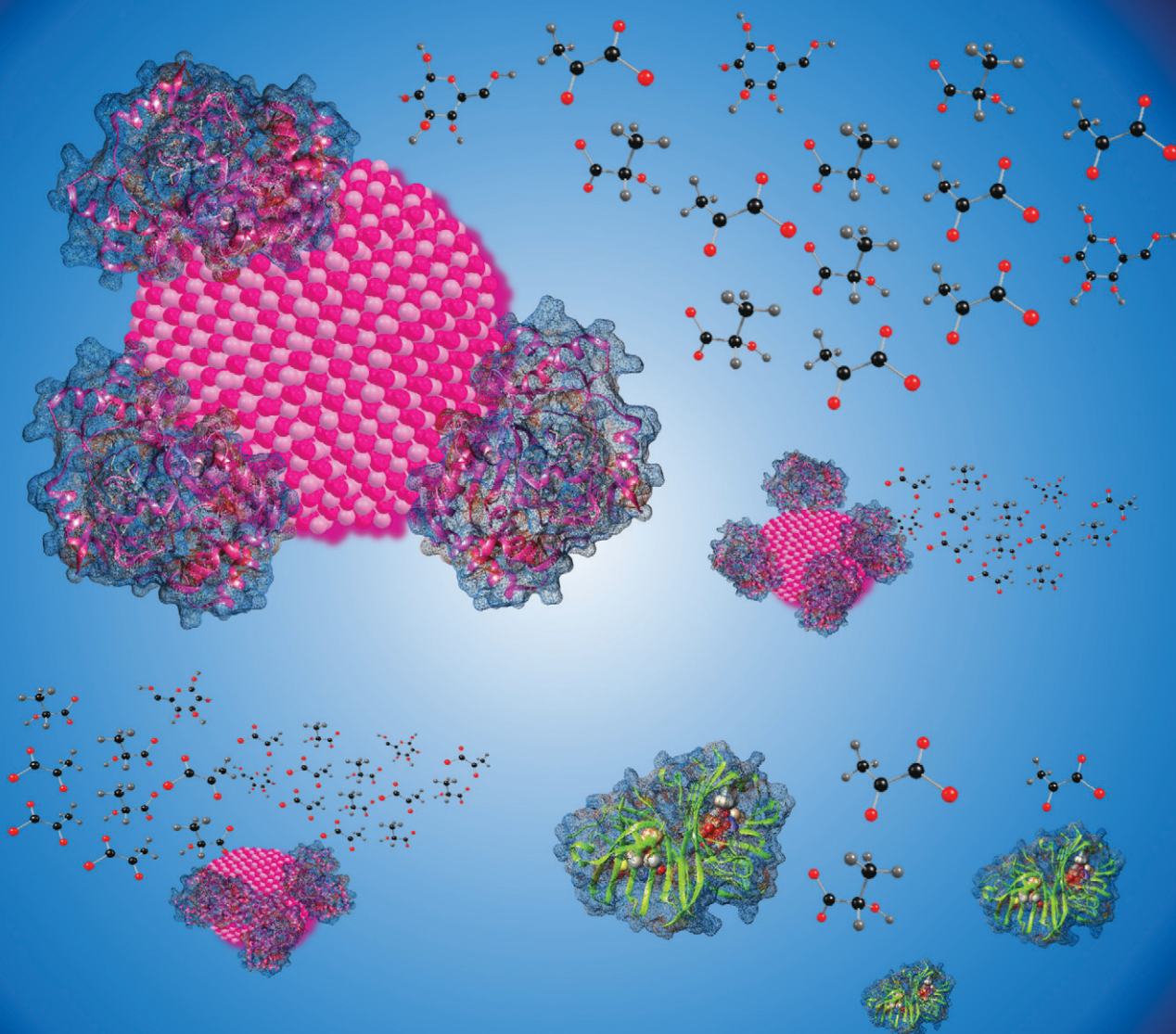


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REVIEW ARTICLE

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Enhancing enzymatic activity with nanoparticle display – an updated compendium and engineering outlook



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Enhancing enzymatic activity with nanoparticle display – an updated compendium and engineering outlook

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Almost all utilization of biocatalysis in the burgeoning field of synthetic biology requires not only enzymes but also that they function with peak efficiency, especially when paired with other enzymes in designer multistep cascades. This has driven concerted efforts into enhancing enzymatic performance by attaching them to macroscale scaffolding materials for display. Although providing for improved long-term stability, this attachment typically comes at the cost of decreased catalytic efficiency. However, an accumulating body of data has confirmed that attaching enzymes to various types of nanoparticle (NP) materials can often dramatically increase their catalytic efficiency. Many of the causative mechanisms that give rise to such enhancement remain mostly unknown but it is clear that the unique structured and interfacial environment that physically surrounds the NP material is a major contributor. In this review, we provide an updated and succinct overview of the current understanding and key factors that contribute to enzymatic enhancement by NP materials including the unique structured NP interfacial environment, NP surface chemistry and size, and the influence of bioconjugation chemistry along with enzyme mechanics. We then provide a detailed listing of examples where enzymes have displayed enhanced activity of some form when they are displayed on a NP as organized by material types such as semiconductor quantum dots, metallic NPs, DNA nanostructures, and other more non-specific and polymeric nanomaterials. This is followed by a description of what has been learned about enhancement from these examples. We conclude by discussing what more is needed for this phenomenon to be exploited and potentially translated in the design and engineering of far more complex molecular systems and downstream applications.

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Design, System, Application

Enzymes are constantly finding utility in biotechnology and especially within the exponentially growing field of synthetic biology. Across this vast application space it is many times desirable to enhance and improve the kinetic activity of the enzymes and especially in configurations when they are paired with other enzymes in designer multistep cascades. Although mutational selection and enzyme evolution are clearly important ways to address the need for catalytic improvements, they still require significant expertise and resources to be properly undertaken. However, a growing body of literature confirms that displaying enzymes on nanoparticles (NPs) can provide for significant kinetic improvements in their performance. In this review, we provide a state of the art compendium of enzymatic enhancement by NP attachment as a function of major NP material classes. We further describe what is currently known about this enhancement phenomenon with a focus that includes contributions from the unique structured interfacial environment that surrounds NPs, NP surface chemistry and size, and the influence of bioconjugation chemistry along with enzyme mechanics. We then look towards the future and discuss what more is needed for this phenomenon to be fully exploited and translated in the design and engineering of far more complex molecular systems and downstream applications.

Introduction

Enzymes are the key catalysts that speed up almost all biochemical reactions including many of those that would occur exceedingly slowly. This has made almost all attempts at biosynthesis without enzymes effectively a non-viable process. However, though a given enzyme may speed up a

catalytic transformation quite significantly by several orders of magnitude, it can be even more desirable to enhance the kinetic activity of that enzyme even further than what can be initially achieved or apparent within the context of several different applications. Such enhancement above an enzyme's native rate of activity allows for increased throughput and yield in an assay or biocatalytic industrial application, decreasing substrate requirements and improving output in research and diagnostic assays or pharmaceutical candidate/activity screening, along with matching kinetic rates between jointly-coupled enzymes in

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designer multienzyme cascaded reactions amongst other potential utility.^{1–5} There are several different approaches to achieving such enhancement, some of which have been applied independently or even jointly, and these include looking for more permissive or active enzyme homologs from other source species, mutational selection and/or evolution for optimized versions of the enzyme itself, and parametric adjustment of overall components and reaction conditions in a multienzyme cascade within a design of experimental framework.^{6–16} Interestingly, attaching enzymes to macroscale scaffolds such as surfaces, beads, and resins can help increase an enzyme's structural stability and, in turn, it's viable lifetime providing for long-term application along with potential reuse by allowing for the attached enzymes to be removed with the scaffolding and added to another reaction.^{17,18} However, the latter is not normally pursued to enhance enzyme activity such as the catalytic rate or k_{cat} , for example, primarily because chemical attachment to macroscale scaffolding materials is usually achieved with a concomitant decrease in that enzyme's kinetic properties.^{17–19} This is believed to arise as a result of the linkage chemistry decreasing the enzyme's overall freedom of movement, or by blocking or limiting access to its active site, along with the possibility of chemically modifying key residues needed for catalysis during attachment or association with the scaffold.^{20–22} Another aspect to appreciate is that, in many cases, enzyme structures are somewhat metastable, which allows them to better sample their substrates, and any structural impediment to this can be functionally deleterious.^{23–26}

Somewhat counterintuitively, a growing number of recent reports have confirmed that attaching or displaying enzymes on nanoparticles (NPs) or various other similarly-sized nanoscale scaffolding materials can increase enzymatic activity, sometimes to quite a significant extent, for example by increasing the k_{cat} apparent by 50-fold or more (*vide infra*). These nanomaterials can range from those that are primarily inorganic such as metallic NPs to those that are predominantly biological such as dense DNA constructs. We note that there exist almost no current review articles or other sources that bring together a detailed listing of which enzymes have been enhanced and what NP materials this has occurred with. Although an excellent conceptual reference, the main source material in this vein originates from 2012 and this reference does not include anything that occurred subsequently in the past 12 years.²⁷ The only other partial sources in this vein include ref. 28–30. Our focus in this review is to provide a brief discussion of some of the key factors associated with achieving NP enhancement of enzymatic activity based on what is understood about this phenomenon. See Fig. 1 for a schematic of many of the inter-related factors that are found at the NP–enzyme interface and which can influence enzymatic activity and especially kinetic enhancement. We then provide a state of the art compendium of examples where such an enhancement phenomenon has been observed across different nanoparticulate materials. We conclude with a discussion of how this phenomenon can be exploited and potentially translated during the design and engineering of more complex molecular systems. The latter can help drive unique

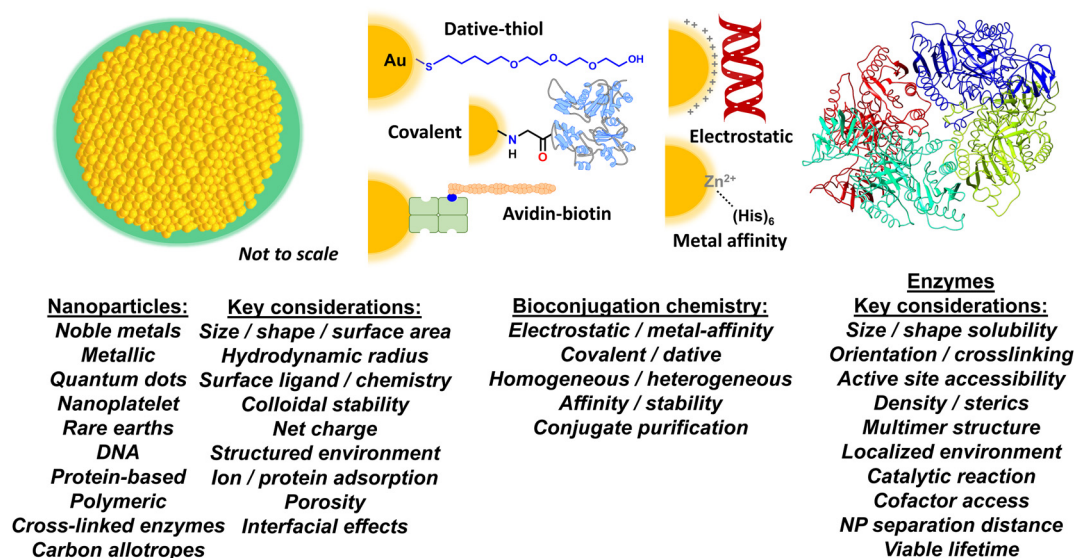


Fig. 1 Nanoparticle–enzyme bioconjugation and key considerations for accessing enhanced enzymatic activity. Schematic depicting a prototypical NP structure (left), different bioconjugation chemistries (middle), and an enzyme (right). The NP (gold) is surrounded by a layer of surface ligands (green), which provide it with colloidal stability. Bioconjugation chemistries shown include: a poly(ethylene glycol) or PEG attached to a gold NP surface via dative thiol–gold interactions; covalent amide bond formation between an aminolated NP and a carboxyl group on a protein; binding of a biotinylated peptide to an avidin modified NP; electrostatic assembly of a negatively charged DNA to a positively charged NP surface; and MAC between the (His)₆ motif and the surface ions of a Zn-overcoated quantum dot (QD). The listings underneath include some examples on NP material classes and bioconjugation chemistry approaches along with key considerations that can effect or are implicated in the generalized phenomenon of enzymatic acceleration when displayed on a NP surface.



enzymatic applications in synthetic biology and other related biotechnological fields.

