aldehyde</td>
<td><img src="image" alt="Diagram" /></td>
<td>4.2.1.150</td>
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### TABLE 1C

**Summary of pathway reactions and enzymes**

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<td>Trans-2-C(n + m)-acyl-CoA → 3-hydroxy-C(n + m)-acyl-CoA</td>
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<td>4.2.1.17; 4.2.1.135; 1.1.1.35; 1.1.1.36</td>
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**TABLE 1C-continued**

Summary of pathway reactions and enzymes

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<td>Acetyl-CoA ↔ Acetaldehyde</td>
<td><img src="image1.png" alt="Reaction Illustration" /></td>
<td>1.2.1.10 Acetyl-CoA reductase</td>
<td><em>E. coli</em> exE, <em>Salmonella enterica</em> exE</td>
<td><em>E. coli</em> mhpF</td>
<td>NP_416959.1, AAA80299.1, NP_414888.1</td>
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**TABLE 1D**

Summary of pathway reactions and enzymes

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<td>Acyl-CoA ↔ Carbonylic acid</td>
<td><img src="image2.png" alt="Reaction Illustration" /></td>
<td>3.1.2.- Acetyl-CoA + acyl-CoA thioesterase 2.8.3.8 Acetyl-CoA:Acetyl-CoA thioesterase transacylase 2.3.1.-; 2.7.2.1; Phosphotransacylase</td>
<td><em>E. coli</em> ydiA, <em>E. coli</em> ybgC, <em>Alcaligenes lwoffii</em> ydiB, <em>Bacillus subtilis</em> ydiB</td>
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Termination module

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<td>Acetyl-CoA ↔ Alcohol</td>
<td><img src="image3.png" alt="Reaction Illustration" /></td>
<td>1.2.1.84 Alcohol-forming coenzyme-A thioester reductase</td>
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<td><em>Arabidopsis thaliana</em> At3g1980, <em>Arabidopsis ssp</em></td>
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<td>Acyl-CoA → Aldehyde</td>
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<td>Aldehyde forming CoA thioester reductase</td>
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Thr Met Asp Ile Gly Arg Thr Val Leu Gln Asn Tyr Leu Pro Arg His
405  410  415

Arg Leu Asp Ala Gly Thr Phe Gly Thr Met Gly Val Gly Leu Gly Phe
420  425  430

Ala Ile Ala Ala Ala Ala Val Val Ala Lys Asp Arg Ser Pro Gly Glu Trp
435  440  445

Ile Ile Cys Val Glu Gly Asp Ser Ala Phe Gly Phe Ser Gly Met Glu
450  455  460

Val Glu Thr Ile Cys Arg Tyr Asn Leu Pro Ile Ile Leu Leu Val Val
465  470  475  480

Asn Asn Asp Gly Ile Tyr Gin Gly Phe Asp Thr Asp Thr Trp Lys Glu
485  490  495

Met Leu Lys Phe Gin Asp Ala Thr Ala Val Val Pro Pro Met Cys Leu
500  505  510

Leu Pro Asn Ser His Tyr Glu Gin Val Met Thr Ala Phe Gly Gly Lys
515  520  525

Gly Tyr Phe Val Glu Thr Pro Glu Gin Gin Leu Gin Lys Ser Leu Arg Gin
530  535  540

Ser Leu Ala Asp Thr Thr Lys Pro Ser Leu Ile Asn Ile Met Ile Glu
545  550  555  560

Pro Gin Ala Thr Arg Lys Ala Gin Asp Phe His Trp Leu Thr Arg Ser
565  570  575

Ann Met

<210> SEQ ID NO 2
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<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

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Ala Lys Val Ile Ala Ala Leu Lys Thr Gin Asp Val Glu Tyr Ile
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Phe Gly Ile Val Gly Ile Pro Val Thr Glu Ile Ala Ile Ala Gin
35  40  45

Gln Leu Gly Ile Lys Tyr Ile Gly Met Arg Asn Glu Gin Ala Ala Cys
50  55  60

Tyr Ala Ala Ser Ala Ile Gly Tyr Leu Ser Ser Arg Pro Leu Leu Val
65  70  75  80

Ile Gly Gin Ser Gin Gin Gin Thr Met Gly Ala Phe Gin
85  90  95

Glu Phe Pro Gin Val Gin Ala Cys Arg Leu Tyr Thr Lys Phe Ser Ala
100 105 110

Arg Pro Ser Ser Ile Gin Ala Ile Pro Phe Val Ile Gin Lys Ala Val
115 120 125

Arg Ser Ser Ile Tyr Gin Arg Pro Gly Ala Cys Tyr Val Asp Ile Pro
130 135 140

Ala Asp Phe Val Gin Gin Met Asn Val Gin Val Gin Gin Gin Gin Gin Gin Glu
145 150 155 160

