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## Navigating the landscape of enzyme design: from molecular simulations to machine learning

Jiahui Zhou and Meilan Huang \*

Global environmental issues and sustainable development call for new technologies for fine chemical synthesis and waste valorization. Biocatalysis has attracted great attention as the alternative to the traditional organic synthesis. However, it is challenging to navigate the vast sequence space to identify those proteins with admirable biocatalytic functions. The recent development of deep-learning based structure prediction methods such as AlphaFold2 reinforced by different computational simulations or multiscale calculations has largely expanded the 3D structure databases and enabled structure-based design. While structure-based approaches shed light on site-specific enzyme engineering, they are not suitable for large-scale screening of potential biocatalysts. Effective utilization of big data using machine learning techniques opens up a new era for accelerated predictions. Here, we review the approaches and applications of structure-based and machine-learning guided enzyme design. We also provide our view on the challenges and perspectives on effectively employing enzyme design approaches integrating traditional molecular simulations and machine learning, and the importance of database construction and algorithm development in attaining predictive ML models to explore the sequence fitness landscape for the design of admirable biocatalysts.

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### 1. Introduction

Over the past decade, enzyme biocatalysis has become a promising alternative to traditional chemical transformations for the sustainable production of valuable chemicals such as biofuels and pharmaceuticals<sup>1–3</sup> and hence has attracted increasing attention from both academia and industries. In order to meet the requirements of large-scale industrial production, new biotechnologies have been developed to discover novel enzymes or optimize existing enzyme biocatalysts to improve their catalytic activities, substrate specificity, selectivity, stability, *etc.*<sup>4–7</sup> The success of structure-based enzyme design strategies has been exemplified in numerous cases in rational design, semi-rational design and *de novo* design. However, it remains challenging to design novel biocatalysts for specific reactions by navigating the vast protein fitness landscape. Recently, machine learning has emerged as an efficient strategy to harness the available data, accelerating the discovery of enzyme biocatalysts and enabling the accurate prediction of mutation sites to achieve biocatalysts with desirable properties.<sup>8–10</sup>

#### 1.1 Structure-based enzyme design

The semi-rational enzyme design approach is based on the prior knowledge of enzyme structure and function to navigate

the vast theoretical sequence space by screening a small sequence library generated from random mutagenesis or targeted mutagenesis.<sup>11</sup> Efficient procurement of mutant variants with the desired functionalities may be achieved by constructing smart mutant libraries and employing appropriate experimental or computational high-throughput screening methods.<sup>12–14</sup> Rational enzyme design requires detailed knowledge of the enzyme's mechanism of action, *e.g.* how it binds to substrates and catalyzes reactions, to guide enzyme engineering for improved or altered function. In addition to mutations based on existing natural sequences, the functional enzymes can be designed from scratch through pre-construction of catalytic sites and selection of protein scaffolds, followed by atomistic simulations.<sup>15–17</sup>

Structure-based enzyme design requires the identification of active sites and substrate binding pockets, however, many enzymes of interest lack resolved structures, and their sequences often exhibit low homology with the known proteins with available crystal structures, making homology modeling unsuitable for obtaining reasonable starting structures. In the past few years, deep-learning based protein structure prediction tools such as AlphaFold2<sup>18</sup> and RoseTTAFold<sup>19</sup> have shown great success in predicting protein 3D structures. Ligand binding mode and the dynamic properties of protein complexes can be further explored by using molecular docking and molecular dynamics simulations. The functions and catalytic mechanisms of enzymes are highly intricate, and are dependent on binding affinities

*School of Chemistry and Chemical Engineering, Queen's University, David Keir Building, Stranmillis Road, Belfast BT9 5AG, Northern Ireland, UK.*  
*E-mail: m.huang@qub.ac.uk*



of the substrates and the reaction kinetics of the enzymes. Hybrid molecular mechanics and quantum mechanics (QM/MM) enable the prediction of enzyme-catalyzed reaction kinetics. It is worth noting that structure-based enzyme design requires advanced knowledge in molecular modeling and is also computationally prohibitive for screening a large database to identify the enzyme sequences with desirable functions.

## 1.2 ML-accelerated enzyme design

In the era of big data, enzyme sequence and structural and functional data have been accumulated and shared at an unprecedented pace. This provides a wealth of information resources for machine-learning guided enzyme design by learning the inherent patterns from data to make predictions. However, the surge in data also brings about the challenge of efficiently harnessing the data to generate generalized ML models to make accurate predictions for accelerating the design of enzymes with improved properties.<sup>20–22</sup>

In this review, we summarize the techniques and applications of computer-aided enzyme design using molecular simulation approaches and machine learning techniques. We also provide our perspectives on effective enzyme design through the synergetic combination of molecular simulations, machine learning and experimental validations.

## 2 Computer-aided enzyme design tools and applications

### 2.1 Enzyme modelling methods

**2.1.1 Molecular modeling.** The rationale of structure-based enzyme engineering is that the structures of enzymes dictate their functions. Designing biocatalysts with admirable functions, or optimizing specific catalysts to achieve improved catalytic efficiency, selectivity or stability often requires an in-depth understanding of the relationship between their structures and functions. For this, accurate acquisition of enzyme structures is essential.

Compared with the vast protein sequence space in nature (with over 244 million protein sequences in the UniProt database<sup>23</sup> as of May 2024), the number of protein structures is much smaller (with over 220 thousand structures in the Protein Data Bank<sup>24</sup>). Currently characterized structures only account for less than 10% of the total protein sequences, and the capability of structure characterization largely lags behind that of sequence acquisition (Fig. 1a). Experimentally determining the three-dimensional structure of a protein is a costly and time-consuming process and some proteins are highly flexible, which makes structural determination even more challenging. When the 3D structures of proteins are not available, computational methods become powerful tools in predicting protein structures based on their sequences.



Jiahui Zhou

*Jiahui Zhou is a PhD student at Queen's University Belfast, working under the supervision of Dr Meilan Huang. He received his BS degree in Pharmaceutical Engineering from Beijing Institute of Technology in 2020. His research focuses on using molecular modeling and simulation methods to unravel complex phenomena in enzyme catalysis.*



Meilan Huang

*Meilan Huang obtained her PhD from Zhejiang University in physical chemistry in 2003. She worked as a research assistant on organic synthesis of anticancer drugs in Prof Fengling Qing's lab in the Key Laboratory of Organofluorine Chemistry at Shanghai Institute of Organic Chemistry, Chinese Academy of Sciences in 1998. She was a postdoc in the Department of Chemistry at the University of Calgary, Canada in 2003, working with Prof Arvi Rauk on the theoretical study of the oxidation mechanism related to Alzheimer's diseases. Awarded with Wellcome Trust International Fellowship on computer-aided drug design, she spent two years in the Laboratory of Physical and Theoretical Chemistry at the University of Oxford, working with Prof W. Graham Richards during 2004–2006 and then moved to Department of Medicine at the University of British Columbia, Canada in 2006, working with Prof Artem Cherkasov on the rational design of infectious disease therapies. She joined Queen's University Belfast as a lecturer in 2007. The research in the Huang group is focused on molecular modelling and theoretical catalysis at the interface of chemistry and biology.*





**Fig. 1** Molecular modeling in enzyme engineering. (a) Growth rate of the data in the Protein Data Bank and UniprotKB/TrEMBL database. (b) Protein modeling approaches. (c) Modelled structures for a new sesquiterpene synthase JeSTS4 using different protein modeling approaches.<sup>25</sup> HM\*: homology model was built using the crystal structure of the sesquiterpene synthase Copu9 from *coniophora puteana* (PDB: 7OFL<sup>26</sup>) as a template (sequence identity: 25%); *ab initio* models were built using I-TASSER and AlphaFold2, respectively. (d) Modelled structures for the Ga98 variants<sup>27</sup> with three progressed single mutations using ColabFold.

**2.1.1.1 Traditional modeling methods.** When 3D structures of proteins are not available, computational methods have shown their power in predicting protein structures based on their sequences.<sup>28</sup> Structure prediction approaches can be classified into template-based modeling represented by homology modeling and protein threading, or template-free modeling (*ab initio* modeling)<sup>29</sup> (Fig. 1b).

For sequences that share certain homology with crystal structures, their homology models can be built using tools such as Modeller<sup>30</sup> and Swiss-Model.<sup>31</sup>

For sequences with low sequence identity to known crystal structures, the fold recognition method (*e.g.* protein threading) can be used to predict structures by matching the query sequence directly onto the 3D structures of other solved proteins.

For sequences with no structural similarity to any solved proteins, *ab initio* modeling can be used to predict protein structures from scratch.

In principle, the global lowest energy conformation of a protein can be obtained using molecular simulations. In 1998, molecular dynamics simulations (MD simulations) disclosed a marginally stable folded conformation during the folding process of a 36-residue peptide,<sup>32</sup> marking the first simulation-based *ab initio* modeling. Due to the demanding computational cost, it is impractical to predict full length protein structures using simulation-based *ab initio* modeling.

Currently, most of the *ab initio* protein structure prediction tools are composite approaches that combine fold recognition, structure assembly, and structure refinement. For example, I-TASSER developed by Zhang lab<sup>33</sup> utilizes protein threading to identify similar structural motifs from the structure database, to assemble the well-aligned motifs. For the unaligned regions, Monte Carlo based modeling is used to predict the structure. In Rosetta developed by Baker,<sup>34</sup> the target sequence is segmented into a consecutive window of three or nine residues and its structure is predicted by selecting fragments that are then assembled by a Monte Carlo strategy to construct the structure.

**2.1.1.2 Deep learning-based structure prediction methods.** AlphaFold1<sup>35</sup> secured the top ranking in the CASP13 free modeling (FM) category.<sup>36</sup> AlphaFold1 extracts co-evolutionary information and employs neural networks to generate residue contact maps, which are then used to predict protein structures.