Colloidal nanoparticles and their unique interface

The European Commission defines a nanomaterial as “a natural, incidental or manufactured material containing particles, in an unbound state or as an aggregate or as an agglomerate and for 50% or more of the particles in the number size distribution, one or more external dimensions is in the size range of 1 nm up to 100 nm”.³¹ There are, of course, many other definitions, depending upon the source and application.³² These definitions are generally extended to state that the material should not be exclusively naturally occurring; therefore a protein by itself would generally not be classified as a nanomaterial, however, an artificially engineered NP assembled from several naturally occurring proteins would. For these purposes, we define a NP as an engineered nanomaterial that has one of its dimensions <100 nm regardless of what it is constituted from. NPs can also be assembled from chemicals such as polymers or even constituted from carbon allotropes.³²

The current application under discussion, where NPs are interfaced with enzymes, requires that the NP materials be dispersible in aqueous media such as buffers. Since most NPs consisting of metals, semiconductors, and carbonaceous materials are synthesized within organic media or at high temperature, this typically means that the NPs must be further modified post-synthetically to display molecules on their surface, referred to as NP ‘surface ligands’ in the parlance of the field. These ligands impart colloidal stability to the NPs in aqueous media. Ligands can range chemically from small charged molecules to long poly(ethylene glycol) or PEG derivatives along with amphiphilic polymers and many other similarly functional materials such as polyelectrolytes (Fig. 1). The key concept to appreciate is that one part of the ligand interacts with the NP surface while another provides the NP with colloidal stability in the medium (*i.e.*, the particles remain evenly and stably distributed throughout the solution). Obviously, the size and chemical properties of these ligands will all influence the activity of an enzyme that is attached to it or placed in or around it. In contrast to this, biological-based NPs assembled from proteins or DNA, for example, typically have intrinsic colloidal stability in aqueous media given that their constituent materials are naturally soluble in water. The interested reader is referred to several excellent reviews on this subject for more information.^{33–38}

More pertinently, recent work has revealed that colloidal NPs universally structure their surrounding environment through the physicochemical influence of the dense ligand layer that provides the colloidal stability to the NPs.^{39,40} The characteristics of this structured environment remain mostly speculative as the requisite metrology to probe these nanoscale confines does not yet exist.^{41,42} However, it is believed that this structured environment includes pH, ionic, charge, and density gradients that may extend to twice the

NP diameter in some cases. This nanoscale structuring of the surrounding environment and the immediate boundary layer that it influences between the NP–enzyme conjugate and the bulk environment is believed to be a major contributing factor to the enzymatic enhancement described below.^{2,43,44}

Nanoparticles – selected methods for enzyme bioconjugation

How an enzyme is chemically attached to or otherwise displayed on a NP is obviously another major contributing factor to both accessing kinetic enhancement and the magnitude with which it is manifested (Fig. 1). From an idealized perspective, the chemistry used to bioconjugate an enzyme to a NP should allow for control over: i – the ratio of enzyme attached per NP; ii – the distance between the enzyme and NP; iii – the orientation of the enzyme on the NP; iv – the affinity of the enzyme attachment on the NP; v – provide for homogeneous attachment and a homogenous final bioconjugate structure; and, vi – all of these should be replicable with another unrelated NP and enzyme as desired.⁴⁵ The reality is that for most NP–protein bioconjugation reactions currently utilized, the majority of these properties are not achievable in any method let alone even jointly approach a plurality. The breadth of NP–bioconjugation chemistries along with their considerations has been extensively discussed in several focused review articles.^{46–55}

The benefits and liabilities of each bioconjugation chemistry should be carefully considered in terms of the composite material's final application and whether that choice of chemistry is appropriate or compatible for that purpose or not. For example, protein assembly to noble metal NPs and especially gold NPs (AuNPs) often relies on dative bonding between thiols and the gold surface.^{56–58} Many times, this requires that cysteine residues with their thiol side group be recombinantly introduced into the protein in a surface available position to facilitate the interaction. If the enzyme already has a cysteine or a cysteine-based dithiol in its structure, introduction of further cysteine(s) can result in thiol scrambling and loss of activity. Moreover, if a single thiol is used, it can have a strong rate of desorption from the gold surface, which can impinge on a need for extended bioconjugate stability; this can be somewhat alleviated by having multiple thiols participate in the chemistry.^{59,60} In contrast to dative thiol bonding, other bioconjugation chemistries rely on covalent chemistries with use of the common carboxyl–amine linkage by amide bond formation using EDC-NHS (1-ethyl-3-(3-dimethylaminopropyl) carbodiimide – *N*-hydroxysuccinimide ester) reactants being perhaps the most popular.⁵² To facilitate this type of linkage, the NPs often display either the requisite carboxyl or amine group on their surface ligands with the other cognate chemical functionality originating from the enzyme. However, the ubiquitous presence of these groups on most proteins usually results in heterogeneous enzyme display along with cross-linked NP–enzyme clusters blocking active sites. If these same groups are part of the enzyme's active site, this chemistry can

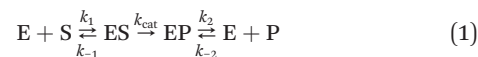


further inactivate the enzyme altogether. Additionally, the large concentrations of reactants used in these reactions lead to a need for extensive purification. Although ubiquitous and certainly powerful, biotin-avidin chemistry also comes with its own set of issues. Unless a labeling site is recombinantly introduced into a protein for site-specific biotinylation during expression,⁶¹ it is typically introduced into the protein using derivatives of the above heterogeneous covalent chemistry targeting amines, thiols or carboxyls. Most avidins are naturally tetravalent and attached to the NP *via* some type of covalent chemistry or electrostatically. This means that, in conjugation with heterogeneous biotinylation, the enzyme or other biomolecule will also be displayed in a variety of heterogeneous orientations that cannot be controlled and may even crosslink the NPs if it displays multiple biotin moieties.⁶²

Our work assembling enzymes to semiconductor QDs has mostly relied on metal affinity coordination (MAC) between (His)₆-motifs introduced at a protein's C- or N-termini and the Zn on a ZnS shell that overcoats the core/shell QD structure.⁶³ The (His)₆-motifs are usually introduced into the proteins recombinantly for subsequent purification using metal-affinity media and are thus already present on many expressed proteins.⁶⁴ Extensive characterization has shown that the imidazole side chains of (His)₆ will coordinate to the Zn on the QD surface with very high affinity ($K_d \sim 1$ nM), as it is available in the appropriate 2⁺ valence state similar to the transition metals (*e.g.* Ni, Co, Cr) commonly found as chelates in metal affinity media such as nitrilotriacetic acid (NTA).^{50,65} Even when access to the QD's ZnS shell may be precluded by a large surface solubilizing ligand, as long as the QD's surface displays multiple carboxyls, the same type of polyhistidine binding can still be accomplished in practice.^{66,67} Exposing these carboxylated QDs to small amounts of transition metals allows the ions to be coordinated by the carboxyls and functionally mimic the NTA group. More pertinently, attaching enzymes to QDs in this manner occurs almost spontaneously and provides for control over enzyme orientation on the QD and the average enzyme ratio per QD, which follows a Poisson distribution and displays an upper limit dictated purely by steric fitting considerations.⁶⁸ One potential issue here is that many enzymes are tetrameric as self-assembled from monomeric units and will thus display multiple (His)₆ at their multiple termini; this, in turn, will crosslink the QDs and enzymes into clusters. As discussed below, this can actually be very desirable when creating channeled multienzyme cascades with QDs.

Methods for quantifying enzyme kinetics and changes thereof

For appropriate context towards understanding the enhancement phenomenon, it is helpful to first consider enzyme activity within the formalism of the simplest-generalized version of the Michaelis-Menten (MM) model of enzymatic activity as given in eqn (1):⁶⁹



Here, the enzyme (E) and substrate (S) interact by a first order mechanism in a reaction where S is converted to product (P) after formation of an enzyme-substrate (ES) complex and where substrate concentration is made significantly higher than the Michaelis constant (K_M), which reflects the E's affinity for that substrate, *i.e.* $[S] \gg K_M$. For simplicity, these reaction conditions assume no reversible back-conversion of P to S. k_{cat} delineates the catalytic rate and is usually given in units of turnover per second or minute depending upon the observed rate. k_1 and k_{-1} are the rates of E-S association and dissociation, respectively, while k_2 and k_{-2} represent the respective rates of E and P release along with back formation into an EP complex, respectively. The latter is a common source of product inhibition when high concentrations of P are formed.⁶⁹ The mechanism and rate of EP release or k_2 play an important putative role in enzyme enhancement when attached to NPs (discussed more below). The methodology most commonly used to determine K_M and k_{cat} is to assay the initial reaction rate (or enzyme velocity V) as a function of substrate concentration in an experimental format that keeps the enzyme concentration fixed and very low relative to S, whereby $[E] \ll [S]$. This format is undertaken so as to meet Briggs-Haldane conditions and satisfy the initial steady-state reaction approximation.⁶⁹ The resulting data is then fit using the MM model and some derivation of eqn (2), which can also provide for estimation of the corresponding ES association and dissociation rates (k_1 and k_{-1}).

$$v = \frac{d[S]}{dt} = \frac{V[S]}{K_M + [S]} = \frac{k_{cat}[E]_0[S]}{k_1^{-1}(k_{-1} + k_{cat}) + [S]} \quad (2)$$

A second order rate constant can also be derived from the ratio of k_{cat}/K_M ; this is sometimes referred to as the specificity constant, and is also used as a metric for describing enzyme efficiency.⁶⁹ The data shown in Fig. 2E and F represents a sample of how enzyme kinetic data is commonly plotted. Obviously, more complex enzymatic mechanisms will necessitate far more complex models than this.

Some additional methods for quantifying enhancement in the observed activity of an enzyme include monitoring the overall amount of product formed under a given set of conditions and specific activity. Specific activity provides a metric for the activity of an enzyme per milligram of total protein and is usually given in units of $\mu\text{mol min}^{-1} \text{mg}^{-1}$. In its classical usage, specific activity provides a measurement or indirect reflection of enzyme purity in the mixture. Important to note is that in the current context where enzymes are displayed on NPs, a change in specific activity between free enzymes or as attached to NPs does not necessarily mean that it is now purer when attached to the NP in the strict sense of the definition, rather that it has become more active.



