Structure prediction and activity analysis of human heme oxygenase-1 and its mutant

Zhen-Wei Xia, Wen-Pu Zhou, Wen-Jun Cui, Xue-Hong Zhang, Qing-Xiang Shen, Yun-Zhu Li, Shan-Chang Yu

INTRODUCTION

Heme oxygenase (HO) is responsible for the physiological breakdown of heme into equimolar amounts of biliverdin, carbon monoxide, and iron. Three isoforms (HO-1, HO-2, and HO-3) have been identified. HO-1 is ubiquitous, its mRNA levels and activity can be increased several-fold by heme, other metalloporphyrins, transition metals, and stress-inducing stimuli. In contrast, HO-2 is present chiefly in brain and testes and is virtually uninducible. HO-3 has very low activity; its physiological functions probably include heme binding. The HO system has been strongly highlighted for its potential significance in maintaining cellular homeostasis. Nevertheless the physiological correlations of the three isoforms and their reciprocal interrelation have been poorly understood.

HO-1 regulates the levels of serum bilirubin as the rate-limiting enzyme in heme degradation pathway. Recent reports showed that HO-1 was identified as an ubiquitous stress protein and it had important physiological roles. Overexpression of HO-1 gene resulted in protection from cytokine-induced oxidative stress[5], inflammation[6-9], apoptosis[10-17] and proliferation[18-25]. It has been reported that histidine (His) residues at positions 25, 84, 119 and 132 in HO-1 sequence are conserved in rat, human, mouse and chicken. These histidines may be important for heme-binding[26]. His25 and His32 mutants were reported for the proximal heme iron ligand in rat heme oxygenase-1 (rHO-1)[27]. The unambiguous spectroscopic demonstration that His25 is the proximal iron ligand leaves the role of His32 uncertain. His 25 is essential for heme deactivation activity of the enzyme. The research on human HO-1 (hHO-1) structure has been shown that hHO-1 embodies a novel protein fold that consists primarily of α-helices, and the heme is held between two of these helices[28].

It is unclear whether hHO-1 mutant has the same characteristics and displays catalytic inactivity but binding heme as Ala replacing His 25. In this study, the characteristics of wild hHO-1 (whHO-1) and its mutant were predicted by bioinformatics method. On the basis of the results, the truncated hHO-1 cDNA mutant was constructed by site-directed mutagenesis. Two expression plasmids, pBH1-0 and pBH1-0(M) containing whHO-1 and hHO-1 His25Ala mutant (△H)HO-1, respectively, were constructed and expressed in E.coli, and then isolated, purified and analyzed of their activities.
Methods
Structure analysis of hHO-1 and its mutant

Swiss-PdbViewer and Anthropt 5.0 were used to analyze the structure diversity and physical-chemical changes between whHO-1 and ΔhHO-1. The phi prime structure was refined by energy minimization. Gromos96 was used as force field. The molecular surfaces and electrostatic potential were calculated. The rationality of the resulted model was validated by Ramachandran plot.

Construction of ΔhHO-1 expression vector pBHO-1 (M)

EcoRI/Sall digested fragment of pBHO-1, which is from expression vector containing the truncated hHO-1 cDNA (804 bp), was cloned into the EcoRI/Sall sites of plasmid pBSKS II to construct plasmid pBSHO-1. hHO-1 His25Ala mutant cDNA could be amplified by PCR using pBHO-1 as template. The mutation primer (5'-GACAGCATGCCTCGAGGTGAGCTCAGGCGCTTTAGGCAGCGATCC-3') and reverse primer (5'-AACAGCTATGACCATG-3') were chosen, which have changed the His 25 codon CAC into Ala codon GCC. The reaction conditions were: at 94 °C for 5 min, followed by 94 °C for 1 min, at 55 °C for 1 min, 72 °C for 2 min, with amplification repeated for 30 cycles, and finally at 72 °C for 10 min. The mutant cDNA was gel-purified and digested with Sphl/Sall and then cloned into Sphl/Sall sites of plasmid pGEM3Z. hHO-1 His25Ala mutant cDNA was screened by restriction digestion and confirmed by sequencing in pGEM3. hHO-1 His25Ala mutant cDNA was obtained from Sphl/Sall digested fragment of pGEM3ZHO-1 (M) and then cloned into pBHO-1, thus the expression vector containing ΔhHO-1 was designated as pBHO-1 (M) (Figure 1).

