Engineering globins for efficient biodegradation of malachite green: two case studies of myoglobin and neuroglobin†

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Malachite green (MG)-contaminated wastewater resulting from industrialization causes a global problem because of its toxicity and widespread usage. Compared with traditional physical and chemical approaches, biodegradation provides a new route for the degradation of MG. As promising candidates for native enzymes, artificial enzymes have received tremendous attention for potential applications due to unlimited possibilities based on precise design. In this study, we rationally engineered artificial enzymes based on myoglobin (Mb) and neuroglobin (Ngb). We introduced an aspartic acid (H64D mutation) in the heme pocket of Mb. A distal histidine (F43H mutation) was further introduced into H64D Mb to obtain a double mutant of F43H/H64D Mb. Moreover, we used A15C/H64D Ngb as designed recently for comparison studies. The H64D Mb, F43H/H64D Mb, and A15C/H64D Ngb were found to catalyze MG degradation efficiently, with activities much higher than those of native enzymes, such as dye-decolorizing peroxidase and laccase (83–205-fold). The crystal structure of H64D Mb was solved and the interactions of MG and H64D Mb and A15C/H64D Ngb were investigated by using both experimental and molecular docking studies. The biodegradation products of MG were also revealed by ESI-MS analysis. Therefore, these artificial enzymes have potential applications in the biodegradation of MG in textile industries and fisheries.

Introduction

With the vigorous development of textile industries, more than 100 000 commercial dyes have been developed and more than 10% of the dyes are discharged without proper treatment, leading to environmental pollution and causing health risks.1–4 Organic dyes typically have stable aromatic structures, which are difficult to be biodegraded, resulting in widespread environmental contaminations. Malachite green (MG), as a synthetic triphenylmethane dye, is extensively used in the textile industry for its excellent dying capacity. The use of MG has been banned in several countries,5 because MG can irritate the skin and cause permanent damage to the eyes of humans and animals, as well as other health risks.6,7 Meanwhile, MG is still used in aquaculture as a parasiticide and fungicide due to its low cost and good antibacterial effect. Therefore, it is urgent and significant to develop methods for the treatment of MG in the environment.

To date, several physicochemical methods have been utilized for the removal of MG from the environment, such as acidification, lime flocculation, ozone oxidation, oxidative electrolysis, and adsorbents.8–11 However, each of these methods has limitations, such as high cost,11 more energy consumption, secondary pollution,12 and low efficiency.13 Moreover, tertiary treatments such as photochemistry and ozone oxidation are often involved in this process.16

Remarkably, biological treatment has drawn considerable attention for MG degradation due to the characteristics of being environmentally-friendly, efficient, and low-cost. For example, Hao et al. purified a monomeric laccase and showed that it was able to decolorize MG.17 Recently, Guan et al. expressed a mutant of CoTα-laccase in E. coli cells and showed that it could decolorize MG under neutral and alkaline conditions.19 Moreover, heme enzymes such as lignin peroxidase, manganese peroxidase, and horseradish peroxidase were applied for the decolorization of MG.18–21,41 Meanwhile, the catalytic efficiency of these native enzymes needs to be improved, and it is still
required more efficient enzymes for biodegradation. For example, in a recent study,\textsuperscript{23} we have realized MG degradation by an artificial heme enzyme engineered in the protein scaffold of myoglobin (Mb), F43H/H64A Mb, with modification of the heme active site.

In addition to Mb, neuroglobin (Ngb) also belongs to the family of globins. Ngb has a similar protein fold to Mb, whereas it has bis-His coordination (His64/His96).\textsuperscript{23,24} Ngb has been shown to exhibit catalytic functions by alteration of the heme coordination states. For example, Tejero and colleagues showed that mutants of Ngb were able to catalyze reactions involving H2O2.\textsuperscript{25} Recently, we have shown that a double mutant of A15C/H64D Ngb exhibits multiple peroxidase activities.\textsuperscript{26}

In this study, we constructed more efficient artificial enzymes based on the globins including both Mb and Ngb by engineering new mutants such as H64D Mb, F43H/H64D Mb and A15C/H64D Ngb. These new artificial enzymes were designed to process a key Asp or His residue in the heme distal site that is responsible for H2O2 activation. We then applied them for the biodegradation of MG. The catalytic efficiencies of these enzymes were further compared to those of other published native and artificial enzymes.

