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Introduction

Alternative design strategies to help build the enzymatic retrosynthesis toolbox

Shelby L. Hooe, 回 ab Gregory A. Ellis 回 and Igor L. Medintz 问 *a

Most of the complex molecules found in nature still cannot be synthesized by current organic chemistry methods. Given the number of enzymes that exist in nature and the incredible potential of directed evolution, the field of synthetic biology contains perhaps all the necessary building blocks to bring about the realization of applied enzymatic retrosynthesis. Current thinking anticipates that enzymatic retrosynthesis will be implemented using conventional cell-based synthetic biology approaches where requisite native, heterologous, designer, and evolved enzymes making up a given multi-enzyme pathway are hosted by chassis organisms to carry out designer synthesis. In this perspective, we suggest that such an effort should not be limited by solely exploiting living cells and enzyme evolution and describe some useful yet less intensive complementary approaches that may prove especially productive in this grand scheme. By decoupling reactions from the environment of a living cell, a significantly larger portion of potential synthetic chemical space becomes available for exploration; most of this area is currently unavailable to cell-based approaches due to toxicity issues. In contrast, in a cell-free reaction a variety of classical enzymatic approaches can be exploited to improve performance and explore and understand a given enzyme's substrate specificity and catalytic profile towards non-natural substrates. We expect these studies will reveal unique enzymatic capabilities that are not accessible in living cells.

The desire to both understand and control the synthesis of complex molecules began as early as 1828, when Friedrich Wöhler ushered in the era of modern organic chemistry via the first ever synthesis of urea.¹ However, it was not until the early 1900s that the elucidation of chemical structure and synthetic processes started to revolutionize the field of organic chemistry. One of the leaders in organic chemistry during this time (ca. 1910-1950) was Sir Robert Robinson who is remembered for his research on anthocyanins, alkaloids, and the famous Robinson annulation. Robinson was one of many early organic chemists who exploited a unique ability to recognize the relationships and patterns which existed between chemical structures and their reactivity. Advances by early organic chemists like Robinson paved the way for later chemists including R. B. Woodward, E. J. Corey, K. C. Nicolaou, and countless others to build on this mechanistic understanding and further grow the field of organic chemistry.² Seminally, Corey brought retrosynthetic organic chemistry into the mainstream for assembling complex molecules in the 1960s, and chemical synthesis itself has, in turn, contributed to expanding the number of proven retrosynthetic pathways.^{3,4}

Retrosynthesis provides an efficient tool for mitigating problems in planning the synthesis of complex organic molecules. It starts from the complex molecule to be synthesized and works in the reverse direction to sequentially break down the complex molecule into a series of elementary steps of substrate combinations that lead to the desired product. The 'Network of Organic Chemistry' now contains almost every reaction published in the literature and hosts 35 million chemicals typically allowing for derivation of multiple suggested pathways towards the synthesis of any new molecule.⁵

Although incredibly powerful and representing the current 'state-of-the-art', multistep organic synthesis typically relies on specific catalysts and requires a sequential and stepwise process (assemble reactants \rightarrow synthesis \rightarrow purify product from reactants and undesired byproducts \rightarrow determine yield/purity/enantiomeric excess (ee) \rightarrow repeat with next reaction step) that is characterized by diminishing yield as the number of steps and time needed increase, ubiquitous organic waste, sometimes nontrivial product isolation, along with relying on requisite expertise. In contrast, eons of evolution has allowed nature to evolve enzymes and improve their properties allowing them to efficiently catalyze complex multistep syntheses. Enzymes possess the ability to carry out a variety of multistep reactions inside the cell, in a one-pot manner if you will, achieving unparalleled regio- and stereospecificity within the cellular cytosol. Cells have numerous membrane-bound organelles where reactions take place, as well

^a Center for Bio/Molecular Science and Engineering Code 6900, U.S. Naval Research

