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Proton Nuclear Magnetic Resonance Investigation of the Native and Modified Active Site Structure of Heme Proteins

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PROTON NUCLEAR MAGNETIC RESONANCE INVESTIGATION OF THE
NATIVE AND MODIFIED ACTIVE SITE STRUCTURE OF HEME PROTEINS

A dissertation submitted in partial fulfillment of the
requirements for the degree of
DOCTOR OF PHILOSOPHY
in
CHEMISTRY
by
Zhonghua Wang

2011
To:  Dean Kenneth Furton  
College of Arts and Sciences

This dissertation, written by Zhonghua Wang, and entitled Proton Nuclear Magnetic Resonance Investigation of the Native and Modified Active Site Structure of Heme Proteins, having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this dissertation and recommend that it be approved.

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Date of Defense: October 5, 2011

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University Graduate School

Florida International University, 2011
DEDICATION

I dedicate this dissertation to my wife, Taiyi Chen, my mother, Fenglan Jiang and my father, You Wang. Without their understanding, encouragement, support, and love, the completion of this work would not have been possible.
ACKNOWLEDGMENTS

I am deeply grateful to many people that have helped me during my delightful and rewarding graduate study at Florida International University in the past few years. First, I would like to thank my research advisor, Dr. Xiaotang Wang, who has been not only an intelligent and respectable mentor, but also a close and thoughtful friend to me. His patience, guidance, and encouragement strongly supported me through my graduate studies and have offered me a great foundation to build a career in science. I would also like to thank all my committee members, Dr. Watson Lees, Dr. Fenfei Leng, Dr. Kathleen Rein and Dr. Lidia Kos, for all of their constructive advice and professional assistance. Thanks to all the members of Dr. Wang’s group, especially Wei Lin, Zheng Wang, Lin Jiang, Hui Tian, Rui Zhang, Hua Ling, Dr. Hedong Bian, and Dr. Yuchang Jiang, for all of the inspiring discussions, selfless help, and for creation of a pleasant lab environment. I would like to extend my appreciation to Yali Hsu and Dr. Yaru Song for their enthusiastic help and careful maintenance of the NMR and MS instruments. Special thanks go to Dr. Palmer Graves and Dr. Uma Swamy for all their kindness help and guidance to teach me how to be a good teaching assistant. I would like to especially acknowledge Dr. Chengxiao Zhang and Dr. Peng Lu for all their care, support and encouragement to me during the years. I also want to thank all of my family and friends for their continuous support, understanding, and encouragement. Finally, I would like to thank the Department of Chemistry and Biochemistry, Florida International University, the National Institutes of Health, and the National Science Foundation for the assistanship support and the funding support during my graduate studies.
ABSTRACT OF THE DISSERTATION

PROTON NUCLEAR MAGNETIC RESONANCE INVESTIGATION OF THE
NATIVE AND MODIFIED ACTIVE SITE STRUCTURE OF HEME PROTEINS

by

Zhonghua Wang

Florida International University, 2011

Miami, Florida

Professor Xiaotang Wang, Major Professor

Hemoproteins are a very important class of enzymes in nature sharing the essentially same prosthetic group, heme, and are good models for exploring the relationship between protein structure and function. Three important hemoproteins, chloroperoxidase (CPO), horseradish peroxidase (HRP), and cytochrome P450cam (P450cam), have been extensively studied as archetypes for the relationship between structure and function. In this study, a series of 1D and 2D NMR experiments were successfully conducted to contribute to the structural studies of these hemoproteins.

During the epoxidation of allylbenzene, CPO is converted to an inactive green species with the prosthetic heme modified by addition of the alkene plus an oxygen atom forming a five-membered chelate ring. Complete assignment of the NMR resonances of the modified porphyrin extracted and demetallated from green CPO unambiguously established the structure of this porphyrin as an $N_{III}$-alkylated product. A novel substrate binding motif of CPO was proposed from this concluded regiospecific N-alkylation structure.

Soybean peroxidase (SBP) is considered as a more stable, more abundant and less expensive substitute of HRP for industrial applications. A NMR study of SBP using 1D
and 2D NOE methods successfully established the active site structure of SBP and consequently fills in the blank of the SBP NMR study. All of the hyperfine shifts of the SBP-CN$^-$ complex are unambiguously assigned together with most of the prosthetic heme and all proximal His170 resonances identified. The active site structure of SBP revealed by this NMR study is in complete agreement with the recombinant SBP crystal structure and is highly similar to that of the HRP with minor differences.

The NMR study of paramagnetic P450cam had been greatly restricted for a long time. A combination of 2D NMR methods was used in this study for P450cam-CN$^-$ complexes with and without camphor bound. The results lead to the first unequivocal assignments of all heme hyperfine-shifted signals, together with certain correlated diamagnetic resonances. The observed alternation of the assigned novel proximal cysteine $\beta$-CH$_2$ resonances induced by camphor binding indicated a conformational change near the proximal side.
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LIST OF ABBREVIATIONS