In contrast, AlphaFold2<sup>19</sup> employs a completely new architecture, differing significantly from previous methods which relied on residue contact maps to indirectly predict protein tertiary structures. The approach to predict protein structures is to learn the three-dimensional structure of proteins directly from their amino acid sequences, a so-called “end-to-end” learning method. AlphaFold2 has significantly advanced the development of “end-to-end” structure prediction, wherein the





predict the structure of GB98-T25I but failed to predict the structures of Gb98 and GB98-T25I/L20A correctly.<sup>43</sup>

Structure prediction tools can serve as initial points for structural and functional analysis of enzymes, however careful inspection has to be conducted for the structure model obtained. Moreover, understanding the subtle mutation effects, particularly single mutations on enzyme properties such as enhanced stability or activity requires more precise structural simulations and sampling.

## 2.1.2 Molecular dynamics simulations

**2.1.2.1 Classical MD simulation method.** In structure-based drug discovery, protein targets are usually treated as fixed to allow large scale virtual screening to identify potential hits, by evaluating the binding affinities of small ligands in the binding pocket of the drug target, which can then be processed for bioassay. However, in biocatalysis, due to the promiscuity of enzyme's catalytic pocket induced by mutations or ligand binding, it is inappropriate to neglect the dynamic conformations of enzymes, which cannot be obtained by experimental X-ray, NMR or the *ab initio* models. Molecular dynamics provides an effective way to describe the dynamic properties of enzymes at the atomic level to interpret their functions.<sup>51</sup> The development of molecular dynamics (MD) methodology tailored for biological macromolecules such as GROMACS,<sup>52</sup> AMBER,<sup>53</sup> CHARMM<sup>54</sup> and OpenMM<sup>38</sup> and acceleration of simulations by graphics processing units (GPU) on high-performance computing (HPC) has enabled accurate and fast prediction of protein structures as well as the binding modes of protein–ligand or protein–protein interactions.

CHARMM is one of the most widely used MD software packages and the CHARMM force field has been developed along with the software since the 1980s.<sup>55</sup> A user-friendly graphic interface CHARMM-GUI<sup>56</sup> was developed to prepare the input of simulations interfaced with widely used MD simulation packages such as CHARMM, GROMACS, AMBER and OpenMM. GROMACS<sup>52</sup> is known for its highly optimal computing efficiency and open-source code and has become one of the most popular MD software packages for biomacromolecules. It is interfaced with different forcefields including AMBER99SB,<sup>39</sup> CHARMM36,<sup>57</sup> GROMOS<sup>58</sup> and OPLS-AA/M.<sup>59,60</sup> Benchmark studies on the commonly used MD simulation packages showed that GROMACS was optimal for biomolecular simulations of medium-sized systems at the microsecond level.<sup>61,62</sup> The AMBER package<sup>53</sup> includes the AMBER simulation software with the AMBER force-field. The program assembly package AmberTools is freely accessible and convenient for preparing the input and result analysis. The input files generated by AmberTools can also be converted by third-party scripts such as ParmEd (<https://github.com/ParmEd/ParmEd>) and acpype (<https://github.com/alanwilter/acpype>) so as to be readable by other MD software packages like GROMACS. Other efforts have been reported to automate the process of preparing the AMBER inputs and conducting result analysis.<sup>63</sup> OpenMM<sup>38</sup> is an open-source MD simulation package with a layered and modular architecture, making it easily integrable with other applications. It is highly extensible, allowing for the implementation of various plugins.

**2.1.2.2 Enhanced sampling methods.** Depending on the software, hardware and molecular system, the timescale of MD usually ranges from tens to hundreds of nano seconds. It has been demonstrated by a number of MD simulation case studies that the properties of protein–ligand complexes can be captured using simulations at the nano second time scale. However, it is difficult to observe large conformational changes for enzyme complexes *e.g.* from the reactant to product states of the enzyme by traditional MD simulations, because high energy barriers need to be overcome for the transitions between different conformations to take place, making it challenging to extensively sample free energy landscape.

Potential of mean force (PMF)<sup>64</sup> is a modern statistical method commonly used to characterize the energetics of transitions in biomolecules. However, it is impractical to compute PMF directly from MD simulations because of the large configurational space of proteins and also a large energy barrier along the reaction coordinate. Various sampling techniques have been developed to effectively and accurately compute PMF. An effective technique in enhanced sampling to gain large-scale conformational changes is enhanced sampling<sup>65</sup> including the umbrella sampling method,<sup>66</sup> metadynamic method,<sup>67</sup> accelerated molecular dynamics method (AMD)<sup>68</sup> and replica exchange molecular dynamics, REMD.<sup>69</sup>

Umbrella sampling<sup>66</sup> is one of the most widely used enhanced sampling methods in MD.<sup>70</sup> The conformations between the thermodynamic states are sampled in a set of umbrella windows along the reaction coordinate  $\xi$ . At each window  $\xi_i$  ( $i = 1, 2, 3, \dots, N$ ), MD simulations are conducted with a bias potential (umbrella potential) added to restrain the system around a narrow space around  $\xi_i$  so as to enable more efficient conformational sampling in this region.

The bias potential is usually calculated using a harmonic function

$$V_i^b(\xi) = \frac{1}{2}k_i(\xi - \xi_i)^2 \quad (1)$$

where  $k_i$  is the force constant.

The free energy at the position  $\xi_i$  is calculated with the bias potential added onto the unbiased total energy of the state  $U(R)$ , which is a function of the coordinate  $R$

$$U_i^b = U(R) + V_i^b(\xi) \quad (2)$$

For each umbrella window, the probability distribution  $P_i(\xi)$  along the reaction coordinate is represented by an umbrella histogram  $h_i(\xi)$ . The weighted histogram analysis algorithm (WHAM) is a widely used technique in umbrella sampling to calculate PMF from the histogram, to resume the unbiased free energy profile by umbrella integration to obtain the complete free energy landscape along the minimum free energy pathway.

Umbrella sampling is traditionally combined with the post-analysis process. Following the MD runs for a number of biased window simulations, the neighbouring overlapping windows are combined, which allows the system to transit from one conformation state to another and generate the free energy over a large range of reaction coordinates. Adaptive umbrella



sampling<sup>71</sup> constructs a good biasing potential to counterbalance the free energy barrier, so as to allow self-consistently determining the bias potential with less human intervention to achieve a uniform distribution.

Metadynamics is also a bias potential-based method.<sup>72,73</sup> Bias potential is placed on the Hamiltonian of the system thus the system would skip the transition barrier provided the growing bias potential counterbalances the transition barrier. This strategy can escape local minimum and allows for navigating free energy landscape as a function of a few collective variables (*e.g.* bond to be formed or broken, bond angle or dihedral) related to enzyme-catalyzed reactions with accelerated sampling. The choice of independent collective variables is crucial for those reactions for which prior knowledge of reaction coordinates is not available.<sup>67</sup>

Both umbrella sampling and metadynamics methods require prior knowledge on the degree of freedom for the motion of interest, based on either reaction coordinates or collective variables. The accelerated molecular dynamics method (aMD) does not need prior knowledge of potential energy wells or saddle points to explore the rare events that are related to the reaction. A bias potential is added to the true potential such that it is easier for the system to escape from the potential well and move from one low-energy basin to another. This strategy accelerates the sampling of the conformational landscape while converging to correct probability distribution. Replica exchange molecular dynamics based on a replica-exchange method (REM) also does not need knowledge of reaction coordinates. It generates an ensemble consisting of multiple copies (replicas) at different temperatures, and the copies are exchanged to overcome high-energy barriers so as to effectively explore the transitions among different states and conformational space.

These enhanced sampling methods have largely sped up the conformational sampling, however, they may still be slow processes while sampling irrelevant states so that not suitable to be used to refine the large scale predicted *ab initio* models. The Bayesian-based modeling employing limited data (MELD)<sup>74,75</sup> method applies restraints to incorporate data in MD simulations with coarse physical insight, which harnessed weak information and generated multiple-funnel landscape, and sped up the sampling by up to five orders of magnitude. Recently, MELD combined with REMD (MELD × MD) was employed to predict the *ab initio* models of Ga98 and its variants (Fig. 1d)<sup>50</sup> and accurately predicted all of the four structures.

The advancement of deep learning algorithms has also contributed to the development of enhanced sampling techniques.<sup>76,77</sup> For example, Tao *et al.* developed a deep learning enhanced adaptive sampling method that can predict larger conformational changes efficiently.<sup>78</sup> Tiwary *et al.* developed an enhanced sampling method that combined AlphaFold2 with deep learning enhanced MD to generate a collection of Boltzmann-weighted protein conformations from sequences, using the structures predicted by AlphaFold2 as the initial inputs.<sup>79,80</sup> Combining deep learning with statistical

mechanics, Noé *et al.* developed an adaptive sampling method that generated unbiased equilibrium samples of protein conformations using Boltzmann generators initialized by metastable states, without the need of prior knowledge of reaction coordinates.<sup>81</sup>

**2.1.2.3 Binding free energy calculations.** The catalytic efficiency of enzyme biocatalysts is dependent on both the thermodynamic binding free energy and reaction kinetic activation energy of the enzymes. The binding affinities of substrates in enzymes can be estimated by binding free energy calculations. The commonly used methods are MM/PB(GB)SA.<sup>82–84</sup>

In MM/PB(GB)SA, the MD simulation is run for the system solvated in a periodic box with water and counterions. Then the binding free energy between the enzyme and its substrate can be calculated for MD simulated structures processed by stripping the solvent and counterions, according to eqn (3):

$$\Delta G_{\text{Binding}} = G_{\text{ES}} - G_{\text{E}} - G_{\text{S}} \quad (3)$$

where E denotes the enzyme and S the substrate. In turn,  $\Delta G_{\text{Binding}}$  can also be represented as eqn (4):

$$\Delta G_{\text{Binding}} = \Delta H - T\Delta S = \Delta E_{\text{MM}} + \Delta G_{\text{sol}} - T\Delta S \quad (4)$$

