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Citation: Dhamankar, Himanshu, and Kristala LJ Prather 2011 Microbial Chemical Factories: Recent Advances in Pathway Engineering for Synthesis of Value Added Chemicals. *Current Opinion in Structural Biology* 21(4): 488–494.

As Published: <http://dx.doi.org/10.1016/j.sbi.2011.05.001>

Publisher: Elsevier B.V.

Persistent URL: <http://hdl.handle.net/1721.1/79702>

Version: Author's final manuscript: final author's manuscript post peer review, without publisher's formatting or copy editing

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Microbial Chemical Factories: Recent Advances in Pathway Engineering for Synthesis of Value Added Chemicals

Himanshu Dhamankar and Kristala L. J. Prather*

Department of Chemical Engineering, Synthetic Biology Engineering Research
Center (SynBERC), Massachusetts Institute of Technology, Cambridge, MA 02139

* Corresponding author,

77 Massachusetts Avenue, Room 66-454

Cambridge, MA 02139

617-253-1950

kljp@mit.edu

Keywords: *de novo* pathway design, microbial chemical factories, pathway engineering, enzyme
promiscuity, protein engineering, cofactor balances

Running title: **Pathway Engineering for Synthesis of Value Added Chemicals**

1 **Abstract**

2 The dwindling nature of petroleum and other fossil reserves has provided impetus towards microbial
3 synthesis of fuels and value added chemicals from biomass-derived sugars as a renewable resource.
4 Microbes have naturally evolved enzymes and pathways that can convert biomass into hundreds of
5 unique chemical structures, a property that can be effectively exploited for their engineering into
6 Microbial Chemical Factories (MCFs). *De novo* pathway engineering facilitates expansion of the
7 repertoire of microbially synthesized compounds beyond natural products. In this review, we visit some
8 recent successes in such novel pathway engineering and optimization, with particular emphasis on the
9 selection and engineering of pathway enzymes and balancing of their accessory cofactors.

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1 Introduction

2 We rely heavily on the dwindling resource of crude petroleum to supply not only fuels but also
3 an array of valuable chemicals. Globally, roughly 20% of crude petroleum consumed is used for the
4 synthesis of products other than transportation and industrial fuels, a sizeable fraction of which are
5 petrochemicals [1]. Petrochemicals are used as raw materials in the manufacturing of a variety of
6 products such as polymers, textiles, paints, solvents, pharmaceuticals, detergents, waxes and lubricants
7 that find applications in our day-to-day lives. Reducing our dependence on petroleum by utilizing
8 biomass as a renewable resource requires an integrated approach towards engineering of
9 microorganisms into 'microbial chemical factories' (MCFs) that can be used in a 'biorefinery' for the
10 conversion of biomass into both fuels and value added biochemicals. Biosynthesis of high value chemical
11 compounds from biomass using natural or engineered pathways in microorganisms also serves as a
12 promising alternative to chemical synthesis processes that employ expensive, hazardous and non-
13 renewable raw materials and reagents as well as harsh processing conditions.

14 The evolution of the field of metabolic engineering has led to the development of principles and
15 tools that enable construction and optimization of MCFs by tapping into naturally occurring pathways in
16 specific host organisms, heterologous expression of non-native pathways in well-characterized hosts, or
17 engineering *de novo* biosynthetic pathways for synthesis of various natural and non-natural products. *De*
18 *novo* pathway engineering refers to the design and construction of novel pathways (hitherto unknown in
19 nature in any single organism) by assembling multiple existing partial pathways from different organisms
20 or using promiscuous or engineered enzymes as biocatalysts to catalyze a series of biotransformations
21 with non-natural substrates [2*]. This capability is of particular importance for truly expanding the
22 repertoire of value added products that can be synthesized microbially.

23 Figure 1 outlines the steps involved in the design of novel pathways for MCFs and various tools
24 and approaches used for their optimization. Once a pathway has been designed and selected for

1 experimental exploration [3*], suitable pathway enzymes need to be selected to catalyze the reaction
2 steps. Pathway enzymes are the tireless machines of the MCFs that sequentially process raw materials
3 into desired value added products and govern the pathway rates, selectivity, yield and overall
4 productivity. Whenever possible, enzymes known to specifically catalyze reactions with the respective
5 pathway intermediates are selected. If no enzyme has been documented to catalyze the required
6 biochemical reaction with the specific pathway intermediate, other enzymes exhibiting the required
7 biocatalytic activity known to act with structurally and chemically similar substrates may be selected to
8 take advantage of natural enzyme promiscuity exhibited by many enzymes. The selected enzymes are
9 then expressed in a suitable organism grown in a culture medium supplemented with the required
10 starting materials for the microbial synthesis of the desired value added product.

