Biochemical and Cellular Investigation of *Vitreoscilla* Hemoglobin (VHb) Variants Possessing Efficient Peroxidase Activity

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Peroxidase-like activity of *Vitreoscilla* hemoglobin (VHb) has been recently disclosed. To maximize such activity, two catalytically conserved residues (histidine and arginine) found in the distal pocket of peroxidases have successfully been introduced into that of the VHb. A 15-fold increase in catalytic constant (k_{cat}) was obtained in P54R variant, which was presumably attributable to the lower rigidity and higher hydrophilicity of the distal cavity arising from substitution of proline to arginine. None of the modifications altered the affinity towards either H\textsubscript{2}O\textsubscript{2} or ABTS substrate.

Spectroscopic studies revealed that VHb variants harboring the T29H mutation apparently demonstrated a spectral shift in both ferric and ferrous forms (406–408 to 411 nm, and 432 to 424–425 nm, respectively). All VHb proteins in the ferrous state had a \( \lambda_{soret} \) peak at ~419 nm following the carbon monoxide (CO) binding. Expression of the P54R mutant mediated the downregulation of iron superoxide dismutase (FeSOD) as identified by two-dimensional gel electrophoresis (2-DE) and peptide mass fingerprinting (PMF). According to the high peroxidase activity of P54R, it could effectively eliminate autoxidation-derived H\textsubscript{2}O\textsubscript{2}, which is a cause of heme degradation and iron release. This decreased the iron availability and consequently reduced the formation of the Fe\textsuperscript{2+}-ferric uptake regulator protein (Fe\textsuperscript{2+}-Fur), an inducer of FeSOD expression.

**Keywords:** *Vitreoscilla* hemoglobin, peroxidase-like activity, enzyme kinetics, catalytic constant, proteomic analysis, iron superoxide dismutase

*Vitreoscilla* hemoglobin (VHb) is a bacterial globin originated from the obligate aerobic bacterium *Vitreoscilla* sp. The organism resides naturally in oxygen-limited environments such as stagnant ponds or decaying vegetative matters. The VHb has been thought to maintain an essential level of intracellular dissolved oxygen [44]. Nowadays, it has become a versatile tool in current biotechnology, especially for large-scale fermentation, since shortage of oxygen supply during cultivation can be overcome by heterologous expression of VHb. Cell growth and productivity are enhanced particularly under oxygen-depleted conditions [15, 23, 24]. Thus, VHb technology has been applied for industrial-scale production of many valuable compounds [45, 50].

Beside its role in O\textsubscript{2} delivery, VHb has been involved in various cellular activities, such as protein translation efficiency [3, 36], cellular metabolism [21], and alleviation of nitrosative stress [18, 22]. Recently, peroxidase-like activity has been disclosed in VHb [25, 38]. The steady-state kinetic study has revealed that VHb catalyzes the oxidation of various aromatic substrates via a ping-pong mechanism, as normally found in horseradish peroxidase (HRP) [25]. However, this catalytic activity seems to be limited possibly as a result of the improper nature of its active site. In comparison with HRP, two catalytically important residues (distal histidine and arginine) are missing in VHb, as illustrated in Fig. 1, which may account for the low activity. The distal histidine is believed to be a catalyst by firstly acting as a base to withdraw electron from H\textsubscript{2}O\textsubscript{2} substrate and then turning to be an acid to donate hydrogen atom to heme-bound peroxy [28, 42]. This facilitates the cleavage of the peroxide bond yielding the release of a water molecule. Additionally, the positive charge of conserved arginine provides pulling force to this bond dissociation, which in turn accelerates compound I formation [35, 42].

In the present work, VHb variants possessing distal histidine and arginine residues have been constructed and determined the peroxidase kinetics in the assay reaction with H\textsubscript{2}O\textsubscript{2} and aromatic substrate. Equivalent positions of
VHb, Tyr29 and Pro54, as found in HRP, have been selected for mutagenic modifications. Proteomic study has also been exploited to observe the consequence of VHb expression on protein profiling of E. coli. Such finding would suggest an additional role of this oxygen-binding protein when it behaves as a H$_2$O$_2$ detoxifying enzyme.