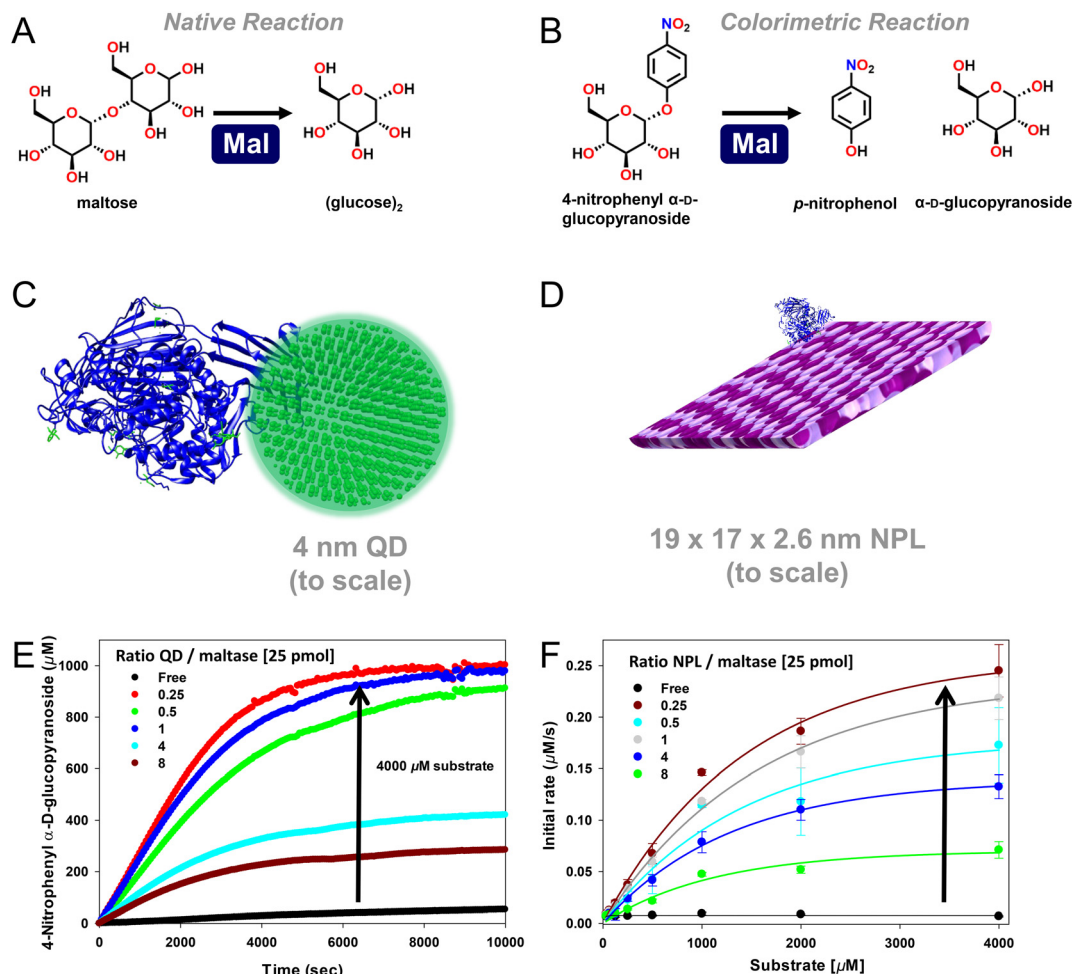


Fig. 2 Enhanced maltase activity when attached to semiconductor quantum dots (QDs) or nanoplatelets (NPLs). (A) Native maltase catalytic reaction and its activity on the (B) colorimetric 4-nitrophenyl α -D-glucopyranoside substrate, which is used to monitor activity in the assays shown. At scale schematic depiction of maltase as assembled to (C) \sim 4 nm diameter 520 nm emitting CdSe/CdS/ZnS core/shell/shell QDs²²⁴ and (D) *ca.* $19 \times 17 \times 2.6$ nm (LWH) 585 nm emitting CdSe/ZnS core/shell NPL materials. Both the QDs and the NPLs were surface functionalized with the zwitterionic CL4 ligand (see Fig. 3).¹⁴⁹ Protein structure coordinates obtained from the protein structure of human maltase-glucoamylase PDB # 3CTT, originally derived from ref. 225. (E) Enzymatic activity progress curves for a constant concentration of 25 picomoles of maltase as assembled at the indicated increasing ratios to 4 nm diameter QDs in the presence of 4000 μ M substrate. (F) Initial rate of 25 picomoles of maltase enzyme as assembled to the indicated increasing ratios of the NPL materials *versus* increasing concentrations of the colorimetric substrate. Maltase k_{cat} apparently increased \sim 50 \times and then \sim 125 \times when displayed on the QDs and NPLs, respectively, as indicated by the arrows. Maltase concentration held constant while QD/NPL concentration varied to achieve the indicated ratios. Data in panels E and F drawn from ref. 2.

Compendium of enzymatic enhancement examples

In this section, we provide a somewhat detailed, but certainly not exhaustive listing, of examples where enzyme activity has been enhanced when attached to some type of NP material. To qualify, the NP material has to have at least one of its dimensions within a size ≤ 100 nm. We further separate the data by NP material types including semiconductor QDs and related nanoplatelet (NPL) materials in Table 1, AuNPs in Table 2, metallic NPs in Table 3, other NP materials in Table 4, and DNA-based NPs in Table 5. The activity itself also has to have some reported quantifiable metric of enhancement. Some pertinent and/or relevant details are also provided on the NP surface and bioconjugation chemistries used for each example along with how the enzyme activity was assayed.

In Table 1, we list representative results collected for different individual enzymes displayed on the surface of semiconductor QDs. Notably, all the examples mentioned using QDs rely exclusively on the MAC of the enzyme's pendant (His)₆-motifs to the QD's ZnS shell for NP bioconjugation. The pendant coordination to one of the enzyme's end termini suggests that minimal to no perturbations were being contributed to these systems from heterogeneous enzyme assembly. The reported range of different QD sizes, which have been used to study the effect of immobilization on the enhancement of an individual enzyme's activity, is moderately narrow, ranging from \sim 4.0 up to 10 nm in QD diameter. Although QD emission is known for being size-tunable, the inherent chemistry that gives rise to this quantum confined property limits the size



Table 1 Enhanced enzymatic activity when displayed on semiconductor quantum dots and nanoplatelets

Material (size-diameter)	Enzyme	NP-attachment chemistry	Reaction	Results ^a	Ref.
CdSe/ZnS core-shell 520 nm QDs (~4.0 nm)	Horseradish peroxidase (HRP)	HRP-(His) ₆ - QD ZnS shell metal affinity coordination (MAC)	Oxidation of TMB	>2× <i>k</i> _{cat}	94
CdSe/ZnS core-shell 525 nm QDs (~4.2 nm)	Phosphotriesterase (PTE)	PTE-(His) ₆ - QD ZnS shell MAC	Paraoxon to <i>p</i> -nitrophenol	~4× initial rate; ~2× <i>k</i> _{cat} / <i>K</i> _M	72
	Benzaldehyde lyase (Bal)	Bal-(His) ₆ - QD ZnS shell MAC	Benzaldehyde + acetaldehyde to (<i>R</i>)-2-hydroxy-1-phenylpropan-1-one	~30% increase in <i>k</i> _{cat} > 3× <i>k</i> _{cat} / <i>K</i> _M	95
CdSe/ZnS core shell (~4/9 nm)	Engineered PTE trimer	PTE-(His) ₆ - QD ZnS shell MAC	Paraoxon to <i>p</i> -nitrophenol	~2× <i>k</i> _{cat} ; ~2× <i>k</i> _{cat} / <i>K</i> _M	96
CdSe/CdS/ZnS core/shell/shell 520 nm QDs (~4.0 nm) NPLs (LWH of ~19.2 × 17.3 × 2.6 nm)	Lactate dehydrogenase (LDH)	LDH-(His) ₆ - QD ZnS shell MAC	Pyruvate to lactate	>50× in turnover on QD/~10× in <i>k</i> _{cat} / <i>K</i> _M on QD and NPL	97, 2
	Maltase (Mlt)	Mlt-(His) ₆ - QD/NPL ZnS MAC	Maltose to glucose	~50× <i>k</i> _{cat} on QD ~125× <i>k</i> _{cat} on NPL	
	Glucokinase (Glk)	Glk-(His) ₆ - QD/NPL ZnS MAC	Glucose to glucose-6-phosphate	~12× <i>k</i> _{cat} on QD ~7× <i>k</i> _{cat} on NPL	
	Phosphofructokinase (PFK)	PFK-(His) ₆ - QD/NPL ZnS MAC	Fructose-6-phosphate to fructose-1,6-bisphosphate	~4× <i>k</i> _{cat} on NPL	
	Glyceraldehyde-3-phosphate dehydrogenase (GPD)	GPD-(His) ₆ - QD/NPL ZnS MAC	Glyceraldehyde-3-phosphate to 1,3-bisphosphoglycerate	~5× <i>k</i> _{cat} on QD ~3× <i>k</i> _{cat} on NPL	
	Phosphoglycerate kinase (PGK)	PGK-(His) ₆ - QD/NPL ZnS MAC	1,3-Bisphosphoglycerate to 3-phosphoglycerate	~3× <i>k</i> _{cat} on QD 2.3× <i>k</i> _{cat} on NPL	
	Phosphoglycerate mutase (PGM)	PGM-(His) ₆ - QD/NPL ZnS MAC	3-Phosphoglycerate to 2-phosphoglycerate	~6× <i>k</i> _{cat} on QD and ~4× <i>k</i> _{cat} on NPL	
CdSe/ZnS core-shell 523 nm QDs (~4.3 nm)	Glucose dehydrogenase (GDH)	GDH-(His) ₆ - QD ZnS shell MAC	D-Glucose to D-glucono-1,5-lactone	~5× <i>k</i> _{cat} / <i>K</i> _M	98
CdSe/ZnS core-shell 525 nm QDs (~4.2 nm)	Beta-galactosidase (β-gal)	β-Gal-(His) ₆ - QD ZnS shell MAC	2-Nitrophenyl β-D-galactopyranoside to 2-nitrophenol and galactose	~4× <i>k</i> _{cat}	70
	Alkaline phosphatase (AlkP)	AlkP-(His) ₆ - QD ZnS shell MAC	Conversion of 4-methylumbelliferyl phosphate to 4-methylumbelliferyl	25% increase in <i>k</i> _{cat}	99
CdSe/ZnS core-shell 630 nm QDs (size not reported)	Lysozyme (Lyz)	Lyz-(His) ₆ - QD ZnS shell MAC	Hydrolysis of glycosidic bonds	~2–2.5× activity	100
CdSe/ZnS core-shell 545 and 605 nm QDs (~5 and 10 nm)	Endoglucanase and exoglucanase	Enzyme (His) ₆ - QD ZnS shell MAC	Cellulose digestion	4.3–4.9× initial rate	101
CdSe QDs (20 nm)	Cellulase catalytic domain	Biotinylated-enzyme to streptavidin QDs	Cellulose digestion	>7× sugar yield	102

Notes: MAC = metal affinity coordination; NPLs = nanoplatelets (585 nm emitting CdSe/ZnS core/shell four monolayers CdSe); QDs = quantum dots; TMB = tetramethylbenzidine; LWH = length × width × height. ^a Reported relative to the freely diffusing enzyme (without immobilization).

of the final QD material to a relatively small range in contrast to other materials such as AuNPs, see below. Almost half of the examples (7 out of 15) in Table 1 originate from ref. 2, where QD-displayed enzymes were allowed to form self-assembled nanoclusters to engage in channeled activity. In that study it was important to profile the kinetic activity of each enzyme independently when displayed on the QD to better numerically simulate and match their relative activities during cascaded assays by controlling their ratios to each other when present in these clusters. Although not listed in the current table, this same reference also has data for two other QD sizes of 9.7 and 13.4 nm diameter, where the same 7 enzymes also displayed some enhancement but not of the same magnitude as when attached to the smallest ~4.0 nm diameter QDs.