Here,  $\Delta H$  represents the binding enthalpy and  $-T\Delta S$  accounts for the conformational entropy change upon ligand binding.  $\Delta H$  can be decomposed into different terms: the gas phase free energy contributions  $\Delta E_{\text{MM}}$  (eqn (5)) and the solvation free energy contributions  $\Delta G_{\text{sol}}$  (eqn (6)).

$$\Delta E_{\text{MM}} = \Delta E_{\text{bond}} + \Delta E_{\text{angle}} + \Delta E_{\text{dihedral}} + \Delta E_{\text{ele}} + \Delta E_{\text{vdw}} \quad (5)$$

In eqn (5),  $\Delta E_{\text{MM}}$  includes the internal energy ( $\Delta E_{\text{bond}}$ ,  $\Delta E_{\text{angle}}$  and  $\Delta E_{\text{dihedral}}$ ), electrostatic contribution ( $\Delta E_{\text{ele}}$ ) and van der Waals contribution ( $\Delta E_{\text{vdw}}$ ).

$$\Delta G_{\text{sol}} = \Delta G_{\text{pol}} + \Delta G_{\text{non-pol}} = \Delta G_{\text{PB/GB}} + \Delta G_{\text{non-pol}} \quad (6)$$

In eqn (6), the solvation energy can be decomposed into electrostatic term  $\Delta G_{\text{pol}}$ , and non-electrostatic term  $\Delta G_{\text{non-pol}}$ . The PB and GB models estimate the polar component of the solvation.  $\Delta G_{\text{PB/GB}}$  is calculated with the electrostatic component calculated using the Poisson–Boltzmann equation or the generalized Born model.

The nonpolar free energy  $\Delta G_{\text{non-pol}}$  is proportional to the molecule's total solvent accessible surface area (SASA), with a proportionality constant  $\gamma$  derived from experimental solvation energies of small non-polar molecules (eqn (7)).

$$\Delta G_{\text{non-pol}} = \gamma \text{SASA} + b \quad (7)$$

To decide the minimum free energy pathways between states of an enzymatic system, the free energy pathway can be explored by umbrella sampling breaking down the distance along the reaction coordinates into a series of very small coupling parameter  $\lambda$  ( $\lambda$  varies from 0 to 1). MD simulations are run at the fixed reaction coordinates along the reaction



pathway and then the free energy change at each point is calculated by integrating the mean values of the derivatives (eqn (8)).

$$\Delta G = \int_{\lambda=0}^{\lambda=1} \left\langle \frac{\partial H}{\partial \lambda} \right\rangle_{\lambda} d\lambda \quad (8)$$

Another class of methods is alchemical methods, where binding free energy is estimated by the statistical analysis of the simulated thermodynamic pathway between two end states. Free perturbation (FEP)<sup>85</sup> and thermodynamic integration (TI)<sup>64,86</sup> methods are commonly used alchemical methods to explore the enzyme conformation landscape. In free energy perturbation (FEP),<sup>85</sup> the free energy difference between two states of a system is calculated using eqn (9).

$$\Delta G_{\lambda} = -RT \ln \left\langle e^{-(\Delta H_{\lambda'} - \Delta H_{\lambda})/RT} \right\rangle_{\lambda} \quad (9)$$

where the triangular brackets denote an average of thermodynamic windows over a MD simulation run for state A.

In thermodynamic integration (TI),<sup>64</sup> the free energy difference between two states is calculated by the integration of the ensemble average of the derivative of Hamiltonian with respect to  $\lambda$  at different  $\lambda$  values for alchemical reaction pathways.

These robust free energy methods are accurate in principle but require extensive sampling from long MD simulations. They have been combined with conformational sampling techniques such as umbrella sampling and alchemical simulations to speed up the calculations.

### 2.1.3 Quantum mechanics and multiscale simulations.

The catalytic efficiency of enzymes is not only dependent on the binding free energies of reactants, but also the reaction barriers of the catalytic reactions. Quantum mechanics (QM) and hybrid QM/MM methods are commonly used to evaluate the reaction mechanism of enzymes, with the initial structures taken from either crystal structures or MD simulated structures.

**2.1.3.1 QM cluster method.** In the QM cluster method, the active site of the enzymes is calculated by QM methods most commonly density functional theory and the remainder of the enzyme is fixed and treated using the continuum solvent with dielectric constant  $\epsilon = 4$  to reduce the computing cost. The QM region is usually composed of the substrates, cofactors, metals and interacting residues with side chains truncated. The method is usually applied using different sized models; a smaller model to quickly explore possible reaction pathways, and a larger model to study the environment of the active site.<sup>87</sup> With the increasing computing power, QM can contain more than 300 atoms nowadays.<sup>88</sup>

QM-Cluster methods optimize only truncated active site models, eliminating the degree of freedom of the region beyond the active site and hence reducing the complexity of the sampling problem. However, during the geometry optimization of a QM cluster model, geometric constraints have to be introduced to avoid the deformation of the active site in absence of the full protein environment. Dasgupta *et al.*

proposed to apply a harmonic confining potential to the terminal atoms (“anchor atoms”) of the QM model, rather than using fixed-atom constraints adopted in traditional QM-cluster methods. This approach improved optimization efficiency and robustness in locating the transition states,<sup>87</sup> and would be particularly useful for those enzymes with large conformational change during the reaction process involving notable entropic effects.

It is usually impossible to achieve reliable kinetic and thermodynamic results by calculating a small QM cluster model. A “maximal” QM cluster model with a residue interaction network of the entire protein was developed and provided reliable results.<sup>89</sup> QM methods have similar computing costs to QM/MM calculations and are popular to those who are only interested in the overall reaction mechanism; however, they may generate different conformations compared to those predicted by QM/MM methods.

**2.1.3.2 QM/MM method.** Hybrid quantum mechanics/molecular mechanics (QM/MM) methods combine accurate QM methods to study the reactions and classical MM force field methods to capture the conformational energetics and have been widely used to study enzyme-catalyzed reactions.<sup>90–99</sup> The starting structures can be obtained either from experimental X-ray or NMR structures or reliable molecular modeling followed by proper sampling from multiple replicas of MD simulations.

Additive QM/MM is a popularly used scheme based on the following equation:

$$E_{\text{Total}} = E_{\text{QM}(R,r)} + E_{\text{MM}(R)} + E_{\text{QM/MM}(R,r)}$$

The effect of the MM region on the QM region is calculated using either electrostatic embedding or mechanical embedding. For accurate QM/MM studies, the polarization effect of MM estimated using the Drude oscillator (DO) model is insignificant for enzyme systems that involve no significant charge transfer.<sup>100</sup> Appropriate choice of the QM region in the QM/MM calculations is crucial for attaining meaningful results.

Bím *et al.* recommended a mechanism-based practice for predicting the mutation effect on enzyme kinetics,<sup>101</sup> which was in good agreement with the experimental value. It combined QM/MM and QM, where QM/MM is used to optimize the geometries of reactants, transition states, intermediates and products and QM is used to estimate the energies.

**2.1.3.3 QM/MM MD method.** QM cluster and QM/MM methods are suitable for exploring the potential energy surface of reactions. Since the enzymatic reaction process involves conformational dynamics, a combination of QM/MM and MD can be employed to extensively sample the potential energy surface. However, QM/MM MD simulations are computationally very expensive because the QM energy and forces are computed from a converged SCF at every step. For example, a QM/MM MD simulation with a QM region containing 49 atoms, using B3LYP density functional with the 6-31G\* basis set and on an NVIDIA V100, can achieve only 1.86 ps per day.<sup>102</sup> The scalable QM/MM



MD calculation framework MiMiC<sup>103</sup> enables running several ps per day in a single simulation using thousands of standard CPU cores.

Alternatively, a less expensive semiempirical method has been adopted in QM/MM MD to reduce the computing cost. For example, the PM3 semiempirical method was employed in a steered QM/MM MD in the hydride transfer mechanism study of zinc-dependent hydrogenase/reductase.<sup>104</sup>

The steered QM/MM MD method<sup>105</sup> has been used to study the enzymatic reactions at an affordable time scale. This method applies harmonic forces on selected atoms to the reaction mechanism along the reaction coordinate and has been used for the design of industrial catalysts such as glycosyltransferases,<sup>106</sup>  $\omega$ -transaminase,<sup>107</sup> and MHETase.<sup>108</sup>

In enzyme engineering, it is useful to know the binding free energy contribution from individual residues. Recently, an *ab initio* QM/MM<sup>109</sup> method was reported to obtain the electrostatic, polarization and van der Waals contributions from each residue to the activation barrier, as well as the contributions from different collective variables along the reaction coordinate to explore the possible reaction mechanism. This was achieved through a mean force integration along the free energy pathway and the reaction coordinate by analyzing the MD simulation trajectories.

For tutorial and practical guidance on the QM cluster, QM/MM and QM/MM MD multiscale simulations on biomolecules, we recommend reading recent reviews.<sup>110–112</sup>

## 2.2 Enzyme design applications

There are perennial challenges in enzyme design to identify the active site related to the reaction mechanism and fine-tune enzymes to improve their properties. The enzyme fitness landscape describes the relationship between the enzyme variants and fitness, which measures how well a given enzyme can perform a target function (Fig. 2a). However, the potential protein sequence space is vast, necessitating effective strategies to search through it and identify sequences with desired functions. Common strategies include random mutagenesis, semi-rational design, rational design, and *de novo* design.

Random mutation is conducted when structures are not available and is often combined with high-throughput screening. Hence, we will not discuss this strategy in our review. Compared to high-throughput screening, rational and semi-rational enzyme design strategies demonstrate significant promise due to their reduced cost and efficiency.

The semi-rational design strategy is based on structures and prior knowledge of enzyme functions. It constructs small libraries by performing site-directed mutagenesis on several specific residues, which are identified around the catalytic site of the enzyme.