Laboratory, Washington, DC 20375, USA. E-mail: igor.medintz@nrl.navy.mil

^b National Research Council, Washington, DC 20001, USA

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as molecular crowding that create localized regions of function within the cytosol. Further, there are certainly biochemical processes that begin at one location in a cell and end at another. Our use of the descriptive 'one-pot' for reaction site refers to the cell as a total synthetic entity here. Depending upon substrain type, it is estimated that an E. coli bacterial cell can carry out >1400 different enzymatic reactions with more than a plurality running concurrently.⁶ Moreover, most of the complex molecules made by nature still cannot be emulated by current organic chemistry.⁷ Given the vast number of enzymes that exist in nature, many of which are still waiting to be described but which will nevertheless contribute to expanding the repertoire of available reactions, and the incredible potential of directed evolution, the field of synthetic biology contains perhaps all the necessary building blocks to bring about the realization of applied enzymatic retrosynthesis.8 This would allow the power of enzymatic synthesis to be harnessed in a similar manner to its analogous chemical system while bringing with it remedies for many of the issues that plague organic synthesis. Current thinking anticipates that enzymatic retrosynthesis will be primarily implemented using conventional cell-based synthetic biology where requisite designer and evolved multi-enzyme pathways hosted by chassis organisms carry out an 'optimized' designer synthesis.8-11 In relying primarily on cell-based approaches, a significant amount of potential synthetic chemical space is precluded simply due to cellular toxicity issues.

In this perspective, we suggest that there are several other complimentary approaches that can not only contribute to building the enzymatic retrosynthesis toolbox, but which can also help expand its capabilities significantly beyond that of what cells can provide. Beyond enzyme evolution and redesign, we look to see if there are other ways to get Nature's currently available repertoire of enzymes to perform in a more efficient manner ex vivo and to catalyze reactions that they normally would not undertake in a cell. In conjunction with this, we highlight key areas of research and viable alternatives to cell-based systems that are needed to populate the enzyme retrosynthetic database such that it can incorporate substrates that would be toxic to cells along with, perhaps more importantly, those considered non-natural or xenobiotic. We begin by looking at the promise of cell-based synthetic biology along with its limitations as the latter are the key areas we wish to address and overcome.

Synthetic biology and enzymatic synthesis

Synthetic biology is being touted as the next industrial revolution due to its potential to address the many existing socioeconomic problems associated with the exponentially growing demands of an industrially and technologically-dependent planet.^{12–14} This potential is real and is certainly not under dispute here. For these purposes, the salient points are that the expected way forward will engineer cellular chasses (*e.g.* bacteria or yeast) to host designer enzymatic pathways to produce the desired molecules from precursors or even generic carbon–nitrogen sources in an efficient manner.⁸⁻¹¹ If a particular chemical reaction/step is not available in Nature, then it is hoped that new enzymes can be evolved to accomplish this step.¹⁵⁻¹⁷ Cellbased approaches come with many inherent benefits, which also serve to highlight the current interest and enticement in exploiting them, these include: cells replicate themselves; cells can be engineered to tolerate focused enzymatic pathways that produce a given molecule while removing competing pathways; recombinant DNA technology and molecular biology now provide a powerful toolbox for engineering cells and pathways; cells can make and continuously replenish all the enzymes they are directed to use; the confines of a cell can act as a highly concentrated synthetic vessel to facilitate efficient catalysis; and industrial fermenters can host thousands of gallons of an actively producing culture.¹⁸

So what is it that cell-based synthetic biology cannot do (at least currently)? Cells typically cannot tolerate the vast majority of unnatural or xenobiotic substrates, especially, if they inhibit a key pathway or adversely alter key cellular molecules or structures downstream. This point is exemplified by amino acid analogs, many of which are highly toxic to cells as they cannot be incorporated into nascent proteins and can poison the cell.^{19,20} This does not mean that enzymes in a given cascade cannot act on that substrate; in fact, the enzymes may be tolerant or promiscuous to the analog and the chemical transformations still potentially useful, just not in the context of a cell if they are to be exploited. Another key point to be appreciated is the interdependence amongst the vast number of enzymatic pathways present in a cell. Nature is nothing if not efficient and frugal, and so one enzymatic pathway can crossfeed into multiple products from a common intermediary rather than requiring multiple redundant pathways to make the same intermediary. For example, cellular glycolysis feeds into or is otherwise intertwined with amino acid, nucleic acid, and fatty acid synthesis along with many other metabolic processes. Unfortunately, along with these pathways competing with each other, which can be detrimental from a production standpoint, this also now provides multiple points to poison a cell. There may also not be a pathway for cellular internalization of a non-natural analog and similarly, no pathway for export before it builds up and becomes toxic. Moreover, if a natural version of an analog is required for growth, this can make for complex final product separation.