Assembly of individual enzymes onto semiconductor QDs led to improvements in the enzyme's catalytic rate (*k*_{cat})

ranging from 2–50 times greater than what was obtained with the freely diffusing enzyme. Similarly, 2–10× improvements in enzyme efficiency (*k*_{cat}/*K*_M) were also reported. Of the 14 reported enzymes in Table 1, half have also been immobilized onto ~585 nm emitting CdSe/ZnS core/shell NPLs with an average *L* × *W* × *H* of ~19.2 × 17.3 × 2.6 nm. Utilizing these NPLs with their larger size and greater dimensionality instead of semiconductor QDs led to 2–125× improvements in *k*_{cat} and ~10× improvements in *k*_{cat}/*K*_M. It is worth highlighting that of all the different enzymes that have been immobilized onto either semiconductor QDs or NPLs, maltase (Mlt) showed the greatest improvement in activity when immobilized relative to the freely diffusing enzyme. As shown in Fig. 2, maltase was being studied for the conversion of maltose to glucose as part of a 10 enzyme joint saccharification and glycolytic cascade and displayed a remarkable 50× and 125× improvement in *k*_{cat} when



Table 2 Enhanced enzymatic activity when displayed on gold nanoparticles

Nanoparticle material (size-diameter)	Enzyme	NP-Attachment chemistry	Reaction	Results ^a	Ref.
AuNPs (20 nm)	Glucose oxidase (GOx)	Thiol bonding at the cysteine-rich side of GOx	Glucose conversion to gluconic acid and H ₂ O ₂	3× specific activity	103
AuNPs (3 nm)	Laccase	Hybrid immobilization <i>via</i> adsorption or amide coordination	Oxidation of ABTS	2× $k_{\text{cat}}/K_{\text{M}}$	104
AuNPs (50 nm)	Nitroreductase (NTR)	Amino groups of Cys-tagged NTR	Reduction of CB1954 prodrug	22% decrease in K_{M} 112% increase in k_{cat} 512% increase in $k_{\text{cat}}/K_{\text{M}}$	105
AuNPs (30 nm)	Pepsin	Enzyme immobilization <i>via</i> adsorption	Hydrolysis of peptide bonds	79% decrease K_{M} 118% increase in V_{Max} 110% increase in $k_{\text{cat}}/K_{\text{M}}$	106
AuNPs of varying size (1.5–100 nm)	PTE	PTE-(His) ₆ conjugation to Ni ²⁺ -NTA AuNP surface	Conversion of paraoxon to <i>p</i> -nitrophenol	3 to 10× k_{cat} 2× $k_{\text{cat}}/K_{\text{M}}$	107
HS-PEG ₇ -COOH functionalized AuNPs (10–30 nm)	Pepsin	EDC cross-linking	Hydrolysis of peptide bonds	73% decrease in K_{M} and 107% increase in $k_{\text{cat}}/K_{\text{M}}$	108
Polyelectrolyte functionalized AuNPs (36.4 nm)	Papain	Amide coupling between amino groups on papain and the COOH AuNPs (EDC/NHS)	Conversion of BAPNA to <i>p</i> -nitroaniline	59% decrease in K_{M} 4211% increase in k_{cat} 6667% increase in $k_{\text{cat}}/K_{\text{M}}$	71
Cysteamine surface functionalized AuNPs (25 nm)	Lipase	Carboxyl group of enzyme and AuNPs EDC/NHS cross-linking	Conversion of <i>p</i> -nitrophenyl palmitate to <i>p</i> -nitrophenol	41% decrease in K_{M} 181% increase in $k_{\text{cat}}/K_{\text{M}}$	109
Citrate modified AuNPs of varying size (5–20 nm)	PTE	PTE-(His) ₆ conjugation to Ni ²⁺ -NTA AuNP surface	Conversion of paraoxon to <i>p</i> -nitrophenol	~17× V_{max}	110
Citrate modified AuNPs (47 nm)	Rhamnulose-1-phosphate aldolase	Adsorption	Aldol addition	~4× initial rate	111
Au nanostars (37 and 83 nm Feret diameter)	Amylase	(His) ₆ conjugation to Ni ²⁺ -NTA AuNP surface	4-Nitrophenol colorimetric substrate	76% increase in k_{cat}	77
	Glucokinase		NADH <i>via</i> coupled enzyme assay	156% increase in k_{cat}	

Notes: ABTS = 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt; BAPNA = *N*-benzoyl-DL-arginine-4-nitroanilide hydrochloride; EDC = 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride; NHS = *N*-hydroxysuccinimide; NPs = nanoparticles; NTA = nitrilotriacetic acid. PTE = Phosphotriesterase. ^a Reported relative to the freely diffusing enzyme (without immobilization).

immobilized onto either QDs or NPLs, respectively.² This result directly speaks to the improvements in kinetic activity that can be potentially accessed by this phenomenon.² Studies with beta-galactosidase (β -gal) attachment to QDs revealed further and somewhat counterintuitive findings about the enhancement mechanism.⁷⁰ Although expressed as a monomer (~116.3 kDa) from a single gene during *E. coli* production, the functional form of this enzyme is an obligate tetramer of *ca.* 465 kDa displaying 4×(His)₆, one at each of the four monomers' termini, and has a length that is slightly greater than 10 nm. This large size means that, in practice, this enzyme is not displayed at a QD surface but rather displays up to 4 QDs around its surface. Here, selective display of QDs around β -gal's surface resulted in a further 3× enhancement in k_{cat} . Table 1, as well as the subsequent tables, also reveals a wide range of activity improvements

reported across a variety of different enzymes from different sources and manifesting a variety of different catalytic processes suggesting that such enhancement is not limited to only certain enzymes or catalytic mechanisms.

Table 2 outlines results reported for the assembly of individual enzymes onto AuNPs. Here, a wide range of differentially-sized AuNPs, ranging from around *ca.* 1.5 up to 100 nm in NP diameter, were utilized to study the effect of NP assembly on enzyme activity. Additionally, a variety of surface-modifications for the reported AuNP systems were explored including HS-PEG₇-COOH functionalized AuNPs, polyelectrolyte functionalized AuNPs, and cysteamine functionalized AuNPs with the rest assumed to be citrate modified AuNPs. The type of enzyme attachment strategy also varies across the AuNPs possessing these different surface functionalizations including, adsorption, thiol-bonding, MAC,



Table 3 Enhanced enzymatic activity when displayed on selected metallic nanoparticles

Material (size-diameter)	Enzyme	NP-Attachment chemistry	Reaction	Results ^a	Ref.
CLEA – amino-functionalized iron oxide NPs (50–100 nm)	Pectinase (containing xylanases and cellulases)	Cross-linking <i>via</i> glutaraldehyde addition	Hydrolysis of wheat straw	1.8× half-life	112
Iron oxide with polyacrylic acid – gallic acid (13.1 nm)	Laccase	EDC cross-linking	Oxidation of ABTS	4.4-Fold increase in initial rate	87
Cu(OH) ₂ nanocages (170 nm)	Laccase	Enzyme amines to APTMS-functionalized nanocage surface	Oxidation of 2,6-DMP or ABTS	14- to 18× k_{cat} 8- to 14× k_{cat}/K_M	89
Nano-sized Fe ₃ O ₄ modified with APTES (12 nm)	Lipase	Enzyme immobilization <i>via</i> the aldehyde activated magnetite	Hydrolysis of either <i>p</i> -nitrophenol butyrate or <i>p</i> -nitrophenol palmitate	9.2× <i>p</i> -nitrophenol butyrate 32× <i>p</i> -nitrophenol palmitate	113
Ni-NTA/H ₂ N-SiO ₂ @Fe ₃ O ₄ NPs (50 nm)	Bienzyme hydroxylase monooxygenase (HpaBC)	HpaB-(His) ₆ and HpaB-(His) ₆ to Ni-NTA	<i>ortho</i> -Hydroxylation of 4-hydroxyphenylacetate	2.6× increase in product formation	114
Cu ₃ (PO ₄) ₂ nanoflowers (3 μm/2 μm/5–8 μm) ^b (15–20 μm) ^b (20 μm) ^b (Size not reported) (20–40 nm) (100 nm)	Laccase	Immobilization <i>via</i> amide groups of enzyme and Cu ²⁺	Oxidation of epinephrine or ABTS	1.5× and 3.6× initial rate/6.5× product	115–117
	HRP		Oxidation of <i>o</i> -phenylenediamine	5.1× initial rate	118
	Lipase		Conversion of 4-nitrophenyl acetate	4.6× V_{max}	119
	Carbonic anhydrase		Conversion of paraoxon	2.9× product formation	120
Cu ₂ O NP (<350 nm)	Lipase		Lauric acid 1-dodecanol esterification	51× specific activity	84
	Hydroxylase		Conversion of 2,4-dichlorophenol to 3,5-dichlorocatechol	1.6× specific activity	121
Cu ₂ O NP (<350 nm)	Laccase	Immobilization <i>via</i> amide groups of enzyme and Cu ²⁺	Oxidation of syringaldazine	4.0× increase in product formation	122
	Organophosphorus hydrolase	Co ²⁺ binding to allosteric site of enzyme	Paraoxon conversion to <i>p</i> -nitrophenol	3× increase in product formation	123
Co ₃ (PO ₄) ₂ ·8H ₂ O (1 μm × 450 nm)	Organophosphorus hydrolase	Co ²⁺ binding to allosteric site of enzyme	Paraoxon conversion to <i>p</i> -nitrophenol	3× increase in product formation	123
Co ₃ (PO ₄) ₂ nanoflower (7 μm × 200 nm)	D-Psicose 3-epimerase	Enzyme binding to Co ²⁺	D-Fructose conversion to D-psicose	7.2× specific activity	124
Cu ₂ O nanowire mesocrystal (120 nm pore size, dia ~90 nm)	Laccase	Enzyme binding to Cu ²⁺	Oxidation of syringaldazine	10× specific activity	88
Fe ₃ O ₄ nanoring (70 nm outer diameter height 50 nm)	Beta-galactosidase	Immobilization <i>via</i> enzyme binding to Fe ²⁺	<i>o</i> -Nitrophenyl-β-galactoside conversion to <i>o</i> -nitrophenol	1.8× increase in product formation	125
Cu ₃ (PO ₄) ₂ ·3H ₂ O (30 nm)	Urease	Enzyme binding to Cu ²⁺	Ureas conversion of urea to CO ₂ NH ₃	40× specific activity	126
Membrane/Cu nanoflower (2 μm pore size, 4 μm diameter)	Laccase	Enzyme amide groups and Cu ²⁺	Oxidative coupling of phenol to 4-aminopyrine to form antipyrine dye	2.0-fold increase in product formation	127

Notes: ABTS = 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt; APTMS = (3-aminopropyl)-trimethoxysilane; APTES = 3-aminopropyltriethoxysilane; 2,6-DMP = 2,6-dimethoxy-phenol; EDC = 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride; CLEA = cross-linked enzyme aggregate; NPs = nanoparticles NTA = nitrilotriacetic acid. ^a Reported relative to the freely diffusing enzyme (without immobilization). ^b Full 3-D size, not NP size.

and covalent chemistry. Assembly of individual enzymes onto AuNPs led to improvements in the enzyme's k_{cat} parameter ranging from 300× improvement to a truly remarkable 4211% greater than what could be achieved with the freely diffusing enzyme. Similarly, 100–6667% improvements in the enzyme k_{cat}/K_M and 22–79% decrease in the enzyme K_M were reported across the different AuNPs. That the K_M values did not undergo similar changes in magnitude again suggests that some of these enhancements may arise from changes in the rate of EP release (*vide infra*). Notably, papain assembly onto

polyelectrolyte functionalized AuNPs displayed the largest improvement in overall enzyme activity compared to the freely diffusing enzyme. For example, the protease papain was studied for the conversion of *N*-benzoyl-DL-arginine-4-nitroanilide hydrochloride (BAPNA) to *p*-nitroaniline. When assembled onto polyelectrolyte functionalized AuNPs (36.4 nm in diameter), the papain enzyme displayed a 59% decrease in K_M , a 4211% increase in k_{cat} , and a 6667% increase in k_{cat}/K_M relative to the non-assembled enzyme. However, these results were attributed less to the AuNP aspect of the material and



Table 4 Enhanced enzymatic activity when displayed on other nanoparticles and nanoparticle-like materials