**Materials and Methods**

**Bacterial Strain and Chemical Reagents**

Cloning and expression of each VHb variant were conducted in *Escherichia coli* strain TG1 (lac-pro), Sup E, thi I, hsd DS5F' tra D36, pro A B', lacI, lacZ, M15 (amp', dut'). PCR-based site-directed mutagenesis of the VHb gene was carried out using pV6h as a template [25]. Mutagenic primers were purchased from MWG Biotech (Ebersberg, Germany). 2,2'-Azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) and hydrogen peroxide [30% (v/v) H$_2$O$_2$, $c_{25\text{cm}}$=72.8 M$^{-1}$ cm$^{-1}$] [20] were obtained from Sigma and Aldrich, respectively. Before use, the H$_2$O$_2$ solution was standardized and freshly prepared using the corresponding molar adsorption coefficient described above. Other reagents were of analytical grade and commercially available.

**Site-Directed Amino Acid Substitution of Distal Residues of VHb**

Selected residues (Tyr29 and Pro54) in the distal pocket of VHb were substituted with histidine and arginine, respectively, in order to simulate the active site of typical peroxidase. The *vhb* gene was amplified by using single-mutation-containing primers as follows: (Y29H_F: 5'-GAATCTTTGGAGCAGCATATAAAAACCTGTTGCC-3'; P54R_F: 5'-GAATCTTTGGAGCAGCGTTGCGCAGGCGTGGGCGATG-3', mutations shown in bold). Complete digestion of parental plasmid DNA was enabled by addition of *DpnI* restriction enzyme to the PCR product (37°C for an hour). DNA with desired mutation was then transformed into *E. coli* TG1 competent cells and cultured overnight at 37°C on Luria–Bertani (LB) agar supplemented with 100 µg/ml ampicillin (Amp). Colonies were selected randomly for plasmid isolation. Verification of the correct mutation on each plasmid was carried out by DNA sequencing (BM Labbet, Lund, Sweden). A double mutant, Y29H/P54R, was generated by introducing the P54R mutation into the pV6H_Y29H mutant plasmid.

**Expression and Purification of VHb Mutants**

*E. coli* TG1 cells carrying plasmids coding for native VHb and its variants (Y29H, P54R, and Y29H/P54R) were cultivated in modified LB medium (10 g tryptone, 10 g yeast extract, and 5 g NaCl, pH 7.2–7.4) supplemented with 100 µg/ml ampicillin and 0.3 µM ß-aminolevulinate acid (ß-ALA; heme precursor) [7], at 37°C, 150 rpm. Once cell density (OD$_{\text{600}}$) reached 0.6–0.8, recombinant VHb proteins were actively produced as a consequence of gene induction by isopropyl-ß-D-1-thiogalactopyranoside (IPTG) at 0.5 mM final concentration. Then, cultures were further incubated at 30°C, 150 rpm for 14–16 h. Reddish cells indicating the hemoglobin expression were collected, washed, and ruptured by ultrasonic disintegration. Crude cell extracts were centrifuged at 13,000 ×g, 4°C for 15 min and the supernatant was collected. Before loading onto an affinity column, the supernatant was dissolved in lysis buffer (100 mM Tris-HCl, pH 8.0, to remove imidazole). Protein was quantified by the Bradford dye-binding assay [11] with bovine serum albumin (BSA) as a standard and its homogeneity was determined by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE). Purified protein (>95% purity) was utilized in all experiments.

**Spectroscopic Measurements of VHb Proteins**

UV–Vis spectra of VHb variants (5 µM in 50 mM sodium phosphate buffer, pH 7.5) were recorded at room temperature on a DU 800
UV/Vis spectrophotometer (Beckman Coulter, U.S.A.). In addition, changes in the Soret peak were monitored following chemical reduction or oxidation (by excess amount of sodium dithionite or ammonium persulfate, respectively) as well as CO-binding in the same buffer.