**Results and discussion**

**Structural characterization and substrate binding**

To provide structural information for the introduced Asp64 in the heme distal pocket, we crystallized H64D Mb and solved the X-ray crystal structure (PDB entry 7XCQ and Table S1). As shown in Fig. 1A, the structure revealed that the single mutant provides an environment at the heme distal site, which is similar to that in the native chloroperoxidase, with a distal glutamate responsible for the activation of H2O2 (Fig. 1B).\textsuperscript{27} Moreover, the introduction of Asp64 in Mb was shown to form a specific binding site for substrates and act as a general acid-base catalytic system to generate an oxoferryl species in reaction with H2O2.\textsuperscript{28,29}

It is of vital importance for substrate binding to the catalytic site of enzymes. To probe whether Mg could bind to the protein scaffold, we first performed molecular docking studies for Mg binding to H64D Mb (Fig. 1C) and A15C/H64D Ngb (Fig. 1D), respectively. The overall structure of the Mg–H64D Mb complexes revealed that the molecule of Mg, like the shape of a fan, binds to the protein cavity between helices E and F. The distance between the heme iron and the central carbon of Mg is approximately 8.0 Å (Fig. 1C), which may facilitate the oxidative cleavage of Mg upon the formation of oxoferryl species. The calculated ten most favorable Mg–H64D Mb complexes have similar conformations with the lowest binding energy calculated to −5.92 kcal mol\(^{-1}\) (Table S2).

The overall structure of the Mg–A15C/H64D Ngb complexes was shown in Fig. 1D. It showed that Mg bound to similar positions to the protein scaffold whereas it adopted a different conformation concerning that of Mg binding to H64D Mb, with a short distance of ~9.4 Å to the heme iron center. The lowest binding energy of Mg–A15C/H64D Ngb complexes was calculated to be −5.42 kcal mol\(^{-1}\) (Table S3), which is only −0.05 kcal mol\(^{-1}\) higher than that of the Mg–H64D Mb complex. These theoretical results suggest that Mg may bind to the scaffold of both Mb and Ngb close to the heme active site, albeit with a different conformation.

**Experimental studies of Mg binding to H64D Mb and A15C/H64D Ngb**

The binding of Mg to ferric H64D Mb and A15C/H64D Ngb was further suggested by EPR studies as shown in Fig. 2. The EPR spectrum of ferric H64D Mb exhibited the characteristic high-spin signals of Fe\(^{3+}\) (g\(_z\) = 6.0; g\(_\perp\) = 2.0, Fig. 2a). The intensity of the g\(_\perp\) signal decreased slightly upon the addition of Mg to H64D Mb (Fig. 2b), suggesting a slight alteration of the heme coordination. Meanwhile, the ferric A15C/H64D Ngb exhibited typical 6-coordinated high-spin heme signals with g tensor calculated to g\(_z\) = 6.0 and g\(_\perp\) = 2.0 (Fig. 2c), which were similar to other globins with H2O2His coordination.\textsuperscript{30,31} A small fraction of low-spin species were also detected with g = 3.0 and g = 2.2,
due to the presence of small amounts of low-heme species. These heme signals almost remained the same in the presence of MG (Fig. 2d), which suggests that the binding of MG does not affect the heme active site.

To further probe the effect of MG binding on the protein secondary structure, we collected the far UV-CD spectra of H64D Mb and A15C/H64D Ngb in the absence and presence of MG. The spectrum of H64D Mb exhibited two major negative absorptions at 209 nm and 222 nm, which indicates that the protein has more α-helix content than β-sheets (Fig. 3A). Upon the addition of MG, only slight increases in ellipticity were observed for the characteristic regions of the secondary structure, suggesting that the binding of Mb almost has little effect on the secondary structure. Moreover, similar results were observed for the spectra of A15C/H64D Ngb (Fig. 3B), suggesting the binding of MG to protein with only slight alteration of the secondary structure.

**MG degradation catalyzed by H64D Mb and A15C/H64D Ngb**

With the information on MG–protein interactions, we then evaluated the dye-decolorizing activity of H64D Mb and A15C/H64D Ngb toward the substrate, MG. The effects of H₂O₂ concentration, pH values and temperature on the catalytic ability of H64D Mb and A15C/H64D Ngb were studied. As shown in Fig. 4A and B, the blue solution almost faded completely in the presence of 0.2 mM H₂O₂ for both enzymes. Meanwhile, the reaction was inhibited obviously for H64D Mb compared with A15C/H64D Ngb at higher concentrations of H₂O₂, which indicates that A15C/H64D Ngb has a high tolerance of H₂O₂.