1D  One dimensional
2D  Two dimensional
AB  Allylbenzene
ABTS 2,2’-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) diammonium salt
cam  Camphor
CcO  Cytochrome c oxidase
CcP  Cytochrome c peroxidase
CMP  Coprinus macrorhizus peroxidase
CN  Cyanide
CooA  CO-sensing transcriptional activator
COSY  Correlation spectroscopy
Cpd 0  Compound 0
Cpd I  Compound I
CPO  Chloroperoxidase
CW  Q-band continuous wave
CYP  Cytochrome P450
DEAE  Diethylaminoethyl (cellulose)
DMSO  Dimethyl sulfoxide
DTT  Dithiothreitol
EDTA  Ethylenediaminetetraacetic acid
ENDOR  Pulsed electron nuclear double resonance spectroscopy
EPR  Electron paramagnetic resonance
<table>
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<tr>
<th>Acronym</th>
<th>Description</th>
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<tr>
<td>FPS</td>
<td>Fructose Peptone Salts</td>
</tr>
<tr>
<td>HPLC</td>
<td>High-performance liquid chromatography</td>
</tr>
<tr>
<td>HRP</td>
<td>Horseradish peroxidase</td>
</tr>
<tr>
<td>HRP-C</td>
<td>Horseradish peroxidase C isoenzyme</td>
</tr>
<tr>
<td>IPTG</td>
<td>Isopropyl β-D-1-thiogalactopyranoside</td>
</tr>
<tr>
<td>KCN</td>
<td>Potassium cyanide</td>
</tr>
<tr>
<td>MALDI</td>
<td>Matrix-assisted laser desorption/ionisation</td>
</tr>
<tr>
<td>MCD</td>
<td>Monochlorodimedone</td>
</tr>
<tr>
<td>MnP</td>
<td>Manganese peroxidase</td>
</tr>
<tr>
<td>N-ABPP</td>
<td>N-allylbenezene-protoporphyrin adduct</td>
</tr>
<tr>
<td>NADH</td>
<td>Nicotinamide adenine dinucleotide, reduced</td>
</tr>
<tr>
<td>NADPH</td>
<td>Nicotinamide adenine dinucleotide phosphate, reduced</td>
</tr>
<tr>
<td>N-GfPP</td>
<td>N-griseofulvin-protoporphyrin</td>
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<tr>
<td>N-MePP</td>
<td>N-methylprotoporphyrin</td>
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<tr>
<td>NMR</td>
<td>Nuclear magnetic resonance</td>
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<tr>
<td>NOE</td>
<td>Nuclear Overhauser effect</td>
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<tr>
<td>NOESY</td>
<td>Nuclear Overhauser enhancement spectroscopy</td>
</tr>
<tr>
<td>NOS</td>
<td>Nitric oxide synthase</td>
</tr>
<tr>
<td>P450</td>
<td>Cytochrome P450</td>
</tr>
<tr>
<td>PDA</td>
<td>Potato Dextrose Agar</td>
</tr>
<tr>
<td>PMSF</td>
<td>Phenylmethylsulfonyl fluoride</td>
</tr>
<tr>
<td>rSBP</td>
<td>Recombinant SBP</td>
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<tr>
<td>Rz</td>
<td>Reinheitzahl value</td>
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<tr>
<td>SBP</td>
<td>Soybean peroxidase</td>
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<td>------------------------------------------------</td>
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<tr>
<td>SDS</td>
<td>Sodium dodecyl sulfate</td>
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<tr>
<td>TOCSY</td>
<td>Total correlation spectroscopy</td>
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<tr>
<td>TOF</td>
<td>Time of flight</td>
</tr>
<tr>
<td>Tris</td>
<td>Tris(hydroxymethyl)aminomethane</td>
</tr>
<tr>
<td>WT</td>
<td>Wild type</td>
</tr>
<tr>
<td>Ala</td>
<td>Alanine</td>
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<tr>
<td>Arg</td>
<td>Arginine</td>
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<td>Asn</td>
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<td>Trp</td>
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<td>Val</td>
<td>Valine</td>
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CHAPTER I. LITERATURE REVIEW

1.1 Introduction to Hemoproteins

Hemoproteins (EC 1.11.1.7) are one of the largest classes of metalloproteins that are ubiquitous and essential for every organism and are involved in a wide range of biological processes while retaining a basically unaltered prosthetic group consisting of an iron-protoporphyrin IX (heme). Hemoproteins are therefore good models for exploring the relationship between protein structure and function. However, after over eighty years of research, the relationship between the various porphyrin architectures and the biochemical function of hemoproteins remains poorly defined.

The biological functions of hemoproteins include: electron transfer (b- and c-type cytochromes), oxygen transport (hemoglobin) and storage (myoglobin), oxygen reduction (cytochrome oxidases), organic substrates oxygenation (cytochromes P450), small molecules sensing (CooA), and peroxides reduction (catalases and peroxidases) (1-2). Studies of hemoproteins have suggested that the flexibility in hemoprotein functions normally arises from a combination of differences in both heme constituents and the polypeptide of the various hemoproteins (3).

Concerning the prosthetic heme group, there are four common representatives, that contain a planar iron-tetrapyrrole ring system with different side chains (Figure 1.1). Heme \( a \) provides cofactors for the cytochrome \( a_{a3} \) of cytochrome \( c \) oxidase (CcO), an important respiratory protein (4). Heme-\( b \) (also termed iron-protoporphyrin IX) is the most abundant and important among these four hemes, since heme \( b \) forms the prosthetic group of hemoglobin, myoglobin, catalase, most peroxidases, the b-type cytochromes and the cytochrome P-450s. Heme \( c \) is the prosthetic group for the c-type
cytochromes, which are the most extensively studied family of electron transfer proteins (5). Heme $d_i$ is found as a prosthetic group in the heme-containing nitrite reductases designated as cytochromes $cd_i$ (6). Since heme $b$ is the only prosthetic group of all the hemoproteins related to my work, I will limit my discussion to enzymes containing a heme $b$.

Heme $b$ has eight side chains which are conventionally numbered as position 1-8 in a clockwise sense as shown in Figure 1.1 B: four methyl groups located at positions 1, 3, 5, and 8, two vinyl groups at positions 2 and 4, and two propionate groups at positions 6 and 7. The carbon atoms of the methine bridges are named separately from the side chains as $\alpha$, $\beta$, $\gamma$, and $\delta$, which results in the hydrogen on each methine bridge carbon named as meso-$\alpha$, $\beta$, $\gamma$, and $\delta$ proton, respectively. The four pyrrole rings are, clockwise, named as pyrrole ring I, II, III and IV. The iron atom is placed in the center of the heme with the four pyrrole nitrogens equatorially ligated. In hemoproteins, the heme iron normally has two axial ligands binding sites which are located above (termed distal side) and below the porphyrin plane (termed proximal side). The importance and function of these two axial ligands will be discussed later.

Heme $b$ in hemoproteins can exist as two rotational isomers called isomers A and B that differ by a $180^\circ$ rotation of the heme group about the $\alpha$-$\gamma$ axis, and leads to different protein-heme contacts. For isomer A, the porphyrin substituents are arranged in a clockwise sense when viewed from the proximal side of proteins, such as chloroperoxidase (CPO) and horseradish peroxidase (HRP). For isomer B, the porphyrin substituents are arranged in a counterclockwise sense, such as cytochrome $c$ peroxidase (CcP) and manganese peroxidase (MnP). Two-fold rotation about the
Figure 1.1 Hemes commonly found in biological systems.
\(\alpha-\gamma\) axis interchanges the 2- and 4-vinyl groups with the 3- and 1-methyl groups, respectively. Studies by NMR have shown that the equilibrium ratio of A to B form depends on the protein sequence, and the oxidation state and the ligands of the heme (7).

Concerning the polypeptide part of the molecule, it is obvious that the protein structure around the heme center is of great importance in controlling the type of reaction catalyzed, and directing the heme to a prescribed function specifically and efficiently. The environment surrounding the heme iron in hemoproteins dictates their activity and makes them a diverse group of enzymes. In particular, axial coordination to the central iron atom, covalent attachment of the heme to the protein, H-bonding interactions, conformations, and amino acid side chains in the immediate vicinity of the active site, are all of dominant importance in governing the redox potentials and enzymatic functions.

The presence and the nature of the fifth and sixth axial ligands to the iron will be crucial in determining the reactivity of hemoproteins. For example, in oxidases or oxygenases which need to bind oxygen or hydrogen peroxide, it is obviously of paramount importance to have a coordination site (distal site) available for exogenous ligands. At the same time, the proximal site will be occupied by heteroatoms from nucleophilic amino acid residues. The distal position in these hemoproteins is normally the catalytic center and its coordination state varies during the catalytic cycle. Meanwhile the proximal ligand is highly conserved and is used as an important indication to differentiate each hemoprotein families. For example, in all heme peroxidases except chloroperoxidase, the proximal ligand is always a histidine nitrogen; while in catalase, it is a tyrosine oxygen, and in all P450s and CPO, it is a cysteinate sulfur serving as the proximal ligand. In contrast, the hemoproteins involved in simple electron transfers,
which do not directly bind exogenous ligands but need to avoid reorganization at the heme iron site, are therefore almost invariably hexa-coordinated with the iron in the low-spin state with both axial sites occupied by heteroatoms from the peptide backbone amino acids (8). Because of the variety and complexity of hemoproteins, I will only further introduce several hemoproteins which are closely related to my researches in this chapter.