The current solution to some of these issues comes in the form of either major metabolic reengineering or cell-free biosynthesis, where the cellular cytosol and its contents are appropriated, or recombinantly constituted, to do the catalysis *in vitro*.²¹ Although clearly useful, this brings with it some of the same issues and a significant financial burden. For example, a single 25 μ L reaction from the PURExpress *In vitro* Protein Synthesis Kit manufactured by New England Biolabs has a current cost of ~ US\$25.^{22,23} In-between the boundaries of purely cell-based approaches *versus* purely cell-free methods for synthetic biology exist many different potentially viable variants where cell culture, cell lysates, and designer mixtures of enzymes and other components can still be used to carry out a set of reactions of interest.^{24–29} In cell lysates, the enzyme has been grown and harvested within the cell and subsequent lysis has been performed to release the enzyme from the cell. Therefore, the cell lysate represents a material containing an abundance of the cell-free enzymes of interest along with other cellular components. One undisputed advantage of this approach is the reduction in cost relative to reconstituted/artificial cell-free systems since the cell lysate requires fewer purification steps for production. However, a disadvantage of using cell lysates is that mixture complexity is increased relative to purified enzymes since lysis induces a release of all enzymes present in the cell, such as periplasmic nucleases and peptidases, which then remain in the cell extract potentially reacting with intermediates and/or substrates present.³⁰ A variety of possible hybrid systems exist for performing multi-step reactions such as adding purified enzymes to cell lysates, combining chemical catalysts with enzymes, supplementing cell-based systems with purified enzymes, etc.³¹⁻³⁵ If needed, there is no reason why the first part of a reaction sequence couldn't be done outside a cell with a toxic but cell impermeable substrate or intermediate. Further, two-pot reactions with a cell-free reactor and cell-based reactor can even be linked through chemical engineering methods. It must be appreciated that with each additional component or custom configuration there exists a consequent increase in mixture complexity and a series of benefits and liabilities that will need to be carefully considered.

The true power of synthetic biology will only be unleashed when it can access the chemical space now precluded from cells and move from 'natural' substrates and products to those that are not found in nature; in essence, when we can harness enzymes to make almost any molecule we desire in a designer fashion. This will be especially true if we can do this with the knowledge and power of unfettered retrosynthesis driving the field. So how do we get from these issues to a form of enzymatic retrosynthesis that can accommodate such non-natural chemistry and even build on it? We suggest several avenues of research that are needed, which cumulatively have strong potential for positive contribution here. Namely,

(i) Developing focused minimalistic multi-enzyme synthetic systems.

(ii) Undertaking substrate tolerance and specificity studies for major classes of enzymes.

(iii) Optimizing desired catalytic activity outside a cell.

(iv) Determining relevant thermodynamic-kinetic properties.

The concepts behind each of these ideas are briefly developed in the following sections. We note that many of these ideas are not unique to us and have been previously iterated in various different forms by others.^{8,30}

Minimalist multi-enzyme synthetic systems

Broadly speaking, the minimum number of components needed to implement a multi-enzyme biosynthetic cascade are the requisite enzymes, their cofactors, and substrates. The driving concept behind 'minimalist' here is to remove all manner of competitive pathways and the presence of any substrate inhibitors such that the only possible reaction is the one desired; this, of course, does not account for reversible or back-reactions. The number of molecules and their derivatives which are known to either be recalcitrant to diffusion through cell walls and/or induce cellular death are both abundant and diverse.³⁶ The net result of this is a vast synthetic chemical space, which simply cannot yet be explored within the constraints of cellular-based systems, and which make this concept of minimalism attractive within the field of synthetic biology.

Within cellular-based systems, an enzyme's promiscuity cannot surmount what the cell itself cannot tolerate. Outside a cell, an individual enzyme's promiscuity can and will increase overall flexibility in the design of biosynthetic pathways.³⁷ By isolating an enzyme which carries out a desired function from the cell, the ability to identify all the promiscuous attributes for a specific enzyme can now be realized (vide infra). In the construction of multistep one-pot enzymatic reactions, working outside the cell also greatly simplifies pathway design to synthesize a product of interest and this is especially true if all manner of competition is removed. The ability to combine enzymes isolated from different organisms to function in the same biosynthetic pathway also offers the potential to truly harness all the reactivity nature has to offer. This also obviates the need, for example, to back engineer eukaryotic enzymes for expression in bacterial systems.³⁸ Further, minimalist approaches allow for rapid detection and identification of bottlenecks and/or degradation pathways.³⁷ By simplifying the number of components within a system to include only what is required, the ability to identify sources to problems such as slow conversion, incomplete conversion, or off target activity/degradation within a biosynthetic pathway become much easier. A great deal of leeway is also available in how such reactions are implemented with easy adjustment to variables such as volume, concentration, temperature, and the like. Moreover, concerted efforts are currently underway to create multifunctional nanoscale scaffolds that stabilize the enzymes, control the placement, order, and stoichiometry of the enzymes, and even allow them to access phenomena such as channeling.39