11 Once a novel or natural pathway has been established and the synthesis of the product of
12 interest has been demonstrated, enhancing pathway yields is critical to attain industrially viable product
13 titers. Protein engineering can be used for improving the selectivity and activity of the pathway enzymes
14 and can effectively complement conventional metabolic engineering approaches such as increasing the
15 precursor supply by varying pathway enzyme expression levels or knocking out competing pathways to
16 enhance productivity. Further, pathway enzymes often require cofactors for catalysis. Imbalances and
17 limitations arising out of an inability to effectively recycle cofactors can hamper pathway productivity,
18 warranting specific efforts for cofactor regeneration.

19 A number of research groups have employed combinations of these strategies towards
20 developing novel pathways and enhancing productivities of already established pathways for the
21 microbial synthesis of a number of value added biochemicals including the polymer and pharmaceutical
22 building blocks putrescine [4], cadaverine [5], succinic acid [6,7], 3-hydroxybutyric acid [8] and 3-
23 hydroxyvaleric acid [9], native silk protein [10], the high value flavoring agent natural vanillin [11,12],
24 pharmaceutical drug precursors such as taxadiene [13], amorpho-4,11-diene [14] and D-glucaric acid

1 [15] and plant secondary metabolites such as flavanoids, stilbenoids [16] and isoprenoids [14,17]. In this
2 review, we visit some recent studies focusing on the design, engineering and optimization of MCFs for
3 synthesis of value-added products, with particular emphasis on exploiting enzyme promiscuity for *de*
4 *novo* pathway construction and engineering of pathway enzymes and cofactor balances to enhance
5 pathway productivity.

6 **Exploiting enzyme promiscuity for design and engineering of novel pathways**

7 All living cells metabolize a limited set of common metabolites into a wide range of molecules
8 essential for the sustenance of various life processes. The key to the generation of such molecular
9 diversity is the array of enzymes coded by the genomes, capable of catalyzing a wide variety of
10 biochemical transformations, often with more than one substrate. Enzyme promiscuity [18,19*] refers
11 to the ability to catalyze the same reactions with a variety of structurally and chemically similar
12 substrates (substrate promiscuity) or multiple different chemical reactions (catalytic promiscuity)
13 sharing certain mechanistic features and is critical for expanding natural cell metabolism for the
14 biosynthesis of valuable new chemicals. While the use of catalytic promiscuity for development of novel
15 biocatalysts has been extensively reviewed [20-22], exploiting natural substrate promiscuity for catalysis
16 in novel pathways has received less attention. Indeed, many enzymes naturally exhibit a broad
17 substrate range by accommodating alternative molecules in their active site, a property that can be
18 effectively employed for catalysis with non-native substrates in novel pathways.

19 The Liao group at UCLA [23**,24**,25] has effectively employed such promiscuity for the
20 production of various non-natural alcohols and amino acids in *E. coli* from various 2-ketoacid
21 intermediates in the native branched chain amino acid synthesis pathways. The inherent ability of the
22 enzymes LeuA, LeuC and LeuD of the Leu operon to initiate chain elongation with their natural
23 substrate, 2-ketoisovalerate during L-leucine synthesis, was extended to the structurally similar natural

1 metabolite 2-keto-3-methylvalerate (precursor to L-isoleucine) followed by decarboxylation by the
2 broad substrate range enzyme 2-ketoisovalerate decarboxylase (KIVD) from *Lactococcus lactis* and
3 reduction by alcohol dehydrogenase VI (Adh6) from *Saccharomyces cerevisiae* Adh6 for synthesis of the
4 novel branched chain alcohol (S)-3-methyl-1-pentanol. In a separate study, the authors explored the
5 ability of the *E. coli* transaminase IlvE and valine dehydrogenases from various *Streptomyces* species to
6 catalyze amination of 2-ketobutyrate for the synthesis of the non-natural amino acid L-homoalanine
7 [24**]. Our group has effectively used enzyme promiscuity exhibited by enzymes of the butanol and
8 polyhydroxyalkanoate synthesis pathways from different organisms for the microbial synthesis of
9 enantiopure forms of the biodegradable polymer precursor 3-hydroxyvaleric acid as a value-added
10 product[9].