Glu Gin Cys Gin Ser Pro Gin Ile Ser Gin Ile Gin Glu Thr Ser Ala
Val 165 170 175
-continued-

Cys Thr Ala Ala Ser Val Ile Arg Asn Ala Lys Gin Pro Leu Leu Ile
180 185 190

Ile Gly Lys Gly Ala Ala Tyr Ala His Ala Glu Glu Ser Ile Lys Lys
195 200 205

Leu Val Glu Gin Tyr Lys Leu Pro Phe Leu Pro Thr Pro Met Gly Lys
210 215 220

Gly Val Val Pro Asp Asn His Pro Tyr Cys Val Gly Ala Ala Arg Ser
225 230 235 240

Arg Ala Leu Gin Phe Ala Asp Val Ile Val Leu Phe Gly Ala Arg Leu
245 250 255

Asn Trp Ile Leu His Phe Gly Leu Pro Pro Arg Tyr Gin Pro Asp Val
260 265 270

Lys Phe Ile Gin Val Asp Ile Cys Ala Glu Glu Leu Lys Asn Asn Val
275 280 285

Lys Pro Ala Val Thr Leu Leu Gly Asn Ile His Ala Val Thr Lys Gin
290 295 300

Leu Leu Glu Glu Leu Asp Lys Thr Pro Thr Gin Tyr Pro Pro Glu Ser
305 310 315 320

Lys Trp Thr Lys Thr Leu Arg Glu Lys Met Lys Ser Asn Glu Ala Ala
325 330 335

Ser Lys Glu Leu Ala Val Ser Lys Ser Leu Pro Met Asn Tyr Tyr Thr
340 345 350

Val Phe Tyr His Val Gin Glu Gin Leu Pro Arg Cys Phe Val Val
355 360 365

Ser Glu Gly Ala Asn Thr Met Asp Ile Gly Arg Thr Val Leu Gin Asn
370 375 380

Tyr Leu Pro Arg His Arg Leu Asp Ala Gly Thr Phe Gly Thr Met Gly
385 390 395 400

Val Gly Leu Gly Phe Ala Ile Ala Ala Val Val Ala Lys Asp Arg
405 410 415

Ser Pro Gly Gin Thr Ile Ile Cys Val Glu Gly Asp Ser Ala Phe Gly
420 425 430

Phe Ser Gly Met Glu Val Glu Thr Ile Cys Arg Tyr Asn Leu Pro Ile
435 440 445

Ile Leu Leu Val Asn Asn Asn Gly Ile Tyr Gin Gly Phe Asp Thr
450 455 460

Asp Thr Thr Lys Gin Met Leu Lys Phe Gin Asp Ala Thr Ala Val Val
465 470 475 480

Pro Pro Met Cys Leu Leu Pro Asn Ser His Tyr Gin Val Met Thr
485 490 495

Ala Phe Gly Lys Gly Tyr Phe Val Gin Thr Pro Glu Glu Leu Gin
500 505 510

Lys Ser Leu Arg Gin Ser Leu Ala Asp Thr Thr Lys Pro Ser Leu Ile
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Trp Leu Thr Arg Ser Asn Met
545 550

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<211> LENGTH: 496
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<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 3

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Ala Lys Val Ile Ala Gin Ala Leu Lys Thr Gin Asp Val Glu Tyr Ile
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Phe Gly Ile Val Gin Ile Pro Val Thr Glu Ile Ala Ile Ala Ala Gin
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Gln Leu Gly Ile Lys Tyr Ile Gly Met Arg Asn Glu Gin Ala Ala Cys
50     55     60

Tyr Ala Ala Ser Ala Ile Gly Tyr Thr Ser Arg Pro Gly Val Cys
65     70     75     80

Leu Val Val Ser Gly Pro Gly Leu Ile His Ala Leu Gly Gly Met Ala
85     90     95

Asn Ala Asn Met Asn Cys Trp Tyr Met Glu Arg Cys Met Ser Pro Pro
100    105    110

Ile Ser Met Ala Glu Thr Ser Ala Val Cys Thr Ala Ala Ser Val Ile
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Arg Asn Ala Lys Gin Pro Leu Leu Ile Ile Gly Lys Gly Ala Ala Tyr
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Ala His Ala Glu Gin Ser Ile Lys Leu Val Gin Thr Lys Leu
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Pro Phe Leu Pro Thr Pro Met Gly Gly Val Val Pro Asp Asn His
165    170    175