Material (size-diameter)	Enzyme	NP-Attachment chemistry	Reaction	Results ^a	Ref.
Graphene oxide (GO) – MgNP (GO size 0.5–5 μm)	α-Amylase (amy)	Amy-Mg affinity GO cross-linking <i>via</i> glutaraldehyde	Cleavage of 1,4-α-D-glycosidic bond in linear amylose and amylopectin	2.3×/4.3× (8 °C/90 °C) V_{max} ; 2.5× product formation (18 °C)	128
GO-GQDs (2.3 nm × 0.9 nm)	HRP	Covalent attachment	Oxidation of TMB	1.9× K_M	129
Magnetic Fe ₃ O ₄ NP CLEAs (30–50 nm)	Lipase	AOT-activated CLEA lipase immobilized to NP with APTES	Transesterification of 2-phenylethanol and vinyl acetate	20× increase in product formation	130
MWCNTs (size not reported)	Lipase	EDC/NHS cross-linking	Hydrolysis of <i>p</i> -nitrophenol palmitate	10× specific activity	131
Polydopamine coated AgNP (20 nm)	Trypsin	Enzyme immobilization <i>via</i> catechol groups on AgNPs	Hydrolysis of proteins	1.6× hydrolysis casein, 8–17% increase ovalbumin and BSA	132
Polycaprolactone (PCL) nanofibers (324 nm in diameter)	Lipase	Immobilization <i>via</i> PCL backbone	Hydrolysis/transesterification of <i>p</i> -nitrophenyl palmitate	14× specific activity	133
Carbon dot (3 nm)	Laccase	Carbon dot phosphate backbone	Oxidation of ABTS	1.9× specific activity	134
Cu ²⁺ -adsorbed pyrene-PAA/PPEGA (50 nm)	Laccase	Enzyme amide groups and Cu ²⁺	Oxidation of ABTS	4.5× specific activity 3× k_{cat}	135
SWCNT (diameter: 1–2 nm, length: 5–30 μm)	Laccase	Enzyme adsorption	Oxygen reduction	6× increase in electrocatalytic current	136
Mesoporous silica NP (6 nm)	Laccase	Enzyme adsorption	Oxidation of ABTS	1.2× half life	137
Carbon nanotube (diameter: 20–40 nm; length: 5–15 μm)	Lipase	Enzyme adsorption	Lauric acid 1-dodecanol esterification	68× specific activity	84
ZIF-8 MOF (300 nm with pore sizes of 5–20 nm)	Cytochrome c	PVPP-modified enzyme assembly to Zn ²⁺ of MOF	Amplex red conversion resorufin	10× specific activity	138
Pluronic polymer (30 nm)	Lipase Cytochrome c	Amphiphilic grafting	Hydrolysis of 4-nitrophenyl butyrate Conversion of 2,2'-azinobis-(2-ethylbenzthiazoline-6-sulfonate)	67× specific activity 670× specific activity	90
Fe ₃ O ₄ NP hydrogel (size not reported)	L-2-HAD _{ST} dehalogenase	Conjugation to acrylic acid <i>via</i> the protein's lysines	Dehalogenation of L-2-haloalkanoates to D-2-hydroxyalkanoates	2.0× k_{cat}/K_M	139
Siliceous mesocellular foam (mesoporous dia. ~36 nm)	Lipase	Enzyme adsorption	Tributyryl conversion to butyric acid	25× specific activity	140

Notes: ABTS = 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt; AOT = sodium bis-2-(ethylhexyl) sulfosuccinate; APTES = 3-aminopropyltriethoxysilane; BSA = bovine serum albumin; CLEA = cross-linked enzyme aggregate; EDC = 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride; MWCNT = multiwalled carbon nanotubes; NHS = *N*-hydroxysuccinimide; NPs = nanoparticles; PAA/PPEGA = block copolymer [poly(acrylic acid)/poly(poly(ethylene glycol) acrylate)]; PTE = phosphotriesterase; PVPP = polyvinyl pyrrolidone; SWCNT = single-walled carbon nanotube. TMB = tetramethylbenzidine. ^a Reported relative to the freely diffusing enzyme (without immobilization).

more to the charged polymer surface and high papain loading capacity achieved with the polymer chains of the polyelectrolyte surface.⁷¹ It may be that the high charge density of the polyelectrolyte surface also contributed in other ways such as providing for localized substrate sequestration. See also related discussion on DNA structures below.

The unique material chemistry available with AuNPs has also provided useful properties for these types of studies. The above-mentioned study where AuNPs ranging over almost 2 orders of magnitude in size (from 1.5 to ~100 nm diameter) were used to ascertain the effect of NP size on PTE catalytic enhancement would probably have not been possible with other individual NP types.⁷² This is because the unique

chemistry of AuNPs allows for this size range to be achieved, although synthesis of some of the larger sizes usually necessitates using smaller AuNP particulates as seeds for further growth. There are also a variety of other AuNP shapes synthetically available including triangles, nanostars, and nanoflowers for evaluation with enzymes.^{73–76} Díaz and coworkers assayed the activity of amylase, maltase, and glucokinase as assembled to both large and small gold nanostars as part of a study looking at their cascaded activity.⁷⁷ Due to their shape, these NPs are characterized by their Feret diameters meaning that they were measured along a specific plane that could include the largest end-to-end distance between the stars tips. Díaz found that both amylase



Table 5 Enhanced enzymatic activity when displayed on DNA nanostructures

DNA nanostructure (size or structure)	Enzyme	NP-Attachment chemistry	Reaction	Results ^a	Ref.
DNA triangle (120 nm length per side)	HRP	Covalent attachment to free lysine amine side chains	Oxidation of a series of phenol derivatives	>3× specific activity	141
DNA origami triangle (120 nm length per side)	β-Amylase	NTA-modified 22 bp-oligomer guide strand bound to enzyme's His ₆ in presence of Cu(II) ions	Hydrolysis of every second α-1,4 glycosidic linkage in starch	4× k_{cat}	44
	Maltase		Hydrolysis central α-1,4 glycosidic maltose bond yielding 2 glucose	3× k_{cat}/K_M >35× k_{cat}	
	Glucokinase		Phosphorylation of glucose to glucose-6-phosphate	2× k_{cat}/K_M >3× k_{cat} 11.5× k_{cat}/K_M	
3D octahedral DNA scaffold (12 six-double helix bundles and 120 staples)	Glucose oxidase	Sulfo-EMCS treated enzymes assembled directly to activated oligonucleotides	Conversion of glucose to gluconic acid	3.5× k_{cat}	142
DNA nanocage (~54 nm × 27 nm × 26 nm; inner cavity: 20 nm × 20 nm × 17 nm)	Glucose oxidase	Oligonucleotide-conjugated enzyme annealed directly to complementary DNA nanocage	Conversion of glucose to gluconic acid	5.4× k_{cat}	143
Rectangle (70 × 100 nm ²) on microbeads	HRP (R)-Selective alcohol dehydrogenase	Halotag based oligonucleotide binder protein.	Oxidation of ABTS Convert 5-nitrononae-2,8-dione 1 to (R)-syn/anti-hydroxyketones (60:40)	9.5× k_{cat} 2.2× k_{cat}	144
2D DNA triangle 25 nm	HRP	Covalent chemistry with bifunctional linker	Various substrates including <i>p</i> -aminophenol	100–300× activity	141
Square pyramidal DNA scaffold ~35–45 nm	Xylose reductase Xylitol dehydrogenase	Leucine zipper/Halo-tag enzyme fused to DNA binding proteins	NADH cofactor production or consumption	3–4× turnover frequency	145
48.5 kbp lambda phage DNA	β-Lactamase	Streptavidin-biotin chemistry	Chromogenic nitrocefin substrate	~2× k_{cat} 1.7× k_{cat}/K_M 3.8× activity	146

Notes: ABTS = 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt; bp = base pair; NTA = nitrilotriacetic acid; Sulfo-EMCS *N*-ε-maleimidocaproyl-oxysulfosuccinimide ester-. ^a Reported relative to the freely diffusing enzyme (without immobilization).

and glucokinase k_{cat} increased between 76% and 156%, respectively. In contrast to what was seen with QDs above, maltase activity actually decreased when displayed on these NPs by almost 50%. It is not readily apparent why this enzyme would now decrease in activity after increasing by almost 150× when attached to NPLs above.⁷⁷ This is a prime example of where differences among inherent properties such as NP size, shape, surface ligand/solubilization approach, and bioconjugation chemistry could all potentially contribute to this stark difference. We also note that considerable research has been undertaken using lasers to plasmonically heat AuNPs, which, in turn, can transduce the heat to the localized surroundings including an enzyme on its surface in a very rapid manner before the heat dissipates into the bulk.^{78–82} This may represent a possible tool for probing the narrow confines of the NP–enzyme interface by selectively activating or speeding up its activity in an effort to understand how enzymatic enhancement is manifest.

The data compiled in Table 3 summarizes results reported from the assembly of individual enzymes onto metallic NPs other than gold including those synthesized from iron (Fe), cobalt (Co), and copper (Cu). Unlike the AuNPs above, these NPs will exist mostly as a variety of complex oxides based upon how they were synthesized.⁸³ The NPs in this table include a variety of NP shapes each with its own unique dimensionality including nanocages, nanowires, nanorings,

and nanoflowers. They also vary widely in size ranging from 12 nm to 20 μm with the latter reflecting the overall dimensions of the materials and not their nanoscale components, which include those that are <100 nm in size. Enzyme assembly to this class of materials relies heavily on enzyme functionalization *via* the metal (M^{2+} ; M = Fe, Co, or Cu) center of the produced nanomaterial. Assembly of individual enzymes onto these metallic nanomaterials led to improvements in their specific activity with observed increases ranging from 1.6 up to 51× greater than what was obtained from the freely diffusing enzyme.

Similarly, a range of 2.6× up to 6.5× improvements in enzyme product formation and 1.5–5.1× increases in the enzyme initial rate were reported across the different metallic nanomaterials. Of these materials, the greatest enhancement was observed with a lipase enzyme assembled onto a $Cu_3(PO_4)_2$ nanoflower (20–40 nm in size). This lipase was studied for the esterification of lauric acid 1-dodecanol and when assembled onto a $Cu_3(PO_4)_2$ nanoflower demonstrated a 51× improvement in specific activity (*i.e.*, improved activity) relative to the freely diffusing enzyme.⁸⁴ Lipases are interesting enzymes, some of which are known to have hinges over their binding pockets and undergo enhancement or more correctly display a 'turn on' in activity at organic–inorganic interfaces and cell membranes.^{85,86} This is thought to help them better access substrates with partial solubility.



Still not known is how this mechanism would be affected by NP display, although the unique environment around the NP is also commonly considered an interface. We also note that data on the activity of the enzyme laccase as attached across different NP materials was reported. Here, it is seen that attachment to iron oxide NPs (13.1 nm dia.) resulted in a >4-fold increase in specific activity with the same caveat given here about interpreting this metric as above.⁸⁷ Similarly, 10× increases in specific activity are reported for attachment to Cu₂O nanowires (90 nm) while *ca.* 14–18× improvements in k_{cat} and $k_{\text{cat}}/K_{\text{M}}$ are achieved for laccase display on Cu(OH)₂ nanocages (170 nm).^{88,89} Along with some other examples of increased laccase activity on other NP materials given in Table 3, the similarity of the activity increases with their magnitude being almost all within 10-fold of each other suggests that a common enhancement mechanism is responsible for what is observed with this particular enzyme.

Table 4 summarizes results reported for the assembly of several different enzymes onto other NP and NP-like materials that do not fall into the same scaffolding material categories as the systems presented previously. This listing comprises a variety of different materials, including metal organic frameworks (MOFs), hydrogels, polymer materials, cross-linked enzyme aggregates (CLEAs), silica-based materials, and carbon/graphene or carbon allotrope-derived materials. The NP materials vary in overall size ranging from ~3 nm to 5 μm with the caveat that the latter have some dimensional property in a size range of <100 nm. Additionally, there is a significant amount of variation in the enzyme to NP assembly strategies utilized for these materials including covalent attachment, adsorption, grafting, and direct chemical enzyme cross-linking. The latter are again generally undertaken with heterogeneous conjugation chemistries leading to functionally-mixed and heterogeneous ensemble materials. NP–enzyme assembly in this case led to improvements in the enzyme's product formation rates ranging from 2.5 up to 20× and increases in specific activity ranging from 1.9 up to 670× greater than what could be achieved with the freely diffusing enzyme. Of these materials, the greatest enhancement was observed with cytochrome c as assembled onto a pluronic polymer material (30 nm in size). The pluronic polymer was composed of a central hydrophobic polypropylene oxide block attached to two hydrophilic poly(ethylene) oxide side blocks. This enzyme was assayed with the conversion of 2,2'-azinobis-(2-ethylbenzthiazoline-6-sulfonate) and when conjugated to the pluronic polymer material demonstrated an impressive 670× improvement in specific activity relative to the freely diffusing enzyme.⁹⁰ As above, the increase in specific activity probably does not arise from an improvement in purity per the strict definition, but rather should be interpreted as an increase in apparent activity when attached to the NP material.

It should be noted that CLEAs are a popular approach to exploiting the activity of multiple linked enzymes in the context of a multienzyme cascade within both research and industrial applications.^{91,92} They are also many-times formed

using heterogeneous covalent crosslinking chemistries with reactive agents such as glutaraldehyde.⁹³ This can and will lead to an exacerbation of the aforementioned covalent chemistry issues that affect the catalytic activity of a given enzyme across the resulting ensemble CLEA material. More pertinently, the CLEAs that are formed in this way are typically evaluated in the context of the intended final application, *i.e.* cascaded activity by a linked multienzyme assembly, along with product formation and not in terms of changes in individual enzyme rates. This makes collecting requisite kinetic data for each enzyme's individual performance in this configuration rather challenging.