11 In each of these studies, enzymes chosen to be tested for promiscuity were selected based on
12 knowledge of catalytic activity on structurally similar native substrates or functional groups. Screening of
13 multiple possible enzymes exhibiting the required biocatalytic activity from different organisms can be
14 an effective strategy for sampling different naturally evolved enzyme activities and specificities to
15 identify an enzyme exhibiting activity towards a required pathway intermediate. For example, in the
16 synthesis of non-natural alcohols, the Liao group screened 7 different ketoacid decarboxylases from four
17 different organisms before identifying KIVD as a broad substrate range ketoacid decarboxylase [25].
18 Computational approaches may be effectively employed to comb through the extensive genomic,
19 metabolic and enzyme databases such as KEGG [26], BRENDA [27], BioCYC [28]and NCBI
20 (<http://www.ncbi.nlm.nih.gov/guide/>) that classify well-characterized natural enzymes on the basis of
21 the specific and generalized reactions they catalyze, in order to identify such enzyme candidates for
22 screening [29,30]. BLAST searches (blastp) using a protein sequence of known activity as a query may be
23 used to identify homologous enzymes from various organisms capable of exhibiting similar biocatalytic
24 functions with different substrate specificities, an approach effectively employed for identifying a PHA

1 synthase enzyme capable of acting on lactate for the synthesis of polylactic acid (PLA) [31]. Knowledge
2 of the catalytic active site or specific motifs important for catalytic activity and specificity can facilitate
3 narrowing these searches towards isolation of enzyme variants exhibiting differences in order to sample
4 alternative activities and specificities. Kim and coworkers [32*] used this strategy to identify new
5 fructosyl peptide oxidase enzymes that find applications in diagnostics for diabetes. Molecular docking
6 is yet another valuable *in silico* tool to predict promiscuity *a priori* by studying interactions of a given
7 enzyme active site with a range of different substrates of interest and estimating activities and
8 specificities when sufficient information about enzyme crystal structures and active sites is available
9 [18,33].

10 Indeed, identification of candidate enzymes for novel pathways relies on continued investigation
11 of enzyme crystal structures and identification of catalytic active sites as well as development and
12 validation of reliable bioinformatics approaches for utilizing this information for *a priori* prediction of
13 enzyme activity and specificity. Simultaneously, the information garnered from the experimental
14 exploration of enzymes in novel pathways can in turn offer useful insights into enzyme catalytic activity
15 and serve to validate and refine the developed computational tools. Further, such substrate promiscuity
16 also serves as an excellent starting point for engineering novel biocatalysts with higher activity and
17 specificity towards the non-native substrates.

18 **Enhancing pathway productivity through protein engineering**

19 Improving pathway productivity requires selective diversion of the precursor metabolites and
20 pathway intermediates away from natural metabolism or competing side reactions and specifically
21 towards the pathway of interest, without hampering cell growth and viability. Conventionally, this is
22 achieved by enhancing precursor metabolite pools by knocking out host genes encoding enzymes
23 catalyzing competing reactions and controlled overexpression of pathway enzymes to enhance flux

1 along the pathway and reduce metabolic burden [34*,35*]. Overexpression of pathway enzymes and
2 enhancement of precursor metabolite pools cannot, however, overcome inherent limitations associated
3 with activities and specificities of pathway enzymes. Protein engineering allows one to enhance these
4 properties by altering key amino acid residues and can effectively complement metabolic engineering
5 efforts for selective and efficient synthesis of the desired products. Leonard and Ajikumar and
6 coworkers demonstrated the implementation of such an integrated metabolic and protein engineering
7 approach for effectively enhancing titers of levopimaradiene (a valuable diterpenoid precursor to
8 pharmaceutical ginkgolides) over 2600-fold in *E. coli* by engineering the key rate limiting pathway
9 enzymes levopimaradiene synthase (LPS) and geranylgeranyl diphosphate synthase (GPPS) for increased
10 activity and specificity, in addition to amplifying fluxes towards the precursor metabolites isopentenyl
11 diphosphate (IPP) and dimethylallyl diphosphate (DMPP) [36**]. For evolution of GPPS exhibiting higher
12 activity towards geranylgeranyl diphosphate (GPP) synthesis from IPP and DMPP, random mutagenesis
13 using error prone PCR was used to create a library of GPPS mutants. Lycopene biosynthesis from GPP
14 was used as a colorimetric reporter for high throughput screening of this library for isolating an
15 improved GPPS variant. Yang and coworkers used directed evolution of the enzyme propionyl-CoA
16 transferase (Pct) from *Clostridium propionicum* to effectively activate lactate to lactyl-CoA to serve as a
17 substrate for microbial synthesis of the valuable polymer, polylactic acid (PLA) and other lactic acid co-
18 polymers using an engineered PHA synthase [31,37*].