Pro Tyr Cys Val Gly Ala Ala Arg Ser Arg Ala Leu Gin Phe Ala Asp
180    185    190

Val Ile Val Leu Phe Gly Ala Arg Leu Asn Trp Ile Leu His Phe Gly
195    200    205

Leu Pro Pro Arg Tyr Gin Pro Asp Val Lys Phe Ile Gin Val Asp Ile
210    215    220

Cys Ala Glu Glu Leu Gly Asn Asn Val Lys Pro Ala Val Thr Leu Leu
225    230    235    240

Gly Asn Ile His Ala Val Thr Lys Gin Leu Leu Glu Leu Asp Lys
245    250    255

Thr Pro Trp Gin Tyr Pro Pro Glu Ser Lys Trp Trp Lys Thr Leu Arg
260    265    270

Glu Lys Met Lys Ser Asn Glu Ala Ala Ser Lys Glu Leu Ala Ser Lys
275    280    285

Lys Ser Leu Pro Met Asn Tyr Tyr Thr Val Phe Tyr His Val Gin Glu
290    295    300

Gln Leu Pro Arg Asp Cys Phe Val Ser Gin Asp Gin Ala Asn Met Thr Met
305    310    315    320

Asp Ile Gly Arg Thr Val Leu Gin Asn Tyr Leu Pro Arg Arg Ile Thr
325    330    335

Asp Ala Gly Thr Phe Gly Thr Met Gin Val Gin Leu Gly Phe Ala Ile
340    345    350

Ala Ala Ala Val Val Ala Lys Gin Arg Ser Pro Gin Lys Gin Thr Ile
355    360    365

Cys Val Glu Gly Asp Ser Ala Phe Gly Phe Ser Gin Met Glu Val Glu
370    375    380
Thr Ile Cys Arg Tyr Asn Leu Pro Ile Ile Leu Leu Val Val Asn Asn
385 390 395 400
Asn Gly Ile Tyr Gin Gly Phe Asp Thr Asp Thr Trp Lys Glu Met Leu
405 410 415
Lys Phe Gin Asp Ala Thr Ala Val Val Pro Pro Met Cys Leu Leu Pro
420 425 430
Asn Ser His Tyr Gin Val Met Thr Ala Phe Gly Gly Lys Gly Tyr
435 440 445
Phe Val Gin Thr Pro Gin Leu Gin Ser Leu Arg Gin Gin Ser Leu
450 455 460
Ala Asp Thr Thr Lys Pro Ser Leu Ile Asn Ile Met Ile Gin Pro Gin
465 470 475 480
Ala Thr Arg Lys Ala Gin Asp Phe His Trp Leu Thr Arg Ser Asn Met
485 490 495

<210> SEQ ID NO 4
<211> LENGTH: 518
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

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Ala Lys Val Ile Ala Gin Ala Leu Lys Thr Gin Asp Val Gin Tyr Ile
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Phe Gin Ile Val Gin Gin Gin Gin Pro Val Thr Glu Ile Ala Gin
40  45
Gln Leu Gin Gin Gin Tyr Gin Gin Gin Gin Gin Gin Gin Gin Gin
65  70  75  80
Val Ala Ser Ala Cys Gin Tyr Gin Gin Gin Gin Gin Gin Gin Gin
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Tyr Ala Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Val Leu Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
115
Ile Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
125
Glu Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
135
Asn Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Gln Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Met Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Val Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Val Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Ile Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Val Leu Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Lys Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Lys Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Arg Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Arg Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
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Phe Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin Gin
255
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Val Lys Phe Ile Gln Val Asp Ile Cys Ala Gln Leu Leu Gly Aen Aen 275 280 285
Val Lys Pro Ala Val Thr Leu Leu Gly Aen Ile His Ala Val Thr Lys 290 295 300
Gln Gln Leu Ala Ser Lys Lys Ser Leu Pro Met Aen Tyr Tyr Thr Val 310 315 320 325
Phe Tyr His Val Gln Gln Leu Pro Arg Asp Cys Phe Val Val Ser 330 335
Glu Gly Ala Aen Thr Met Asp Ile Gly Arg Thr Val Leu Gln Aen Tyr 340 345 350 355
Leu Pro Arg His Arg Leu Asp Ala Gly Thr Phe Gly Thr Met Gly Val 360 365
Gly Leu Gly Phe Ala Ile Ala Ala Val Val Ala Lys Arg Arg Ser 370 375 380 385
Pro Gly Gin Trp Ile Ile Cys Val Gly Asp Ser Ala Phe Gly Phe 390 395 400
Ser Gly Met Glu Val Glu Thr Ile Cys Arg Tyr Aen Leu Pro Ile Ile 405 410 415
Leu Leu Val Val Aen Aen Aen Gly Ile Tyr Gln Gly Phe Asp Thr Asp 420 425 430 435
Thr Trp Lys Glu Met Leu Lys Phe Gin Asp Ala Thr Ala Val Val Pro 440 445 450
Pro Met Cys Leu Leu Pro Aen Ser His Tyr Glu Gin Val Met Thr Ala 455 460 465
Phe Gly Gly Lys Gly Tyr Phe Val Gin Thr Pro Glu Gin Thr Leu Gin Lys 470 475 480 485
Ser Leu Arg Gin Ser Leu Ala Asp Thr Thr Lys Pro Ser Leu Ile Aen 490 495
Ile Met Ile Glu Pro Gin Ala Thr Arg Lys Ala Gin Aen Phe His Thr 500 505 510
Leu Thr Arg Ser Aen Met 515

