Engineering the biomimetic cofactors of NMNH for cytochrome P450 BM3 based on binding conformation refinement†

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Cytochrome P450 BM3 (BM3) is an important oxidoreductase that is widely used in drug synthesis, chemical synthesis, and other industries. However, as BM3 unquestionably increases costs by consuming a natural cofactor that unstably provides electrons, an alternative biomimetic cofactor with simpler structures represented by nicotinamide mononucleotide (NMNH) has been utilized. Currently, few reports exist on artifically modified BM3 enzymes using NMNH, especially regarding theoretical simulation and calculation. With the cognition of the mechanism in mind, we propose a strategy that optimizes and refines catalytic conformation. Based on constrained molecular dynamics simulation, the distance between N-5 of FAD flavin and C-4 of NMNH is used as a cue for the determination of improved conformation, and the potential positive mutants are subsequently screened virtually in accordance with binding free energy requirements. As a result, the $K_{cat}/K_M$ values of the favorable mutant S848R increased to 205.38% compared to the wild-type BM3 with NMNH. These data indicate that our strategy can be applied for the specific utilization of biomimetic cofactors by oxidoreductases represented by BM3.

Introduction

Oxidoreductase, which catalyzes biological oxidation/reduction reactions, is widely used for industrial drug synthesis, chemical synthesis, and food processing, among other industries. Most oxidoreductases require natural cofactors in the catalysis process, such as nicotinamide adenine dinucleotide (NAD$^+$) and nicotinamide adenine dinucleotide phosphate (NADP$^+$), whose reduced forms are NADH and NADPH, respectively. However, these natural cofactors are structurally complex and unstable, which undoubtedly increases the cost and obstacles of industrial production. Nowadays, many biomimetic cofactors with simpler structures, such as nicotinamide mononucleotide (NMN$^+$/NMNH)$^5$, have been developed and utilized. However, most wild-type enzymes either cannot utilize biomimetic cofactors, including NMNH, or exhibit low activity when such cofactors are utilized.

Cytochrome P450 BM3 (BM3, EC1.14.14.1) is a soluble monoxygenase derived from Bacillus megaterium.$^7$–$^9$ On the one hand, it is a typical 120 kDa natural fusion enzyme, with two parts comprising a 55 kDa P450 heme domain and a 65 kDa P450 reductase (CPR) domain, which contains both the FMN-binding domain (FMN domain) and the FAD/NADPH-binding domain (FAD domain).$^8$ As in CPR, electrons pass from NADPH to FAD and thence to FMN before being shunted to the heme iron. On the other hand, BM3 formerly hydroxylates long-chain fatty acids, amides, and alcohols at the $\omega$-1, $\omega$-2, and $\omega$-3 positions.$^{11}$ Due to its self-sufficiency in electron transfer, incomparable regioselectivity, and stereoselectivity, various researchers have employed protein engineering strategies to obtain BM3 mutants that exhibit specific substrate selectivity.$^{12}$ Munro’s group demonstrated that A82F, F87V, and F87V/A82F mutants of BM3 can catalyze diverse azole antifungal drugs more efficiently with voriconazole, omeprazole, and esomeprazole.$^{13,14}$ The BM3 variant C400S designed by Arnold’s group could complete NAD(P)H-driven cyclopropanation with high conversion in whole-cell catalysis.$^7$ However, the wild-type (WT) BM3 consumes only a natural cofactor that provides electrons, which is a disadvantage. Consequently, research on biomimetic cofactors.
has exploded, and finding alternative biomimetic cofactors is the key to resolving this issue. Biomimetic cofactors, such as NMNH, N-4-methoxybenzyl-1,4-dihydrionicotinamide and nicotinamide cytosine dinucleotide, have been confirmed to be used by oxidoreductase. Positive mutants were obtained by randomized and irrational strategies, and the mechanism of enzymes using biomimetic cofactors and further rational artificial modification strategies have not yet been explained in detail.

Currently, few studies have been conducted on artificially modified BM3 enzymes using NMNH, especially regarding theoretical simulation and calculation. Recently, our group has obtained several modified mutant strains of Gox2181, changing its exclusive preference for the cofactor NADPH into dual utilization activity. This rational design method largely completes the specific utilization of biomimetic cofactors by oxidoreductase.