Rational design strategies typically utilize molecular modeling and structural sampling methods to explore enzyme–substrate binding modes. Additionally, dynamic structures are considered through molecular dynamics simulations and the reaction mechanism is explored by employing quantum



**Fig. 2** Enzyme design approaches. (a) The fitness landscape map of an enzyme shows the relationship between different variants of an enzyme and their fitness (such as catalytic efficiency, thermal stability, substrate specificity, etc.). Each variant corresponds to a point on the map and the height of the point represents the fitness of the variant. (b) Directed evolution mimics the natural evolution process to improve the function of proteins through multiple rounds of random mutation, screening and selection. (c) In the semi-rational design approach, the key sites identified based on enzyme structures are mutated with saturation mutagenesis to improve the enzyme function. (d) In the rational design approach, the sites identified based on the dynamic structures and catalytic mechanism of enzyme are mutated to improve protein function. (e) *De novo* design methods are used to construct protein backbones from scratch to generate protein structures with new functions.



mechanical calculations, thereby greatly reducing the search space on the fitness landscape.

Both semi-rational and rational approaches focus on modifying natural enzymes to alter or confer new catalytic functions, while *de novo* enzyme design strategies aim to generate novel enzymes usually by incorporating the active site of the reaction into a simplified artificial protein scaffold.

There are many structure-based enzyme design/engineering studies. Here we focus on recent computer-aided enzyme design cases that were guided by semi-rational and rational design strategies to improve the enzyme properties, such as enhancing enzyme's activity, controlling regio- or enantioselectivity preferences, broadening substrate scope and altering enzyme function.

**2.2.1 Improving activities.** Crystal structures can serve as a basis for semi-rational design strategies. Several studies have reported the successful application in enhancing enzyme catalytic activity by combining site-directed mutagenesis. For example, based on the X-ray solved crystal structure and docking studies of Leucine dehydrogenase (LeuDH, EC 1.4.1.9), which can catalyze  $\alpha$ -keto acids and free ammonia to produce  $\alpha$ -amino acids, Mu *et al.* selected 6 key residues and mutated them into hydrophobic residues of different sizes for pocket reshaping.<sup>113</sup> The designed variants with double mutations increased the catalytic efficiency toward the natural and non-

natural substrates. Based on the crystal structure of flavin-dependent halogenase, Chaiyen *et al.* engineered the intermediate (HOX) transfer tunnel that connects two active sites, as a result, to reshape the tunnel, so that the engineered enzyme showed the improved catalytic efficiency (Fig. 3a).<sup>114</sup>

Multichemical state analysis (MCSA) is an enzyme design method developed for the redesign of enzymes with multiple substrates. Large structure ensembles were abstracted from MD simulation to model each of the chemical states, and library design was performed by sub-designs comprising overlapping subsets of the total designed positions, thus the sequence space was explored effectively. The enzyme sequences were optimized and a ranked list, which is based on Boltzmann-weighted sequence energies averaged over the structural ensembles, was used to generate a position probability matrix (PPM) for each sub-design. Screening a designed small combinatorial library for aminotransferase gave promising variants with up to 200-fold improvement in catalytic efficiency.<sup>116</sup>

In the absence of a crystal structure, different modeling methods can be used to generate enzyme structures. Qin *et al.* constructed the structure of L-lysine hydroxylase from *Niastella koreensis* (NkLH4) through homology modeling and achieved a 24.97-fold increase in activity for L-lysine by employing semi-rational combinatorial active-site saturation test (CAST) on four positions.<sup>117</sup>



**Fig. 3** Hotspot region identification in semi-rational design approaches. (a) Engineering the tunnel (shown in green) passing through the FADH<sup>-</sup> binding site and the tryptophan binding site. The structure is produced based on the crystal structure of flavin-dependent halogenase (FDH) (PDB ID: 7CU2<sup>115</sup>). (b) Structural modeling of Wild Type JeSTS4 by I-Tasser and AlphaFold2. (c) The two hotspot regions were identified for JeSTS4 by combining coevolution and the structural information obtained from MD simulations. Reproduced with permission.<sup>25</sup> Copyright 2022, American Chemical Society.





to industry.<sup>133</sup> Based on the crystal structure and homology models of variants, the preference toward the admirable (*R*)-enantioselectivity was achieved for both *E*- and *Z*-citrals isomers, by only introducing one or two mutations for a NADPH-dependent OYE enzyme OYE3.<sup>134</sup> Site-directed mutagenesis based on the crystal structural analysis of two stereocomplementary OYE enzymes GsOYE and BfOYE4 gave stereodivergent products.<sup>135</sup>

Cytochrome P450 enzymes are a superfamily of enzymes that are important for the synthesis of complex bioactive molecules such as natural products and drug metabolism. Based on the crystal structure, the regioselectivity of P450 BM3 was tailored to give hydroxylated derivatives at different positions of a sesquiterpene lactone compounds parthenolide (PTL) and micheliolide (MCL).<sup>136,137</sup> Based on the analysis of the crystal structures of two P450 enzymes IkaD and CftA, it was suggested that the structural difference at the polar moieties of the two enzymes accounts for the regioselectivity and chemselectivity for PoTeM,<sup>138</sup> and the regioselectivity of a P450 enzyme IkaD for a polycyclic tetramate macrolactams (PoTeM) ikarugamycin was altered by fine-tuning the catalytic pocket.<sup>138</sup>

In the search for stereocomplementary serine lipase CALB, all four stereodivergent variants of serine lipase CALB were obtained by only screening an ultra-small variant library constructed based on the MD simulated structures preferable to

the four respective stereoisomer products.<sup>139</sup> By employing a workflow combining Rosetta enzyme design and MD simulation-based free energy ranking, Delgado-Arciniega *et al.* introduced 6–8 simultaneous mutations in a ketoreductase and altered the enantioselectivity. They experimentally characterized only four variants and found three variants exhibited inverted enantioselectivity in the reduction of acetophenone-like substrates and an  $\alpha$ -keto ester, significantly reducing the experimental screening workload.<sup>140</sup>

Based on the substrate binding mode of wild type cyclohexanone monooxygenase (WT-CHMO) studied from MD simulations, we found that the substrate is sandwiched between the top or bottom of the binding site featured by two residues F434 and L437 (Fig. 5a). A single mutation at either position led to a complete reversal of enantiopreference towards 4-alkyl and 4-phenyl substituted cyclohexanones.<sup>141</sup> However, there is still room for further improvement in reversing the enantioselectivity for cyclohexanone with short substituents like a methyl or ethyl group. Therefore, we designed the F434I/L437A/T435L triple mutation to reconstruct a smaller binding pocket and achieved complete reversal of enantiopreference for cyclohexanone with short substituents.<sup>142</sup> Furthermore, we found that replacing F279, located in the second sphere near the active site and forming hydrophobic interactions with F434, with a larger residue like tryptophan, would achieve a marked improvement



Fig. 5 Mutations of Baeyer–Villiger monooxygenases (BVMOs) for improved properties. (a) Single mutation at two active residues F434 or L437 surrounding the substrate reversed the natural enantiopreference of WT-CHMO.<sup>141</sup> The crystal structure of CHMO (PDB ID: 4RG3<sup>143</sup>) was used. (b) Engineering the second sphere residue F279 into smaller residues like Valine reversed the enantioselectivity of CHMO toward diverse substrates.<sup>125</sup> (c) Expanding substrate scope of PAMO by engineering the bulge region that is present in PAMO but absent in CHMO.<sup>144</sup> (d) Improving the thermal stability of CHMO by creating additional disulfide bonds between two adjacent cysteine residues.



in enantio- or regioselectivity across a wide range of substrates. Conversely, replacing it with smaller residues would achieve a complete reversal of enantiopreference (Fig. 5b).<sup>125</sup>

For the design of terpene synthases, the water flow regions identified from MD simulations provided guidance on reshaping the active site of a sesquiterpene synthase to catalyze the synthesis of a valuable terpenoid product while avoiding the hydroxylated product.<sup>145</sup> A single mutation of another sesquiterpene synthase, pentalenene synthase, diverted the reaction pathway to give different products, because of the reshaped binding pocket disclosed by molecular docking and MD simulations.<sup>146</sup>

QM/MM and MD simulations disclosed the reversed regioselectivity of thermostable CHMO (TmCHMO) for 4-phenyl-2-butanone to give the abnormal product attributed to the conformational changes in the Criegee intermediate and transition states in the reaction pathway.<sup>147</sup> MD simulations and QM/MM calculations elucidated the catalytic mechanism of PAMO toward its native substrate phenylacetone and the alkyl migration mechanism of the Criegee intermediate decay.<sup>148</sup> Furthermore, based on MD simulations of PAMO, we proposed the requirements for a catalytic pocket favourable for non-native linear substrate 2-octanone, which provides structural insight for further engineering the enzyme to accommodate linear substrates.<sup>149</sup> QM cluster calculations disclosed that the change in the chirality of the Criegee intermediates and transition states accounts for the regioselectivity so as to give the normal or abnormal products by the WT-TmCHMO and its variants, respectively.<sup>150</sup>

**2.2.3 Broadening the substrate spectrum.** Bayer–Villiger monooxygenases (BVMOs), comprising many subfamilies of enzymes depending on their respective substrates, such as cyclohexanone monooxygenases (CHMO), phenylacetone monooxygenase (PAMO) and cyclopentanone monooxygenase (CPMO), catalyze the insertion of an oxygen atom in ketones to give esters or lactones. There are universal hotspot regions in different Bayer–Villiger Monooxygenase (BVMO) subfamilies that are responsible for the enzymes' properties such as substrate scope, enantio- and regio-selectivities and stability.<sup>151</sup>