1-72. (canceled)

73. A genetically modified microorganism comprising a DNA molecule encoding a polypeptide that catalyzes a substrate to product conversion resulting in the condensation of an initial C(n) aldehyde with formyl-CoA to produce a 2-hydroxy-C(n+1)-acyl-CoA, wherein the 2-hydroxy-C(n+1)-acyl-CoA is one carbon longer than the initial C(n) aldehyde, and wherein n refers to the number of carbons in the initial aldehyde.

74. The genetically modified microorganism of claim 73, wherein said polypeptide is a 2-hydroxyacetyl-CoA lyase or a naturally occurring homolog thereof.

75. The genetically modified microorganism of claim 73 further comprising:

a. at least one DNA molecule encoding a polypeptide that catalyzes the conversion of a single carbon substrate to formyl-CoA,
b. at least one DNA molecule encoding a polypeptide that catalyzes conversion of the 2-hydroxy-C(n+1)-acyl-CoA to a C(n+1)-aldehyde, and
c. at least one DNA molecule encoding a polypeptide that catalyzes conversion of an intermediate of the prior step to a desired product, wherein the desired product contains more than one carbon atom, and
d. optionally comprising at least one DNA molecule encoding a polypeptide that catalyzes regeneration of the initial C(n) aldehyde molecule.

76. The genetically modified microorganism of claim 75, wherein:

a. the polypeptide that catalyzes the condensation of the formyl-CoA and the C(n)-aldehyde to the 2-hydroxy-C(n+1)-acyl-CoA is overexpressed, and wherein the polypeptide is selected from the group consisting of a 2-hydroxyacetyl-CoA lyase, an oxalyl-CoA decarboxylase and a benzaldehyde lyase; and wherein the microorganism further comprises:
b. a DNA molecule that encodes a 2-hydroxyacyl-CoA dehydratase that catalyzes the conversion of the 2-hydroxy-C(n+1)-acyl-CoA to a trans-2-(C(n+1))-enoyl-CoA; and

c. a DNA molecule that encodes a 2-enoyl-CoA reductase that catalyzes the conversion of the 2-enoyl-(n+1)-enoyl-CoA to a (C(n+1))-acyl-CoA; and

d. a DNA molecule that encodes an acyl-CoA reductase that catalyzes the conversion of the C(n+1)-acyl-CoA to the (C(n+1))-aldehyde.

77. The genetically modified microorganism of claim 76, wherein:

a. the 2-hydroxyacyl-CoA lyase is an amino acid sequence encoded by a gene selected from Homo sapiens hac11 (Q9UJ83), Rattus norvegicus hac11 (Q7TCH7), Dictyostelium discoideum hac11 (Q54DA9), Mus musculus hac11 (Q9QXW0), or homologues thereof, and wherein the oxalyl-CoA decarboxylase is an amino acid sequence encoded by a gene selected from Mycobacterium sp. MOTT36Y W7S-25140 (YP_006308704.1), Mycobacterium tuberculosis TKK-01-0051 K875_00487 (KBZ76942.1), Poly-nucleobacter necessarius subsp. asymptoticus QLW-P1DMWA-1 PmcB_0637 (YP_001155419.1), E. coli oxc (NP_416874.1), or homologues thereof;

b. the 2-hydroxyacyl-CoA dehydratase is an amino acid sequence encoded by a gene selected from Clostridium propioniciucus ldcABC (G3KIM3, G3KIM4, G3KIM5), Clostridium difficile ldcABC (AAV-40818.1, AAU-40819.1, AAV-40820.1), or homologues thereof; and