**Experimental**

**Preparation of protein structure**

The crystal structure of the FAD/NADP-binding domain (FAD domain) of P450 BM3 in the presence of the ligand NADPH has been resolved (PDB ID: 4DQL). Given that the ribose-nicotinamide mononucleotide of NADPH is highly disordered, this structure (4DQL) does not contain the exact conformation of ribose nicotinamide. Fortunately, the crystal structure of NADPH-cytochrome P450 oxidoreductase (CYPOR) from rats contains detailed information on the ligand NADPH (PDB ID: 1J9Z). The FAD domains from 4DQL and 1J9Z are very similar (approximately 49.0% sequence similarity). Therefore, the structure of NADPH was obtained by aligning 1J9Z to 4DQL. The extra R group of NADPH was then deleted to generate the structure of NMNH, as shown in Fig. 1(A) and (B).

**MD simulation**

The AM1-BCC method was used to calculate the charge of NADPH and NMNH. The ff14SB and GAFF2 force fields of AMBER18 (ref. 24) software were employed to provide parameters for protein and substrate, respectively. The TIP3P water box with a buffer distance of 10 Å (about 171,000 water molecules) was used as the solvent environment, and 17 sodium ions were added to maintain neutrality of the system. Next, energy minimization is carried out in two steps using the steepest descent and the conjugate gradient methods to remove bad contacts of atoms. In the first step, the solvent water molecules are optimized, and the solute molecules are constrained under a force of 500 kcal (mol Å^2)^{-1}. In the second step, the entire system is optimized without any constraints. Then, the system was slowly heated from 0 to 300 K for 300 ps under the weak constraint of 10 kcal (mol Å^2)^{-1}. The SHAKE algorithm was used to constrain all chemical bonds involving hydrogen atoms, and the particle mesh Ewald method was used to handle long-range electrostatic interactions. Finally, MD simulation was performed for 10 ns under the NPT ensemble.

**Computer virtual mutation**

All residues (except the catalytic triad) within 8 Å of the substrate were considered by the computer virtual saturation mutation strategy to screen for favorable mutants. The leap module of AMBER18 was used to replace amino acid side chains to achieve amino acid mutations. The structure of the protein after mutation is optimized by the steepest descent and the conjugate gradient methods. During the optimization, the distance between N-5 of FAD and C-4 of NMNH atoms is constrained so that the substrate is in a catalytic conformation. The structure of the wild type is optimized using the same method. The molecular mechanics/generalized Born surface area (MM/GBSA) method is subsequently adopted to calculate the binding free energy of protein and substrates based on the optimized structure obtained after mutation. The free energy of binding is used as the objective function to select mutations with enhanced affinity.

**MM/GBSA method**

The MM/GBSA method was used to calculate the binding free energy (ΔG_{bind}) of protein and substrate molecules as follows:

\[ ΔG_{bind} = ΔG_{gas} + ΔG_{sol} \] (1)

where ΔG_{gas} and ΔG_{sol} represent the gas phase binding free energy and solvation free energy, respectively. Among these variables, ΔG_{gas} can be divided into two parts:

\[ ΔG_{gas} = ΔE_{ele} + ΔE_{vdW} \] (2)
where \( \Delta E_{\text{ele}} \) and \( \Delta E_{\text{vdw}} \) represent electrostatic interactions and van der Waals (vdW) interactions, respectively. \( \Delta G_{\text{sol}} \) can be divided into two parts:

\[
\Delta G_{\text{sol}} = \Delta G_{\text{pb}} + \Delta G_{\text{np}}
\]

(3)

where \( \Delta G_{\text{pb}} \) represents the polar solvation free energy, which is calculated by the GB model\(^{12,13} \) with "igb = 2" \( \Delta G_{\text{np}} \) representing the non-polar solvation free energy, which is calculated using the following equation:

\[
\Delta G_{\text{np}} = \gamma \times \text{SASA} + \beta
\]

(4)

where SASA represents the solvent-accessible surface area, as calculated by the MSMS program.\(^{32} \gamma \) and \( \beta \) represent 0.005 kcal (mol \( \text{Å}^2 \))\(^{-1} \) and 0.00 kcal (mol)\(^{-1} \), respectively.