As shown in Fig. 4C, at fixed concentrations of MG (10 μM) and H₂O₂ (0.2 mM), the maximal activity was observed at pH 6.0 and pH 7.0 for H64D Mb and A15C/H64D Ngb, respectively. The temperature had a large influence on the activity of H64D Mb compared to that of A15C/H64D Ngb (Fig. 4D). When the temperature increased from 15 °C to 85 °C, both enzymes exhibited the highest rate of decolorization efficiency at 33–55 °C. Meanwhile, when the temperature was above 55 °C, an inhibition effect was observed for H64D Mb. These observations further indicate the high thermal stability of A15C/H64D Ngb.

Under the optimal degradation conditions, we monitored the UV-vis spectral changes of MG catalyzed by H64D Mb and A15C/H64D Ngb, respectively (Fig. 4E and F). The absorbance of 617 nm of MG was decreased and blue-shifted gradually, suggesting the generation of degradation products, and the solution color was changed from blue to transparent in 1 h. These observations show that these two heme enzymes are efficient in the degradation of MG.

**Catalytic efficiency toward MG**

To obtain the kinetic parameters (k_{cat} and K_{m}) for degradation of the substrate by the enzymes, we carried out steady-state kinetic studies with different concentrations of MG and monitored the absorption changes at 617 nm in the UV-vis spectra under the optimal reaction conditions. Moreover, we evaluated the catalytic ability of F43H/H64D Mb by mutation of Phe43 to His43. The values of k_{obs} and the concentrations of MG were then plotted and fitted to the Michaelis–Menten equation (Fig. 5A). The parameters of k_{cat} and K_{m} are listed in Table 1. The

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**Fig. 3** Comparison of the CD spectra of (A) H64D Mb (1 μM) and (B) A15C/H64D Ngb (1 μM) in the absence and presence of MG (20 μM) in 5 mM KPi (pH 6.0 for H64D Mb and pH 7.0 for A15C/H64D Ngb).

**Fig. 4** Digital photos of MG oxidized by (A) H64D Mb and (B) A15C/H64D Ngb with different H₂O₂ (0–1.0 mM) concentrations at 25 °C for 2 h; effects of different (C) pH and (D) temperature on the catalytic oxidation rate of MG by H64D Mb and A15C/H64D Ngb in the similar conditions; the UV-vis spectra of MG oxidized by (E) H64D Mb and (F) A15C/H64D Ngb.

**Fig. 5** (A) Steady-state rates of oxidation as a function of MG concentrations (0–16 μM) catalyzed by H64D Mb, F43H/H64D Mb and A15C/H64D Ngb using H₂O₂ as an oxidant; (B) the comparison of the catalytic efficiency of Mb and Ngb mutants and the natural enzymes.
results showed that the $k_{\text{cat}}$ value of H64D Mb and F43H/H64D Mb are 0.1218 s$^{-1}$ (87-fold vs. WT Mb$^{22}$) and 0.3148 s$^{-1}$ (225-fold vs. WT Mb), respectively. These results indicate that the aspartic acid at the 64 position and the distal His43 could facilitate the catalytic ability, which agree with previous studies.$^{38-40}$ It showed that the distal Asp64 or His43 in Mb with a suitable distance to the heme iron efficiently acts as acid–base catalysis in activation of H$_2$O$_2$, generating the reactive compound I (an oxoferryl heme π-cation radical) or compound II (oxoferryl heme). These species may be responsible for the substrate degradation by oxidative cleavage through radical or oxygen transfer.

Among these enzymes, A15C/H64D Ngb exhibited the highest catalytic efficiency with $k_{\text{cat}}/K_m$ value up to 94 929 M$^{-1}$ s$^{-1}$, which is 151- and 205-fold higher than that reported for dye-decolorizing peroxidase from Bacillus subtilis (BsDyP$^{35}$) and the mutated CotA-lacase SF enzyme.$^{36}$

**Product analysis of MG degradation**

To identify the degradation products of MG catalyzed by H64D Mb, we conducted ESI-MS studies of the solution after the reaction. As shown in Fig. 5A for the mass of MG (329 m/z), the peak disappeared after the reaction and new peaks were generated. These new peaks included 226, 212, 197 and 134 m/z, which could be assigned the following species, (Fig. 6B(a)) $m/z = 226$ is 4-(dimethylamino)benzophenone (DLBP); (Fig. 6B(b)) $m/z = 212$ is 4-((methylamino)benzophenone (MLBP); (Fig. 6B(c)) $m/z = 197$ is 4-aminobenzophenone (ABP); (Fig. 6B(d)) $m/z = 134$ is 4-(methylamino)benzaldehyde (MLBD), respectively. Moreover, the MG degradation products catalyzed by A15C/H64D Ngb produced small molecules such as those detected with molecular weights of 315, 301, 287, and 165 m/z, which are corresponding to (p-dimethylaminophenyl)[p-methylaminophenyl]phenylmethyl (DM-PM), (p-methylaminophenyl)[p-methylaminophenyl]phenylmethyl (MM-PM), (p-methylaminophenyl)phenylmethyl (M-PM) and 4-(dimethylamino)benzoic acid (DM-BA), respectively.