c. the 2-enoyl-CoA reductase is an amino acid sequence encoded by a gene selected from Rhodobacter sphaeroides Acul (Q3K691.1), E. coli ydhH (NP_417719.1), E. coli fabl (NP_415804.1), Enterococcus faecalis fabk (NP_816503.1), Bacillus subtilis fabl (KF98655.1), Euglena gracilis TER (Q5EU90.1), Treponema denticola TER (NP_971211.1), Vibrio cholera fabH (B1P0R8), or homologues thereof; and

d. the acyl-CoA reductase is an amino acid sequence encoded by a gene selected from Acinetobacter calcoaceticus acr1 (AAC45217.1), Acinetobacter sp Strain M-1 acrM (BA885476.1), Clostridium beijerinckii acrD (AA766436.1), E. coli euE (NP_416950.1), Salmonella enterica euE (AAAA80209.1), E. coli mhpF (NP_414885.1), or homologues thereof.

78. The genetically modified microorganism of claim 75, wherein:

a. the polypeptide that catalyzes the condensation of the formyl-CoA and the (n)-aldehyde to the 2-hydroxy-C(n+1)-acyl-CoA is overexpressed, and wherein the polypeptide is selected from the group consisting of a 2-hydroxyacyl-CoA lyase, an oxalyl-CoA decarboxylase and a benzaldehyde lyase; and wherein the microorganism further comprises:

b. a DNA molecule that encodes an acyl-CoA reductase that catalyzes the conversion of the 2-hydroxy-C(n+1)-acyl-CoA to a 2-hydroxy-C(n+1)-aldehyde; and

c. a DNA molecule that encodes a polypeptide that catalyzes the conversion of the 2-hydroxy-C(n+1)-aldehyde to a 1,2-(C(n+1))-diol, wherein the polypeptide is a 1,2-diol oxidoreductase or an alcohol dehydrogenase; and

d. a DNA molecule that encodes a diol dehydratase that catalyzes the conversion of the 1,2-(C(n+1))-diol to the (C(n+1))-aldehyde.

79. The genetically modified microorganism of claim 78, wherein:

a. the 2-hydroxyacyl-CoA lyase is an amino acid sequence encoded by a gene selected from Homo sapiens hac11 (Q9UJ83), Rattus norvegicus hac11 (Q7TCH7), Dictyostelium discoideum hac11 (Q54DA9), Mus musculus hac11 (Q9QXW0), or homologues thereof, and wherein the oxalyl-CoA decarboxylase is an amino acid sequence encoded by a gene selected from Mycobacterium sp. MOTT36Y W7S-25140 (YP_006308704.1), Mycobacterium tuberculosis TKK-01-0051 K875_00487 (KBZ76942.1), Poly-nucleobacter necessarius subsp. asymptoticus QLW-P1DMWA-1 PmcB_0637 (YP_001155419.1), E. coli oxc (NP_416874.1), or homologues thereof;

b. the acyl-CoA reductase is an amino acid sequence encoded by a gene selected from Acinetobacter calcoaceticus acr1 (AAC45217.1), Acinetobacter sp Strain M-1 acrM (BA885476.1), Clostridium beijerinckii acrD (AA766436.1), E. coli euE (NP_416950.1), Salmonella enterica euE (AAAA80209.1), E. coli mhpF (NP_414885.1), or homologues thereof;

the 1,2-diol oxidoreductase or alcohol dehydrogenase is an amino acid sequence encoded by a gene selected from E. coli betA (NP_414845.1), E. coli dkgA (NP_417884.5), E. coli euG (NP_416948.4), E. coli ficO (NP_417279.2), E. coli ucpA (NP_416291.4), E. coli yahK (NP_414859.1), E. coli yblO (NP_415026.1), E. coli yblE (NP_415152.1), E. coli yigY (YP_026233.1), E. coli yigB (NP_416990.4), or homologues thereof;

d. the diol dehydratase is an amino acid sequence encoded by a gene selected from Klebsiella oxytoca pddABC (Q59470, Q59471, Q59472), E. coli pddCDE (CAS09680, CAS09681, CAS09682), S. enterica pddCDE (NP_456590, NP_456591, NP_456592), or homologues thereof.