Lastly, Table 5 summarizes results reported for the assembly of individual enzymes onto different DNA-based nanostructures including DNA origami triangles, octahedra, rectangles, and nanocages. These materials vary minimally in size, relative to some of the other materials, ranging from ~20–120 nm in length for any side of the DNA nanostructure. Given that each of these nanostructures is assembled from multiple smaller DNA strands that are hybridized together into the final architecture, the definition for having at least one dimension <100 nm is not as rigidly constraining. The enzyme assembly strategies utilized for these DNA-based nanostructures include direct annealing of enzyme-labeled DNA to complementary DNA, (His)₆-MAC, direct attachment by DNA binding proteins when fused to the enzyme, and other covalent attachment methods. Aside from horse radish peroxidase (HRP), which had much higher enhancement activity, assembly of individual enzymes onto DNA nanostructures led to improvements in enzyme k_{cat} ranging from 3–35× greater than what could be achieved with the freely diffusing enzyme and 2–11.5× improvements in the enzyme $k_{\text{cat}}/K_{\text{M}}$. Of these materials, the greatest enhancement was observed with the maltase enzyme assembled onto a DNA origami triangle (120 nm in length per side). This maltase enzyme was implemented for the hydrolysis of a central α-1,4 glycosidic bond in maltose and related maltosidic sugars derived from amylose as part of a saccharification pathway yielding two glucose molecules when assembled onto the DNA triangle. In this context, maltase demonstrated a >35× improvement in k_{cat} relative to the freely diffusing enzyme.⁵ Notably, this same maltase enzyme showed the greatest overall enhancement in activity relative to other reported systems for both QD and NPL-based scaffolding materials (Table 1).

Towards elucidating the mechanism of enzyme enhancement by nanoparticles

As originally suggested by Ansari and others, enzymes commonly display enhanced activity when they are displayed on a NP surface.^{27–29,147} Reports of this phenomenon have been steadily accumulating over the last 10-plus years and indeed many of the examples in this time period are listed in the tables included and further expounded upon in the corresponding discussion. A prototypical example of how such enhancement would



manifest is provided in Fig. 2. Here, the enzyme maltase originating from *Schizosaccharomyces pombe* as expressed in laboratory *E. coli* was attached to both semiconductor QDs and nanoplatelets (NPLs) by metal affinity *via* its terminal (His)₆ motif. NPLs are a new form of quasi 2-dimensional materials synthesized from the same constituents as QDs but having far larger length × width dimensions with a very shallow height.¹⁴⁸ When the 74.5 kDa monomeric maltase protein was attached to ~4 nm diameter QDs at a ratio of <1 per QD, k_{cat} increased *ca.* 50×. When assembled to 19 × 17 × 2.6 nm (LWH) 585 nm emitting CdSe/ZnS core/shell NPL materials at the same ratio, maltase manifests a ~125-fold increase in k_{cat} .² Fig. 2B and C show at scale schematics of maltase as attached to these two different nanomaterials. When considered at face value, these depictions provide strong support for the unique NP environment serving to enhance maltase activity as the NPL clearly has far more of such an environment surrounding it and the relatively smaller maltase enzyme. Steric considerations alone suggest that a range of 4–8 and 21–31 maltase could fit around these two differentially shaped/sized QD and NPL materials, respectively.²

Studies utilizing the enzyme phosphotriesterase (PTE), which originates from *Brevundimonas diminuta*, as assembled to QDs and AuNPs have yielded perhaps the most insight into the underlying mechanisms that give rise to NP-enzymatic enhancement.^{72,107} This 341 residue protein forms an obligate dimer and was expressed with a C-terminal (His)₆ motif for assembly to the ZnS surface of

QDs again by MAC. Since the C-termini in the dimer are in very close proximity to each other, it is assumed that this enzyme does not crosslink QDs when assembled to them as they both should bind to the same NP. Fig. 3A schematically depicts the components of this system including the enzymatic reaction and the structure of the paraoxon substrate, zwitterionic DHLA-CL4 ligand used to make the QDs colloiddally stable, and a triethyl phosphate competitive inhibitor.^{72,149} Fig. 3A compares the initial rates of PTE activity when consuming paraoxon as the PTE concentration incrementally increases and it remains freely diffusing or as assembled to a fixed concentration of 625 nm emitting CdSe/ZnS core/shell QDs with a corresponding increase in ratios per QD. PTE k_{cat} more than doubled from ~45 s⁻¹ to >110 s⁻¹ when displayed on the QDs at ratios of 6–8 per QD. The upper assembly limit of PTE to these 9.2 ± 0.8 nm diameter QDs was estimated at 28 confirming that the ratios utilized in the assays did not approach a steric limit with all enzymes present assumed to be attached to the QDs. Interestingly, PTE k_{cat} increased by 4-fold when displayed on smaller 525 nm emitting QDs (diameter ~4.2 ± 0.5 nm). PTE K_{M} was concomitantly decreased (meaning a lowering of affinity with an increased value of K_{M}) as k_{cat} increased in these examples. This is not unexpected because k_{cat} and K_{M} are usually mechanistically-linked with one changing in the converse direction of the other unless the enzyme undergoes very complex changes in its mechanism of activity.⁶⁹ One of the utilities or potential benefits of such enzymatic enhancement that still remains mostly

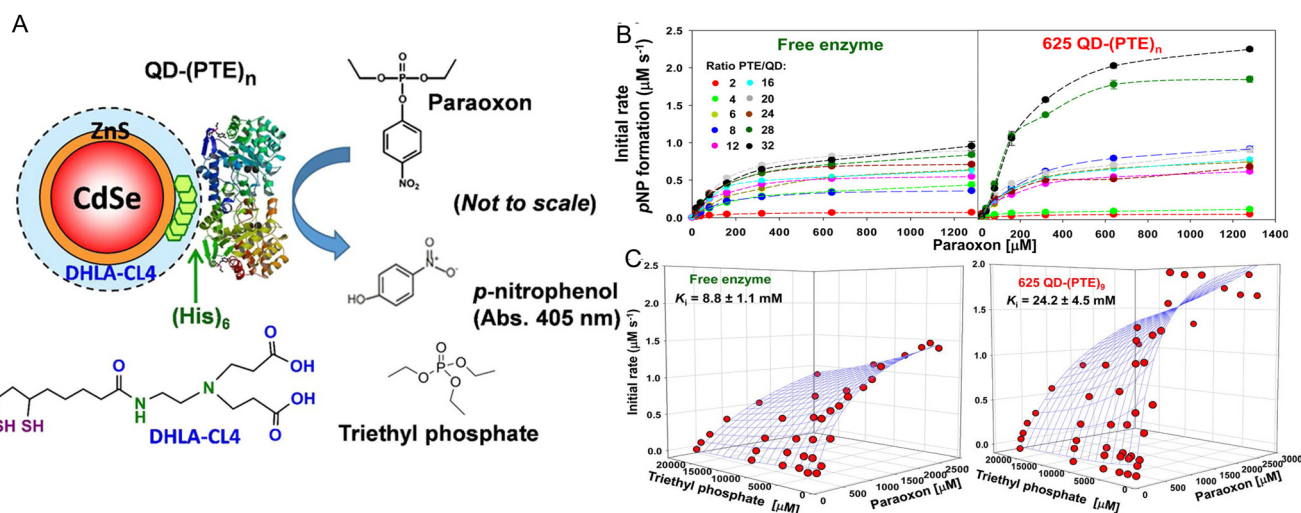


Fig. 3 QD phosphotriesterase bioconjugate and paraoxon hydrolysis. (A) Schematic of a CdSe/ZnS core/shell QD surface-functionalized with the DHLA-CL4 ligand.¹⁴⁹ PTE is ratiometrically self-assembled to the QD surface by its terminal hexahistidine (His)₆ sequence. The average number or valency of PTE per QD is controlled through the molar stoichiometry added during assembly and the conjugates are directly utilized without subsequent purification. PTE hydrolysis of paraoxon substrate to *p*-nitrophenol product, which absorbs at 405 nm, is used to monitor enzyme activity. Structure of the PTE competitive inhibitor triethyl phosphate. Note, not to scale. (B) Initial rates of *p*-nitrophenol product formation for (left) free PTE enzyme and (right) 625 QD-(PTE)_{*n*} bioconjugates assembled at the indicated ratios of *n* when exposed to an increasing concentration of paraoxon substrate. (C) 3D plots of PTE initial rates *versus* increasing paraoxon concentration in the presence of increasing triethyl phosphate inhibitor for (left) free enzyme and 625 QD-(PTE)₉ (right); estimated K_{i} values are included with each. Reproduced with permission from ref. 72 Copyright 2015 American Chemical Society.



unexplored is highlighted in Fig. 3C where it was revealed that PTE display on QDs allows it to remain far more active with a much higher concentration of triethyl phosphate competitive inhibitor present than the free enzyme. Indeed the K_i value tripled from *ca.* 8.8 up to 24.2 mM when the enzyme was attached to the surface of the QDs.

Fig. 4 presents representative data providing insight into the mechanisms that underlie PTE enhancement on QDs. Fig. 4A shows results from assaying PTE both when freely diffusing and as assembled to the above 525 and 625 nm emitting QDs as the temperature is raised in the form of an Arrhenius plot.⁷² Thermally activated changes are seen in all cases with an activation energy of 37.9 kJ mol^{-1} estimated for the free enzyme. That the QD-displayed enzymes show the same slope over temperature change as the free enzyme confirms that their activation energies are the same. Thus, the enzymatic enhancement from QD attachment does not arise from a change or especially lowering of the enzyme's energy barrier, at least for PTE. The possibility of lowering an

enzyme's energy of activation by NP display should not, however, be discounted. For example, Kouassi and colleagues reported that attachment of cholesterol oxidase to 10–56 nm diameter magnetic Fe_3O_4 NPs lowered the enzyme's activation energy by >30% from 13.6 to 9.3 kJ mol^{-1} .¹⁵⁰ Fig. 4B shows results collected in a similar format of assaying free PTE enzyme activity *versus* that of when QD displayed in buffer with increasing concentration of sucrose present. Raushel previously used this same format to show that PTE experienced decreased turnover as the microviscosity of the localized environment increased with increasing sucrose presence in the buffer and concluded that this decreased activity arose from a decrease in k_2 , the rate of E-P release, which was postulated to be the enzyme's rate limiting step similar to that of many other enzymes.^{69,151,152}

In essence, as the viscosity increases it makes it harder for the enzyme to release its product. However, when PTE is assembled to QDs, the opposite effect is seen strongly suggesting that QD display somehow alleviates this rate-