1.2 Heme-Thiolate Proteins

As mentioned above, the proximal heme ligand of hemoprotein is an important indication for classification. Therefore, hemoproteins using a thiolate anion from a deprotonated cysteine residue as the proximal ligand are grouped as heme-thiolate proteins, in contrast with another major hemoprotein category, the heme-imidazole proteins, whose proximal ligand is from a histidine imidazole nitrogen.

The known heme-thiolate proteins currently include all cytochromes P450, CPO, nitric oxide synthases (NOS), cystathionine β–synthase, sensor protein CooA and eIF2α kinase. With the same proximal ligand, heme-thiolate proteins exhibit highly versatile, diverse and essential functions in biological systems. The remainder of this chapter will focus on the structures, functions, and mechanisms of cytochromes P450 and CPO that constitute the major part of my dissertation research.

1.2.1 Cytochromes P450

Cytochromes P450 (CYP) are a family of monooxygenases (mixed function oxidases) named for their characteristic absorption maximum (the Soret peak) at around 450 nm upon binding of carbon monoxide by the reduced protein. The title of “cytochrome” is earned because of the electron transfer functionality and colorful character of the P450 enzymes. Cytochrome P450 monooxygenases require only two electrons and two protons
to reductively cleave atmospheric dioxygen, producing a single water molecule in the process while saving the second oxygen atom for substrate functionalization and formal oxidation (Equation 1). This two-electron reduction process is in contrast with dioxygenases which need the full four-electron reduction of dioxygen to generate two water molecules (9). Electrons that are required for the catalysis were found to come from NADH or NADPH via electron transfer proteins, such as cytochrome P450 ferredoxin and cytochrome P450 reductase.

\[
\text{R-H} + \text{O}_2 + 2e^- + 2\text{H}^+ \rightarrow \text{ROH} + \text{H}_2\text{O} \tag{1}
\]

Ever since the first P450 was discovered in 1958 (10-11), massive amount of effort has been put into studying the ubiquitous P450s for over half a century. Many reviews (9, 12-22) and books (23-28) have been devoted to the superfamily of P450s as catalyst. There are two main features that attracted the interest of chemists: (i) the P450s can catalyze the selective monooxygenation of a wide variety of organic substrate, and (ii) the P450s play a key role in the oxidative metabolism of drugs and other xenobiotics including steroid hormones, which is a crucial step in the adaptation of living organisms to their always changing chemical environment (18). The intense research performed on P450s has revealed more than 11,500 distinct proteins that are spread across almost all living organisms (9). All P450s have the highly conserved cysteine thiolate acting as the heme proximal ligand, which is considered to play critical roles in the catalytic mechanisms. Although the sequence homology between P450 members from different gene sources can be as low as ~20% on the amino acid level, their overall structures have high similarity.
The P450s have revealed a high diversity of biological functions, catalyzed reactions, and acceptable substrates (19, 21-22). The biological functions of P450s include the biosynthesis and biodegradation of important endogenous compounds and the metabolism of xenobiotics as mentioned above. The reactions catalyzed by P450s are listed in Table 1.1. The substrates for P450s include fatty acids, steroids, prostaglandins, as well as a multitude of foreign compounds such as drugs, anaesthetics, organic solvents, ethanol, alkylaryl hydrocarbon products, pesticides, and carcinogens.

Table 1.1 Reactions catalyzed by cytochromes P450 (14, 19, 22, 29).

<table>
<thead>
<tr>
<th>Reaction</th>
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<tbody>
<tr>
<td>Hydrocarbon hydroxylation</td>
<td>Alkene epoxidation</td>
</tr>
<tr>
<td>Alkyne oxygenation</td>
<td>Arene epoxidation</td>
</tr>
<tr>
<td>Aromatic hydroxylation</td>
<td>N- Dealkylation</td>
</tr>
<tr>
<td>S- Dealkylation</td>
<td>O- Dealkylation</td>
</tr>
<tr>
<td>N- Hydroxylation</td>
<td>N- Oxidation</td>
</tr>
<tr>
<td>S- Oxidation</td>
<td>Oxidative deamination</td>
</tr>
<tr>
<td>Dehydratations</td>
<td>Alcohol and aldehydes oxidations</td>
</tr>
<tr>
<td>Oxidative dehalogenation</td>
<td>Reductive dehalogenation</td>
</tr>
<tr>
<td>N- Oxide reduction</td>
<td>Epoxide reduction</td>
</tr>
<tr>
<td>Reductive β-scission of alkyl peroxides</td>
<td>NO reduction</td>
</tr>
<tr>
<td>Isomerizations</td>
<td>Oxidative C-C bond cleavage</td>
</tr>
<tr>
<td>Allylic rearrangement</td>
<td>Sulfoxidation</td>
</tr>
<tr>
<td>Desaturation</td>
<td>Deformylation</td>
</tr>
<tr>
<td>Peroxidase-type oxidation</td>
<td>NO-Synthase-type oxidation</td>
</tr>
</tbody>
</table>
The detailed catalytic cycle of dioxygen activation and substrate hydroxylation derived from the most studied P450, the camphor monooxygenase, P450cam, from *Pseudomonas putida*, is shown in Figure 1.2. The resting state enzyme [1] is in an oxidized ferric low-spin, six-coordinate form with a water molecule serving as the distal ligand. Upon substrate binding, the distal water is removed, leaving the heme iron as a high-spin, five-coordinate form shown as intermediate [2] (19), and meanwhile the reduction potential is increased to allow electron transfer to occur. The water exclusion process initiated the first electron transfer to the heme iron from NAD(P)H via electron transfer proteins, yielding a reduced P450 with a five-coordinate heme (intermediate [3]). The binding of dioxygen forms an unstable ferrous-oxy intermediate [4]. The following second electron transfer step, which is the rate limiting step, forms a ferric peroxo-iron species [5], which is consequently protonated to the formation of the hydroperoxo-iron complex [6], Compound 0 (Cpd 0). Another proton is transferred into the active site to the distal oxygen to initiate the formation of a highly reactive Fe^{IV}=O (ferryl-oxo) complex [7], Compound I (30), which is the most widely accepted key intermediate in P450-catalyzed reactions. The Compound I will hydroxylate the bound substrate and finish the reaction cycle by releasing the product and adopting a water molecule to regenerate the resting state enzyme.