19 Completely combinatorial strategies towards altering protein activity and specificity, such as
20 random mutagenesis or directed evolution do not require detailed enzyme crystal structure or active
21 site information, but are time consuming, laborious and often require several rounds of mutagenesis
22 and selection. They may also be impractical in the absence of high throughput screens or selection
23 strategies. While technologies such as MAGE [38] , TRMR [39] and SCALES [40] have thus far focused on
24 modifying regulatory regions to affect protein expression, they offer excellent practical solutions

1 towards accelerated directed evolution across multiple target loci simultaneously within a short period
2 of time. However, they still rely on high throughput screens to isolate improved mutant candidates. On
3 the other hand, site directed mutagenesis of select residues in the active site can limit the size of the
4 mutant library and obviates the need for a high throughput screen, but requires detailed information
5 about the active site and substrate-enzyme interaction to be derived from the enzyme crystal structure.
6 In the absence of such information for an enzyme of interest, constructing homology based crystal
7 structure models by threading residues of the enzyme of interest onto known crystal structures of
8 similar enzymes can allow identification of a putative active site. In the previously mentioned study on
9 levopimaradiene synthesis [36**], the authors used this approach to identify a putative active site for
10 the LPS enzyme and replaced individual residues in this active site with alternative residues from
11 paralogous LPS-type enzymes from other organisms, one at a time to sample alterations in activity and
12 specificity using the resulting distribution of levopimaradiene and by-products as a screen. Mutations in
13 residues that resulted in the most significant productivity enhancements were subjected to saturation
14 mutagenesis and were combined to engineer an LPS enzyme with 10-fold enhancement in
15 levopimaradiene productivity.

16 Protein engineering can also be employed for building on and improving naturally promiscuous
17 activities employed in novel pathways. Zhang and coworkers adopted this approach to enhance the
18 specificities of the enzymes LeuA and 2-ketoisovalerate decarboxylase (KIVD) toward the non-natural
19 substrates 2-keto-3-methylhexanoate and 2-keto-3-methylvalerate respectively by enlarging the
20 substrate binding pockets by replacing bulky residues with smaller ones to facilitate interaction,
21 resulting in 20-fold enhanced synthesis of the non-natural alcohol (*S*)-3-methyl-1-pentanol [23**]. Yang
22 and coworkers used site directed mutagenesis of key residues identified to confer specificity in 5
23 different broad substrate range PHA synthase enzymes to evolve higher specificity towards
24 incorporation of lactate [31]. These numerous examples underscore the immense potential of

1 engineered enzymes to enhance yields by reducing by-product formation and accelerating rate limiting
2 steps, and inherent challenges in engineering enzymes stemming from the limited available information
3 about enzyme crystal structures and catalytic mechanisms and lack of convenient high throughput
4 screens.

5 **Enhancing productivity through cofactor balancing**

6 Pathways for synthesis of value-added products often employ oxidation-reduction reactions
7 catalyzed by enzymes using the nicotine amide dinucleotide cofactors NAD(H) and NADP(H) that are as
8 critical for catalysis as the enzymes themselves. In the course of the biocatalytic transformations, these
9 cofactors are oxidized or reduced and need to be recycled back to their active forms via natural cell
10 metabolism or a subsequent pathway step for continued catalysis. In the absence of such recycling, the
11 pathway can come to a grinding halt. These cofactors also play a critical role in redox reactions in natural
12 cell metabolism where NAD^+ is often the cofactor of choice for oxidation of substrates in catabolic
13 reactions with NADPH serving as the reductant in biosynthetic reactions. Thus, pathway enzymes often
14 have to compete with natural cellular enzymes for these essential cofactors. Decreased active cofactor
15 availability due to competition with natural metabolism as well as rapid depletion of cofactor pools due
16 to an imbalance between pathway demands and natural cofactor recycling can pose serious limitations
17 on pathway productivity [41]. For example, NADH and NADPH are oxidized to NAD^+ and NADP^+ ,
18 respectively, during the synthesis of reduced fermentation products such as ethanol, lactate, butanol,
19 succinate, 3-hydroxybutyrate and polyhydroxybutyrate and their availability can influence both the
20 productivity and product profile of pathways [9,41,42].