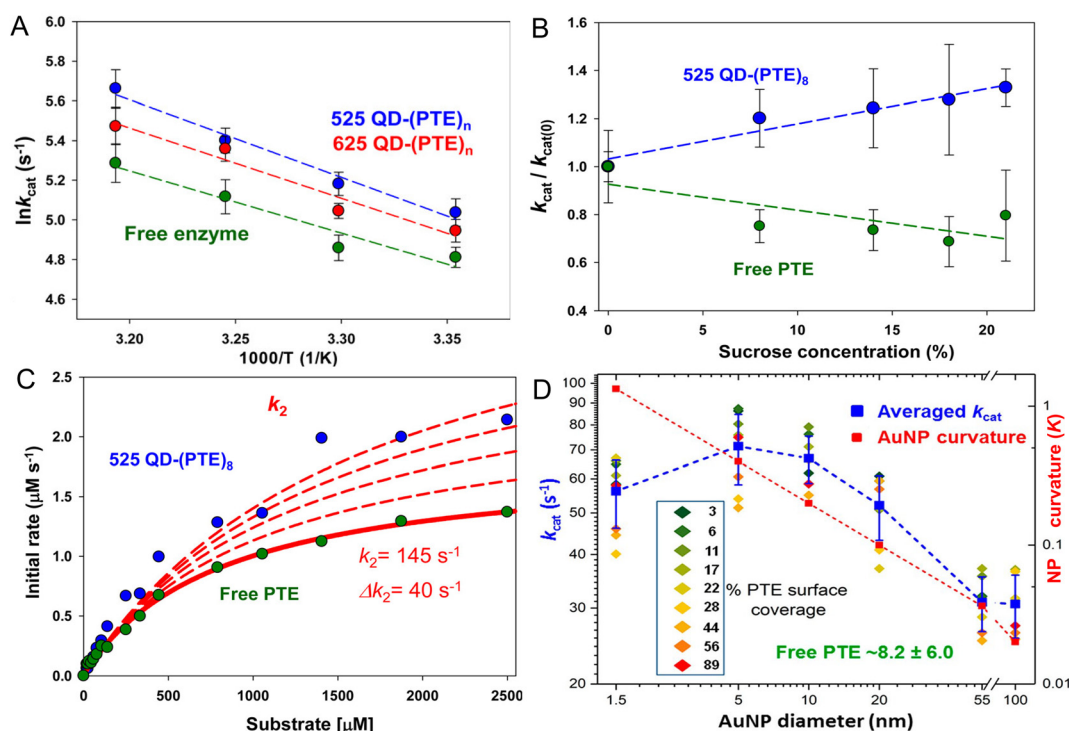


Fig. 4 Analyses of QD phosphotriesterase bioconjugate activity. (A) Arrhenius plot of averaged $\ln k_{\text{cat}}$ values *versus* inverse temperature (Kelvin) for QD–PTE assemblies and free enzyme. Slopes of the fitted data were -3.9 ± 0.4 , -3.5 ± 0.5 , and -3.1 ± 0.6 (average = -3.5 ± 0.4) for the 525 QD, 625 QD, and free enzyme, respectively. (B) Analysis of PTE k_{cat} on and off QD *versus* increasing sucrose concentration. Plots of normalized PTE k_{cat} values from 525 QD–(PTE)₈ conjugates (blue) and equivalent amounts of free PTE *versus* increasing sucrose concentration. Linear fits added to each data series. (C) Plot comparing the effect of potential changes in k_2 on initial PTE rates. The experimental rates of free PTE (green) and 525 QD–(PTE)₈ (blue) *versus* substrate concentration are plotted. An initial k_2 value of 145 s^{-1} was derived from the experimental data. The effect on initial rates of increasing the k_2 value by increments of 40 s^{-1} is then estimated with the red dashed lines. Note the overall qualitatively good fit between free and on QD experimental formats. (D) Comparison of averaged PTE k_{cat} values as AuNP surface coverage increases for the AuNP size series following the indicated color-coding (inset). The average value (of individual replicates) is shown in blue along with its standard deviation. The dashed blue line joins the average values together. Unassembled free PTE k_{cat} value ($8.2 \pm 6.0 \text{ s}^{-1}$) indicated in green. AuNP curvature ($\kappa = 1/\text{radius}$) values in log scale are plotted in red with a line of best fit. Percent PTE coverage was based on ratios of 2.5, 8, 32, 128, 969, and 3202 PTE being assembled to the 1.5-, 5-, 10-, 20-, 55-, and 100 nm-diameter NPs as the 100% coverage maxima. PTE concentration held constant while NP concentration varied to achieve the indicated ratios. Panels A–C reproduced with permission from ref. 72 Copyright 2015 American Chemical Society. Panel D reproduced with permission from ref. 107 Copyright 2019 American Chemical Society.



limiting step even though the localized viscosity should be just as high and the enzyme should encounter the same difficulty in releasing P. By being displayed on the QD surface, the PTE enzyme is now somehow protected from this viscosity increase – presumably by the localized environment around the QD, which is radically different from that of bulk solution. Supporting this notion, Fig. 4C shows simulations fitting the experimentally derived data for both free and QD attached PTE enzymes from Fig. 4B where the rate of k_2 alone was modified while all other variables were kept constant. This excellent fit to the experimental data was not seen when attempting the same exercise while varying k_1 . This exercise suggests that by just increasing the product dissociation rate and effectively alleviating this rate-limiting step, one could potentially explain this effect of QD-enzyme enhancement.

Lastly, Fig. 4D draws from a study where a series of increasingly larger AuNPs were used to ascertain the effect of NP size on PTE catalytic enhancement.⁷² This study showed that PTE k_{cat} increased with NP size from 1.5 nm up to the range of ~5–10 nm AuNP diameter and then steadily decreases as the NP size increases and the NP curvature decreases although the catalytic rate at the largest 100 nm diameter size is still 3–4× that of the free PTE. PTE enhancement is also consistently greater with a lower percentage of surface coverage across all NP sizes tested. These PTE studies all utilized (His)₆-MAC for NP bioconjugation, thus all the enzymes were in the same orientation. That the enhancement was highest with the lowest NP display ratio or lowest surface coverage argues that enzyme freedom of movement, even on a NP surface, is still important. Attaching this enzyme to a bulk flat surface using the same chemistry would orient it on the surface in the same manner, yet enzymatic enhancement is normally not seen in this configuration. It is believed that the reason behind this is that such bulk surfaces lead to stagnation layers near the enzyme where the local substrate is rapidly depleted.^{5,28,30} Cumulatively, the PTE data shows that enhancement is dependent on NP size and curvature with smaller NPs performing better, does not arise from a change to the enzyme's thermodynamics, and most likely arises from a putative change or alleviation of k_2 , the enzyme's rate limiting step. We postulate and some of the experiments discussed above support the notion that the latter arises in part from the unique and structured environment that is found around colloidal NPs (*vide supra*).

Results from displaying QDs around β -gal's surface, which provided for a further 3× enhancement in that enzyme's k_{cat} , are also quite informative even if somewhat counterintuitive (Table 1). β -Gal is considered to be a diffusion limited enzyme meaning that it already functions at its optimal rate and should demonstrate a hard-ceiling towards increased activity arising purely from continuously increasing the concentration of the substrate. This suggested that either a rather unlikely super-diffusional rate of substrate accessibility was now contributing to activity in the presence of QDs or some other change in activity was taking place. Assuming a

similar overall rate-limitation arising from k_2 , the rate of E-P release as discussed above, it seems likely that an alleviation of this step would again be responsible for the observed enhancement. Supporting this, the largest increase in k_{cat} was noted for the β -gal enzyme assemblies displaying the maximum of 4 QDs rather than lower valencies as was seen above with maltase (Fig. 2). We hypothesize that the larger size of this enzyme now requires more QDs to be displayed around it to reap the full benefits of the unique NP interfacial environment. Alternatively, some evidence suggests that proton transfer during β -gal's catalysis is part of the rate-limiting step for k_{cat} .¹⁵³ This would then suggest that the unique QD environment acts to alleviate this limiting mechanism somehow by allowing for an increased rate of proton transfer.

We also note that aside from results reported with NPLs, enzyme enhancement appears to generally increase when smaller NP materials are utilized.^{2,70,72,95,98,99,107,110} Further support for the importance of smaller NP size on achieving enhancement comes from the work of Mukai *et al.*,¹⁵⁴ who assembled the same 7–10 glycolytic enzymes (*albeit* cloned from mice testis) as Breger *et al.*,² via His₆ MAC to 500 nm diameter Ni-NTA silica NPs in different cascaded combinations. In contrast to Breger's finding of kinetic enhancement with half of the enzymes in the cascade, they found that enzyme activity decreased in all cases when tethered to their significantly larger NPs. Cumulatively, this suggests that the enzyme activity should achieve its maximum enhancement for NP materials with zero dimensionality – or the no NP present control. That this also does not occur again argues for the unique NP environment contributing to this phenomenon.

Important to mention is that a frequent alternative explanation put forth is that of localized substrate sequestration around the NP-enzyme bioconjugate.^{155–161} This should not be discounted as it may actually be a contributor to some of the examples seen with DNA-based NP structures. Here, the high-localized concentration of DNA in conjunction with its high charge density may actually be able to build up significant localized concentrations of substrates especially if they are oppositely charged and this, in turn, can increase the rate of substrate binding and catalysis. For the specific case of horseradish peroxidase, HRP (Table 5), Lin and Wheeldon found that attachment of this enzyme to a 2-dimensional 25 nm DNA triangle led to 100–300× increases in activity depending upon what type of substrate was used in the subsequent assay.¹⁴¹ By performing kinetic assays using a library of different HRP chemical substrates, it was revealed that the DNA scaffolding itself could enhance the HRP activity in a manner that is akin to the so called Sabatier principle. This principle is drawn from the field of heterogeneous catalysis and posits that the binding between the substrate and the catalyst should be “just right” and should be neither too weak nor too strong. If substrates bind too weakly, they will fail to associate with the catalyst and result in no net reaction while strongly bound substrates will be conversely too slow to



dissociate and prevent subsequent reactions *via* product inhibition by blocking of the active sites. Lin and Wheeldon concluded that here the substrates bound to the DNA scaffold and not the enzyme but the overarching trend expected for the Sabatier principle remained in place where the weakly and strongly binding substrates manifested no kinetic enhancement in their activity rates while that of the intermediately bound or 'just right' substrates resulted in the highest >300% enhancement in HRP activity. Presumably, this binding led to high-localized concentrations of given substrates around the DNA scaffold, which, in turn, effectively allowed for increased enzyme activity by promoting substrate binding. Hess has posited similar pH-driven changes being responsible for enhancement of HRP and other enzymes when attached to DNA scaffolding.¹⁶² There is also evidence for this effect being related to having high localized concentrations of DNA in a NP assembly. For example, Zhao reported that direct conjugation of the enzymes sarcosine oxidase and HRP to single oligonucleotide strands significantly reduced their activity.¹⁶³ However, it should also be noted that the same report also confirms that inserting the DNA-tethered enzymes into a tetrahedral DNA structure significantly increased their activity to beyond that seen for the free unmodified enzyme.¹⁶³

Wheeldon drew upon these results and some other similar studies that examined enzymatic rate changes in the context of localized physicochemical characteristics to suggest a set of design rules for enzyme–DNA nanostructures where enhanced catalysis is desired.¹⁵⁶ A mixed approach of molecular simulations and kinetic analysis was able to confirm that interactions between enzyme substrates and DNA-scaffolded enzyme nanostructures could increase local substrate concentrations. The enhancement would manifest as a reduction in the apparent K_M and increased enzyme efficiency. The primary determinants to accessing this effect reduce primarily to exerting control over the local physical and chemical environment to obtain the required increases in the local concentration of substrates. Although scaffolding of enzymes on DNA may allow them to manifest enhancements in their activity, the reasons that give rise to this may be amongst the most complex to elucidate mechanistically.^{141,144,164} As Wheeldon elegantly demonstrated above,¹⁵⁶ the physicochemical characteristics of the enzyme, substrate, DNA scaffold itself, the buffering environment, and especially that of the buffer's pH will all interact in a complex manner that can be hard to predict. Given its strongly-charged nature and the way that this is displayed in an almost uniform manner, it would not be unexpected for DNA to universally structure its localized environment as well in a manner akin to that predicted for the colloidal NPs above.^{39,40} This would certainly add another confounding variable to the determinative mix. Fortunately, many of these properties can be isolated and changed independent from the others (*e.g.* buffer pH) and therefore may allow for parametric testing to elucidate some of the contributions and underlying roles of each. There have also been examples reported of mixed NP–DNA–enzyme systems that also yielded some form of enzymatic improvement.^{165–167} The

material complexity here suggests that parsing out and understanding the enhancement mechanism may be incredibly complicated in this situation as localized substrate sequestration from DNA and NP interfacial structuring could both be simultaneously present and contribute to different extents.