**Measurement of Peroxidase Kinetics of VHb Variants**

The oxidations of reducing substrate, ABTS, by mutant proteins were determined using the DU 800 UV/Vis spectrophotometer. Initial rates for the one-electron oxidation of ABTS substrate in the presence of H$_2$O$_2$ were measured over the first 30 s at ambient temperature. It is worth to note that concentrations of H$_2$O$_2$ and reducing substrate were in enormous excess compared with VHb concentration to ensure the pseudo-first-order reaction. A 1-ml reaction mixture in 50 mM sodium phosphate buffer, pH 7.5 containing 0.5 µM VHb proteins, 3–75 mM H$_2$O$_2$, and 100–500 µM ABTS was used. Triplicate measurements were performed for each point of substrate concentration. The initial velocity for the formation of ABTS$^+$ radical was obtained by Beer’s law using its extinction coefficient of ε$_{420}$=14.7 mM$^{-1}$ cm$^{-1}$ [1]. Consequently, it was used for plotting and calculation of kinetic parameters including $K_m$ and $V_{max}$ for both H$_2$O$_2$ and ABTS substrates.

Moreover, the rate of compound I formation ($k_+$) was measured at ambient temperature [26] in 50 mM sodium phosphate, pH 7.5. The concentration of H$_2$O$_2$ substrate was varied over the range of 3–75 mM at 200 µM ABTS reducing substrate and 0.5 µM VHb protein. Rate constants were obtained by fitting the recorded data to pseudo-first-order reaction using the epsilon value of the ABTS$^+$ radical described above.

**Preparation of Protein Samples for Proteomic Experiment**

*E. coli* TG1 expressing wild-type VHb or its variants was grown aerobically at 37°C, 150 rpm in 5 ml of LB medium supplemented with 100 µg/ml ampicillin, and then a 100 dilution of the pre-culture was made into 50 ml of LB/Amp broth. This culture was incubated aerobically at 37°C, 150 rpm in 5 ml of LB medium supplemented with 100 µg/ml ampicillin, and then a 100 dilution of the pre-culture was made into 50 ml of LB/Amp broth. This culture was incubated further for 16 h at the same condition. To induce VHb expression, 1 mM IPTG was added into the bacterial culture during the exponential phase (OD$_{600}$=0.6–0.8). After collecting cells by centrifugation (4,500 × g, room temperature, 10 min), cells were washed three times using 40 mM Tris-HCl, pH 8.0, to remove residual salts and other chemicals. Cells were then dissolved in 1 ml Tris-HCl buffer, pH 8.0. Lysis buffer (250 µl), freshly prepared before use by mixing 10 mg/ml dithiothreitol (DTT) and 10 µg/ml protease inhibitor cocktail into 7 M urea, 2 M thiourea, and 4% CHAPS, was added into the cell suspension. A clear supernatant containing total proteins was then obtained by ultrasonic disintegration (Branson Sonifier Model 450) followed by centrifugation at 13,000 × g for 30–60 min at room temperature. Bradford’s method was used for determination of protein concentration using 5-bromo-2’-deoxyuridine (BrdU) as a standard. Subsequently, the cellular extract was mixed with 1 M acrylamide (1:10 of total volume), left at room temperature for 10 min, and then stored at −20°C until use.

**Two-Dimensional Gel Electrophoresis (2-DE)**

Experiments were performed according to procedures described previously [21]. Briefly, protein sample was examined using a 2D electrophoresis set purchased from Atto Corporation, Japan (IEF disc gel and mPAGE; Models AE-6541 and AE-6531). All procedures were done in accordance with the manufacturer’s recommendation with minor modifications. One hundred µg of cellular proteins was carefully loaded on the top of pre-casting IEF rod gels with pH range of 3–10 (Atto Corporation). The gel was overlaid with 150 µl of upper electrode buffer (0.2 M NaOH). This first-dimension electrophoresis was operated at 300 volts for 3.5 h. Diffusion of proteins from the agarose gel was prohibited by soaking the 1D gel in 2.5 g% trichloroacetic acid (TCA) for 3 min. Fixed gels were then washed 3 times with distilled water. Before performing the second-dimension electrophoresis, gels were immersed in SDS equilibration buffer [50 mM Tris-HCl, 6 M urea, 30% (v/v) glycerol, 2% SDS and bromphenol blue, pH 8.8] for 10 min. After that, rod gels or Whatman filter paper dotted with standard protein markers were placed onto the pre-casting gradient (5–20% acrylamide) SDS–PAGE gel (Atto Corporation). All components were connected to one another by sealing with 1% agarose gel. Second-dimension gels were electrophoresed initially at 10 mA for 10 min and subsequently run at 40 mA for 90 min. Colloidal Coomassie blue was used to stain gels with mild agitation (20–25 rpm) for 2–3 h. Gels were then washed several times with distilled water to remove unstained dye and finally photographed.