As previously mentioned, P450s from different sources are found to preserve a highly similar secondary and tertiary structure in spite of their low amino acid sequence homology (9, 17). Crystal structures of four P450s: P450cam, P450BM-3, P450eryF and P450terp, which are shown in Figure 1.3, exhibit very similar topology. The two
Figure 1.2 The cytochrome P450 catalytic cycle (23). RH is substrate and ROH is product; Scys is the cysteinate ligand; the heme porphyrin is represented as a circle. Decoupling pathways resulting from oxidase activities of P450, in competition with the monoxygenase activity, are called autoxidation, and peroxide and oxidase shunts.

Hydrophobic helices which sandwich the heme prosthetic group from the proximal and distal direction, respectively, are similarly positioned in all P450s, leaving no part of the heme directly exposed to bulk solvent. The highly conserved cysteine ligand is situated at the N-terminal end of the proximal helix.

Cytochrome P450_{cam} (CYP101) is the first to be completely purified (31) and analyzed by X-ray diffraction (32-33) and the most studied among all P450 members.
Figure 1.3 Schematic structures obtained from the crystallographic results of: A. P450\textsubscript{cam} (PDB code: 2CPP), B. P450\textsubscript{BM-3} (PDB code: 1JPZ), C. P450\textsubscript{eryF} (PDB code: 1EUP) and P450\textsubscript{terp} (PDB code: 1CPT). The helix-rich domain is shown on the right side and the helix-poor domain on the left side.

A large amount of crystal structure studies of substrate/inhibitor complexes of P450\textsubscript{cam} provided important insights into the relationship between substrate binding, spin state, redox potential and stereospecificity. However, although P450\textsubscript{cam} is the best characterized among the P450 superfamily, a fundamental question remains to be addressed: How does
the substrate fit into the active site of the enzyme? This mystery has troubled researchers for a long time because all crystal structures of P450<sub>cam</sub>, no matter substrate-free or substrate-bound form, are always in a static conformation with no obvious path into the active site and provided only limited information regarding the dynamics of the binding process. The “invisible” structural and conformational changing process upon substrate binding is partly revealed by my NMR research of P450<sub>cam</sub> and is described in detail in Chapter 4.

1.2.2 Chloroperoxidase

Chloroperoxidase (CPO) was first discovered by Hager and coworkers as a chlorinating enzyme from a marine fungus, *Caldariomyces fumago* (34). During the last half century of study, CPO has been found to be one of the most diverse and versatile catalysts among the hemoprotein superfamily. In addition to its biological role of halogenating organic molecules (Equation 2) (35), CPO was also found to possess peroxidase (Equation 3) and catalase (Equation 4) activities (36), together with P450-like oxygen insertion reactions (Equation 5) without any requirement for NADH or coenzyme (37). In general, CPO can utilize hydrogen peroxide or organic hydroperoxide as the oxidant to catalyze a wide variety of one- and two-electron oxidations, such as the oxidation of alcohols to aldehydes, aldehydes to acid, and amines to nitroso compounds, propargylic oxidation (38-40), N-demethylation and dealkylation (41), dimerization of phenols, stereospecific hydroxylation and chiral sulfoxidations (39, 42, 43). Chloroperoxidase can also catalyze enantioselective epoxidation of cis-disubstituted alkenes with high turnovers in concert with high enantiomeric excess (ee) for a range of aliphatic and aromatic olefins in this class (38, 44-45). This capability makes CPO a
promising candidate to be an important industrial catalyst because chiral epoxides are very useful to form bifunctional compounds via stereo specific ring-opening.

Halogenation (normal function): \[ \text{RH} + \text{H}_2\text{O}_2 + \text{Cl}^- + \text{H}^+ \rightarrow \text{R-Cl} + 2\text{H}_2\text{O} \]  

Dehydrogenation (peroxidases): \[ 2\text{RH} + \text{H}_2\text{O}_2 \rightarrow \text{R-R} + 2\text{H}_2\text{O} \]  

Peroxide decomposition (catalase): \[ 2\text{H}_2\text{O}_2 \rightarrow \text{O}_2 + 2\text{H}_2\text{O} \]  

Oxygen insertion (P450): \[ \text{RH} + \text{H}_2\text{O}_2 \rightarrow \text{R-OH} + \text{H}_2\text{O} \]  

In addition to catalyzing a wide range of reactions, CPO also serves as a unique and precious structure model for the study of the mechanism of hemoproteins. Chloroperoxidase, composed of 299 amino acid residues, is a typical heme enzyme containing ferric protoporphyrin IX as its prosthetic group. Although CPO shares a number of physiochemical functions with cytochromes P450, the distal pocket of CPO possesses a hydrophilic environment similar to those of heme peroxidases and catalases (46-47). However, in contrast to other peroxidases, the proximal side of the heme active site in CPO has the same feature as cytochromes P450 with a thiolate cysteine as the proximal heme ligand rather than a histidine as in other peroxidases (48-49). The electron-rich thiolate ligand has been generally believed to be critical to the cleavage of the peroxide oxygen-oxygen bond in the absence of the assistance provided by polar active site residues as in horseradish peroxidase. Combined with a P450-like proximal heme environment and a peroxidase-like distal environment, CPO can be considered as a P450-peroxidase hybrid. The hybrid architecture may explain why CPO can catalyze the transformation of such a broad range of substrates and exhibits both P450-type and peroxidase-type activities.
Although shares functional and structural similarities with other hemoproteins, CPO holds some distinctive features revealed by its crystal structure shown in Figure 1.4 (50). The overall tertiary structure of CPO is dominated by eight helical segments, which bears little resemblance to peroxidases or P450s, implying a separate evolutionary origin. Although the distal pocket environment of CPO is polar as in other peroxidases, a glutamic acid rather than a histidine is positioned directly adjacent to the peroxide-binding site (Figure 1.5). It has been proposed that this glutamic acid serves as the acid–base catalyst to help cleaving the peroxide bond in the formation of Compound I (Cpd I), an oxyferryl porphyrin π-cation radical, in CPO (51). The peroxide binding site is flanked by two phenylalanine residues, Phe103 and 186, whose benzene rings are almost parallel with the heme plane. These phenylalanine residues are suggested to play a key role in the binding of hydrophobic substrates. The substrate-binding pocket of CPO more closely resembles P450s in that a small opening above the heme allows access to the ferryl-oxo center in Cpd I. The crystal structure of CPO also reveals that, as the result of post-translational modification, CPO is heavily glycosylated and the carbohydrate account for about 19% of the total molecular weight (52).

All reactions catalyzed by CPO are ping-pong reactions in which Compound I is formed in the initial reaction and subsequently reacts with a second substrate to return the protein to the native state (53). Accidentally, it was discovered that when incubated with allylbenzene (AB) or other selected terminal unsaturated substrates in the presence of hydrogen peroxide, CPO will be rapidly and quantitatively converted to a green species with a complete vanishing of enzymatic activity (Figure 1.6) (54). This phenomenon
Figure 1.4 Schematic structure obtained from the crystallographic result of chloroperoxidase (PDB code: 2J5M).