The potential in this approach is perhaps epitomized by work from the Bowie and Lou Labs towards the synthesis of molecules such as monoterpenes and D-glucaric acid, respectively.40,41 Monoterpenes are a class of isoprenoids widely used within the pharmaceutical, cosmetic, and agricultural industries.42-44 However, within cells, monoterpenes are highly toxic and previous cell-based studies have found that specific types of monoterpenes can cause damage to the plasma membrane leading to decreased downstream ATP concentrations.45 This property has hindered the industrial application of cell-based terpene biosynthesis due to the low inhibitory concentrations that must be maintained in order to sustain cell viability.46 The Bowie Lab reported the design of a multi-enzyme cell-free system to produce monoterpenes from glucose at production yields which were over an order of magnitude greater than cellular toxicity limits.40 They combined 27 enzymes to process glucose into monoterpenes and by switching between specific terpene



Fig. 1 Multienzymatic pathway converting glucose into monoterpenes. Schematic of the synthetic enzyme system converting glucose to monoterpenes. Glycolytic and mevalonate enzymes are highlighted in blue and orange respectively. Auxiliary enzymes constituting the purge valve, (mGap and NoxE) and phosphate recycling by pyrophosphatase are highlighted in red, purple, and cyan. Different monoterpenes can be made by using alternative terpene synthases (green). Abbreviations: Hex – hexokinase, Pgi – glucose-6-phosphate isomerase, Pfk – phosphofructokinase, Fba – fructose-1,6bisphosphate aldolase, Tpi – triose phosphate isomerase, Gap – gald-3-P dehydrogenase, mGap – gald-3-P dehydrogenase, Pgk – phosphoglycerate kinase, dPgm – 2,3-bisphosphoglycerate-dependent phosphoglycerate mutase, Eno – phosphoenolpyruvate hydratase, PykF – pyruvate kinase, PDH – pyruvate dehydrogenase, PhaA – acetyl-CoA acetyltransferase, Hmgs – β -hydroxy β -methylglutaryl (HMG)-CoA synthase, Hmgr – HMG-CoA reductase, Mvk – mevalonate kinase, Pmvk – phosphomevalonate kinase, Mdc – mevalonate-PP decarboxylase, Idi – isopentenyl-PP isomerase, Fpps – farnesyl-PP synthase, NoxE – NADH oxidase, PPase – pyrophosphatase, Gald-3-P – glyceraldehyde 3-phosphate, DHAP – dihydroxyacetone phosphate, P – phosphate, PEP – 2-phosphophenolpyruvate, HMG – 3-hydroxy-3-methylglutaryl, PP – diphosphate. Schematic drawn from ref. 40.

synthase enzymes at the final step of the pathway, they could vary the product from limonene to pinene and sabinene; see the enzymatic pathway schematic in Fig. 1. This pathway utilized enzymes drawn from 13 different species/strains many of which

were optimized mutants. Later work demonstrated a cascaded system that produced the bioplastic polyhydroxybutyrate from glucose along with a 'purge valve' node for preventing the buildup of excess reduced nicotinamide adenine dinucleotide phosphate (NADPH) which would become inhibitory.30 Similarly, p-glucaric acid is a targeted high-value molecule derived from biomass due to its potential application in biodegradable detergents and polymers.⁴⁷ Several cell-based methods for its biosynthesis have been previously described in both yeast and E. coli. However, these strategies are limited in industrial fermentation methods due to low conversion rates, unbalanced pathway fluxes, and issues of cellular acid toxicity.48,49 Cell-free methods were similarly developed by Lou for the production of p-glucaric acid from sucrose whereby addition of a single NAD⁺ regenerative enzyme was able to significantly increase the overall efficiency of this pathway.⁴¹ Beyond the benefits and capabilities described here, such minimalist enzymatic systems also provide a potent format for implementing the next area of discussion.