**Protein Identification by Peptide Mass Fingerprinting (PMF)**

Protein spots on gels were taken out manually and placed into 96-well plates filled with 50% methanol and 5% acetic acid. Each of the protein spots was digested by high-grade trypsin (Promega, U.K.), mixed with spotter solution (10 mg/ml α-cyano-4-hydroxyphenylacyl acid dissolved in 66% acetonitrile and 0.1% trifluoroacetic acid), and then spotted onto a 384-well target plate (Bruker Daltonics, Germany). Protein digestion and spotting were carried out using a Spot Handling Workstation (GE Heathcare, U.S.A.). Digested proteins were shot with approximately 500 laser beams for acquisition of mass spectra, which were then matched in the NCBInr database using BioTool 2.0 (Bruker Daltonics) and MASCOT 2.2 software (MatrixScience, http://www.matrixscience.com). Permission of only 1 incorrect cleavage per peptide and a score of ≥1.0 KDa initial mass tolerance were applied in all searches. Protein identification was confidently decided if the score of matched proteins was greater than 80 (p<0.05).

Fig. 2. SDS–PAGE showing purified proteins of all VHb variants: molecular mass marker (M); wild-type VHb (1); and Y29H (2); P54R (3); and Y29H/P54R (4) variants.
RESULTS

Construction of VHb Variants and Investigation of Their Spectral Properties

Three VHb variants (designated as Y29H, P54R, and Y29H/P54R) were successfully constructed in which the target residues were initially identified with the aid of molecular modeling served by PyMOL [12]. Supplementation of δ-ALA, a heme precursor, to cultures significantly improved heme content, which could be readily observed by the increase of pellet reddishness [7] (data not shown). After purification by Ni²⁺-IDA affinity chromatography, the wild-type VHb and its variants showed approximately >95% homogeneity as located at ~16 kDa (Fig. 2). To investigate the spectral properties of VHb proteins in ferric, ferrous, and carbonyl forms, ammonium persulfate and sodium dithionite were added in excess to the hemoglobin solution prior to acquiring the spectra. For CO-bound form, VHb protein was treated with dithionite followed by exposure to CO gas for a few seconds. Fig. 3 shows that the UV–Vis spectra of wild-type VHb and P54R variants were nearly identical, and their untreated proteins were likely to be in a ferric state, with Soret peaks positioned at 406 and 408 nm, respectively. Y29H and Y29H/P54R variants revealed unusual absorption properties in which the Soret maxima were shifted to 411 nm. These positions were usually close to the oxy-form ($\lambda_{Soret}=414$ nm) but they lacked typical peaks at 543 and 576 nm known as $\alpha$-bands [27]. Instead, they showed a broad minor peak at 534 nm. To verify whether they were in oxygenated form, both protein samples were bubbled with CO gas and the spectra were then recorded. However, interconversion of oxygenated protein, if formed, to CO-bound protein was not detected (data not shown) since exposure to CO gas of oxygenated VHb has been proven to change the absorption peak to 419 nm [27]. Therefore, these findings indicated that all protein samples were present in the ferric form. Other confirmative evidences revealed that further addition of ammonium persulfate did

![Graphs showing spectral properties of VHb variants](image-url)

Fig. 3. UV–Vis spectra of VHb samples (5 µM) in 50 mM sodium phosphate buffer, pH 7.5. Ferrous or reduced form (dashed line) was obtained following the addition of excess dithionite into ferric-VHb (solid line); and CO complex (dotted line) was obtained by exposing the ferrous-VHb to CO gas for a few seconds. The absorption spectra in the visible region starting at 450 nm were magnified, and the scale is indicated on the right.
Measurements were performed in 50 mM sodium phosphate buffer, pH 7.5, at room temperature.

Table 1. The steady-state kinetic properties of VHb variants in reaction with aromatic substrate ABTS in the presence of H$_2$O$_2$.