Figure 1.5 The top view (A) and side view (B) of the heme active site of chloroperoxidase. The Glu183 and His105 can form hydrogen bond and function as acid-base catalyst. The two phenylalanine residues may play a role in the binding of hydrophobic residues.
suggested that CPO was inactivated in a P450-type reaction involving the mechanism-based formation of N-alkylporphyrins during the oxidation of allylbenzene. Most amazingly, the inactive CPO (green CPO) can slowly return to native CPO upon standing with a concomitant regain of greater than 80% of that of the original enzyme’s activity (55).

![Diagram](image)

**Figure 1.6** The chloroperoxidase catalytic cycle including the allylbenzene inactivation and auto-restoration. RH is substrate and X’ is Cl’, Br’, or I’.

The suicide inactivation of CPO is shown to be another common reactivity shared with cytochromes P450 in addition to the structural and functional similarity. The mechanism-based inactivation of P450s also results in “green pigments” with the active site heme being modified by addition of the unsaturated substrate skeleton plus an oxygen atom (56-57). It was reported that the olefin can predominately alkylate the
nitrogen of pyrrole ring IV of the prosthetic heme, but the acetylenes prefer to alkylate pyrrole ring I (58). The reversible N-alkylation of the prosthetic heme was also reported for a P450 2E1 mutant T303A (59-60).

Q-band continuous wave (CW), pulsed electron nuclear double resonance spectroscopy (ENDOR), Mössbauer and electron paramagnetic resonance (EPR) studies of allylbenzene-inactivated CPO identified that the green heme species resulted from the N-alkylation of the heme with a five-membered chelate ring formed (55, 61). However, the techniques used in previous studies were not able to identify which of the four pyrrole ring nitrogen was alkylated. Therefore, a study using NMR was carried out to characterize the modified heme of CPO and the results are described in Chapter II of this dissertation.

1.3 Heme Peroxidases

Heme peroxidases (donor: H2O2, oxidoreductase: EC 1.11.1.7) are a large ubiquitous superfamily of b-type hemoproteins that catalyze the oxidation of a number of substrates in the presence of hydrogen peroxide. The reactions catalyzed by heme peroxidases are important to biological systems, ranging from oxidation of aromatic and heteroatom compounds, through free radical oligomerizations and polymerizations of electron-rich aromatics, to electron transfer, sulfoxidation, epoxidation, demethylation and oxidative degradation of lignin (62-63). Nonmammalian heme peroxidases are often grouped with P450s because of having similar intermediates during their catalytic cycles. Nonmammalian heme peroxidases can be divided into three distinct classes (64). Class I are the intercellular peroxidases, such as yeast cytochrome c peroxidase, gene-duplicated bacterial peroxidases and ascorbate peroxidases. Class II are the extracellular fungal peroxidases and class III are the well-known extracellular plant peroxidases. Although
the primary sequence homology may greatly vary, all none-mammalian heme peroxidases share a common helical-hold overall structure. As mentioned before, all heme peroxidases utilize histidine as the proximal ligand of the heme iron except for chloroperoxidase. In this chapter, I will briefly introduce three important heme peroxidases related to my research: Horseradish peroxidase (HRP), cytochrome c peroxidase (CcP) and soybean peroxidase (SBP).

1.3.1 Horseradish Peroxidase

Horseradish peroxidase is one of the most important enzymes obtained from the plant horseradish, *Armoracia rusticana*, and has been extensively studied for more than a century. Most of our current knowledge about heme peroxidases comes from the studies of HRP. Horseradish peroxidases can utilize hydrogen peroxide to oxidize a wide variety of organic and inorganic compounds. Horseradish peroxidase has been exploited extensively in applications and commercial uses spanning from bioscience to biotechnology, including organic synthesis (65), bioremediation (66), biocatalysis (67), diagnostics (68), biosensors (69), transgenics (70), protein engineering (71) and even to therapeutics (72). Comparing with other popular alternatives such as alkaline phosphatase, horseradish peroxidase has advantages for industrial applications, such as enzyme size, stability, and price aspects. It also has a high turnover rate that allows the generation of strong signals in a relatively short time span. Although the term “horseradish peroxidase” is used generically, the root of the horseradish plant actually contains a number of distinctive peroxidase isoenzymes, among which the C isoenzyme (HRP-C) is the most abundant and consequently most widely used in studies and applications.
The HRP-C is a 44 kD glycoprotein containing a single polypeptide of 308 amino acid residues, with two calcium ion binding sites and four disulfide bonds between cysteine residues 11–91, 44–49, 97–301 and 177–209. The two calcium binding sites are located proximal and distal to the heme plane, respectively, and are linked by a network of hydrogen bond to the heme-binding region. The presence of the calcium ions is significant for the structural and functional integrity of HRP-C. Removal of calcium ions results in a decrease in both enzymatic activity and thermal stability (73), and causes subtle changes in the heme environment (74). The calcium ions are also essential for correct folding of denatured or recombinant HRP (75).

Most reactions catalyzed by HRP can be expressed by Equation 6, in which one molecule of hydrogen peroxide is reduced to water and two molecules of radical product, RH•, are generated (70). The catalytic mechanism of HRP-C, has been investigated extensively (Figure 1.7) (65), which involves the formation of the well established and powerful oxidant intermediates: compounds I and II.

$$2\text{RH}_2 + \text{H}_2\text{O}_2 \rightarrow 2\text{RH•} + 2\text{H}_2\text{O}$$ (6)

Similar as in the peroxide shunt pathway of P450s, the first step in the catalytic cycle of HRP is the reaction that \(\text{H}_2\text{O}_2\) bind to the resting state of the enzymatic heme Fe\(^{III}\) as the distal ligand. The bound \(\text{H}_2\text{O}_2\) is heterolytically cleaved to generate the high oxidation state intermediate compound I, an Fe\(^{IV}\) ferryl-oxo center and a porphyrin-based cation radical. Compound I stores two oxidation equivalents, which are used subsequently for substrate oxidation, thereby leading to crucial functions. A putative transient Fe\(^{III}\)-hydroperoxy intermediate (compound 0) formed prior to compound I has been detected in reactions between HRP-C and \(\text{H}_2\text{O}_2\) at low temperatures (76). The highly conserved
distal histidine and arginine residues (Figure 1.8A) are suggested to cooperatively play important roles through acid-base catalysis in the cleavage of the O-O bond (Figure 1.9) (77-78). The second step in the catalytic cycle is conducted by the first one-electron reduction, which requires the participation of a reducing substrate and leads to the generation of compound II, an Fe$^{IV}$ ferryl-oxo species in which the Fe$^{IV}=O$ species remains intact but the porphyrin radical has been reduced (79-80). The catalytic activity of HRP-Compound II is suggested to link to the protonated distal His42 (81). The remaining oxidation equivalent stored in compound II is released consequently in the second one-electron reduction step, which returns the enzyme to its resting state.