80. The genetically modified microorganism of claim 75, wherein the single carbon substrate is carbon dioxide, and wherein the at least one DNA molecule encoding a polypeptide that catalyzes the conversion of the single carbon substrate comprises:

a. a DNA molecule encoding a carbon dioxide reductase that catalyzes the conversion of carbon dioxide to formate;

b. a DNA molecule encoding an acetate or formate kinase that catalyzes the conversion of the formate to formyl-phosphate; and

c. a DNA molecule encoding a phosphatase acetyl-transferase that catalyzes the conversion of the formyl-phosphate to the formyl-CoA.

81. The genetically modified microorganism of claim 80, wherein:

a. said carbon dioxide reductase is an amino acid sequence encoded by a gene selected from Acetobacterium woodii f0liF2, lycC2, lycB3, hydA2 (YP_002658050.1, YP_002658053.1, YP_002658050.1, YP_002658050.1), or homologues thereof;

b. said formate kinase is an amino acid sequence encoded by a gene selected from Salmonella typhimurium sAeC (P63441), E. coli AAcE (NP_416799.1), or homologues thereof; and
c. said phosphate formyl-transferase is an amino acid sequence encoded by a gene selected from E. coli endD (NP_416953.1), E. coli pta (NP_416800.1), Salmonella typhimurium PduL (Q9KDN5), or homologues thereof.

82. The genetically modified microorganism of claim 75, wherein the conversion of the intermediate to the desired product is selected from the group consisting of:

a. the conversion of an acyl-CoA to a carboxylic acid;

b. the conversion of an acyl-CoA to a terminal alcohol;

c. the conversion of an acyl-CoA to an aldehyde;

d. the conversion of an aldehyde to a terminal alcohol;

e. the conversion of an aldehyde to a terminal alkyl; and

f. the conversion of an aldehyde to a terminal amine; and wherein:

i. the substrate to product conversion of (a) is performed by a thioesterase, acyl-CoA:acyl-CoA transferase, or phosphotransacetylase and carbonyl kinase; ii. the substrate to product conversion of (b) is performed by an alcohol-forming coenzyme-A thioester reductase;

iii. the substrate to product conversion of (c) is performed by an aldehyde-forming coenzyme-A thioester reductase;

iv. the substrate to product conversion of (d) is performed by an alcohol dehydrogenase;

v. the substrate to product conversion of (e) is performed by an aldehyde dehydrogenase; and

vi. the substrate to product conversion of (f) is performed by a transaminase.

83. The genetically modified microorganism of claim 82, wherein:

a. said overexpressed thioesterase is an amino acid sequence encoded by a gene selected from E. coli tesA (NP_415027.1), E. coli tesB (NP_414986.1), E. coli yciA (NP_415769.1), E. coli fisD (NP_414977.1), E. coli ydiF (NP_416201.1), E. coli ybgC (NP_415624.1), Alcanivorax borkumensis tesB2 (YP_692749.1), Fibrobacter succinogenes Fas2108 (YP_005822012.1), Prevotella ruminicola Pr655 (YP_003574018.1), Prevotella ruminicola Pr1687 (YP_003574982.1), or homologues thereof;

b. said overexpressed acyl-CoA:acyl-CoA transferase is an amino acid sequence encoded by a gene selected from E. coli aotD (NP_416725.1), Clostridium kluveri cat2 (AA92344.1), Clostridium acetobutylicum etfAB (NP_149326.1, NP_149327.1), E. coli ydiF (NP_416209.1), or homologues thereof;

c. said overexpressed phosphotransacetylase is an amino acid sequence encoded by a gene selected from Clostridium acetobutylicum pbtB (NP_349676.1), Enterococcus faecalis pbt (AAD55374.1), Salmonella enterica pduL (AAD39011.1), or homologues thereof;

d. said overexpressed carbonyl kinase is an amino acid sequence encoded by a gene selected from Clostridium acetobutylicum buk (AAK81015.1), Enterococcus faecalis buk (AAD55375.1) or Salmonella enterica pduW (AAD39021.1), or homologues thereof;

e. said overexpressed alcohol-forming coenzyme-A thioester reductase is an amino acid sequence encoded by a gene selected from Clostridium acetobutylicum adhE2 (YP_009076789.1), Arabidopsis thaliana ATh2a (NP_415089.1), Arabidopsis thaliana ATh344560 (AEE77915.1), Arabidopsis thaliana ATh356700 (AEE79553.1), Arabidopsis thaliana ATh522250 (AED93034.1), Arabidopsis thaliana CER4 (AEE86278.1), Marinobacter aquaeolei VT8 maqu_2220 (YP_959486.1), Marinobacter aquaeolei VT8 maqu_2507 (YP_959769.1), or homologues thereof;