**Construction of strains and plasmid**

Cytochrome P450 BM3 (GenBank J04832.1) was amplified from the plasmid pET-28a(+) with an N-terminal His-tag. Site-directed mutagenesis, using WT BM3 as a template, was performed using PrimeSTAR Max DNA Polymerase (Takara) for the PCR process. All primers were synthesized by Generay Biotechnology (Shanghai, China), which was used for mutagenesis, as shown in Table S1.\(^{†} \) According to the manufacturer’s instructions of the Blunting Kination Ligation Kit (Takara), phosphorylation was carried out at 37°C for 20 min, after which the phosphatase was deactivated by heat at 70°C for 5 min, and ligation was allowed to take place for 60 min. Next, the ligation mixtures were used to transform E. coli DH5ax cells to select positive clones. According to reports in the literature, the glucose dehydrogenase from *Bacillus subtilis* (Bs GDH, UniProt ID: A0A1B2ATD9_BACIU) of the mutant I195R\(^{4} \) was cloned into pET-22b(+) with an N-terminal His-tag, which was used to reduce NMN\(^{+} \) (Aladdin) and generate NMNH. The gene sequences of all the mutants were determined using Shanghai Sunny Biotechnology.

**Protein expression and purification**

The WT/mutant plasmids were transformed into *E. coli* BL21 (DE3) for the heterogeneous expression of enzymes. Recombinant *E. coli* BL21 (DE3) cells were cultured in Luria–Bertani medium with 50 \( \mu \)g mL\(^{-1} \) kanamycin. After incubation at 37°C and shaking at 220 rpm, cells were grown to an OD\(_{600}\) of 0.8–1.0. The cultures were then induced by adding IPTG to a final concentration of 0.1 mM at 20°C. After a further 12 h of incubation, the cells were harvested by centrifugation at 8000g for 5 min at 4°C.

The cell pellet was resuspended in lysis buffer (50 mM Tris–HCl buffer, 10 mM imidazole, pH 7.5) and the cells were lysed by sonication. Cell debris was removed by centrifugation at 18 000g for 15 min at 4°C. The supernatant was filtered through a 0.22 \( \mu \)m filter. Next, the lysate was subjected to a HisTrap HP affinity column using an AKTA Purifier system. The WT/mutant sample was eluted with a gradient of 10–500 mM elution buffer (50 mM Tris–HCl buffer, 10–500 mM imidazole, pH 7.5), and the target protein was eluted with 100 mM elution buffer (50 mM Tris–HCl buffer, 100 mM imidazole, pH 7.5).

Enzyme purity was analyzed by 8% SDS-PAGE, and active fractions were also buffer exchanged into 50 mM Tris–HCl (pH 7.5). The enzyme was concentrated using Amicon Ultra (Millipore) with a 30 kDa molecular weight cut-off, and the three centrifugation steps were allowed to take place at 4800g at 4°C for 20 min. The concentration of the target protein after purification was determined using the BCA Protein Quantification kit, which was purchased from Yeasen Biotechnology (Shanghai, China).

**Enzyme activity and kinetics investigation**

The enzyme activity was detected using a Multiskan Sky Microplate Spectrophotometer (Thermo Scientific). The enzyme activity reaction mixtures (0.1 mL) containing 0.75 mg mL\(^{-1} \) purified enzyme, 0.3 M glucose, 1 M NaCl, 2 mM NMNH, and 0.2 M PBS buffer (pH 7.5) were placed into a 96-well plate. The plates were pre-incubated at 30°C for 2 min, then 80 \( \mu \)M cytochrome c (Sigma-Aldrich) was added, which was monitored at 550 nm (\( \varepsilon_{550} = 21.1 \) M\(^{-1} \) cm\(^{-1} \)). NMNH was prepared according to previously described methods.\(^{a} \) The activities of the WT/mutant enzymes were expressed as the difference in the ability to reduce cytochrome c between the experimental and control groups. One unit (U) of enzymatic activity was defined as the amount of enzyme required to catalyze the reduction of 1 nmol cytochrome c per min at 30°C (pH 7.5). Enzyme activity data are shown as means with standard deviations.