![Figure 1.7](image)

**Figure 1.7** The catalytic cycle of horseradish peroxidase (HRP-C). The rate constants $k_1$, $k_2$ and $k_3$ represent the rate of compound I formation, the rate of compound I reduction and the rate of compound II reduction, respectively. RH$_2$ and RH• represent a reducing substrate and its radical product, respectively.

Because of the presence of attached carbohydrates which hinders crystal growth, the first crystal structure of HRP-C using X-ray crystallography appeared in the literature relatively late (82) as a recombinant enzyme expressed in *Escherichia coli* in non-glycosylated form. The structures of HRP-C contain three α-helices that are
Figure 1.8 The highly conserved active site structures of (A) horseradish peroxidase C, (B) soybean peroxidase, and (C) cytochrome c peroxidase. The heme group and the catalytic residues Arg, distal His and proximal His are in atom colors.

Figure 1.9 Mechanism of compound I formation with the distal His42 plays the main role as both proton acceptor from one of the peroxide oxygen and proton donor to the other peroxide oxygen. The distal Arg38 facilitate the heterolytic cleavage of the O-O bond. A very unstable intermediate enzyme complex is formed before compound I and is referred in the literature as compound 0.

additional to the conserved “core” peroxidase fold. The distal residues, His42 and Arg38 are emphasized to play critical roles in substrate stabilization, catalytic oxidation and ligand binding through an extensive hydrogen bonding network, in addition to the cooperation with the proximal His170 to accomplish the “push-pull” mechanism (14).

Although HRP has been used in a vast number of applications such as organic, enzyme assays, chemiluminescent assays, immunoassays and the treatment of waste waters, it should be pointed out that HRP is relatively less stable than other enzymes commonly used in industrial applications (e.g., hydrolases). Furthermore, the relatively
expensive price of HRP is another obstacle preventing the usage of HRP in applications such as peroxidase-catalyzed enantioselective oxidations at the industry level. Solutions to these problems include better process management of hydrogen peroxide to avoid HRP inactivation, use of engineered or alternative enzymes with improved stability, catalytic efficiency and price points. The latter solution leads attentions to a peroxidase, soybean peroxidase (SBP), extracted from soybean seed hull, with ideally improvement from every aspect mentioned above. A brief introduction about SBP is given in Chapter 1.3.3.

1.3.2 Cytochrome c Peroxidase

Cytochrome c Peroxidase (CcP), a class I b-type heme peroxidase from yeast, is the first heme peroxidase whose X-ray crystal structure was elucidated. It is anomalous in that CcP can utilize ferro-cytochrome c, a hemoprotein, as substrate, in addition to small substrates such as guaiacol, styrene, phenylhydrazine, etc. The metabolic role of CcP is not well defined, but was suggested to relate to detoxification of hydrogen peroxide formed in the intermembrane space of mitochondria using ferro-cytochrome c as the electron donor. Despite the little interest of study from a metabolic perspective, CcP has been extensively used as a model system and archetype in the study of peroxidase catalytic mechanism, protein electron transfer, protein engineering, and structure-function interactions.

The catalytic cycle of CcP is similar to that of HRP and other peroxidases, with compound I and II formation and the two-electron reduction of peroxide to water. The highly conserved distal His52 and Arg48 (figure 1.8C) play the same role as the corresponding residues in HRP through acid-base catalysis. From an analysis of the crystal structure of CcP (83), it was proposed that binding of the peroxide to the iron is
facilitated by transfer of one of its hydrogens to the imidazole nitrogen of His52. It is notable that studies of CcP 1.2 Å crystal structure (84) revealed that Arg48 occupies two positions: one termed “out” and the other “in”. When Arg48 occupies the “out” position, a water molecule take up the space vacated by the side chain of Arg48 and functions as the sixth axial ligand of heme iron in the resting state of the enzyme. Conversely, in Compound I, Arg48 occupies the “in” position permanently where it is able to H-bond with the ferryl oxygen atom for stabilization purpose. The major difference between CcP and other peroxidases is that CcP does not form a porphyrin cation radical but instead a cationic Trp191 radical compound I (85). The CcP compound I, is stable with a half-life of several hours in the absence of reductant (86) but rapidly returns to the ferric resting state via sequential reactions with two molecules of ferro-cytochrome c. The Trp191 residue is suggested to be essential for activity and responsible for the high stability of CcP compound I (85).

1.3.3 Soybean Peroxidase

Soybean peroxidase (SBP) is a class III heme peroxidase expressed in the seed hull of soybean ∼20 days after anthesis (87). Soybean peroxidase shows 57% amino acid sequence identity with the most widely studied member of this class of peroxidases, horseradish peroxidase.

As most other glycoprotein peroxidases, the native SBP encountered difficulties in obtaining suitable single crystals for X-ray diffraction. Alternatively, the non-glycosylated recombinant SBP (rSBP) produced in Escherichia coli was used for crystal structure determination (88). The rSBP and HRP-C share a very similar three-dimensional structure (Figure 1.10), active-site environment (Figure 1.8 A, B), and
catalytic mechanism. They have many structural factors in common, such as heme prosthetic group, two calcium ions, four disulfide bonds, a single tryptophan and eight glycosylation sites (89). The crystallographic structure of rSBP revealed three features of mechanistic implications: a bound Tris buffer molecule at the binding site of small phenolic substrates, a highly exposed heme edge, and the contact of one heme vinyl group with a sulfur atom from methionine.

![Figure 1.10 Schematic structures of (A) rSBP (PDB code: 1FHF) and (B) HRP-C (PDB code: 1ATJ) obtained from crystallographic results.](image)

As mentioned in the end of section 3.1 of this chapter, horseradish peroxidase is not an ideal peroxidase for industrial applications but needs improvement for its activity, efficiency, stability, cost, abundance, and large-scale production. Soybean peroxidase, however, has many advantages in these aspects as an attractive alternative to HRP for industrial use. Catalytic activity of SBP is reported to be ~1.5-fold higher than that of HRP-C (90). A ~20 fold higher catalytic efficiency for SBP over HRP-C at their pH optima was also reported (89). The thermal and kinetic stabilities of SBP are better than those of HRP (91-93). SBP has also been found to be greatly resistant to denaturation induced by high concentration of urea and guanidine hydrochloride (90). The above
attributes together with the low cost and high abundance of SBP make it a very attractive candidate for industrial and medical uses, and in fact, SBP has been patented for a broad range of uses, such as biocatalyst, biosensor and wastewater treatment (94-106).

With most of the previous investigations focusing on the physiological property, biological functions and applications of SBP, the structural and chemical properties of the enzyme were somewhat neglected. Structural studies of SBP, other than the crystal structure of recombinant enzyme, include electronic absorption, resonance Raman, and tryptophan fluorescence studies (90, 107-109). It has been suggested that the higher stability of SBP compared to HRP-C may stem from the heme binding to the apoprotein, whereas a contact between heme 8-vinyl group and the sulfur atom of Met37 is observed in the SBP structure (88, 93).