Expression and purification of whHO-1 and ΔhHO-1

A total of 5.0 mL inoculum was set up from plates with fresh colonies of transformed E. coli DH5α and incubated at 37 °C for 12-16 h. From the fresh cultures 400 µL was used to inocuate into 40 mL cultures of the same media. The cells were grown at 37 °C until A₅₆₂ reached 0.3-0.5, then added with 0.5 mmol/L IPTG for 18 h. The expression products were harvested and analyzed with sodium dodecyl sulfate-polyacrylamide gel (SDS-PAGE) and Western blotting at the time points of 7, 9 and 11 h, respectively, after addition of IPTG. Then 500 mL inoculum in medium was incubated under same condition. The harvested cells were centrifuged at 6000 rpm for 20 min at 4 °C, washed once in PBS, and then centrifuged at 6000 rpm for 10 min at 4 °C. The cells were lysed in 50 mmol/L Tris buffer (pH 8.0) containing 1 mmol/L dithiothreitol, 1 mmol/L EDTA, 1 mmol/L phenylmethanesulfonyl fluoride (PMSF) and sonicated for 10 min. Then the cells were centrifuged at 13 000 rpm for 1 h at 4 °C. The supernatant was collected and ammonium sulfate was added to a final concentration of 30%, and the solution was stirred for 60 min. Following centrifugation (13 000 rpm for 20 min), ammonium sulfate concentration was raised to 60% of saturation. The pellets precipitated by ammonium sulfate were collected and resuspended in 0.1 mol/L potassium phosphate (pH 7.4) and then dialyzed against 1 g/L NH₄HCO₃ for 4 h (×4-times). Protein 130 mg was applied to a Q-Sepharose Fast Flow anion column and eluted with a step gradient of 50 mmol/L Tris-HCl (pH 7.4) (buffer A) containing 0-0.5 mol/L NaCl (buffer B). The fractions containing hHO-1 protein were pooled together and applied to Q-Sepharose Fast Flow anion column again. The protein was eluted with a step gradient of 50 mmol/L Tris-HCl (pH 8.4) (buffer C) containing 0-0.5 mol/L NaCl (buffer B). The gradient was increased linearly from 0-100% buffer B. Fractions containing hHO-1 protein were pooled and dialyzed against 1 g/L NH₄HCO₃ for 4 h (×4-times).

Western blotting
Blot analysis was carried out as previously described[29]. Microsomal protein samples were fractionated by SDS-PAGE under denaturing conditions. The separated proteins were electrophoretically transferred to a nitrocellulose membrane. Western blotting was carried out using monoclonal antibody to hHO-1 and the immunoreactive bands were visualized by staining (1 mol/L Tris-HCl pH 9.5, 1 mol/L MgCl₂, 0.1 mg/mL nitro-blue tetrazolium, 0.1 mg/mL 5-bromo-4-chloro-3-indolylphosphate-toluidine salt).

Analysis of whHO-1 and ΔhHO-1 activities

Protein samples were incubated with heme (50 µmol/L), rat liver cytosol (5 mg/mL), MgCl₂ (2 mmol/L), glucose-6-phosphate dehydrogenase (1 unit), glucose-6-phosphate (2 mmol/L), and NADPH (0.8 mmol/L) in 0.5 mL of 0.1 mol/L potassium phosphate buffer (pH 7.4), for 60 min at 37 °C. Reaction was stopped by putting the tubes on ice, and reaction solution was extracted with chloroform. The rate of bilirubin formation was monitored at 464 nm by a spectrophotometer and then calculated using an extinction coefficient of 40.0 mmol/L (L.cm)[29].

RESULTS

Whole structure comparison between whHO-1 and ΔhHO-1

In heme-heme oxygenase-1 complex, heme is sandwiched in whHO-1. When Ala 25 replaces His 25, the complex structure does not change. In the heme-binding pocket, Ala 25 loses contacting with heme as His 25 does. The molecular surface has a catalytic reaction pocket, which includes Thr 21, His 25, Ala 28, Glu 29, Gly 139, Asp 140, Gly 143. After binding heme, His 25 still lies on the surface but Gly139, Asp140 and Gly143 are covered by heme. When Ala 25 replaces His 25, Ala 25 still lies in the surface (Figure 2). In activity domain, the bond angle, dihedral angle and chemical bond appear differently after Ala 25 replacing His 25 (Table 1, Figure 3).
Table 1  Bond angle, dihedral angle and the distance between atoms of $\Delta$hHO-1