<table>
<thead>
<tr>
<th>VHb proteins</th>
<th>$k_m$ (s$^{-1}$)</th>
<th>$K_m$ (H$_2$O$_2$) (mM)</th>
<th>$K_c$ (ABTS) (µM)</th>
<th>$k_c$ (M$^+$ s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VHbH$_6$</td>
<td>0.81±0.03</td>
<td>11.72±1.23</td>
<td>55.38±10.11</td>
<td>30.73±0.69</td>
</tr>
<tr>
<td>Y29H</td>
<td>0.2±0.03</td>
<td>30.83±5.14</td>
<td>160.54±47.25</td>
<td>4.89±0.63</td>
</tr>
<tr>
<td>P54R</td>
<td>11.8±2.79</td>
<td>22.2±1.33</td>
<td>3.238.79±623.07</td>
<td>23.08±0.24</td>
</tr>
<tr>
<td>Y29H/P54R</td>
<td>0.18±0.06</td>
<td>9.13±0.87</td>
<td>1.686.81±699.13</td>
<td>1.17±0.14</td>
</tr>
</tbody>
</table>

Measurements were performed in 50 mM sodium phosphate buffer, pH 7.5, at room temperature.

not alter the spectra (data not shown). Meanwhile, treatment of VHb proteins with excess dithionite produced a 432-nm peak for the wild type and the P54R variant, and a 425-nm peak for Y29H and Y29H/P54R variants. As expected, CO complexes of all proteins appeared at 419 nm. However, slightly different peaks in α-bands among each protein were observed, as shown in Fig. 3, in which two minor peaks were found at 534–537 and 556–563 nm.

Steady-State Kinetics of VHb Variants

Initial rates of formation of ABTS radical upon reacting with H$_2$O$_2$ were determined and calculated by Beer’s law as mentioned earlier. Results of the enzymatic characterization are summarized in Table 1. Expectedly, the P54R variant possessed approximately 15-fold increase in catalytic constant ($k_m$) for the peroxidase-catalyzed reaction compared with that of the wild-type VHb (11.80 and 0.81 s$^{-1}$, respectively). More importantly, such catalytic constant was approximately 60-fold higher than those of the Y29H and Y29H/P54R variants (0.20 and 0.18, s$^{-1}$, respectively). However, a 2-fold lower $K_m$ for H$_2$O$_2$ was observed, suggesting that the substituted positively charged amino acid could not potentially afford binding to the H$_2$O$_2$ substrate. The rate of compound I formation ($k_c$) also revealed that this substitution slightly decreased the reactivity towards the H$_2$O$_2$ molecule. As evidenced in the other two variants, the Y29H mutation reduced the catalytic constant by a factor of 4 as well as the $K_m$ for H$_2$O$_2$ and ABTS substrates by a factor of 1–3 and 3–30, respectively. Likewise, $k_c$ constants for compound I formation of variants Y29H and Y29H/P54R were decreased by factors of ~6–26, respectively, compared with the wild-type VHb. Thus, the Y29H mutation seemed to considerably abolish the formation of compound I intermediate. This finding correlated well with the lower affinity to H$_2$O$_2$ substrate as mentioned previously.

Effects of VHb Variants on Protein Expression Profiles of E. coli

To further investigate whether the high efficiency to catalyze the peroxidase reaction of the VHb variants rendered any alterations or consequent effects on the cellular responses of host cells, total protein extract was analyzed by a proteomic approach. After 2-DE experiment, two areas of each gel displaying protein spots with either down- or upregulation were zoomed, and those of the wild type VHb are displayed in Fig. 4A. Several protein spots (Fig. 4B) were selected and identified by mass spectrometry and the results are summarized in Table 2. As shown in Fig. 4C, molecular chaperones DnaK and GroEL were highly expressed and are usually stated as landmarks in several proteomic analyses (nos. 1, 2, 12, 17, 24, and 3, respectively). The levels of alkyl hydroperoxide reductase (AhpC), thiol peroxidase (Tpx), rotamase B, and DNA protection protein (Dps) remained unchanged. The monomeric subunit of bacterial hemoglobin (represented as spots no. 9, 10, 11, 16, 21, 22, and 25) could readily be visualized at molecular mass of ~16 kDa and pI value of ~5.3. However, notification has to be made that expression levels of each VHb variants were not equal. Single amino acid substitution adversely affected the expression level of VHb proteins as determined by the intensity of protein bands (VHbH$_6$>P54R>Y29H>Y29H/P54R). Obviously, the double mutant (Y29H/P54R) was poorly expressed in comparison with others. This suggests that the position of the amino acid residue in the distal pocket is sensitive for mutation. Interestingly, iron superoxide dismutase (FeSOD), one of the most important antioxidant enzymes, was expressed in two populations, where one displayed an acidic character (nos. 4, 5, and 13) and another was more basic (nos. 14 and 23). It should be mentioned that the acidic (major) form of FeSOD was not found in E. coli TG1 host cells (data not shown). These major isoforms (nos. 4, 5, and 13) completely disappeared in cells expressing P54R variant, possibly due to their distinct properties. By contrast, they could be easily detected in other samples (nos. 4, 5, 13, 14, and 23). Transformation of acidic FeSOD isoform to a more basic form was observed only upon expression of the P54R mutant. In addition, flagellin protein (no. 18) emerged in cells expressing this variant, but it was undetectable in the others. No significant difference in the overall protein expression profile was disclosed among cells expressing wild-type VHb, Y29H, and double mutant.