The method to determine kinetic parameters is similar to that for enzyme activity. In the reaction mixture, the enzyme concentration was 6.25 \( \mu \)M, and the concentration of cytochrome c varied in the range of 3.125 \( \mu \)M to 160 \( \mu \)M. Each set of data was tested in parallel at least in triplicate. The \( K_{\text{cat}} \) and \( K_{m} \) values were fitted to Michaelis–Menten using non-linear regression (OriginPro 9.0).

**Results and discussion**

**Structural comparison of NMNH and NADPH**

Before MD simulations, the electron transport mechanism of BM3 and the difference between NMNH and NADPH should be understood in detail. As shown in Fig. 1(C), the first transfer of electrons in BM3 from NADPH to FAD form FADH2. Next, the electrons pass from FADH2 to FMN, leading to the simultaneous formation of FMNH and FADH semiquinone. The final step is to provide electrons from FMNH semiquinone to iron in the heme domain.\(^{7,31,34} \) Fig. 1(A) and (B) show the structures of FAD, NMNH, and NADPH. The structural difference between NMNH and NADPH is the lack of the R part, but both have a nicotinamide moiety in common. The nicotinamide C-4 of NMNH or NADPH donates a proton to the isooalloxazine N-5 of FAD. It was found that a shorter distance between N-5 (FAD) and C-4 (NMNH or NADPH) atoms can provide a better conformation.

Both NMNH and NADPH can be catalyzed by wild enzymes, but they have a lower preference for NMNH. To study the reasons for different preferences between NMNH and NADPH, MD simulations of FAD domain–NMNH and FAD domain–NADPH were performed for 10 ns. Given the stability of the protein crystal
structure and cofactors, the root mean square deviation (RMSD) of the protein binding pocket backbone atoms, FAD, NMNH, and NADPH relative to the initial structure was calculated to measure the overall stability of the structure. As shown in Fig. 2, the RMSD of the FAD domain–NMNH and FAD domain–NADPH simulations tended to be stable, without major fluctuations, which indicates that the simulations have reached convergence within 10 ns and can be used for further analysis.

According to the electron transport mechanism, the closer the distance between the N-5 and C-4 atoms, the greater the probability of the electron transfer. When the distance is large, electrons cannot be transferred from N-5 to C-4, and the catalytic reaction cannot occur. Therefore, the distance between the N-5 and C-4 atoms is an important factor that determines the influence of the catalytic reaction. The distances and their frequency distributions during the FAD domain–NMNH and FAD domain–NADPH simulation are plotted in Fig. 3. It can be found that the distance of the FAD domain–NMNH system is evidently farther than that of the FAD domain–NADPH, which causes the electron transfer of the FAD domain–NMNH system to be more difficult.

This phenomenon explains the low catalytic efficiency of the BM3 with NMNH cofactor. At the same time, it shows that distance is a prerequisite for catalysis, which can be used as a research basis for constructing rational designs that improve catalytic activity.

In addition, a longer time (100 ns) MD simulation on FAD domain–NMNH complex was performed to verify the convergence of simulation and the reliability of the results. The RMSD and distance between the N-5 and C-4 atoms were calculated and presented in Fig. S4.† As can be seen, there is no significant difference between the average distance of 100 ns and 10 ns simulation. This result demonstrates the convergence of our simulation and the reliability of the results.

Screening of mutants

The structural comparison found that the distance between the N-5 (FAD) and C-4 (NMNH) atoms is a prerequisite for catalysis. To improve the catalytic activity of BM3 with NMNH cofactor, the distance of N-5 (FAD) and C-4 (NMNH) is used as a constraint condition, and the binding free energy of BM3 and NMNH is used as an objective function to select favorable mutants. All residues (49 with the exception of the catalytic triad) within 8 Å near the NMNH performed the computer virtual saturation mutation. The binding free energies of 931 single point mutants are presented in Table S2.† Based on the principle of the scattered mutation points, ten mutations with lower binding energy, as shown in Table 1, were selected for further experimental verification.