21 Effective recycling of cofactors is hence essential and may be achieved by specific metabolic
22 engineering efforts targeting natural host pathways that can bring about such recycling or via enzyme
23 mediated cofactor regeneration using a suitable sacrificial substrate or transhydrogenases. For example,

1 Chelmer and coworkers employed a stoichiometric model to identify gene knockouts for improving the
2 NADPH availability for the biosynthesis of the natural flavanoid polyphenols (+)-catechins and
3 leucoanthocyanidins from dihydroquercetin, targeting natural *E. coli* metabolic pathways for effective
4 recycling of NADP to achieve a four-fold improvement in the yields [43]. The disruption of the
5 phosphoglucose isomerase gene and simultaneous over-expression of NAD⁺ kinase or soluble
6 transhydrogenases can be employed for diversion of glucose flux away from the Embden-Meyerhof-
7 Parnas (EMP) pathway towards the pentose phosphate pathway (PP) to enhance NADP recycling to
8 NADPH as demonstrated in the enhanced synthesis of thymidine [44]. This, however, comes at the cost
9 of loss of valuable carbon in the decarboxylation step in the PP pathway that can decrease pathway
10 yields. Transhydrogenases bring about the reversible transfer of reducing equivalents between NADH
11 and NADP and can be employed to replenish pools of NADPH or NADH (at the cost of the other) to
12 restore the cofactor balance, an approach effectively employed to enhance pathway productivity in
13 polyhydroxybutyrate synthesis [45]. Hwang and coworkers employed enzyme mediated cofactor
14 recycling using glucose dehydrogenase and glucose as a sacrificial molecule to regenerate NADPH
15 consumed in the simultaneous synthesis of the valuable fragrance compound 2-phenylethanol and the
16 non-natural amino acid L-homophenylalanine, enhancing the yields four-fold over the strains without
17 such cofactor regeneration [46]. Altering cofactor specificity via protein engineering is yet another
18 approach to achieve effective cofactor recycling [47,48]. These various examples again highlight the
19 importance of cofactor balancing for enhancing pathway productivity.

20 **Conclusions**

21 Design, engineering and optimization of novel pathways expands the array of value added products that
22 can be synthesized from biomass as alternatives to petroleum derived chemicals. Successful
23 development of MCFs is essential to cope with our dwindling fossil resources but poses a number of
24 challenges, including lack of well characterized enzymes, poor activity of selected pathway enzymes, low

1 product titers, poor yield and selectivity, metabolic burden and unfavorable cofactor balance. Advances
2 in the fields of metabolic engineering, biochemistry, protein engineering and synthetic and molecular
3 biology have produced experimental and *in silico* tools to address these challenges. In the absence of a
4 large number of well-characterized enzymes, exploiting natural enzyme promiscuity is critical for design
5 of novel pathways. Protein engineering can further improve on such promiscuous activity and enhance
6 specificity and activity in natural and novel pathways to improve productivity. Cofactor balancing is
7 essential and can further result in several fold improvement in titers. Indeed, for synthesis of value
8 added chemicals using MCFs to be economically feasible and to compete favorably with chemical
9 synthesis, these approaches need to effectively complement conventional metabolic engineering efforts
10 in the design and engineering of pathways.

11 **Acknowledgements**

12 This work was supported by the Synthetic Biology Engineering Research Center
13 (SynBERC) funded by the National Science Foundation (Grant Number EEC-0540879). We thank Kevin V.
14 Solomon for a critical reading of the manuscript.

15

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1 **Figures**

2

3 **Figure 1 Caption:**

4 **Design and Engineering of Pathways for Microbial Chemical Factories (MCFs):** The first step in
5 engineering novel or natural pathways for MCFs is to identify potential natural cell metabolites or
6 biomass derived feedstocks that can serve as starting materials and the series of biochemical reactions
7 required to convert these into the desired product. Martin and co-workers [3*] have reviewed some of
8 the computational tools available for identifying and selecting from the multiple possible pathways
9 connecting different starting materials to a product of interest. Once a pathway is selected, appropriate
10 natural enzymes expected to catalyze pathway reactions need to be selected using enzyme information
11 from various databases. *In silico* approaches such as protein blast searches and molecular docking may
12 help in such enzyme selection. Further pathway optimization to enhance product titers relies on an
13 integrated approach composed of 1) Metabolic Engineering to enhance precursor metabolite availability
14 using gene knockouts and enzyme expression level manipulation 2) Protein Engineering to enhance
15 pathway enzyme specificity and activity and 3) Cofactor Balancing via effective cofactor recycling.

16

Figure

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