PAMO is a thermostable enzyme with high industrial value. However, it has a narrow substrate acceptance range compared to CHMO. Structural comparison showed a bulge (S441–S444), which is present in PAMO, but absent in CHMO (Fig. 5c). Deleting the bulge in PAMO turned the enzyme into a phenylcyclohexanone (PCHMO), which showed a broadened substrate spectrum.<sup>144</sup> Saturation mutagenesis of the bulge region in PAMO using codon degeneracy was conducted and variants that accept 2-aryl cyclohexanone were attained.<sup>152</sup> Mutating a second sphere residue P440 around the bulge achieved the acceptance of a range of substrates.<sup>153</sup> In another work, structure-guided rational design altered the functionality of CHMO to allow it to reduce a range of substituted aromatic  $\alpha$ -keto esters. With high catalytic activity and stereoselectivity. The created reductive activity was attributed to shortened reaction coordinates favourable for hydride transfer in the ketoreductase-like variants in comparison with

the WT enzyme, as observed from docking and MD simulations.<sup>154</sup>

The types of tunnels in metalloenzymes catalyze the reductive or oxidative transport and positioning of small gaseous substrates such as H<sub>2</sub>, N<sub>2</sub>, NH<sub>3</sub>, CH<sub>4</sub>, O<sub>2</sub>, CO, CO<sub>2</sub>, *etc.* dictates the substrate preference, and therefore reshaping the gaseous tunnels would affect substrate selectivity and enzyme functions.<sup>155</sup> The substrate tunnel of a soluble methane monooxygenase (sMMO) hydroxylase has been revealed based on different approaches such as crystallography, MD simulations and mutagenesis of the tunnel-lining residues.<sup>156</sup>

Engineering the composition residues lining the access tunnel of P450<sub>B<sub>5</sub>P</sub> changed the substrate preference.<sup>157</sup> Hotspot identified by MD simulations of haloalkane dehalogenase for the catalytic transformation of linear and branched substrate disclosed the requirements for substrate specificity.<sup>158</sup>

**2.2.4 Tailoring enzymes' function.** The biosynthetic pathway of many enzymes involves multiple reaction steps due to the promiscuity of the enzymes. Engineering enzymes by reshaping the active sites may control the reaction to change the product distribution or change enzyme functions.

Ergothioneine sulfoxide synthase from *Candidatus Chloracidobacterium* (EgtB<sub>Cth</sub>) possesses both EgtB- and Egt1-type activities with the EgtB-type feature more prominent than the Egt1-type; however, the latter is more industrially valuable. By leveraging active site information from EgtB<sub>Cth</sub> crystal structures, EgtB<sub>Cth</sub> variants were designed using Rosetta enzyme design<sup>159</sup> and three mutants were tailored to exhibit Egt1-type characteristics.<sup>160</sup>

Comparison of the key active-site residues in the crystal structures of MPD and MDD that are involved in the bifurcated mevalonate (MVA) pathway, combined with sequence analysis, disclosed the key active-site residues that confer substrate specificity, which facilitated distinguishing enzyme classes involved in two MVA metabolic pathways.<sup>161</sup> In another example, sequence comparison and structural analysis of the homology models of two homologous maize terpene synthases TPS4 and TPS10 disclosed the difference in the key active site residues that determined product specificities, and combined mutation of the different residues in the first and second sphere turned TPS4 into TPS10.<sup>162</sup>

5-Methylene-3,5-dihydro-4*H*-imidazol-4-one (MIO)-enzyme family comprises two classes of enzymes with different functions, *i.e.* aromatic amino acid ammonia lyases (ALs) and 2,3-aminomutases (AMs). Based on the crystal structure of an AL, the substrate binding tunnel of AM was engineered, and the resulting variant showed enzyme function of AL.<sup>163</sup>

Based on the homology model of a sesterterpene synthase SmTS1 and multiple sequence alignment, engineering the substrate binding site residue displayed the function of diterpenes synthase.<sup>164</sup> Similarly, in a semi-rational design based on the crystal structure of a diterpene synthase VenA, VenA was changed to a sesterterpene to accommodate larger substrates.<sup>165</sup>

**2.2.5 Changing the pH-activity profiles.** Modifying the polarity of amino acids near the substrate binding site can



significantly impact the pH-activity profile of an enzyme. Numerous studies have shown that changing the polarity of the catalytic site residues can shift the optimal pH, as exemplified in engineering xylanase,<sup>166</sup> glycosidase,<sup>167</sup> phytase,<sup>168</sup> amylase,<sup>169,170</sup> dehydrogenase<sup>171</sup> and phytase.<sup>172</sup> Further MD simulations may provide insight into the effect of mutations on the dynamic residue-residue interaction network in the active site and hence the pH-activity.

The surface charge of enzymes also plays a crucial role in determining their pH-activity profile.<sup>173–175</sup> For example: the NADH Oxidase from *Bacillus subtilis* exhibits maximum activity at pH 9.0, whereas the pH of its coupled enzyme dehydrogenase is close to 7.0, making the practical industrial application challenging.<sup>176</sup> Introducing negatively charged residues on the enzyme surface using Rosetta design lowered the optimal activity pH to 7.0.<sup>177</sup> In industrial production, vanillin is produced from waste biomass resources and then vanillin is converted to vanillic acid by vanillin dehydrogenase (VDH) under alkaline conditions; however, VDH displayed poor activity at alkaline pH. By mutating non-conserved, negatively charged surface residues to positively charged arginine, the optimal activity was shifted from pH 7.4 to pH 9.0.<sup>171</sup> The comparison of the crystal structures of two SGNH family esterases CrmE10 and AlinE4 showed that the two enzymes have different electrostatic potentials on enzymes' surfaces. Engineering the charge of CrmE10 surface residues from acid to basic improved the alkaline adaption and therefore increased the enzyme's activities (Fig. 6).<sup>178</sup>

**2.2.6 Improving thermostability.** The most common secondary structures of proteins are alpha helices, beta sheets, beta turns and loops, among which alpha helices are more tolerant to multiple mutations than beta sheets,<sup>179</sup> and hence engineering helices would be more liable than engineering beta strands. An enzyme engineering strategy to improve the thermostability of enzymes is replacing the glycine or proline in alpha helices into alanine, which is beneficial to improve the thermostability of helices and hence the overall enzyme thermostability.<sup>180</sup> Zhou *et al.* improved the thermostability of an alkaline pectate lyase (PelN) from *Paenibacillus sp.* by replacing glycine at position 241 on a helical structure with alanine or valine. Additionally combining mutations at positions on beta sheets and the resulting double mutant K93I/G241A retained the high thermostability with improved enzyme activity,<sup>181</sup> which potentiates its industrial applications.

Highly flexible residues may be responsible for protein unfolding and denaturation, leading to decreased thermostability. The highly flexible residues in levansucrase were identified by root mean square fluctuation (RMSF) for MD simulations of the enzyme crystal structure and these residues were mutated to improve the thermostability.<sup>182</sup> The difference in free energy ( $\Delta\Delta G$ ) between the mutant and wild-type enzyme was calculated to assess the stability of mutants and experimental evaluation shows that the designed K82H/N83R mutant is more thermostable than the wild type. A similar design strategy combining MD simulations and  $\Delta\Delta G$  calculations has been used to guide the design of carrageenase,<sup>183</sup>



Fig. 6 Effect of surface electrostatic potential on activity. (a) Protein surface electrostatic potential of two homologous enzymes of the esterase family CrmE10 (top right, PDB: 7C23<sup>178</sup>) and AlinE4 (bottom right, PDB: 7C82<sup>178</sup>). (b) Superimposition of CrmE10 and AlinE4 with the key polar residues on the surface shown in stick mode. (c) The pH/activity profile of CrmE10 and AlinE4.



lipase,<sup>184</sup> and tyrosinase,<sup>185</sup> which attained variants with improved thermostability.

Disulfide bonds can reduce the configurational entropy of the unfolded polypeptide to stabilize the structures of protein.<sup>186</sup> Disulfide bonds can be introduced at non-catalytic residues using MODIP,<sup>187</sup> DbD2<sup>188</sup> or BridgeD<sup>189</sup> server and the effect of designed disulfide bonds on thermostability can be evaluated by calculating  $\Delta\Delta G$  between the designed mutants and the WT enzyme. Two disulfide bonds (S61C–S115C and E190C–E238C) were designed for *Rhizopus oryzae* lipase (ROL) to rigidify the enzyme, and the thermal stability of the enzyme successfully increased by 5.0 °C and 6.9 °C, respectively.<sup>190</sup> The introduction of disulfide bonds near the binding site of divalent cations (*e.g.* Ca<sup>2+</sup>, Mg<sup>2+</sup>) effectively improved the thermostability of polyethylene terephthalate (PET) hydrolase.<sup>191</sup> A simultaneous improvement of stability against oxidation of and thermostability of CHMO was achieved by introducing new disulfide bonds guided by a computational study<sup>192,193</sup> (Fig. 5d). In some enzymes, cysteine and methionine are liable to be oxidized and therefore hamper enzyme activity. Mutating the cysteine and methionine into non-polar residues or serine may enhance oxidative stability and hence thermal stability.<sup>194</sup>

### 3. Machine learning-accelerated enzyme design

Molecular dynamics simulations and the QM/MM method provide valuable insight for atomic level conformational dynamics mechanisms, and the enzymatic reaction mechanism; therefore, they have been widely used to explore conformational space and structure–function relationship. Furthermore, the advances in computer hardware along with the development of accurate force fields and highly efficient sampling methods have enabled employing molecular simulations for enzyme design.<sup>195–198</sup> For example, modulating the protein stability guided by MD<sup>199</sup> and enzyme engineering for natural product biosynthesis aided by QM/MM.<sup>200</sup>

With the dawn of the big data era, various biological databases have become available and machine learning methods have been applied in enzyme engineering.<sup>21,201–205</sup> The advent of a tremendous amount of data from the literature or databases enables us to build machine learning models and implement them into the screening protocol, for example, machine learning guided protocols were reported to predict the properties of mutants so as to reduce the screening demands by traditional experimental high throughput screening.<sup>206,207</sup>

Machine learning (ML) benefits from molecular modeling and accumulated experimental data. It has been implemented in molecular modeling based on atomistic MD and quantum mechanics and facilitated the effective multiscale or coarse-grained modeling, and therefore enabled exploration of the vast space of functional enzyme sequences speeding up the screening of functional enzyme variants.<sup>208–210</sup> The three-pronged atomistic simulations, machine learning and

experimental validation, can be synchronized, functioning just like a troika, and would speed up the efficient screening of potential mutants in the enzyme design protocol, with enhanced accuracy in predicting the effect of mutations.