Substrate tolerance and specificity studies

An enzyme's specificity refers to its ability to activate a specific substrate amongst a pool of structurally similar analogs, while enzyme promiscuity is an enzyme's ability to tolerate a substrate and catalyze a reaction with it, especially when it is not normally part of its functional repertoire.⁵⁰ In directed evolution, it has been shown that promiscuity simplifies the search for an enzyme that will act on a specific target.^{16,17} However, in the design of new biosynthetic pathways, past studies suggest that in multistep cascades the extent of enzyme promiscuity may be more inhibitory than helpful because, as reaction pathways become increasingly complex, so too does the requirement for specificity of the enzymes involved.^{16,17} Accordingly, previous studies have even focused on methods for increasing an enzyme's specificity for a target substrate.^{51,52} To date, this remains a problem for applications of directed evolution in multistep cascade design.8 In terms of reactivity profiles for different enzymes, the most common approach is to generate a substrate scope for a particular enzymatic pathway and provide the corresponding yields for each substrate.53,54 By determining this for major enzyme classes, and variants of a given enzyme - using panels of substrate analogs that include diverse unnatural or xenobiotic derivatives - enough information can be gleaned to be able to predict which substrate(s) a given enzyme will tolerate and which they will not. This is obviously part of the critical information needed to populate a retrosynthetic enzyme database. However, this only provides information on whether a specific reaction will proceed. More valuable, would be the additional understanding of how the initial rate or the total turnover number (TTN) of an enzyme changes across a series of systematically selected substrates. This information is relevant because it provides insight into the robustness of an enzyme and its overall lifetime, which aids in the selection of enzymes for multistep cascades.⁵⁵ Similarly, a thorough understanding of the factors which dictate the initial rate of an enzyme are needed in order to iteratively enhance the performance of an individual enzyme.⁵⁶ It is important to appreciate that the suite of commercial, and in many cases, automated, 'design, test, build, learn' tools developed for bioprospecting in synthetic biology can be brought to bear here and allow such studies to be undertaken in a massively-parallel and automated fashion.^{57,58} The complexity involved for developing a database to aid in enzyme selection methods can be appreciated from the work of Henrissat and co-workers in the development of the CAZy database which compiles carbohydrate-active enzymes into groups based on enzyme sequence and specificity.^{59–61}

Optimize desired catalytic activity outside a cell

While the removal of an enzyme from a cellular environment is often associated with a decrease in enzyme stability, it also provides the opportunity for simplified testing strategies to optimize its activity and analyze its ability to tolerate de novo substrates. By optimization, we stipulate to techniques that do not involve mutagenesis or evolution, which are already wellestablished ways to optimize enzymatic performance but which are based on different principles and considerable effort.⁶²⁻⁶⁴ For a single step enzymatic reaction within a cell-free system, the environmental conditions of the enzyme can be systematically varied such that the accessible operating conditions of the enzyme can be defined and optimized for a specific activity. Similarly, this same strategy can be applied to multistep enzymatic cascades. Here, the ability to optimize conditions whereby the rate of the slowest enzymatic step increases while conditions which are conducive to all enzymes in the system can be maintained is an invaluable tool to optimize the flux through a cascade.⁵⁵ One elegant example of this approach is the six-enzyme cascade developed for the synthesis of amorpha-4,11-diene, a precursor in the synthesis of the drug artemisinin used to treat malaria. This study demonstrated a methodology to test and analyze enzyme ratios, buffer conditions, pH, and ion choice, which increased the rate limiting step of a multienzymatic cascade such that the overall rate of product formation was also optimized.65

For our focus, the goal is to increase the efficiency with which an enzyme catalyzes the reaction of both native and non-native substrates. There are many proven ways to manipulate an enzyme's reaction environment such that some desirable functionality is increased. These include altering buffer composition, pH, ionic concentration, temperature, viscosity, pressure, adding organics and the like, along with adding other exogenous materials.^{11,30,66} See Table 1 for some representative examples. One of the more prominent successes in this regard is the use of lipases and esterases in reactions with different organic solvents added; this has allowed these enzymes to increase their substrate and prochiral selectivity and, in direct contrast to their expected hydrolytic activity, even catalyze synthetic reactions such as C–C and C– heteroatom bond formation along with Michael addition.^{67–69} Clearly, some enzymes are latently capable of doing one reaction Table 1 Examples of changes to enzyme activity and substrate tolerance obtained by altering select reaction conditions