Analysis of expression and purification

According to the results of theoretical calculations and simulations, ten mutations that may increase the enzyme activity of WT BM3 for cytochrome c were screened. According to experimental procedures, WT and mutant proteins have been constructed and expressed. Fig. S1† summarizes the results of the expression procedure. Except for D943M, the molecular mass of the other nine mutations all showed a protein band at 120 kDa on SDS-PAGE, which verifies soluble protein expression. Then, nine mutations were purified by HisTrap HP, and the purity of

![Fig. 2](image-url)

**Fig. 2** The root mean square deviation (RMSD) of the backbone of the FAD domain (wild-type) binding pocket, FAD, NMNH and NADPH. (A) The RMSD of FAD domain–NADPH simulations. (B) The RMSD of FAD domain–NMNH simulations.

![Fig. 3](image-url)

**Fig. 3** Unconstrained dynamic simulations. (A) The distance of N-5 and C-4 atoms during the FAD domain–NMNH and FAD domain–NADPH simulation. (B) The frequency distribution of N-5 and C-4 atoms during the FAD domain–NMNH and FAD domain–NADPH simulation. (C) The lowest potential energy structure of NMNH and FAD during the FAD domain–NMNH simulations. (D) The lowest potential energy structure of NADPH and FAD during the FAD domain–NADPH simulations.

| ID   | Mutant | ΔGbind | ΔΔGbind |
|------|--------|--------|---------|
| WT   | GLY    | −41.54 | 0.00    |
| 1001 | ARG    | −46.18 | −4.64   |
| 943  | ARG    | −45.00 | −3.46   |
| 1001 | GLN    | −44.20 | −2.58   |
| 907  | PRO    | −43.83 | −2.30   |
| 848  | ARG    | −43.35 | −1.81   |
| 866  | ARG    | −43.22 | −1.68   |
| 907  | THR    | −43.21 | −1.68   |
| 905  | THR    | −42.49 | −0.95   |
| 943  | MET    | −42.44 | −0.90   |
| 1002 | ASN    | −42.05 | −0.52   |

*All energies are in kcal mol⁻¹. ΔΔGbind = ΔGbind(mut) − ΔGbind(WT).*

Table 1 Ten mutants screened and corresponding binding free energy (ΔGbind)**
the target protein reached 95% (Fig. S2†), preparing for the next determination of enzyme activity and kinetic parameters.

Analysis of enzyme activity and kinetics

To complete the detection of enzyme activity more accurately and quickly, we use the Multiskan Sky Microplate Spectrophotometer to detect the ability of the mutants to reduce substrate cytochrome c at 550 nm with NMNH cofactor. It was found that only four mutants were active, namely D943R, G905T, S848R, and D1002N. Detailed data of enzyme activity are shown in Fig. 4. Among them, S848R has greater enzyme activity towards NMNH cofactor than WT BM3, and relative enzyme activities for WT BM3 and S848R were 486.42 U mg⁻¹ and 1332.88 U mg⁻¹, indicating that the ability of reduced cytochrome c of this mutant is 2.74-fold of the wild type. Compared to WT BM3, the enzyme activity of the mutants D943R, G905T, and D1002N was decreased by 4.46%, 16.15%, and 54.25%, respectively. Five other mutants, S866R, G907T, G907P, G1001R, and G1001Q, did not demonstrate any activity. It is likely that the mutation site destroys the catalytic active center, leading to lost activity. Using NMNH cofactor, we successfully screened the mutant S848R, which has higher enzyme activity than WT BM3.

Enzyme kinetic parameters of WT BM3 and variant S848R were measured for substrate cytochrome c. Kinetic data of purified proteins are shown in Table 2. The \( K_m \) value of the mutant S848R was 15.20 \( \mu \)M, which was lower than that of the WT (23.09 \( \mu \)M). This indicated that S848R showed a higher affinity with NMNH cofactor. The \( K_{cat} \) value of the mutant S848R was 87.08 min⁻¹ and more than 1.35-fold higher than that of the WT BM3 (64.34 min⁻¹). Moreover, compared to the WT enzyme, the \( K_{cat}/K_m \) values of the mutant S848R increased to 205.38%. The kinetic investigations of cytochrome c reduction with NMNH cofactor based on Michaelis–Menten parameters are shown in Fig. S3.† These data indicate that the BM3 mutant of S848R exhibits higher catalytic efficiency with NMNH, and the increase in catalytic efficiency is due to improvement in the \( K_{cat} \) value and affinity.