To enable interdisciplinary collaboration between experimentalists and computational scientists, it is essential to understand how computers store and process data in a way that is understandable by both parties to facilitate collaborations.<sup>211</sup>

In this section, we will introduce the data processing methods, including the methods of generating descriptors from small molecules and proteins, and utilizing various databases as the data resources for machine learning. Model building and evaluation methods will also be introduced. Finally, the latest machine learning research on enzyme engineering will be reviewed.

#### 3.1 Descriptors for small molecules

To retrieve meaningful patterns and rules in machine learning, the databases need to be processed and converted into numerical descriptors. For example, molecular descriptors representing molecular features are developed to predict the biological activities and screen potential lead compounds in QSAR.<sup>212</sup> These molecular descriptors are classified as 1D global property, 2D planar features or 3D stereo features.

**3.1.1 Descriptor selection and combination.** Feature selection is crucial for machine learning, and the molecular representations should not only capture the diversity of chemical space, but also distinguish the subtle differences among molecules.<sup>213</sup> The descriptors should be simple while retaining key information and consistent and interpretable to assure that the pattern learned from the model would reflect the meaningful relationship between the descriptors and properties rather than being affected by noise.

Removing irrelevant descriptors may improve the accuracy of the prediction to develop robust models. Khan *et al.* reviewed descriptor selection methods in different drug design cases,<sup>212</sup> including the filter method that gradually deletes the low-score features by calculating relevance scores of the descriptors and Wrapper method that gradually deletes descriptors guided by the errors in a validation subset using a support vector classifier.

**3.1.2 Global property descriptors.** Global property descriptors are referred to as physicochemical descriptors of small molecule substrates, which are estimated based on the 2D structure of the molecules. *e.g.*, those properties in Lipinski's rule of five including molecule weight, Log $P$ , the number of H-bond donors/acceptors, *etc.* which are essential properties for drug's pharmacokinetics and hence have been widely used in drug development.<sup>214</sup> In addition, atom-type counts, bond-type counts, and molar refractivity are also global descriptors. It should be noted that most of the global descriptors lack information on the molecular structure or atom connectivity.

**3.1.3 Quantum-chemical descriptors.** Quantum-chemical descriptors including atomic charges, molecular orbital energies, Frontier orbital densities and molecular polarizabilities



are also used in machine learning to predict electrostatic interactions, chemical reactivities, physicochemical, biochemical or pharmaceutical properties of molecules.<sup>215</sup> Combining QM descriptors in machine learning may predict molecular interaction fields and chemical reactivities more accurately.<sup>216</sup>

**3.1.4 Molecular fingerprints and graph descriptors.** The chemical structure features and atom connectivity require 2D representation of molecules (Fig. 7). String representation approaches such as SMILES<sup>217</sup> and InChI<sup>218</sup> were used to store the 2D information of molecules, which can efficiently represent molecular graphic information using standardized and machine-readable formats.

Additionally, molecular structures are compressed into library-based 2D representation by a molecular “fingerprint”, which projects the structure information of molecules into binary codes, with each bit representing molecular structure features or the presence/absence of certain structures. The binary representations such as MACCS<sup>219</sup> are compatible for data storage and also liable for comparing the similarity among molecules.

In contrast to library-based fingerprint representation, circular fingerprints<sup>220</sup> such as Morgan fingerprints, extended-connectivity fingerprints (ECFPs) and functional-class fingerprints (FCFPs) take into consideration of the local environment of molecules to generate a bit vector. For example, the Morgan fingerprint with a radius of 2 considers the connectivity of each atom to other atoms which are linked to the first atom by up to two chemical bonds; it assigns a value of 1 if such a

neighborhood is present in the molecule, otherwise, it assigns 0. These fingerprint methods have been implemented in RDKit toolkits.<sup>221</sup> The vectors generated by fingerprint methods are high dimensional and sparse, and often bring about the issue of bit collision. Google Inc. compared the quality of word representations in vector space for a very large dataset in a word similarity task and reported two model architectures with promising prediction accuracy and efficiency.<sup>222</sup>

Convolutional neural network and natural language processing (NLP) techniques have been used in molecular graphic representations. Fuller and Turk *et al.* reported a Mol2vec algorithm<sup>223</sup> to represent the substructures of a molecule as word vectors and the whole molecule as a sentence. Thus each substructure in the molecule can be more efficiently represented.

Molecular structures can also be represented by molecular graphs. With the development of the graph neural network, each atom in a molecule can be considered as the nodes in graphic structures and the connectivity among atoms are defined as edges. The graphic frame can describe the complicated relationship among the substructures by graphs. Utilizing the graph neural network (GNN), molecular graph descriptors have been widely used in predicting drug-target interactions.<sup>224–227</sup>

To evaluate the catalytic efficiency of enzymes, it is important to estimate the enzyme-substrate interactions as well as enzyme-catalyzed reaction kinetics. Skoraczynski *et al.* developed binary classification models for predicting the reaction



Fig. 7 Machine learning for enzyme design. (a) The data used in enzyme engineering modifications mainly consist of small molecules and protein descriptors. (b) Some commonly used algorithms in regression, classification and clustering models. (c) Evaluation metrics in machine learning models. (d) The challenges in achieving a predictive ML model: the imbalanced distribution of data requires manual curation, *i.e.* some error data must be corrected prior to data preprocessing to assure the quality of prediction models; the issues of model underfitting/overfitting addressed by hyperparameter tuning during model optimization.





On the other hand, a number of approaches have been developed for retrieving sequence-based descriptors from amino acid composition (Fig. 7). A commonly adopted natural language processing (NLP) method is one-hot encoding. One-hot encoding represents the sequence by an array of a binary vector (0 or 1) to indicate the presence of a certain type of 20 amino acids at each position of the sequence. Another NLP method is *n*-gram encoding, where a protein sequence is broken into segments of size *n* to represent the local combinations of amino acids. These segments are then stored in an “*n*-gram” dictionary, which can be used to calculate the similarity among mutant strains. Other language embedding models like ProtVec also treat protein amino acid sequences as a series of “words” and map each amino acid to a vector representation in a high-dimensional space.<sup>260</sup> ProtVec can be easily combined with the aforementioned Mol2vec.<sup>223</sup>

The above vectors capture the similarity and functional relevance among amino acids. Another method position-specific scoring matrix (PSSM)<sup>261</sup> considers homology information among sequences, where each element represents the frequency of a certain amino acid (or base) at a given position across different sequences. These frequencies are calculated through multiple sequence alignments and are then converted into scores or probabilities. PSSM embodies information on conservation and variation of specific amino acids at particular positions in the sequences.

Certain overall physicochemical properties are related to enzyme functions, but their interpretability needs scrutiny. Descriptors based on protein sequences typically reveal fundamental characteristics of enzymes. The amino acid composition indicates the relative proportions and frequencies of different amino acid types, which are related to enzyme diversity and specificity. Conservation describes amino acid residue conservation across different species, which reflects the enzyme's evolutionary history and functional conservation. These descriptors could be used in machine learning for different tasks such as predicting substrate scope, enzyme functions, and classifying enzymes according to their properties.

**3.2.2 Structure-based descriptors.** In addition to sequence-based descriptors, structure-based features have also been used in enzyme engineering.<sup>262</sup> Compared to sequence-based machine learning, structure-based machine learning is more computationally expensive. Structure-based machine learning requires the 3D structures of enzymes to generate the inputs (Fig. 7). The advent of protein structure prediction methods such as AlphaFold and RosettaFold has enabled the acquisition of the protein 3D structures. However, it is still very challenging to acquire large-scale complex structures for a large library of predicted variants, due to the restriction by the demanding computational resources for large-scale simulations and the considerable expertise required for analyzing the predicted structures. Laio *et al.* reviewed the applications of unsupervised learning techniques for the analysis of molecular simulation data, by transforming trajectory data into low dimensional collective variables; thus, the “raw” Cartesian coordinates are converted into compact numerical

representations that preserve relevant information of the simulation trajectory.<sup>263</sup>

The research of using structure-based descriptors in machine learning for enzyme engineering is relatively limited compared to that of the sequence-based approaches. Geometric descriptors such as atomic distances, angles, and dihedrals<sup>264</sup> can be used to describe the spatial relationships among active site residues that are functionally important. These features can be represented by distance matrices and used as the inputs to construct machine learning models (*e.g.* sPairs, an AA-index-based aa pairwise contact potential<sup>248</sup> and residue-residue contact map<sup>265,266</sup>). In addition, enzyme structure representations by space filling curves (SFCs) were reported in classification tasks for evaluating substrate selectivity.<sup>267</sup>

Structure-based features can reflect the substrate–enzyme interaction information. However, it is worth noting that replacing sequence-based features by structure-based features would not necessarily lead to improved predictive performance. In practical applications, these enzyme-related descriptors are often combined to construct comprehensive models to predict enzyme properties. For instance, sequence-based descriptors can be combined with structure-based descriptors to enhance model accuracy for predicting enzyme activities. Protein sequence descriptors have been combined with small molecule structural descriptors to model the interactions between compounds and proteins.<sup>268,269</sup> Incorporating protein structural features in the models may further improve the accuracy and interpretability of the predictive models in the design and optimization of biocatalysts.

### 3.3 Databases

Machine learning algorithms highly rely on the quality of the training dataset (Fig. 7). It is important to resource the enzyme databases for the applications of machine learning. There are numerous publicly available databases online with vast amounts of data. The commonly used databases related to enzyme engineering are summarized in Table 2, encompassing protein sequence database, structure database, protein–ligand interaction database, reaction mechanism database, enzyme property database, *etc.*

It is crucial to craft a dataset to tailor it for specific research objectives and computational resources. For instance, AlphaFold2 achieves high accuracy in predicting protein structures, yet current predictions are limited to the apoprotein and do not include interactions with ligands, cofactors, or metal ions. It is reported that the AlphaFill algorithm<sup>270</sup> can be used to transplant cofactors and ions from experimentally determined structures into the prediction models by AlphaFold2, based on the sequences and structural similarity. However, direct transportation of the ligands or metal ions from known structures has limitation for certain enzymes for which the crystal structures have not been resolved and the sequence identity among species is very low, *e.g.* terpene synthases, an important enzyme family to industry for the biosynthesis of natural products.