**DISCUSSION**

Kinetics of ABTS Oxidation Catalyzed by VHb Proteins

The capability of VHb to catalyze peroxidase-like activity has recently been investigated with a variety of aromatic substrates [25, 38]. However, its reactivity is relatively low...
compared with the HRP. This may be attributable to the absence of catalytic prerequisite in its active site. Therefore, in the present study, an active site of typical peroxidase (containing catalytically conserved histidine and arginine residues) was placed into the distal pocket of VHb at positions 29 and 54, respectively (Fig. 1). Among all variants, single P54R substitution can remarkably increase $k_{\text{cat}}$ (~15-fold compared with wild-type protein) in the reaction with the $\text{H}_2\text{O}_2$ molecule in the presence of aromatic substrate. This result infers that the structural rigidity provided by Pro54 hinders the catalytic moiety. Such substitution can therefore make the active site more flexible, which probably favors the peroxidase reaction [6]. Additionally, the hydrophilic nature of residues lining around the distal site has been verified to be crucial for the interaction with polar substrates like $\text{H}_2\text{O}_2$ in all classes of peroxidase enzyme [42]. Introduction of arginine residue may in turn increase the hydrophilicity of the distal cavity where the substrate binding and catalysis take place [34, 37]. However its $K_m$ values for $\text{H}_2\text{O}_2$ and ABTS were reduced, suggesting that the introduced arginine may not be in a proper position for binding to the $\text{H}_2\text{O}_2$ molecule. This is surprising since results from the structure prediction demonstrated that the guanidinium group of Arg54 has been faced towards the heme cavity (Fig. 1). The distance to the iron atom of HRP appears to be ~5.7–6.0 Å [8], whereas it is only ~3.6 Å in the P54R variant. This proline residue has been replaced by alanine, and the results have

**Fig. 4.** Protein expression profile of *E. coli* expressing wild-type VHb electrophoresed at pH range of 3–10. Two individual areas showing differences in protein expression were chosen for protein identification A. Magnified pictures of selected areas from (B) cells expressing wild-type VHb (a), Y29H (b), P54R (c), and Y29H/P54R (d) variants. Some protein spots were picked and identified (numbers of protein spot designated as identified protein shown in Table 2). C. Types of identified proteins were mapped on gels according to their corresponding spot numbers.
shown that this substitution alters the cyanide binding property [43]. Taken together, replacement of the proline residue with a charged and longer amino acid such as arginine can presumably affect the structure of the distal pocket, resulting in the alteration in binding efficiency [34]. Meanwhile, variants harboring Y29H mutations appear to decrease in catalytic performances such as catalytic constant ($k_{cat}$), substrate affinity ($K_m$), and rate of compound I formation ($k_i$). A plausible explanation is that the His29 is unable to bind and stabilize the bound ligand or substrate because its imidazole side chain most likely projects out of the heme group (Fig. 1). Furthermore, the distance between the distal histidine and iron center is normally 5.5–6.0 Å for peroxidases [8, 17], whereas Y29H mutation enlarges this space to approximately 10.6 Å (Fig. 1), which is inappropriate to bind H₂O₂ substrate and unable to efficiently catalyze the reaction. Moreover, Matsui et al. [28] have constructed myoglobin mutants with varied distal histidine and found that the formation of compound I intermediate, catalytic rate, and anion binding capability are dramatically reduced in variant harboring distal histidine with a distance of 6.6 Å [28]. Our findings confirm such reports that the acid–base catalyst of distal histidine can be achieved only at an appropriate distance.