Mechanism of high catalytic efficiency

Experiments found that the S848 mutant caused an increase in catalytic efficiency, while D943R, G905T, and D1002N led to a decrease. To explore the detailed mechanism of difference in catalytic efficiency, MD simulations of WT, S848R, D943R, G905T, and D1002N were performed in parallel under the same parameter settings. The RMSD of the protein binding pocket backbone atoms, FAD, and NMNH in the four mutant simulations were calculated and are shown in Fig. S5.† It shown that the 10 ns simulation for these four mutants reached convergence. The distances between N-5 (FAD) and C-4 (NMNH) atoms and average values as a function of time in the five simulations are plotted in Fig. 5. The average distances of N-5 (FAD) and C-4 (NMNH) atoms in the WT system was 8.61 \( \AA \), and the average distance in the S848R mutant was reduced to 6.81 \( \AA \), and the average distances in the D943R, G905T, and D1002N mutants increased to 9.06, 8.93, and 11.27 \( \AA \), respectively. The order of distance and the order of relative enzyme activity are essentially equivalent, which shows that the distance of N-5 (FAD) and C-4 (NMNH) atoms is a suitable basis to measure enzyme activity.

For the S848R mutant with high catalytic efficiency, the reason for the closer distance of N-5 (FAD) and C-4 (NMNH) atoms is further explored. The binding free energy between BM3 and NMNH cofactor is calculated using the MM/GBSA method and is shown in Table 3. The binding free energies of the WT

Table 2  Kinetic parameters values for cytochrome c by WT BM3 and mutant S848R by using NMNH

| Enzyme    | \( V_{max} \) (nM min⁻¹) | \( K_m \) (\( \mu \)M) | \( K_{cat} \) (min⁻¹) | \( K_{cat}/K_m \) (min⁻¹ \( \mu \)M⁻¹) |
|-----------|--------------------------|----------------------|----------------------|----------------------------------|
| WT BM3    | 414.22 ± 18.05           | 23.09 ± 3.08         | 64.34 ± 1.75         | 2.79                             |
| S848R     | 525.87 ± 41.41           | 15.20 ± 3.73         | 87.08 ± 2.91         | 5.73                             |
and S848R mutant to NMNH are $-12.18$ and $-13.98 \text{ kcal mol}^{-1}$, respectively, and the S848R mutation resulted in a higher affinity between BM3 and NMNH, which is consistent with the experimental $K_{\text{cat}}$ observation. In addition, residue decomposition is performed to calculate the contribution residues S/R848 in terms of binding free energy by the decomposition model of AMBER18. As shown in Table 3, the contribution of S848 in WT is $-2.72 \text{ kcal mol}^{-1}$, and that of R848 in the S848R mutant is $-4.38 \text{ kcal mol}^{-1}$, indicating that mutation of residue 848 directly leads to an increase in affinity with BM3. In addition to the energy term, electrostatic interactions and vdW interactions both contribute to the improvement of binding affinity.

Besides, the root mean square fluctuation (RMSF) of protein backbone and NMNH are calculated as shown Fig. 6 to explore the impact of S848R mutation on the flexibility of protein structure. In Fig. 6, the RMSF of WT and S848R are very similar overall, however, the RMSF of S848R is significantly smaller than that of WT, which indicates that the S848R mutation makes the structure more stable. Special residues and ligands (848, FAD and NMNH) are highlighted in Fig. 6. The RMSF is relatively small, which shows that they are the key residue/ligand and can stably play a catalytic role. At the same time, for these three key residues/ligands, the RMSF of S848R is smaller than that of WT. The results of RMSF analysis show S848R mutation can make key residue and ligand (848, FAD and NMNH) and even the entire protein more stable to improve catalytic efficiency.