f. said overexpressed alcohol dehydrogenase is an amino acid sequence encoded by a gene selected from E. coli beta (NP_414845.1), E. coli dkgA (NP_414785.4), E. coli endG (NP_416948.4), E. coli fucO (NP_417279.2), E. coli ucpA (NP_416921.4), E. coli yahK (NP_414859.1), E. coli yboO (NP_415026.1), E. coli ydhH (NP_415132.1), E. coli yiaV (YP_026233.1), E. coli ygb (NP_416890.4), or homologues thereof;

h. said dehydrogenase is an amino acid sequence encoded by a gene selected from Synochococcus elongatus PCC7942 orf1593 (Q54764.1), Nostoc punctiforme PCC73102 orf381711 (B2J1M1.1), Prochlorococcus marinus MIT9313 orf3817111 (Q7V6D1.1), or homologues thereof;

i. said overexpressed transaminase is an amino acid sequence encoded by a gene selected from Arabidopsis thaliana At3g22200 (NP_001189947.1), Alcaligenes denitrificans AptA (AA92672.1), Bordetella bronchiseptica BB0869 (WP_015041039.1), Bordetella parapertussis BP07064 (WP_010927683.1), Brucella melitensis BAWG_0478 (EEW88370.1), Buthroderma pseudonabalu BP1026B_10669 (AF165333.1), Chromobacterium violaceum CV0252 (AAQ50697.1), Oceanicola granulosus OG2516_07293 (WP_007254984.1), Paracoccus denitrificans PD1222 Pden_2984 (ABL72050.1), Pseudogulbenkiania ferruxoxidans o-Ta (WP_008952788.1), Pseudomonas putida o-Ta (P28629.1), Ralstonia solanacearum o-Ta (YP_000255833.1), Rhizobium melliflori SME01534 (NP_386510.1), Vibrio fluvialis o-Ta (AEA39183.1), Manau musculus abal (AAH155821.1), E. coli gabT (YP_490877.1), or homologues thereof.

84. The genetically modified microorganism of claim 75, wherein the at least one DNA molecule encoding a polypeptide that catalyzes regeneration to a C(4) aldehyde comprises:

a. a DNA molecule encoding an enoyl-CoA hydratase that catalyzes the conversion of a trans-2-enoyl-CoA to a 3-hydroxy-acyl-CoA;

b. a DNA molecule encoding a 3-hydroxyacyl-CoA dehydrogenase that catalyzes the conversion of the 3-hydroxy-acyl-CoA to a 3-keto-acyl-CoA;

c. a DNA molecule encoding a 3-ketoacyl-CoA thiolase that catalyzes the conversion of the 3-keto-acyl-CoA to acetyl-CoA and an acyl-CoA shortened by two carbons;

d. a DNA molecule encoding an acetyl-CoA reductase that catalyzes the conversion of the acetyl-CoA to the acetaldehyde.

85. The genetically modified microorganism of claim 84, wherein:

a. the enoyl-CoA hydrolase is an amino acid sequence encoded by a gene selected from E. coli fisD (NP_416843.1), E. coli fisD (NP_416843.1), Aeromonas
caviae phaJ (O32472.1), Pseudomonas aeruginosa phaJ1 (BA92740.1), Pseudomonas aeruginosa phaJ2 (BA92741.1), Pseudomonas aeruginosa phaJ3 (BAC4834.1), Pseudomonas aeruginosa phaJ4 (BAC4835.1), or homologs thereof.

b. the 3-hydroxyacyl-CoA dehydrogenase is an amino acid sequence encoded by a gene selected from E. coli fadB (NP_416828.1), E. coli fadD (NP_416843.1), Ralstonia eutropha phaB1 (YP_725942.1), Ralstonia eutropha phaB2 (YP_726470.1), Ralstonia eutropha phaB3 (YP_726636.1), or homologs thereof.

c. the 3-ketoacyl-CoA thiolase is an amino acid sequence encoded by a gene selected from E. coli atoB (NP_416728.1), E. coli yqEF (NP_417321.2), E. coli fadA (YP_026272.1), E. coli fadD (NP_416844.1), Ralstonia eutropha bktB (AAC38322.1), Pseudomonas sp. Strain 313 catF (AAU20407.1), E. coli pmJ (NP_415915.1), Rhodococcus opacus pcaF (YP_002778248.1), Streptomyces sp. pcaF (A222035.1), Ralstonia eutropha phaA (AEF80291.1), Clostridium acetobutylicum thA (AAC26023.1), Clostridium acetobutylicum thB (A2C6026.1), or homologs thereof.

d. the acetyl-CoA reductase is an amino acid sequence encoded by a gene selected from E. coli catE (NP_416950.1), Salmonella enterica catE (AA80209.1), E. coli mhpF (NP_414885.1), or homologs thereof.