Typically, the effect of mutations around the catalytic sites of enzymes is predicted by analyzing their configurations.









Table 3 (continued)

Task	Dataset	Model and framework	Availability	Citation
Predict sequence EC number	Sequences from the UniProt database	CLEAN, a contrastive learning model using ESM-1b embeddings	Code/Web server available	<i>Science</i> , 2023, <b>379</b> , 1358–1363
Generate protein sequences for specific reactions	16 898 sequences from the UniProt database	Generative adversarial network (GAN)	Code available	<i>Nat. Mach. Intell.</i> , 2021, <b>3</b> , s324–333
Generate protein sequences for specific reactions	281 million sequences from the UniParc, UniProtKB, Pfam and NCBI taxonomy databases	Transformer	Code available	<i>Nat. Biotechnol.</i> , 2023, <b>41</b> , 1099–1106
Generate protein sequences for specific reactions	Luciferase-like protein sequences from InterPro	Variational autoencoder (VAE)	Code available	<i>PLoS Comput. Biol.</i> , 2021, <b>17</b> (2), e1008736
Generate protein sequences for specific reactions	MSAs for different protein families generated by PFAM and HMMER	VAE	Code available	<i>Nat. Commun.</i> , 2023, <b>14</b> (1), 2222

The applications of machine learning in enzyme design can be classified into the improvement of properties for specific enzymes or the development of general predictive models. Considering the interest of different readers, we will discuss these two types of tasks respectively in the following section.

### 3.5.1 Practical ML tasks

**3.5.1.1 The predictivity of ML for small datasets.** Building a reliable machine learning model requires a large amount of data. However, for specific enzymes, usually only a limited number of experimental data points are available such that the predictivity of the ML model developed from the small database is insufficient.

By iteratively evaluating ML-predicted sequences and feeding the new experimental data points with improved properties into the training set, the predictivity of the model can be improved. For example, Ohnishi *et al.* used Bayesian optimization to screen a Xylene monooxygenase (XylM) variant library. The library was generated by codon-randomization at five residue positions located close to the ion coordination site at the catalytic site.<sup>281</sup> The iterative predictions after two rounds gave a mutant that increased 3-methylsalicylic acid production by 15 fold compared to that of WT-XylM. In another work, Liu *et al.* disclosed that five positions located at the substrate access channel of the P450 enzyme are critical for the loop stability and hence enzyme's activity.<sup>282</sup> 165 variants were created by simultaneous saturation mutagenesis on these five positions for machine learning. The representation of the sequences was generated by AAindex. The best mutant A86T/T91H/M108G/A109M/T111P showed a 15-fold improvement in the activity compared to the WT.<sup>283</sup>

Transfer learning starts with pre-training a model on a large dataset, then the model is fine-tuned to generate a new model for a smaller dataset, thus the knowledge learned from the large dataset is transferred to improve predictivity performance of the new model for the small dataset. For example, Engqvist *et al.* constructed a dataset of 3 million optimal growth temperatures (OGTs) from the BRENDA database to train a model called DeepET.<sup>284</sup> DeepET is based on a residual neural network using the fast one-hot encoding method to retrieve enzyme thermal adaptation features from enzyme sequences. Transfer learning was then employed to predict two temperature properties related to the thermal stability, *i.e.* optimal catalytic temperature  $T_{opt}$  and melting temperatures ( $T_m$ ) for small datasets. DeepET showed more accurate predictions on these two datasets compared to other feature extraction methods like iFeature<sup>285</sup> and UniRep.<sup>286</sup> However, it is worth noting that if the small dataset is very divergent from the large dataset, the knowledge transferred may not be relevant, leading to poor performance.

In the case of an extremely small data set (9 single mutation variants). Frédéric Cadet *et al.* reported a sequence–activity relationship method called innov'SAR and improved the enantioselectivity of an epoxide hydrolase guided by machine learning. This method numerically encodes the protein sequence by AAindex and then applies the Fast Fourier transform to convert the encoded sequence into protein spectra. The protein spectra



and protein activity are used as learning datasets to build a partial least squares regression (PLS regression) model to predict the activity.<sup>287</sup> In addition, by adding 28 mutants into the training set, the prediction model includes some information about the epistasis between mutations, thereby improving the accuracy of the prediction model.

**3.5.1.2 Impact of dataset construction on predictivity.** Machine learning can guide directed evolution in exploring sequence space. Would the composition of training data affect the predicted results of machine learning?

Guided by ML models, two series of directed evolution for Sortase A were performed. The dataset for one ML model contained a highly positive variant 5 M, whereas the other excluded 5 M.<sup>274</sup> The ML models were trained using the Bayesian optimization method and used to evaluate the probability of a variant being positive and promising variants (with activity 2.2–2.5 times higher than that of 5 M) were predicted by both ML models. However, it is worth noting that the regions for advantageous mutation on the sequence fitness landscape identified by the two ML models are different, indicating that diverse positive variants may be attained by using divergent datasets.

**3.5.1.3 Structure-based machine learning.** Structure-based representations of proteins have been developed to describe the substrate–enzyme interactions. For example, Ran *et al.* reported a deep learning framework EnzyKR, which used complex structures constructed from docking to encode the hydrolase–substrate interactions between hydrolases and the enantiomer products,<sup>288</sup> and showed good performance in predicting activation free energy as well as in predicting enantiomeric excess ratios. Using structure-based ML enzyme engineering, Alper *et al.* obtained a PET hydrolase variant that was generated by combining the triple mutation predicted by ML and double mutation from the scaffold, which showed promising hydrolytic activity and thermostability.<sup>289</sup> The machine learning architecture used in the study employed the 3D CNN method MutCompute<sup>290</sup> proposed by Ross Thyer.

Lu *et al.* represented a structure-based graph convolutional network that denotes the protein–ligand interaction energetics (generated using Rosetta<sup>159</sup>) and successfully predicted the specificity of proteases for two noncanonical substrates.<sup>291</sup>

The dynamic properties of enzymes are crucial for their activities. Tidor *et al.* selected descriptors from 68 geometric parameters including atom distances, planar angles, dihedral angles, *etc.* that represent the local conformation of the active site and accurately predicted the reactivity of ketol-acid reductoisomerase (KARI).<sup>292</sup>

### 3.5.2 General ML models

**3.5.2.1 Predicting enzyme–substrate pairs.** Machine learning has been widely used for predicting protein–ligand interactions based on the datasets constructed from various databases. However, most databases only contain the active substrates that enzymes can catalyze (positive instances) and lack data on non-active substrates (negative instances). The imbalance of

the dataset can lead to models with poor generalization ability. To address this issue, Alexander Kroll *et al.* reported an enzyme–substrate pair prediction model by constructing a database composed of the experimentally validated enzyme–substrate pairs derived from the gene ontology (GO) annotation database and randomly generated negative samples similar to the real substrates using data augmentation.<sup>293</sup> They constructed a gradient-boosted decision tree model for predicting enzyme–substrate pairs. By only using the sequence-based representation for enzymes and GNN-generated fingerprints for small molecules, a general model was achieved with high accuracy that can be applied across enzyme families and a broad range of small molecules.

Hu *et al.* designed a dataset comprised of 606 555 corresponding enzyme–substrate pairs, with the ratio of negative data to positive data around 10 times. They developed a positive unlabeled learning-based enzyme promiscuity prediction (PU-EPP) model for predicting the substrate promiscuity and specificity by extracting substrate features using GNNs and encoding protein sequences using continuous bag-of-words. The model showed good robustness on the test set and successfully identified 15 new enzymes for Mycotoxin Detoxification. It also allowed us to identify the important key residues attributed to the catalytic activity of the enzyme.<sup>294</sup>

**3.5.2.2 Predicting catalytic efficiency.** Binding free energy and reaction energy barrier are closely related to enzymes' catalytic efficiency. However, the intricate and diverse catalytic mechanisms of enzymes pose challenges, especially for those with unknown structures particularly, the experimental and computational simulations methods such as QM/MM or QM/MM MD on these properties are expensive, which has limited their applications in evaluating the large-scale mutations to screen highly efficient enzymes.

$K_M$  (Michaelis constant) and  $k_{cat}$  (turnover number) are two important parameters directly related to the catalytic efficiency of enzymes and therefore it is important to predict these properties to evaluate the effect of mutations.

Nielsen *et al.* reported a deep learning package DLKcat for predicting genome-scale  $k_{cat}$ , using an enzyme-constrained metabolic model which is solely based on substrate structures and protein sequences, combining a graph neural network (GNN) for substrate molecule graphs and a convolutional neural network (CNN) for extracting protein n-gram properties.<sup>268</sup> DLKcat has been used to predict the enzyme activity of  $\beta$ -ketothiolase<sup>295</sup> and thiolase.<sup>296</sup>

Kroll *et al.* predicted  $K_M$  using molecular fingerprints as a numerical representation of substrate molecules.<sup>297</sup> They further predict  $k_{cat}$  values for natural reactions of wild-type enzymes taking into consideration of numerical fingerprints for substrates and products, representing enzymes using transformer networks.<sup>298</sup> The model is able to make meaningful predictions for enzymes that are less than 40% homologous to the data in the training set.