NMR spectroscopy has been applied successfully in the active site studies of many peroxidases, such as HRP (110-114) and CcP (113, 115), which encouraged our interest in using this technique to analyze the heme cavity of SBP. Therefore, an introduction about NMR study of hemoproteins, especially heme peroxidases, is made in the following chapter.

1.4 Nuclear Magnetic Resonance of Hemoproteins

Heme prosthetic group is an exquisitely sensitive chromophore whose spectroscopic properties reflect a wealth of structural detail of its environment in a protein. Therefore the heme prosthetic group offers a unique probe for many biophysical techniques, such as resonance Raman, optical absorption, fluorescence and especially, nuclear magnetic resonance (NMR) spectroscopy.
NMR plays critical roles in bioscience and biotechnology in both imaging and complete three-dimensional structure determination. Compared with X-ray crystallography, another major structural method, NMR has its advantages because it can be carried out under solution conditions without the need for protein crystallization, which can be a frustrating task for many proteins such as glycoproteins. Moreover, in NMR experiments, solution conditions such as the pH, temperature and solute types and concentration can be easily adjusted to closely mimic a given physiological environment. Conversely, it can also be changed to any nonphysiological conditions for studies such as protein denaturation. Furthermore, in addition to protein structure determination, NMR can be applied in investigations of protein dynamic features such as domain movements, as well as studies of protein-substrate, protein-ligand, or protein-protein interactions for structural, thermodynamic and kinetic aspects.

The limitations of NMR spectroscopic study of protein structures in solution, however, are the inherent low sensitivity and poor resolution of the technique. Additionally, the slow protein tumbling increases the spin-spin lattice relaxation time ($T_2$) and therefore increases the resonance linewidth. The increased linewidth coupled with the number of protons in proteins result in serious spectral overlap problems.

New developments in NMR partially alleviated these problems. The NMR sensitivity and resolution have been increased by progress in both theoretical and practical spectrometer technology, which leads to an increasingly efficient utilization from the information content of NMR spectra. The effects of line broadening and spin diffusion can be diminished with deuterium substitution for hydrogen. Likewise, NMR spectra
from uniformly or selectively labeled samples can be drastically simplified to reduce the chance of overlap.

From $^1$H NMR of hemoprotein, the large heme ring current near the heme group can generate a small number of well-separated resonance lines in the spectra of folded, globular hemoproteins. Historically, paramagnetic metalloproteins triggered the development of protein NMR spectroscopy. This is because for the paramagnetic states of hemoproteins, additional well-resolved resonances outside of the crowded diamagnetic 0-10 ppm window are resulted from the interactions with the unpaired electrons of the heme iron. Although these hyperfine shifts only represent less than 3% of the hydrogen atoms in most hemoprotein, these signals can provide an impressive amount of information on the heme environment by revealing the paramagnetic influence on chemical shifts in an expanded chemical shift scale. The hyperfine shifts can be used to monitor electronic structural and magnetic properties of the heme. The hyperfine shifts can also be selectively irradiated in 1D $^1$H NMR experiments to facilitate unambiguous resonance assignment in combination with isotope labeling of heme prosthetic groups. Since the paramagnetism offers more valuable NMR information of hemoproteins, I will further focus on introducing hemoproteins NMR under paramagnetic conditions.

1.4.1 NMR Information Content

The general goal of NMR is to detect and assign the heme and protein signals as much as possible, and to arrange the heme and the amino acid residues in their proper spatial relationships. The information content directly given by an NMR spectrum includes chemical shifts and spin relaxations. After calibration to some convenient reference, the observed chemical shift for an arbitrary nucleus in a paramagnetic
hemoprotein can be described as the sum of the diamagnetic shift for the nucleus in the absence of unpaired electrons and the net paramagnetic hyperfine shift caused by the unpaired electron spins on the iron (116). The latter shift can be further divided into two contributions: the contact shift which arises from partial unpaired spin density delocalization from the paramagnetic heme iron onto a ligand nucleus, and the dipolar shift caused by a dipolar or through-space interaction between any residue nuclei and the unpaired spin on the iron. The paramagnetic hyperfine shift is temperature dependent and will be discussed in Chapter 1.4.2.

Different physical processes are responsible for the spin relaxation of a nucleus. According to the corresponding relationship between the nuclear spin magnetization vector and the external magnetic field, two principal relaxation processes are defined as $T_1$ (spin-lattice relaxation time, longitudinal) and $T_2$ (spin-spin relaxation time, transverse). The relative paramagnetic contribution to the intrinsic $T_1$ for two nonequivalent nuclei, $i, j$ on residues not bonded to the heme iron follows the relationship of Equation 7:

$$r_i/r_j = [T_1_i/T_1_j]^{1/6}$$

(7)

with which the relative distances of the two nuclei to the iron, $r_i$ and $r_j$ can be determined. Therefore, it is convenient to indicate the varying effect of paramagnetic relaxivity throughout a protein in relevant of the heme iron.

1.4.2 Temperature Dependence of Hyperfine Shifts

The temperature dependence of hyperfine shifts is complicated depending on different oxidation/spin state of the heme iron. Under a simplest ideal situation for a well-
isolated spin state with no zero-field splitting, the paramagnetic susceptibility tensor $\chi$ obeys Curie’s law (Equation 8):

$$\chi = \frac{\mu_0 N_A g^2 \beta^2 S(S+1)}{3kT}$$

(8)

where $\mu_0$ is the permeability of vacuum, $N_A$ is Avogadro’s number, $\beta$ is the Bohr magneton, $g$ is the spectroscopic splitting tensor, $k$ is the Boltzmann constant and $T$ is the absolute temperature. The hyperfine shift of a ligand nucleus $j$, $\delta_{hf}^j$ can therefore be simplified as Equation 9:

$$\delta_{hf}^j = (A/h)g^j \beta S(S+1)$$

$$\frac{3kT(\gamma/2\pi)}{3kT(\gamma/2\pi)}$$

(9)

where $(A/h)^j$ is the hyperfine coupling constant for nucleus $j$ in Hz, and $\gamma$ is the nuclear magnetogyric ratio. Therefore $\delta_{hf}$ will exhibit strict Curie behavior (linear in $1/T$ with zero intercept at $T \to \infty$). However, with different electronic structure of the heme iron, the hyperfine shifts may exhibit hyper-Curie (intercept at $T \to \infty < 0$), hypo-Curie (intercept at $T \to \infty > 0$), or even anti-Curie (intercept at $T \to \infty > \delta_{hf}$) temperature behaviors.