Enzyme	Native activity	Alteration	Change	Utility	Ref.
Methyl-parathion hydrolase ^a	Organophosphate hydrolysis	Change divalent cation in catalytic site	100-fold increase in activity from Ca^{2+} to Ni^{2+}	Increased activity	70
Alcohol oxidase ^b	Convert short chain alcohols to aldehydes/ketones	High hydrostatic pressure	Stability to thermal deactivation	Increased lifetime and extended reaction conditions	55, 56, 71 and 72
Glucose oxidase	Glucose oxidation to H_2O_2 and D-glucono- δ -lactone	High hydrostatic pressure	Stability to thermal deactivation	Glucose sensing	73
Thymidine kinase ^c	Phosphorylate thymidine	Change temperature from 82° to $37 \ ^{\circ}C$	Increased promiscuity for non-natural substrate	Phosphorylate wide variety of nucleotide analog prodrugs	74
Bovine liver catalase	Breakdown reactive oxygen species	Increase temperature	Tolerated higher H ₂ O ₂ concentration	Utility in reactions with high H_2O_2 present	75
Lipases	Hydrolyze fatty acids	Increase organic solvent	C–C and C–heteroatom bond formation, Michael addition	Fat removal and industrial synthetic applications	69
Lipases	Hydrolyze fatty acids	Change solvent present	Increase in prochiral selectivity	Commercial transformations	76
Polygalacturonase	Hydrolyzes α -1,4 glycosidic bonds of pectic acid	Change temperature and pressure	Increased activity	Increased activity at lower temperatures	77
Alkaline <i>endo</i> -1,4-β- glucanase	Hydrolyze glucosidic bonds of cellulose	pH change from alkaline to slightly acidic (pH 6)	10-fold increase in hydro- lysis of cellohexaose and cellopentaitol	Optimized activity for additional substrates	78
Laccase ^d	Catalyze oxidation reactions by reducing O_2 to water	Addition of water miscible ionic liquid choline dihy- drogen phosphate	451% rate and 4.5 fold increase in laccase stability at room temperature	2,2'-Azino-bis(3- ethylbenzothiazoline-6- sulfonic acid) diammo- nium salt oxidation	79

in one context and a completely different type of chemistry in a different context.

Another method for optimizing enzyme activity is through various immobilization strategies.⁸⁰ As this is often associated



Fig. 2 Nanoparticle enhancement of enzyme activity. (A) Schematic of a CdSe/ZnS core/shell semiconductor quantum dot (QD) surface-functionalized with the DHLA-CL4 ligand to make it colloidally stable in buffer. Phosphotriesterase (PTE) is ratiometrically self-assembled to the QD surface by its terminal hexahistidine (His)₆ sequence. The average number of PTE per QD is controlled through the molar stoichiometry added during assembly. PTE hydrolysis of paraoxon substrate to *p*-nitrophenol product, which absorbs at 405 nm, is also shown schematically. Structure of the PTE competitive inhibitor triethyl phosphate. QD phosphotriesterase bioconjugate triethyl phosphate inhibition assays. Three-dimensional plots of PTE initial rates *versus* increasing paraoxon concentration in the presence of increasing triethyl phosphate inhibitor for (B) free enzyme and (C) 625 QD nm emitting (diameter 9.3 nm)–(PTE)₉ bioconjugates. Estimated K_{ir} , k_{catr} , and K_{M} values shown for each. Figure reproduced with permission from ref. 85 Copyright 2015 American Chemical Society.

with increased stability, it offers a potential solution to counteract the decreased stability of an enzyme once it is removed from a cellular matrix and increase its useful lifetime.81 Several immobilization strategies have been developed including enzyme display on metal nanoparticles, encapsulation within biological or polymeric materials, and chemical crosslinking.^{39,66,81-84} See Fig. 2 for an example of where nanoparticle display on a semiconductor quantum dot (QD) allowed the enzyme phosphotriesterase to function $2\times$ more efficiently (higher catalytic rate $-k_{cat}$) and further do this in the presence of increased concentrations of a competitive inhibitor, *i.e.*, $3 \times$ increase in K_{i} .⁸⁵ Interestingly, enzymatic enhancement in this context appears to be dependent upon NP size with smaller diameter materials somewhat counterintuitively displaying the largest enhancement.⁸⁶ In the specific context of multiple enzymes (two or more) working together in a concerted or coupled biocatalytical cascade, attachment to a NP surface may potentially offer access to other useful phenomena beyond just enzyme stabilization and/or enhancement, in the form of enzymatic channeling. Probabilistic substrate/intermediary channeling can dramatically improve the rates of catalytic flux in multienzyme systems by increasing the rate of intermediary transfer between proximal enzymes in a manner that competes