Wang *et al.* presented a model DeepEnzyme for predicting  $k_{cat}$  values based on the 3D structures of the proteins. By





Fig. 8 Workflow of ALDELE. The model architecture takes protein sequences and substrate SMILES as inputs and processes them through five toolkits to produce features. These toolkits include RDKit and SMILES for compound inputs, "Words" for protein sequences, PSSM for protein sequences, and protein structure-based features. A two-phase attention NN is applied to extract two sets of vectors representing the protein sequence or ligand. A multi-layer perceptron (MLP) is used for prediction. Reproduced with permission.<sup>280</sup> Copyright 2024, American Chemical Society.

leveraging the features from both sequences and 3D structures, the DeepEnzyme model achieved improved prediction than DLKcat on the performance for those with low homology with the training set.<sup>299</sup>

We recently developed a deep learning-based workflow ALDELE.<sup>280</sup> This workflow includes five toolkits (Fig. 8): (1) NN representation of substrates based on whole compound properties, (2) GNN representation of substrates based on molecular graphs, (3) CNN representation of proteins based on N-Gram vectors, (4) CNN representation of proteins based on PSSM, and (5) CNN representation of protein structure-based features. The comprehensive toolkits allow customized combination of the physicochemical and graphic properties of substrates, with the sequence, evolutionary and structural information of enzymes, for predicting the interactions between enzymes and substrates. Benchmark studies for multiple datasets including a  $k_{\text{cat}}$  dataset comprising 16 838 enzyme-substrate interaction pairs show the accuracy of ALDELE for predicting the biocatalytic activities of enzymes.

**3.5.2.3 Annotating enzyme function.** Protein function prediction is critical for discovering and developing new biocatalysts. Following the sequence-structure-function paradigm, the protein sequence dictates the spatial structure and functions of proteins.<sup>300</sup> Protein sequences, structures, functions, and protein-ligand interactions have been deposited in many

databases such as Uniprot,<sup>301</sup> however, the functions for a large number of newly discovered sequences have not been denoted.

Enzymes can be classified by the Enzyme Commission (EC) number, using a coding system consisting of four digits representing the reaction type, substrate type, reaction type, and specific enzyme order. Estimating the EC number of a new sequence allows predicting the function of the enzyme. Several machine learning models have been developed to predict EC numbers, as well as to predict the related substrates and products, among which the random forest and  $k$ -nearest neighbour-based model combining the enzyme sequence and the structural information of substrates and products was shown to be able predict almost all types of reactions.<sup>302</sup>

The contrastive learning-enabled enzyme annotation (CLEAN) method was trained on the UniProt database to assign enzymes' EC number and functions.<sup>303</sup> The method used the Euclidean distance as a metric to reflect the similarity in enzymes' functions by embedding sequences into numerical vectors and was further validated in the uncharacterized halogenase database.

Additionally, a comprehensive review on the protein representation learning methods by language models since 2015 was reported by Doğan *et al.* They evaluated their performance in identifying protein functions by four benchmarking tasks.<sup>304</sup> *e.g.* The function of proteins is annotated using gene ontology



(GO),<sup>305</sup> which contains the information on molecular function, cellular components and biological processes.

Critical assessment of functional annotation (CAFA) is a project where the performance of different learning and representation methods for predicting the GO-annotated functions of target proteins are benchmarked by the accuracy compared with later acquired protein functions.<sup>306</sup> NetGO 2.0 utilizes protein information from various resources to predict the function of annotated proteins, and achieved top performance in CAFA4.<sup>307</sup> Furthermore, NetGO 3.0 integrates self-supervised protein language model (ESM)-1b embedding to represent protein sequences combined with logistic regression (LR-ESM) and has shown improved capability in predicting protein functions.<sup>308</sup>

Protein function prediction was initially based on the assumption that homologous proteins would share similar functions. However, this approach has obvious limitation with the presence of abundant distant and orphan proteins. Hence data-driven machine learning and deep learning techniques based on various representations of sequence, structure and interaction features have emerged. Dhanuka *et al.* presented a comprehensive review and compared feature-based machine learning and algorithm-based methods for protein function prediction.<sup>309</sup>

**3.5.2.4 Generating functional sequences.** Based on known functional proteins, mutagenesis and selection are commonly employed techniques for generating novel sequences with admirable functions. However, due to the vast sequence space of proteins, it remains challenging to predict sequences with new functions, which necessitates directly generating new sequences with admirable functions from the raw sequences.

Insights gained from sequence variations would provide insights on directed evolution. The advances in machine learning techniques enable deep generative models such as generative adversarial networks (GANs), transformers and variational autoencoders (VAEs) to be used to explore protein sequence space efficiently so as to generate protein sequences with specific functions.<sup>310</sup>

The ProteinGAN model, based on self-attention GAN, generates new protein sequences with natural-like functions.<sup>311</sup> It has been successfully applied to generate soluble and catalytically active sequences of malate dehydrogenase, demonstrating the ability of the neural network architecture to generate highly diverse sequences by learning intricate evolutionary dependencies between amino acids and generalizing across the protein sequence space.

Furthermore, the language model ProGen<sup>312</sup> was developed, trained on 280 million protein sequences using a transformer-based self-attention neural network architecture. The generated artificial sequences showed similar functions to natural proteins from diverse families, as demonstrated in the cases of chorismite mutase and malate dehydrogenase.

VAE models trained on 70 000 oxidoreductases were used to generate new bacterial luciferase. The comparison of VAE models trained on aligned sequences and raw sequences

showed that both models are able to capture the amino acid pattern of the enzyme family, whereas the former is able to better capture the long-distance features inferring their constraints on the protein functions.<sup>313</sup>

Latent generative landscape (LGL) was created using VAE sequence space, enabling flexible exploration of diverse protein functional space without labeling, guiding generative protein design and providing insights into evolutionary fitness and functional diversification.<sup>314</sup>

**3.5.2.5 De novo design of artificial enzymes.** In contrast to traditional enzyme design by directed evolution of native enzymes, the *de novo* design of enzymes with new functions from scratch is still in the infant stage, and new methods are rapidly emerging.<sup>15,315,316</sup>

The Rosetta *de novo* enzyme design protocol has been widely used in generating protein scaffolds since it was first reported over a decade ago.<sup>159,317–319</sup> These design cases mostly rely on existing protein scaffold templates from nature. By transplanting the natural enzyme active sites into other unrelated protein structures and redesigning the amino acid sequence around the substrate, the goal is to stabilize the conformational energy of the enzyme's reaction intermediate state. However, due to limitations in energy functions and design accuracy, the designed enzymes often do not match the activity of natural enzymes.<sup>320</sup>

To address the fitness issue of the designed scaffold, Sarel Fleishman *et al.* proposed a CADENZ approach where the structural fragments from homologous but structurally divergent enzymes were recombined to generate diverse protein scaffolds while preserving enzyme catalytic function.<sup>321</sup>

An enumerative algorithm was developed by Baker *et al.* for generating scaffolds, where enzyme pocket scaffolds were constructed by Monte Carlo assembly of secondary structure folds, thus the possible combination of the structural parameters associated with the folds was enumerated. The approach was successfully applied to generate nuclear transport factor 2 (NTF2-like) protein structures with diverse pockets to accommodate diverse active sites.<sup>322</sup> Recently, Baker *et al.* developed a deep-learning based 'family-wide hallucination' approach to generate a large number of NTF2-like scaffolds with diverse binding pockets and introduce the luciferin substrate of luciferases into scaffolds. The designed artificial luciferases exhibited high activity and specificity toward substrates of natural luciferase.<sup>323</sup>

## 4. Summary and perspectives

In this paper, we reviewed the enzyme design methods guided by computational simulations, as well as the revolution brought to traditional computational modeling by integrating machine learning. We also reviewed enzyme design directed by machine learning.

The development of machine learning-based AlphaFold2 has demonstrated its great success in predicting the protein 3D structures from sequences and hence significantly



expanded the size of the protein structure database for structure-based enzyme design. Ligand-bound conformations now can be generated from apo-protein structures directly.<sup>324</sup> MD simulations and multiscale QM/MM calculations are used to explore protein conformational landscape to guide the site-specific mutagenesis. However, these simulation methods are not suitable for high-throughput screening of a large designed sequence database due to the high computing demands. It is important to balance between the extensive sampling and calculation accuracy. The conformational space can be sampled extensively employing multiple short, parallel MD simulations<sup>325</sup> and enhanced QM/MM sampling would allow the exploration of enzyme free energy surface more efficiently.<sup>326,327</sup>

Machine learning methods may be combined with the traditional simulations to sample equilibrium states and rare events<sup>76,328,329</sup> and even sample the catalytically relevant conformations in catalytic reaction space.<sup>330</sup> Furthermore, exploring protein fitness would benefit from a fully automated process,<sup>10</sup> which will largely reduce the efforts compared to the traditional screening. In an automated device, integrating site-specific mutagenesis and machine learning,<sup>331</sup> the data generated from high-throughput screening were used as the input for the ML model to automatically explore the sequence fitness landscape.

High-quality datasets are crucial for the predictivity and generalization of the ML-model used for enzyme design. Numerous ML-guided enzyme design was based on the datasets extracted from the commonly used databases and substantial efforts are required to collect the specific data such as enzyme sequences, structures, substrate specificity, thermostability, kinetic properties, *etc.* from diverse databases. Additionally, many of these databases contain redundant data and irrelevant information, and the data are usually not standardized. Therefore, additional effort involves cleaning the data to construct a high-quality dataset. Furthermore, usually only the enzyme variants with improved properties were reported in the literature, so that the databases generated are biased toward positive samples. To improve predictivity of ML, artificially constructing negative data sets may be a practical strategy.

On the other hand, the dataset available for machine learning in enzyme engineering is usually small, therefore, it is necessary to develop algorithms tailored to improve the predictivity of ML models for these small datasets. The traditional deep learning models like GNN and RNN were originally designed for large datasets for text recognition and image recognition so they may not attain satisfactory results on small datasets.<sup>332</sup> Transfer learning that transfers the “knowledge” learned from a large dataset to the model for a small dataset holds promise for enhancing the predictivity of ML models.

## Data availability statement

The data that support this study are available from the corresponding author upon reasonable request.

## Conflicts of interest

There are no conflicts to declare.

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