Based on the ESI-MS results, we proposed a generation route of the reaction products, which may involve the generation of reactive radical species upon the activation of H$_2$O$_2$ by these enzymes, resulting in different bond cleavages of the substrate.$^{38-40}$ The cleavage of the C–C bond involving the central carbon by removal of one or two benzenes may generate DLBP and DM-BA. N-Demethylation of DLBP was also observed in a stepwise manner by Du et al.$^4$ Given the identification of

**Table 1** Kinetic parameters for degradation of MG catalyzed by Mbs and A15C/H64D Ngb at optimal reaction conditions, with those of native enzymes shown for comparison.

| Proteins                  | $k_{\text{cat}}$ (s$^{-1}$) | $K_m$ (µM) | $k_{\text{cat}}/K_m$ (M$^{-1}$ s$^{-1}$) |
|---------------------------|-----------------------------|------------|----------------------------------|
| WT Mb$^{22}$              | 0.0014 ± 0.0001             | 1.2 ± 0.2  | 1167                             |
| F43H Mb$^{22}$            | 0.0034 ± 0.0001             | 2.4 ± 0.3  | 1416                             |
| F43H/H64A Mb$^{22}$       | 0.0232 ± 0.0013             | 2.7 ± 0.5  | 8593                             |
| H64D Mb                   | 0.1218 ± 0.0036             | 1.5 ± 0.2  | 81 200                           |
| F43H/H64D Mb              | 0.3148 ± 0.0185             | 8.2 ± 0.9  | 38 390                           |
| A15C/H64D Ngb             | 0.1329 ± 0.0037             | 1.4 ± 0.2  | 94 929                           |
| BsDyp$^{35}$              | 0.0456 ± 0.0016             | 72.0 ± 7.0 | 630                              |
| CotA-lac SF$^{36}$        | 18.36                       | 39 600     | 464                              |

**Fig. 6** ESI-MS spectra of (A) MG, (m/z 329) and its degradation products by catalyzed by (B) H64D Mb and (C) A15C/H64D Ngb. (B(a)) DLBP (m/z 226); (B(b)) MLBP (m/z 212); (B(c)) ABP (m/z 197); (B(d)) MLBD (m/z 134); (C(a)) DM-PM (m/z 315); (C(b)) MM-PM (m/z 301); (C(c)) M-PM (m/z 287); (C(d)) DM-BA (m/z165).
MLBP, ABP, MM-PM, DM-PM, and M-PM, the degradation could also be initiated by N-demethylation of MG, followed by oxidation or further N-demethylation.

As shown in Scheme 1, we proposed a plausible biodegradation mechanism of MG catalyzed by H64D Mb and A15C/H64D Ngb. The biodegradation mechanism for MG catalyzed by H64D Mb probably underwent pathway I, where the oxidation reaction occurred firstly, followed by N-demethylation. Conversely, the biodegradation mechanism for MG catalyzed by A15C/H64D Ngb (pathway II) may include three steps of N-demethylation, with a last step of the oxidation reaction. We noticed that the degradation pathways of MG catalyzed by H64D Mb and A15C/H64D Ngb are different, which may due to their different binding models of MG binding to the heme active site. Interestingly, both oxidative cleavage and demethylation were involved in MG degradation catalyzed by H64D Mb and A15C/H64D Ngb.

Conclusions

In this study, we introduced an aspartate residue, by H64D mutation, into the heme pocket of Mb and obtained the crystal structure of the single mutant. We also engineered a double mutant of F43H/H64D Mb by introducing a distal histidine to the single mutant. Moreover, we used the recently designed A15C/H64D Ngb as a control. EPR studies showed that the binding of the enzyme to MG does not affect the heme active site. At the same time, CD studies showed that the binding of the substrate to the enzyme has little effect on the secondary structure of the enzyme. In addition, we found that A15C/H64D Ngb is highly tolerant and thermally stable to H2O2 compared to that of H64D Mb. Kinetic UV-vis studies further showed that H64D Mb, F43H/H64D Mb and A15C/H64D Ngb effectively catalyze the degradation of MG, with much higher activity than natural enzymes such as decolorization peroxidase and laccase (83–205-fold). Finally, by ESI-MS analysis, we proposed a plausible pathway for the generation of MG biodegradation. This study suggests the potential application of artificial enzymes for the biodegradation of MG in the textile industry and fisheries.