An alternative approach to enhancement using substrate–NP attachment

It has long been known that assembling ligands around a NP could significantly improve the observed binding affinity for a receptor specific to that ligand presumably through localized density and avidity effects.^{168,169} Following from this belief in a conceptual manner, enzymatic enhancement at a NP interface was originally observed for a configuration opposite to that focused on above – where NPs were assembled with repeated copies of a substrate in somewhat high density and then exposed to a freely diffusing enzyme.¹⁷⁰ Perhaps the best understanding of these systems is that of proteases acting upon NP-displayed peptidyl substrates and originates from the seminal studies of Algar.¹⁷¹ Using QD donors displaying different densities or valencies of dye-labeled acceptor peptidyl substrates and quantitative monitoring of reaction kinetics by changes to real-time QD-dye Förster resonance energy transfer (FRET) interactions as a prototypical system,¹⁷² Algar showed that trypsin's observed k_{cat}/K_M increased almost 5× when the peptide substrate was displayed on a QD.¹⁷¹ Analyzing these results in the context of several different mechanisms of enzymatic activity at a surface interface, Algar posited a scooting mode of trypsin activity where the enzyme interacts with a given QD-peptide_(n) substrate due to augmented avidity, rapidly consumes all the substrate on that QD, and then diffuses away to another encounter. The Algar group has extensively characterized the activity of different proteases in this configuration and has gone on to develop several multiplexed sensor configurations including portable versions for monitoring simultaneous protease activity.^{173,100,174–180} They, and others, have also looked at the influence of different NP surface ligands and enzyme net charge on this type of activity.^{181–183} Although mechanistically different from the enhancement seen with NP display of enzymes, it is quite possible that the unique structured NP environment could also contribute to some of the kinetic improvements seen here as well. In this scenario, it may be interesting to test the role of E–P dissociation by performing experiments with increasing viscosities to see if it is a contributor.⁹⁷

Future outlook – systems engineering

The depth and breadth of examples listed above, which include 40 different enzymes and >10 types of NP material families, make it clear that enhancement of enzymatic activity by NP display is a real phenomenon and not just a curiosity or misinterpretation of complex results occurring between different experimental configurations. It is also clear that before engaging in designer engineering of molecular systems



geared towards exploiting such enhancements, far more fundamental studies are still needed. Such studies need to be performed across many different NP materials/types and sizes along with different bioconjugation chemistries. More importantly, these studies need to be performed in a systematic and parametrized manner so that both qualitative and quantitative conclusions can be drawn from an analysis of the results. The enzyme maltase presents an interesting example in this vein. Maltase activity in the form of k_{cat} was enhanced by ca. 50 fold when displayed on 4.0 nm dia. QDs, by almost 150× when attached to $19.2 \times 17.3 \times 2.6$ nm NPLs, 35× on a 120 nm per side DNA origami triangle, but then decreased by almost 50% when attached to Au nanostars of 37 and 83 nm Feret dia.^{5,77,107} Pertinently, the enzyme utilized in these examples is the exact same monomeric version and was assembled to the materials using the same approach – namely that of MAC by its terminal (His)₆ precluding confounding effects from any potential NP crosslinking or heterogeneous orientation. This suggests that differences arise not from the enzyme itself but rather from the effect of the materials in the form of the NP dimensionality along with the surface ligand chemistry used to make the NPs colloidally stable. Another important factor to consider if undertaking these types of parametric studies would be to factor in not only many different enzyme types, *i.e.* different catalytic mechanisms, but also enzymes that have different rate-limiting steps beyond the more common EP release. Of course, more complex enzymatic mechanisms including those that have multiple substrates and multiple reaction steps will necessitate more complex studies. It will also be important to gather data on any changes to a given enzyme's energy of activation when attached to a NP. Work with PTE showed that this was not a factor for that enzyme's attachment to QDs,⁷² however, alternate work did show a reduction when cholesterol oxidase was attached to Fe₃O₄ magnetic particles.¹⁵⁰ Given the dearth of information currently available about this, the role of changes in enzymatic activation barriers in this context remains an open question. There is also the possibility that these changes or even increases in enzyme activation energy when attached to NP can be correlated to different types of catalytic transformations based on the enzyme type and mechanism of action.

One fascinating possibility towards performing the type of parametric analysis intimated above without having to engage in exhaustive high-density mechanistic studies is that of drawing data from the literature and subjecting it to artificial intelligence (AI) or machine learning (ML) based analysis.^{184,185} This approach would draw from the existing body of data in the literature and compiles as many quantitative (*e.g.* NP size, k_{cat} enhancement, enzyme molecular weight, *etc.*) and categorical (*e.g.* NP surface ligand type, substrate type, bioconjugation chemistry used, NP component materials, *etc.*) descriptors or attributes as possible for subsequent analysis. Moreover, negative data in the form of decreases to enzyme activity would be extremely useful in this context and do not have to remain in the background as so-called 'dark data'.¹⁸⁶ Indeed, the latter type of data may be especially useful as it can

allow for parsing of factors that correlate negatively with enhancement and not just those that contribute positively to give a more balanced analysis and deeper insight into the underlying mechanisms. This type of AI-based approach has so far proven to be quite useful for analyzing NP toxicity across hundreds of different material preparations and cell-lines along with design of new electrocatalytic NP materials.^{187–189} An important caveat with this type of approach is that to be really useful it needs extremely large data sets that have huge numbers of examples and that are very well curated in terms of attribute type and number.¹⁹⁰ This suggests that the above compendium is not anywhere near meeting a critical baseline requirement for such an analysis.

Turning towards incorporating NP-enzyme enhancement within engineered molecular systems, several applications currently in use can potentially benefit from this phenomenon in the near term. For example, many enzymatic assays in both research and industrial use require key cofactors for use as electron donors or acceptors in a given reaction.^{191–193} The vast majority of these cofactors, including especially the redox-active nicotinamide adenine dinucleotide cofactor(s) NAD(P)⁺/NAD(P)H, for example, are exceedingly expensive to purchase directly and, in many cases, become the most expensive reagent utilized in these assays. Moreover, buildup of the reduced or oxidized form of the cofactor during the reaction can also poison it. This has led towards sustained efforts to recycle these cofactors in a given reaction using recycling enzymes that are driven by some far cheaper substrates such as glucose towards improving economic viability.^{191–194} In a recent report, Breger and co. showed that glucose dehydrogenase driven reduction of NAD⁺ → NADH (and similarly NAD(P)⁺) could be enhanced by up to 10× simply by displaying it on QDs.⁹⁸ This could certainly increase turnover in these reactions along with even further reducing costs by allowing for even less substrate to be used. Following from this example, many diagnostic assays are based on some form of enzymatic readout such as, for example, the ubiquitous enzyme-linked immunosorbent assay (ELISA) and reverse dot-blot hybridization assays.^{195–197} Increasing the apparent catalytic activity of the enzymes utilized in such formats may represent another cost-effective and powerful way to decrease their reaction time along with the amount of enzyme required. More generally, increasing the activity of enzymes used commercially in all manner of stand-alone industrial applications may help increase their cost-effectiveness along with decreasing the requisite reaction times.³ This would be especially beneficial for enzymes sourced from complex eukaryotes, many of which display low endogenous catalytic rates when applied extracellularly in non-native reaction environments.^{198,199}

The potential application where we believe that NP-enzyme enhancement may play an outsized role is within designer minimalistic cell-free multienzyme cascaded reactions.^{1–6,200} Rather than engineering and maintaining a cellular system to produce a desired enzymatic product, as is the current major focus in the growing field of synthetic



biology, minimizing the required reactions to where all that is present are the requisite enzymes, substrates, and cofactors can allow for production of target molecules that are normally toxic to cells or otherwise not favored due to competing intracellular pathways and cross-inhibiting reactions.^{201–205} Moreover, such a format can allow for potential incorporation of xenobiotic or non-natural substrates towards making new molecules enzymatically.^{6,200} The primary issue faced by these types of reaction formats is that of diffusion limitations and enzyme stability and it is here where NP display of enzymes can contribute. Although not focused on here, attaching enzymes to NPs can and, in many cases, does increase their long-term stability and viable lifetime along with increasing their catalytic activity.^{2,72,97,98} More importantly, assembly of multiple linked enzymes that constitute a biomolecular cascade with NPs into cross-linked nanoclusters can allow them to access intermediary or probabilistic channeling. Channeling is a process that occurs when enzymes are brought into close proximity of each other and where the diffusion of an intermediary from one upstream enzyme into the bulk is decreased and it instead encounters the next downstream enzyme in a cascade so as to produce a net increase in flux through the multistep system. This nanoscale diffusion-limited process is the most efficient form of multienzyme catalysis and occurs when the apparent catalytic rate of flux \gg diffusion rate.^{147,161,206,207} Vranish, Breger and others have shown that QDs can crosslink with multimeric enzymes present in a cascade to form QD–enzyme nanoclusters that speed up overall catalytic flux by orders of magnitude *via* channeling *versus* the same concentration of enzyme freely diffusing in solution.^{2,95,97,98} Within these clusters, enzymatic intermediaries have a high probability of finding the next enzyme in the cascade due to the high localized density of enzymes present. Not only could NP display of enzymes enhance the activity of many of the participants in these cascades, but it could also enhance the rate or efficiency of the channeling process. Moreover, by initially measuring the apparent catalytic rates and affinity of each enzyme when present on a NP and enhanced, one can undertake numerical simulations to optimize the relative ratio of each enzyme present in the cluster to the others so as to maximize the rate of channeled flux.² Overall, this represents a promising approach to increasing the productivity of multienzyme cascades while also potentially requiring a lot less enzyme material. A complementary and mostly unmet need in the field of synthetic biology is that of sensors to monitor and report on the pathways being utilized and the concentrations of products or intermediaries in a reaction regardless of whether it is intracellular or in a cell-free format.^{208,209} NP–enzyme and NP–substrate sensors may also find utility here where they would directly benefit from improved performance.²¹⁰ Combining enzymes with QD's intrinsic photoluminescence and FRET or other energy transfer modalities can allow for sensor assemblies that are capable of extremely complex and concatenated Boolean logic functions.^{211–213} This can allow such sensors to perform

rudimentary computational analysis and provide a more complex and nuanced data output on the underlying processes being monitored.

Although our focus here has been on enzymatic enhancement when displayed on a NP surface, we are cognizant that such attachment can also lead to changes in enzyme stability and as mentioned before this may not necessarily be in a detrimental manner. Examining enzymatic channeling in a coupled pyruvate kinase–lactate dehydrogenase (LDH) enzymatic system assembled on QDs, Vranish noted that assembly of the latter LDH significantly increased that enzyme's stability.⁹⁷ LDH is an obligate tetramer that normally dissociates and ceases functioning at around 10 nM concentration, however, displaying it on QDs allowed it to continue catalyzing turnover even when diluted by a further 1.5 orders of magnitude. In this case, the tetrameric enzyme's 4-pendant terminal (His)₆ moieties cross-linked with the QDs into clusters that presumably helped facilitate structural stabilization. This is certainly a promising result and suggests that not only should enzymes be tested for catalytic enhancement when attached to NPs, but they should undergo further experimental evaluation to determine if they are also stabilized in this context. Such parametric analysis could help provide further information on which types of enzyme structures are more amenable to being incorporated into such configurations. The interested reader is directed towards further reviews describing enzymatic stabilization by NP structures.^{214–217}

The long-term outlook also remains promising especially given how quickly the fields of synthetic biology and biotechnology in general are moving. Enzyme evolution coupled with AI offers the prospect of creating new enzymes capable of new transformations and these too should be amenable to enhancement by NP display.^{218–220} There is also the possibility of accessing the same enhancement phenomenon but without the need for an actual NP material to be present. For example, protein–protein binding motifs and fusion/ligase sequences such as the SpyTag–SpyCatcher system and its myriad derivatives have now matured to the point that they can be designed to allow designer multiprotein clusters to form while still providing stoichiometric control over the participants.^{221–223} This could potentially allow self-assembly or formation of designer CLEAs where component enzymes manifest some enhancement due to the structure's NP-like properties. The ability of NP–enzyme enhancement to allow PTE to tolerate and even work better than the free enzyme in the presence of a triethyl phosphate competitive inhibitor is also quite intriguing (Fig. 3C).⁷² This suggests possible utility in drug screening assays where a given chemical compound may inhibit more than one target enzyme. This would also allow for larger amounts of an inhibitor to be used in an assay. Lastly, it is well worth considering that the actual implementation of NP–enzyme enhancement does not have to be perfect or fully optimized to generate a benefit from its effects – any enhancement can be helpful in many cases. In



summary, although a significant amount of work still needs to be undertaken to fully utilize NP-enhancement of enzyme activity in a reliable and engineered manner, the significant benefits from achieving this suggest that much will be gained by this pursuit in the near and long term.

Author contributions

SLH, JCB, and ILM all conceived and prepared the manuscript.

Conflicts of interest

The authors declare no competing interests.

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