with what is more typically expected to be the much faster rate of product diffusion away from the enzyme.⁸⁵ Fig. 3 highlights some representative data from a recent example where the coupled activity between pyruvate kinase (PykA) and lactate dehydrogenase (LDH) were evaluated when assembled onto QDs.⁸⁷ In this example, the tetrameric enzymes were found to crosslink with the QDs into nanoaggregated structures which stabilized LDH and significantly increased the rate of coupled PykA-LDH activity by ca. 100-fold versus that of equivalent free enzyme (see Fig. 3C and D). Another example looked at the interaction of a 3-enzyme pathway when attached to the surface of a single but much larger gold NP.⁸⁸ Here, the enhanced catalytic flux attributable to channeling processes around the same NP was far more modest at \sim 3-fold. In a further tantalizing report that hinted at application of this phenomena to far more complex systems, NP addition also enhanced reaction rates in full transcription-translation cell-free reaction mixtures. 23,39,66,81-84 Beyond this, even DNA structures are being tested as multienzyme scaffolds due to their ability to directly control enzyme number, order, and sequentiality.39,89 We expect studies of these and related phenomena will help reveal the unique enzymatic capabilities that are not manifest in live cells especially if the substrate is non-natural. Again, such studies can be carried out



Fig. 3 Coupled enzymatic and channeling activity when assembled on a QD scaffold. (A) Reaction scheme for coupled PykA and LDH activity. Individual and coupled enzyme activity monitored by NADH consumption with loss of its absorption at 340 nm. (B) Schematic of the coupled PykA–LDH enzyme system colocalized on a QD surface. The propensity of the enzymes to form cross-linked QD dimers and, to a lesser extent, trimers *via* the enzyme's tetrameric polyhistidine tags located at each monomers distal end is also schematically indicated. Note, not to scale. (C) LDH tetramer stability monitored as its concentration is reduced when free in solution and as assembled to QDs. Specific activity of LDH (μ M NADH consumed s⁻¹ μ M LDH⁻¹) determined at various enzyme concentrations in the absence (black) or presence of 605 nm emitting QD (blue, diameter ~10.1 nm) or 525 nm emitting QDs (red, diameter 4.3 nm). Data fit to a dissociation equation. Note how LDH activity when assembled to the QDs significantly extends its activity profile below the 10 nM dissociation constant of the wildtype enzyme. (D) Coupled PykA–LDH enzymatic kinetics. NADH consumption in a combined PykA–LDH reaction monitored with enzymes assembled to 605 (red) or 525 QDs (green) and QD-free (black). The reactions were at a concentration of 8 LDH (2.4 nM)/4 PykA (1.2 nM)/1 QD (0.3 nM). Enzyme-only contained the equivalent amount of free enzyme. Figure reproduced with permission from ref. 87 Copyright 2018 American Chemical Society.

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in a parallel, parametric format to screen for the desired mechanistic information.

Determining relevant thermodynamickinetic properties

For any single step enzyme catalyzed reaction, the reaction's thermodynamic and kinetic factors control the rate of substrate conversion to product. For successful product formation, the overall difference in energy between the product and reactant under a given set of conditions must be negative (*i.e.*, $\Delta G_r' < 0$). Additionally, the energy between the transition state (TS) and the reactant (*i.e.*, ΔG^{\ddagger}) must be low enough such that the kinetic barrier for the rate-limiting step of the reaction can be overcome. The more the thermodynamic and kinetic free energy values decrease for the rate-limiting step of a reaction, the more facile or favored a given reaction becomes. Experimentally optimizing thermodynamic parameters to achieve optimal enzyme kinetics has previously shown itself to be a valuable tool for successful biosynthesis. For example, Aitken and Heck showed that the activity of a peroxidase enzyme was related to the homolytic O-H bond dissociation energy (BDE) by analyzing the activity of a peroxidase enzyme across a series of monosubstituted phenols.⁹⁰ They found phenols with highly electron-donating (hydroxy and amino) substituents led to inactivation of the enzyme with minimal substrate removal via a mechanism that was unrelated to how the enzyme catalyzed other phenolic substrates. Russell and co-workers identified a temperature dependence on the catalytic rate (k_{cat}) for a chymotrypsin enzyme both in its native state and when modified as a polymer based material.91 Increasing hydrostatic pressure has even been directly correlated to the activity of an alcohol oxidase enzyme.⁷² Lastly, the specific activity of a nitrite reductase was found to correlate with the redox potential of the mediator used in the biocatalytic conversion of nitrite to ammonia.92 In each of these examples, a thermodynamic parameter was varied and correlated with the observed effect on the kinetic activity of the enzyme. Driving reactions to completion by lowering the thermodynamic free energy $(\Delta G'_r)$ is most advantageous to the overall enzymatic reaction as long as the kinetic barrier for the reaction (ΔG^{\ddagger}) decreases, or remains unchanged. Therefore, correlating thermodynamic-kinetic parameters with one another is a simple approach to understanding how to systematically decrease the free energy requirements which dictate product formation. One such useful tool for the determination of thermodynamic values is the eQuilibrator program, which provides estimations of thermodynamic values for small molecules adjusted for experimental pH and ionic strength.93 In developing multienzyme cascades, the obvious requirement is that the difference in energy between product and starting material (i.e., initial reactant) again be negative (*i.e.*, $\Delta G_{\rm r}^{\prime} < 0$). Beyond that, the ability to understand how kinetic parameters for the rate-limiting step of a given enzyme correlate with thermodynamic changes can enable efficient determination of the preeminent enzyme, or homolog thereof, that will be most suitable within a given cascade.