To more intuitively explain the origin of the increased affinity caused by mutations, the lowest potential energy structures of the WT and S848R mutants are plotted in Fig. 7. After structural comparison of wild and S848R mutants, the residue mutates from a short chain residue to a long chain residue, and a longer side chain leads to stronger vdW interactions. On the other hand, the R848 residue with a positively charged group can form a salt bridge with the negatively charged phosphate group on NMNH, which leads to stronger electrostatic interactions. The difference in the above structure results in a higher affinity of the S848R mutant for NMNH, and it also simultaneously shortens the distance between N-5 (FAD) and C-4 (NMNH) atoms, rendering catalysis easier and increasing the $K_{\text{cat}}$ value.

### Conclusions

The use of biomimetic cofactors is a promising method for the economic feasibility of oxidoreductase in industrial production. In this study, the oxidoreductase of P450 BM3 was designed by theoretical simulation and calculation strategies that improve enzyme activity with cofactor NMNH. During the optimization of the MD simulation, the distance between N-5 of FAD and C-4 of NMNH is used as a cue to evaluate improved conformation.

The positive mutant S848R was successfully screened. The enzyme activity of S848R is 2.74-fold of the wild type, and the $K_{\text{cat}}/K_{\text{cat}}$, values increased by 205.38% with NMNH cofactor compared to the wild type. Based on the electron transfer mechanism, the utilization of biomimetic cofactors by oxidoreductase can be effectively improved by optimizing the binding conformation. Through MD simulation and computer virtual mutation strategies, ten mutations with better binding efficiency under the perfect conformation were selected for further experimental verification, of which one mutation was not expressed correctly. Among the nine successfully expressed mutants, five exhibited no enzyme activity and three demonstrated decreased enzyme activity. Eventually, a variant that increased enzyme activity was selected. The success rate of the
binding conformation refinement strategy reached 10%. This strategy can be further optimized to become more automated. Furthermore, our strategy presents important cues and can be applied for the specific utilization of various biomimetic cofactors by oxidoreductases.

**Author contributions**

LJ and TZ designed and controlled the research. YL performed all the experiment and wrote the most of manuscript. YC performed all the calculations and wrote the related manuscript. CZ, YP and QL assisted in experiment. JZHZ, BG, BF and CY revised this research and manuscript. All authors read and approved the final manuscript.

**Conflicts of interest**

The authors declare no competing financial interest.