86. The genetically modified microorganism of claim 75, wherein the desired product is a carboxylic acid.

87. The genetically modified microorganism of claim 75, wherein the desired product is a primary alcohol.

88. The genetically modified microorganism of claim 75, wherein the desired product is selected from a group consisting of an unsaturated fatty acid.

89. The genetically modified microorganism of claim 75, wherein the desired product is selected from a group consisting of an unsaturated fatty acid.

90. A method for the production of the desired product from the single carbon substrate comprising the steps of:

a. growing the microorganism of claim 75 in a culture comprising the single carbon substrate and a growth media;

b. converting said single carbon substrate to the formyl-CoA, condensing the initial C(n) aldehyde with the formyl-CoA to produce the 2-hydroxy-C(n+1)-acyl-CoA, and converting the 2-hydroxy-C(n+1)-acyl-CoA to a C(n+1) aldehyde;

c. converting the intermediate to the desired product; and

d. isolating said desired product from said microorganism or said growth media, or both.

91. A method for the production of the desired product from the single carbon substrate comprising the steps of:

a. providing a reaction mixture comprising the isolated polypeptides from the microorganism of claim 75;

b. contacting the reaction mixture with a single carbon substrate under conditions wherein the single carbon substrate is converted to the formyl-CoA, the formyl-CoA is condensed with an initial C(n) aldehyde to produce the 2-hydroxy-C(n+1)-acyl-CoA, and the 2-hydroxy-C(n+1)-acyl-CoA is converted to a C(n+1) aldehyde, wherein the intermediate is converted to the desired product; and

c. isolating said desired product from said reaction mixture;

wherein the single carbon mole molecule substrate is selected from a group consisting of methanol, formaldehyde, formate, carbon monoxide, carbon dioxide, formyl-CoA and combinations thereof.

92. A genetically modified microorganism according to claim 75, wherein the microorganism is selected from the group consisting of Escherichia, Bacillus, Corynebacterium, Streptomyces, Pseudomonas, Enterococcus, Alcaligenes, Klebsiella, Paenibacillus, Arthrobacter, and Hansenula.

93. The genetically modified microorganism of claim 75, wherein the single carbon substrate is formate and wherein the at least one DNA molecule encoding the conversion of formate to formyl-CoA is acetyl CoA synthetase.

94. The genetically modified microorganism of claim 75, wherein the single carbon substrate is methanol and wherein the at least one DNA molecule encoding a polypeptide that catalyzes the conversion of the single carbon substrate comprises:

a. a DNA encoding a methane monoxygenase that catalyzes the conversion of methane to methanol; and

b. a DNA encoding a methanol dehydrogenase that catalyzes the conversion of methanol to formaldehyde.

c. a DNA encoding an acetylating dehydrogenase that catalyzes the conversion of formaldehyde to formyl-CoA.

95. The genetically modified microorganism of claim 93, said acetyl-CoA synthetase is an amino acid sequence encoded by Escherichia coli (NP_418493.1), or homologs thereof.

96. The genetically modified microorganism of claim 94, wherein:

a. said methane monoxygenase is an amino acid sequence encoded by a single gene selected from Methylosinus trichosporium OB3b mmoXYZB, orfY (P27353, P27354, P27355, Q53563, P27356, Q53562), Methylococcus capsulatus Bath mnoXYZB, orfY (P22860, P18798, P11987, P18797, 22868, P22867), or homologs thereof.

b. said methanol dehydrogenase is an amino acid sequence encoded by a single gene selected from Bacillus methanolicus mdh (P31005), Mycobacterium sp. DSM 3803 mno (C5MRT8), Methylobacterium extorquens mox, moxP (P14775, P16027) or homologs thereof; and

c. said acetylating dehydrogenase is an amino acid sequence encoded by a single gene selected from E. coli mhpF (NP_414885.1), Pseudomonas sp. CL600 dmpF (Q52060) or homologs thereof.

97. The genetically modified microorganism of claim 86, wherein the carbonylic acid is selected from the group consisting of carboxylic acids, (2R)-α-hydroxy carboxylic acids, β-keto carboxylic acids, and α,β-unsaturated carboxylic acids.
98. The genetically modified microorganism of claim 87, wherein the primary alcohol is selected from the group consisting of 1,(3R)-β-diol, 1,(2R)-α-diol, α-keto primary alcohols, and α,β-unsaturated primary alcohols.