Overall, it is our opinion that an understanding of relevant thermodynamic-kinetic parameters which dictate activity across a series of enzyme analogues could be an invaluable tool for iteratively optimized enzyme engineering and selection processes. Most simplistically, it would help to provide a framework for which analogs will be tolerated and which steps in a given enzymatic reaction pathway may or may not work along with what conditions are needed to make it move in the desired direction.

Outlook: towards a framework for assembling retrosynthetic enzymatic pathways

Enzymatic retrosynthetic platforms are starting to be developed with beta versions becoming available and issues with their initial accuracy (\leq 50%) has strengthened the call to populate them with as much data about each enzyme and their substrate tolerance as possible.^{94–96} Coming back full circle, contributions from the above ideas become more apparent in the framework of extending enzymatic retrosynthesis to focus on including nonnatural substrates and making non-natural or xenobiotic products. Similar to how chemical retrosynthesis is utilized conceptually, when a target molecule is desired, the enzymatic retrosynthetic platform will suggest the best combinations of enzymes and order of enzymatic steps needed to assemble that molecule. Within a minimalist reaction, the best combination of prokaryotic and eukaryotic-sourced enzymes can be jointly incorporated to carry out the desired chemistry without being challenged by competing pathways. The breadth of substrates that a given enzyme can tolerate will inform the suggested pathways and precursors to be used along with the intermediaries that can be accessed. Consideration of each enzyme's thermodynamic and kinetic properties in conjunction with optimization of reaction conditions, environment, enzyme immobilization, and the like can all be exploited to improve and drive the reactions forward and overcome any bottleneck or unfavorable step. See also Fig. 4, which illustrates this concept schematically to highlight the types of information that are potentially useful in this context and how they can contribute to enzymatic retrosynthesis and especially that of de novo products.

Of course, all of this is meant to happen within the same framework as that used for cell-based synthetic biology. They are not distinct from each other but are rather meant to be complementary. Our approach enriches and expands the attainable product space and provides choices and enzymatic chemistries currently not available within live cell systems. Some enzymatic reactions are just not available and here is where enzyme redesign and evolution will once again be invaluable. Additionally, not every step in a desired synthesis can be accessed enzymatically and so mixed chemo-enzymatic approaches will continue to be a mainstay as well.⁹⁷ Similarly, not every catalyzed multi-step reaction will be viable in a onepot minimalist cell-free biosynthetic approach and therefore flow-based systems will continue to be necessary to separate





Fig. 4 Schematic highlighting potential contributions from the studies suggested here towards augmenting retrosynthetic enzyme synthesis. The concepts behind each are outlined in the text along with how they can contribute to improving the data that will give rise to retrosynthetic enzymatic capabilities. E1 and E2 represent enzymes in an assembled pathway while the subscripts a-c or a-e represent homologs with different substrate and reactivity profiles.

multi-step reactions into viable steps that enable product formation.⁹⁸⁻¹⁰⁰ One-pot and minimalist are not meant to be all encompassing and immobile approaches, rather they can also be philosophical and conceptual if needed. If one or more steps or reactions can be simplified and minimized to make the overall process more efficient and specific, then the minimalist approach is still contributing to some extent. Advances in artificial intelligence (AI) and machine learning (ML) will clearly be critical to help both compile the exponentially growing library of data describing enzyme properties and characteristics along with mining that data for complex properties and relationships that are not overtly available at a first look to the human eye.¹⁰¹ So is all this easy to achieve? Of course not! A great deal of discovery, enzyme prospecting, experimental testing, metaanalysis, and years of hard work along with overcoming an untold number of problems still remain to see if even some of these concepts prove useful. We look forward to contributing here as well and to learning about other ways to apply enzymes in a similar manner.

Conflicts of interest

There are no conflicts of interest to declare.

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