**Absorption Spectra of VHB Variants**

CO-binding spectra are basically employed to evaluate the oxidation state of hemoproteins [5, 27], since CO is able to bind only the ferrous (Fe²⁺) protein whose Soret peak is then shifted. Because of an appropriate electronic nature of low oxidation state iron center, the synergic $\pi^* \text{π}$ back bonding could be formed only between CO and ferrous protein [29], not the ferric or ferrous states [27]. As demonstrated in Fig. 3, all proteins were expressed in ferric form because their Soret bands remained unchanged after the addition of dithionite. Replacement of Tyr29 with a histidine residue caused substantial spectral shift in both ferric and ferrous states to 411 nm and 424 nm, respectively, as evidenced in the Y29H and Y29H/P54R variants. Since the phenolic side chain of Tyr29 is coordinated with the heme-iron through one water molecule and plays imperative roles in ligand binding [10, 40], substitution of this position may alter the coordination architecture in the distal pocket, leading to the shift in absorption spectra. Similarly, substitution of the distal histidine with an aliphatic residue such as leucine in myoglobin has been demonstrated to shift the $\lambda_{Soret}$ resulting from the loss of stabilizing residue to the heme-bound water molecule [28].

**Influence of Divergent VHB Variants on Protein Profilings of E. coli**

Although VHB protein has been shown to possess several functions in biological systems, involvement of this oxygen-binding protein in the cellular antioxidant system has not yet been reported. This study reports for the first time that expression of a high $k_{cat}$ P54R variant significantly down-regulates FeSOD expression, as demonstrated in Fig. 4 and Table 2, whereas no other alterations in protein profile can be detected in the other variants. This inhibitory effect is independent from the expression level of VHB proteins.

<table>
<thead>
<tr>
<th>Spot No.</th>
<th>Accession No.</th>
<th>Description</th>
<th>Calculated pI value</th>
<th>Nominal mass (M₀)</th>
<th>Protein scorea</th>
<th>Sequence coverage</th>
</tr>
</thead>
<tbody>
<tr>
<td>1, 2, 12, 17, 24</td>
<td>gi</td>
<td>170768574</td>
<td>Chaperone protein DnaK</td>
<td>4.83</td>
<td>69,160</td>
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<tr>
<td>3</td>
<td>gi</td>
<td>38491462</td>
<td>Chaperonin GroEL [Escherichia coli]</td>
<td>4.84</td>
<td>57,475</td>
<td>175</td>
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<tr>
<td>4, 5, 13, 14, 23</td>
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<td>640114</td>
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<td>5.58</td>
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<td>6</td>
<td>gi</td>
<td>170172436</td>
<td>Alkyl hydroperoxide reductase C22 protein (AhpC)</td>
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<td>19,250</td>
<td>101</td>
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<td>7</td>
<td>gi</td>
<td>15800262</td>
<td>Peptidyl-prolyl cis-trans isomerase B (rotamase B)</td>
<td>5.52</td>
<td>18,270</td>
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<tr>
<td>8</td>
<td>gi</td>
<td>15800564</td>
<td>DNA starvation/stationary phase protection protein, Dps</td>
<td>5.72</td>
<td>18,684</td>
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<tr>
<td>9, 10, 11, 16, 21, 22, 25</td>
<td>gi</td>
<td>114816</td>
<td>Bacterial hemoglobin (soluble cytochrome o)</td>
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<td>Range 81–89</td>
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<td>42543455</td>
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<td>86, 81</td>
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<td>Co-chaperonin GroES</td>
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<td>83</td>
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</table>

Protein scores were considered to be significant ($p<0.05$) if they were greater than 80. Protein scores and sequences coverage of multiple protein spots were given as the range of values.