Materials and methods

Materials

Sperm whale wild-type (WT) Mb, H64D Mb, F43H/H64D Mb and A15C/H64D Ngb were expressed in BL21 (DE3) cells and purified as previously reported. The genes of F43H/H64D Mb were constructed by using the QuickChange Site-Directed Mutagenesis Kit (Stratagene) using the WT Mb gene as a template, and the mutations were confirmed by a DNA sequencing assay. All other chemical reagents are commercially available and of analytical grade.

Crystal X-ray diffraction of H64D Mb

The protein crystal was shaped by ferric H64D Mb with high purity (A409 nm/A280 nm > 3.0) and was exchanged into 20 mM potassium phosphate buffer. Single-crystal X-ray diffraction data of H64D Mb were collected at the BL16B1 beamline at the Shanghai Synchrotron Radiation Facility (SSRF) using a MAR mosaic 225 CCD detector with a wavelength of 0.9791 Å at 100 K. The diffraction data were processed and scaled with HKL-2000. The structure was determined by the molecular replacement method, and the 1.6 Å structure of WT Mb [PDB entry 1JP6 (ref. 45)] was used as the starting model. Manual adjustment of the model was carried out using COOT and the models were refined by PHENIX and Refmac5. The stereo-chemical quality of the structures was checked using PROCHECK. All the residues were in the favored and allowed regions and none in the disallowed region.

Molecular modeling

The X-ray structure of H64D Mb was used as the initial structure for docking with MG using the Autodock 4.2.3. The heme iron was set to be the center, with a box size of 60 Å × 60 Å × 60 Å, which covered most of the protein surface. MG as a substrate for docking was generated using the Dundee Prodr2 server. Docked conformations were ranked automatically by Autodock 4.2 using a binding-energy scoring function. The docking results with the ten most favorable conformations after 2000 steps were then visualized and analyzed using VMD 1.9.2.

EPR studies

Electron paramagnetic resonance (EPR) spectra of H64D Mb (0.3 mM, in 100 mM KPi buffer, pH 6.0) and A15C/H64D Ngb (0.3 mM, in 100 mM KPi buffer, pH 7.0) in the absence or presence of MG (0.3 mM) were recorded on a Bruker A300 spectrometer (X-band) equipped with Bruker ER4141VTM liquid nitrogen system. The sample was transferred into an EPR tube with a volume of 300 μL. The spectrum was measured at 100 K, with a frequency of 9.27 GHz, centerfield 2750 G and sweep width 4500 G, microwave power 0.595 mW, and modulation amplitude 3.0 G.

CD spectroscopy

The circular dichroism (CD) studies were performed using a J-Model-1500 spectrophotometer (JASCO) equipped with a Peltier temperature control system. The far-UV CD spectra were recorded in the range of 190–250 nm for H64D Mb and A15C/H64D Ngb (1 μM in 5 mM KPi buffer) in the concentration of MG (20 μM). The conformational stability of protein was assessed by recording the spectra at different concentrations (0, 20 μM). The averaged protein spectrum was corrected by reducing the buffer blank baseline spectrum.

UV-vis spectroscopy

UV-vis spectra were recorded on a Hewlett-Packard 8453 diode array spectrometer. The final concentration of proteins was decided by extinction coefficient, like H64D Mb with ε409 nm of 165 mM−1 cm−1, F43H/H64D Mb with ε407 nm of 159 mM−1 cm−1, A15C/H64D Mb with ε406 nm of 160 mM−1 cm−1, MG with ε617 nm of 148.9 mM−1 cm−1.
Kinetic UV-vis studies

The UV-vis spectra of the reaction were recorded using a Hewlett-Packard 8453 diode array spectrometer UV-vis spectra. The reaction was carried out in a 2 mL cuvette, in which, H64D Mb, F43H/H64D Mb (2 mM in 100 mM KPi buffer, pH 6.0) or A15C/H64D Ngb (2 mM in 100 mM KPi buffer, pH 7.0) was added by varying the concentration of MG (0–16 mM). The reaction was initiated by the addition of H2O2 (0.2 mM), and the decrease of absorption at 617 nm was monitored to determine the initial rate.

ESI-MS

The mass of H64D Mb, A15C/H64D Ngb and the products after the reaction H64D, A15C/H64D Ngb and MG were determined on a G2-XS QToF mass spectrometer (Waters), which was then transferred into the mass spectrometer chamber under positive mode. The mass of H64D Mb, A15C/H64D Ngb and the products a

Conflicts of interest

There are no conflicts to declare.

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