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**Notes and references**

1 S. W. May and S. R. Padgette, Nat. Biotechnol., 1983, 1, 677–686.
2 H. Younus, Oxidoreductases: Overview and Practical Applications, Springer International Publishing, Cham, 2019.
3 S. W. May, Curr. Opin. Biotechnol., 1999, 10, 370–375.
4 A. M. Chanique and L. P. Parra, Front. Microbiol., 2018, 9, 194.
5 C. You, R. Huang, X. Wei, Z. Zhu and Y. P. Zhang, Synthetic and Systems Biotechnology, 2017, 2, 208–218.
6 W. B. Black, L. Zhang, W. S. Mak, S. Maxel, Y. Cui, E. King, B. Fong, A. Sanchez Martinez, J. B. Siegel and H. Li, Nat. Chem. Biol., 2020, 16, 87–94.
7 C. J. C. Whitehouse, S. G. Bell and L.-L. Wong, Chem. Soc. Rev., 2012, 41, 1218–1260.
8 L. O. Narhi and A. J. Fulco, J. Biol. Chem., 1986, 261, 7160–7169.
9 T. K. Hyster and F. H. Arnold, Isr. J. Chem., 2015, 55, 14–20.
10 L. O. Narhi and A. J. Fulco, J. Biol. Chem., 1987, 262, 6683–6690.
11 Y. Miura and A. J. Fulco, Biochim. Biophys. Acta, Lipids Lipid Metab., 1975, 388, 305–317.
12 S. Kille, F. E. Zilly, J. P. Acevedo and M. T. Reetz, Nat. Chem., 2011, 3, 738–743.
13 C. F. Butler, C. Peet, A. E. Mason, M. W. Voice, D. Leys and A. W. Munro, J. Biol. Chem., 2013, 288, 25387–25399.
14 L. N. Jeffreys, H. Poddar, M. Golovanova, C. W. Levy, H. M. Girvan, K. J. McLean, M. W. Voice, D. Leys and A. W. Munro, Sci. Rep., 2019, 9, 1577.
15 P. S. Coelho, Z. J. Wang, M. E. Ener, S. A. Baril, A. Kannan, F. H. Arnold and E. M. Brustad, Nat. Chem. Biol., 2013, 9, 485–487.
16 J. D. Ryan, R. H. Fish and D. S. Clark, ChemBioChem, 2008, 9, 2579–2582.
17 Y. Liu, Y. Feng, L. Wang, X. Guo, W. Liu, Q. Li, X. Wang, S. Xue and Z. K. Zhao, ACS Catal., 2019, 9, 1883–1887.
18 D. Cui, L. Zhang, S. Jiang, Z. Yao, B. Gao, J. Lin, Y. A. Yuan and D. Wei, FEBS J., 2015, 282, 2339–2351.
19 D. Cui, L. Zhang, Z. Yao, X. Liu, J. Lin, Y. A. Yuan and D. Wei, J. Biotechnol., 2013, 167, 386–392.
20 M. G. Joyce, I. S. Ekanem, O. Roitel, A. J. Dunford, R. Neeli, H. M. Girvan, G. J. Baker, R. A. Curtis, A. W. Munro and D. Leys, FEBS J., 2012, 279, 1694–1706.
21 M. Wang, D. L. Roberts, R. Paschke, T. M. Shea, B. S. Masters and J. J. Kim, Proc. Natl. Acad. Sci. U. S. A., 1997, 94, 8411–8416.
22 A. Jakalian, B. L. Bush, D. B. Jack and C. I. Bayly, J. Comput. Chem., 2000, 21, 132–146.
23 A. Jakalian, D. B. Jack and C. I. Bayly, J. Comput. Chem., 2002, 23, 1623–1641.
24 D. A. Case, S. R. Brozell, D. S. Cerutti, T. E. Cheatham III, V. W. D. Cruzeiro, T. A. Darden, R. E. Duke, D. Ghoreishi, H. Gohlke, A. W. Goetz, D. Greene, R. Harris, N. Homeyer, S. Izadi, A. Kovalenko, T. S. Lee, S. LeGrand, P. Li, C. Lin, J. Liu, T. Luchko, R. Luo, D. J. Mermelstein, K. M. Merz, Y. Miao, G. Monard, H. Nguyen, I. Omelyan, A. Onufriev, F. Pan, R. Qi, D. R. Roe, L. J. Roitberg, C. Sagui, S. Schott-Verdugo, J. Shen, C. L. Simmerling, J. Smith, J. Swails, R. C. Walker, J. Wang, H. Wei, R. M. Wolf, X. Wu, L. Xiao, D. M. York and P. A. Kollman, AMBER 2018, University of California, San Francisco, 2018.
25 J.-P. Ryckaert, G. Cicotti and H. J. C. Berendsen, J. Comput. Phys., 1977, 23, 327–341.
26 T. Darden, D. York and L. Pedersen, J. Chem. Phys., 1993, 98, 10089–10092.
27 X. Li, Y. Cong, M. Ma, Z.-N. You, B. Gao, J. Z. H. Zhang and L. Zhang, J. Agric. Food Chem., 2020, 68, 5129–5137.
28 D. Bashford and D. A. Case, Annu. Rev. Phys. Chem., 2000, 51, 129–152.
29 M. Feig, A. Onufriev, M. S. Lee, W. Im, D. A. Case and C. L. Brooks, J. Comput. Chem., 2004, 25, 265–284.
30 V. Tsui and D. A. Case, Biopolymers, 2000, 56, 275–291.
31 A. Onufriev, D. Bashford and D. A. Case, Proteins: Struct. Funct., Bioinf., 2004, 55, 383–394.
32 M. F. Sanner, A. J. Olson and J. C. Spehner, Biopolymers, 1996, 38, 305–320.
33 M. B. Murataliev, R. Feyereisen and F. A. Walker, Biochim. Biophys. Acta, 2004, 1698, 1–26.
34 O. Roitel, N. S. Scrutton and A. W. Munro, Biochemistry, 2003, 42, 10809–10821.