| Bond angle   | Prime simulation | Optimized simulation |
|--------------|------------------|---------------------|
| C24-N-CA     | 119.12           | 121.79              |
| CA-C-N26     | 115.92           | 115.12              |
| Distance     | CB-FE            | 5.55                | 5.58                |
|              | CA-FE            | 6.28                | 6.26                |
| Dihedral angle | $\omega$       | 175.40              | 175.61              |
|              | $\phi$           | -40.76              | -34.45              |
|              | $\psi$           | -60.70              | -62.89              |

Though Ala 25 replaces His 25, the molecular surfaces and electrostatic potential changed little (Figure 4). The electrostatic potential of active pocket was still negative. The mutagenesis had no apparent effect on molecular surface. Ramachandran plot showed that dihedral angles $\phi$ (-33.98) and $\psi$ (-64.46) were in rational range.

Antheprot 5.0 analysis also showed that there was no alteration in the secondary structure between whHO-1 and $\Delta$hHO-1. Garnier, Gibrat, DPM and homology predicted the same results. Physical-chemical characteristics showed somewhat alteration. Hydrophobicity increased, while hydrophilicity decreased. There was no change in antigenicity, helical membranous regions and solvent accessibility.

Figure 2  Prime simulated structure of whHO-1 and $\Delta$hHO-1. Red: heme; Black: His 25 and Ala 25. Ala 25 loses contacting with heme as His 25 does in the heme-binding pocket.

Figure 3  Changes of bond angle, dihedral angle and chemical bond.

Figure 4  Molecular surfaces and electrostatic potential of whHO-1 and $\Delta$hHO-1. The electrostatic potential of active pocket is $-1.800$.

Construction of pBHO-1(M) containing $\Delta$hHO-1

hHO-1 cDNA was site mutated at His 25 (to Ala) by PCR with pBSHO-1 as the template. The 866-bp PCR product showed that nucleotide 3-773 sequences encoded hHO-1 domain from 25-265 AA. The mutant cDNA was cloned into the SphI/SalI sites of plasmid pGEM3Z for constructing plasmid pGEM3ZHO-1 (M). hHO-1 His25Ala mutant cDNA was confirmed by sequencing in pGEM32HO-1(M). The verified hHO-1 mutant cDNA from the SphI/SalI digested fragment of pGEM32ZHO-1 (M) was cloned into pBHO-1, thus the expression vector containing $\Delta$hHO-1 was designated as pBHO-1 (M) (Figure 1). Both whHO-1 and $\Delta$hHO-1 cDNAs equally encoded the proteins containing 265 amino acids with a $M_r$ 30 500.

Expression and identification of whHO-1 and $\Delta$hHO-1 in E.coli DH5$\alpha$

E.coli DH5$\alpha$ was transformed by pBHO-1 and pBHO-1 (M), respectively, and treated with 0.5 mmol/L IPTG for 18 h at 37 . Equal quantities of cells transformed with different expression vectors were lysed by protein electrophoresis buffer. Untransformed E.coli sample was used as the negative control. As shown in Figure 5, E.coli DH5$\alpha$ transformed pBHO-1 or pBHO-1 (M) highly expressed whHO-1 and $\Delta$hHO-1 with a $M_r$ 30 500. Meanwhile, E. coli DH5$\alpha$ not treated with IPTG also expressed whHO-1 and $\Delta$hHO-1, but its expression yield
was significantly lower than that of transformed cells. Analysis of cell lysates showed that whHO-1 and ΔhHO-1 were mainly present in the supernatants.

**Figure 5** Western blotting of whHO-1 and ΔhHO-1 expressed in DH5α. Lanes 1, 2: Expression products of pbHO-1 and pbHO-1(M) in DH5α induced with IPTG; lane 3: Control; lanes 4, 5: Expression products of pbHO-1 and pbHO-1(M) in DH5α not treated with IPTG; lane 6: Marker.

**Purification of whHO-1 and ΔhHO-1**

The protein with a Mr 30 500 was purified by 30-60% ammonium sulphate precipitation and analyzed by SDS-PAGE.