**Table 2.** Summary of 2-DE analysis of *E. coli* TG1 expressing wild-type VHB or its variants identified by mass spectrometry and peptide mass fingerprinting (PMF).
FeSOD catalyzes the dismutation of superoxide anion (O\(^{2-}\)) into \(\text{O}_2\) and \(\text{H}_2\text{O}_2\); and its expression is often believed to be positively regulated by the ferric uptake regulator (fur) [13, 16]. Fur is the major regulatory protein whose function is to control many genes involved in iron uptake as well as FeSOD and MnSOD. Fe\(^{2+}\)-Fur, the active form of this transcription factor, is able to bind the promoter region of fur-controlled genes and then drives the down- or upregulation of such genes. The level of iron availability directly correlates with the level of Fe\(^{2+}\)-Fur active form and induction of FeSOD expression [13, 16]. The suppressed expression of FeSOD in cells expressing the VHbH6 P54R variant is likely to be due to the reduction of intracellular level of iron atoms as a result of lower \(\text{H}_2\text{O}_2\)-induced heme degradation and iron release [31, 32, 46]. \(\text{H}_2\text{O}_2\) can be produced by spontaneous dismutation of \(\text{O}^{2-}\), free radicals formed during autoxidation of hemoglobin [46]. A highly catalytic P54R variant is able to effectively eliminate reactive \(\text{H}_2\text{O}_2\) and therefore reduces the intracellular iron level. As a consequence, the active form of ferric uptake regulatory protein (Fe\(^{2+}\)-Fur) cannot be formed, leading to the suppression of FeSOD expression. This explanation coincides with foregoing study that catalase and glutathione peroxidase are capable of protecting red blood cells from \(\text{H}_2\text{O}_2\)-induced heme degradation [30]. In the other two variants containing distal histidine, this residue plays dual roles in the autoxidation process. At acidic condition, it promotes autoxidation by facilitating an active movement of catalytic proton from solvent to the bound dioxygen via its imidazole side chain. This process is known as proton-catalyzed autoxidation. At the same time, it can prevent the accessible FeO\(^{2+}\) center from the incoming water molecule, which readily binds to heme-bound iron atom and causes autoxidation, by forming the hydrogen bond [39]. Thus, if the distal histidine residue is tilted away from the heme pocket, the rate of this reaction is more pronounced, as found in the \(\beta\)-chain of human hemoglobin [41]. As illustrated in Fig. 1, the side chain of His29 is positioned improperly to stabilize dioxygen in the distal cavity, and therefore, its protective effect cannot be achieved. Then, the improperly positioned distal histidine raises the radical-producing autoxidation level, leading to the increased release of iron from heme degradation. This higher iron availability ensures the expression of FeSOD via Fur-mediated regulation. In contrast, the optimal position of an introduced Arg54 can perhaps provide efficient stabilization of bound dioxygen through hydrogen bonding and consequently hinder the autoxidation [2]. The level of superoxide production and heme degradation is thus reduced in cells expressing this variant. This may decrease the proportion of Fur active form and thereby cause suppression of FeSOD expression (Fig. 4). Our findings open up a high feasibility of applying the P54R variant to rescue cells expressing VHb protein from the susceptibility to \(\text{H}_2\text{O}_2\) [19]. Furthermore, this variant seems to be more applicable in large-scale fermentation, because a high rate of oxygen consumption and cellular metabolism provided by VHb probably makes cells produce more reactive oxygen species (ROS) that impede the productivity of desirable compounds. In conclusion, a highly catalytic variant of VHb (P54R) possessing ~15-fold increase in \(k_{\text{cat}}\) has successfully been constructed based on mimicking of peroxidases active sites. Replacement of Pro54 with an arginine residue contributes to the relaxed nature and higher hydrophilicity of the active site, which thereby effectively facilitates enzymatic catalysis [34, 42]. However, neither Y29H nor P54R mutation was able to promote the affinity to both peroxidase substrates. Interestingly, expression of FeSOD is suppressed in cells expressing the P54R mutant, suggesting that oxidative stress may be less produced. Potent peroxidase activity and lower autoxidation of this variant are proposed to decrease the iron availability and consequently control the Fe\(^{2+}\)-Fur level, an active transcription regulator of the FeSOD gene. \(\text{H}_2\text{O}_2\) generated during autoxidation can be degraded rapidly by this catalytically proficient P54R variant, and it is likely that the side chain of arginine may stabilize bound dioxygen to hamper the autoxidation [2]. This superior variant of VHb can be used in a variety of applications such as a metabolic engineering tool in industrial fermentation [45, 50], a labeled enzyme in immunological assay [38], a biocatalyst in depollution of waste water [4, 9, 33], an electrochemical-redox sensor [14], and a biosensor for \(\text{H}_2\text{O}_2\) determination [47–49].

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