After precipitated with ammonium sulphate, whHO-1 sample was applied to Q-Sepharose Fast Flow anion column (pH 7.4). The first peak containing whHO-1 (No. 1-3 tubes) was collected (Figure 6A) and loaded on Q-Sepharose Fast Flow anion column (pH 8.4) again. whHO-1 was shown to be eluted in the second peak (No. 49-63 tubes) (Figure 6B). All samples were analyzed by SDS-PAGE and Western blotting in order to identify the separation efficiency. The activity of whHO-1 after purification was 30-fold higher than that in the initial lysates (Table 2).

**Figure 6** Q-Sepharose Fast Flow column chromatography. A: pH 7.4 buffer; B: pH 8.4 buffer.

Precipitated ΔhHO-1 was further purified in the same way as described above. One purified protein’s Mr was 30 500. The purified whHO-1 and ΔhHO-1 are shown on SDS-PAGE (Figure 7).

**Table 2** whHO-1 activities after different purification

| Supernatant of DH5α lysates | 30-60% (NH₄)₂SO₄ | Q-Sepharose fast flow |
|-----------------------------|------------------|----------------------|
| whHO-1 activity (U·mg⁻¹·h⁻¹) | 0.5              | 1.8                  | 15                   |
| Purification fold           | 1.0              | 3.6                  | 30                   |

**Figure 7** SDS-PAGE analysis of purified whHO-1 and ΔhHO-1. Lane 1: Marker; lane 2: whHO-1; lane 3: ΔhHO-1.

**DISCUSSION**

An understanding of the structure and function of hHO-1 is of importance in the context of human diseases, but due to limited accessibility and difficulties associated with their purification from microsomal membranes, little direct information has been available until the expression of rHO-1 in E. coli was achieved. The protein thus obtained is soluble, easily purified, and fully active. Encouraged by this result, the research on HO-1 including hHO-1 has been thriving.

hHO-1 is anchored to the endoplasmic reticulum membrane via a stretch of hydrophobic residues at the C-terminus. While full-length hHO-1 consists of 288 residues, a truncated version with residues 1-265 has been expressed as a soluble active enzyme in E. coli. This recombinant enzyme precipitated from ammonium sulfate solution but without a high purity for studies.

In the present studies, we acquired the structures of hHO-1 with residues 1-265 and its mutant using Swiss-PdbViewer. Due to the high similarity between hHO-1 and its mutant, the hHO-1 structure was used as template. Through homology modeling, template selection and target-template alignment, a molecular backbone and side chains were built in one step. Results demonstrated that the refined model, as well as the entire assay, was valid. Predictions of the structure showed that hHO-1 and its mutant had similar electrostatic potential suggesting that they have similar affinity to heme in electrostatic potential. Ala is a hydrophobic amino acid without chemical activity, and its side chain is short. So it has no interference with binding through steric hindrance.

Rat His25Ala HO-1 provides a new approach to construct hHO-1 mutant. hHO-1 mutants were prepared in which residue His25 was replaced by Ala using site-directed mutagenesis. Due to the high similarity between hHO-1 and its mutant, the hHO-1 structure was used as template. Through homology modeling, all templates were selected and target-template alignment, a molecular backbone and side chains were built in one step. Results demonstrated that the refined model, as well as the entire assay, was valid. Predictions of the structure showed that hHO-1 and its mutant had similar electrostatic potential suggesting that they have similar affinity to heme in electrostatic potential. Ala is a hydrophobic amino acid without chemical activity, and its side chain is short. So it has no interference with binding through steric hindrance.

Our study shows that precipitation with ammonium sulfate can enrich whHO-1 and ΔhHO-1 in the lysates. Compared with the whHO-1 activity, ΔhHO-1 activity was reduced by
91.21%. It suggests that His25 is crucial for the catalytic activity of \(\text{whHO-1}\), the proximal His25 ligand is critically required for normal h\(\text{HO-1}\) catalysis. The results are similar to previous reports.

\(\text{hHO-1}\) has anti-inflammatory\(^{6-9}\), anti-apoptotic\(^{10-17}\), and anti-proliferative\(^{18-25}\) effects, and salutary effects on neonatal hyperbilirubinemia\(^{30}\). The results show a valid way to control antiproliferative effects, and salutary effects on neonatal reports\(^{31-35}\).

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Edited by Chen WW and Zhu LH. Proofread by Xu FM