For low-spin (LS) ferriheme group of hemoproteins, the four methyl groups are divided into the 1-CH$_3$, 5-CH$_3$ and the 3-CH$_3$, 8-CH$_3$ pairs because of their relative shifted positions shown on the NMR spectrum. Normally the methyl groups exhibit two strongly (~30 ppm) and two weakly (~5 ppm) low-field contact shifted resonances with either the 1-CH$_3$, 5-CH$_3$ or the 3-CH$_3$, 8-CH$_3$ shown together in the low-field region, depending on the orientation of the proximal ligand with respect to the heme plane. It is generally observed that the two strongly shifted heme methyls exhibit strong hyper-Curie and the other two weakly shifted methyls exhibit anti-Curie behavior (117-124). For the
axial His in peroxidase-CN complexes, the contact shifts normally show strong hyper-Curie behavior.

1.4.3 Nuclear Overhauser Effect

The nuclear Overhauser effect (NOE) is a manifestation of the dipolar interactions between different nuclei. Although the NOE is a common phenomenon for all combinations of closely spaced nuclear spins, $^1$H-$^1$H NOEs are of prime interest for structural studies because it is related to the “through-space” distance between a pair of hydrogens that are in proximity to each other without the need for spin-coupling through covalent linkage. The intensity of the NOE is in first approximation proportional to $1/r^6$, in which $r$ is the distance between the protons, and normally for two adjacent protons, a NOE signal is only observed if their distance is smaller than 5 Å.

However, the NOE is highly restricted in large proteins or paramagnetic systems because spin diffusion in large systems obscures selective primary NOEs. This problem can be solved by saturating a paramagnetically relaxed nucleus $i$ for a very short time $t$, yielding $\eta_{i,j}$, the NOE to proton $j$, becomes independent of the intrinsic relaxation time of the nucleus as shown in equation 10:

$$\eta_{i,j}(t) = \sigma_{ij} t$$  \hspace{1cm} (10)

where $\sigma_{ij}$ is the cross-relaxation rate. The above equation indicates that NOEs should be applicable in paramagnetic systems if resonances can be saturated quickly. Therefore, even steady-state NOEs can be observed in large paramagnetic hemoprotein systems where $\sigma_{ij}$ is large, allowing unambiguous assignment of hyperfine signals in hemoproteins at different oxidation states. The paramagnetism diminishes the NOEs but
at the same time weakens the spin diffusion, so that the resulting NOEs reflect mostly an internuclear distance of ~3 Å (125-126).

The 1D NOE and 2D NOESY experiments are both useful and essential to the determination of protein structure. The 1D NOE is relatively more sensitive and selective for hyperfine shifts but is less effective for signals buried in the diamagnetic envelope. In contrast, the NOESY spectrum offers information of the entire 1H NMR spectrum of a protein with greatly improved resolution of the individual peaks, which is especially valuable for resonances in the crowded diamagnetic window.

A series of NMR studies on HRP-CN– complex using 1D steady state NOE and 2D NOESY (110-114, 126-133), combined with isotope labeling method, have provided unambiguous heme assignments, identification of catalytic residues, and information on the spatial disposition of such residues in the active site. Results from HRP have been used as the benchmark and model systems for developing successful approaches to the NMR studies of other paramagnetic hemoproteins.

1.4.4 Low-Spin Cyanide Complex of Hemoprotein

The low-spin ferric (S=1/2) state of the heme iron is the structurally most informative oxidation/spin state, in which all the electrons are, to first order, in π–bonding or nonbonding iron orbitals. Therefore, electron spin can only be delocalized into π ligand orbitals. For NMR spectra of low-spin state hemoproteins, both linewidth (determined by the electronic spin relaxation) and hyperfine shift dispersion (determined by the magnetic anisotropy) depend critically on the axial ligand because of the orbital splitting gap. In hemoproteins having a histidine proximal ligand, the cyanide inhibited complexes can provide well-resolved 1H NMR spectrum with the narrowest lines and the largest
magnetic anisotropies by utilizing CN\(^{-}\) as the distal ligand. In addition to affecting the side chains on heme, the His/CN\(^{-}\) coordination also produces strong spectral dispersion for nonligated amino acid residues around the heme iron because of significant dipolar shifts.

However, in the case of cyanide inhibited hemoproteins using cysteine as the axial ligand, the Cys/CN\(^{-}\) coordination causes less efficient iron electron spin relaxation, and hence results in larger linewidths, much smaller hyperfine-shift dispersion, and strongly relaxed heme signals, making it much more difficult to extract useful information from the NMR results for nuclei in proximity to the heme iron (123).

In conclusion, with a combination of model compounds or proteins, isotope labeling, differential relaxation study and proper 1D/2D NMR methods, resonances from the heme, the axial ligands, and amino acids in the heme cavity can be assigned. The resulted full scope of the NMR investigation can provide valuable information about the three-dimensional structure of the active site of the hemoprotein in solution.

1.5 References


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CHAPTER II. ISOLATION AND STRUCTURE DETERMINATION OF A NOVEL HEME IN ALLYLBENZENE-MODIFIED CHLOROPEROXIDASE

2.1 Introduction

Chloroperoxidase (CPO) from the ascomycetous fungus, *Caldariomyces fumago* ([1]), is one of the most diverse and versatile catalysts among hemoproteins. CPO is the most extensively studied halogenating enzyme. It is heavy glycosylated (~20 % carbohydrates) and has a molecular mass of 42 kDa. In addition to halogenation, CPO can catalyze a variety of oxidations including epoxidation of olefins ([2]), oxidation of alcohols and aldehydes ([3-4]), hydroxylation of benzylic and propargylic compounds ([5-7]), oxidation of amino into nitroso groups ([8]), N-dealkylation ([9-10]), sulfoxidations ([11]), as well as classic peroxidase ([12]) and catalase ([13]) reactions. Except for the classical heme peroxidase reactions, most catalytic activities of CPO are also common to cytochromes P450. The x-ray crystal structures of CPO ([14]) reveals a cysteine thiolate as the proximal ligand to the heme, which is in marked contrast with other heme peroxidases dominated with a proximal histidine ligand.

The catalytic cycle of CPO is initiated by the heterolytic cleavage of a loosely bound H$_2$O$_2$ molecule, which leads to the formation of a typical heme peroxidase intermediate, compound I. All CPO catalyzed reactions are subsequently facilitated by this highly reactive compound I through direct redox reaction or the formation of other intermediates, such as compound II and the protonated compound II ([15]).

A unique mechanism-based “suicidal” inactivation of CPO was reported that when incubated with allylbenzene (AB) or other selected terminal unsaturated substrates in the presence of hydrogen peroxide, CPO will be rapidly and quantitatively converted into a
green protein species that is devoid of any enzymatic activity (16). Surprisingly, the green CPO can slowly auto-recover to the native CPO upon standing with more than 80% recovery of the original enzyme’s activity (Figure 1.6) (17).

Similar mechanism-based heme modifications have also been reported for cytochromes P-450 (18-39), hemoglobin (40), sulfmyoglobin (41), horseradish peroxidase (42), and heme model systems (43-53). Of all known hemoproteins that undergo mechanism-based inactivations, only CPO can spontaneously regain activity. However, mechanism-based inactivations of a P450 2E1 mutant (34-35) and P450 2B4 as well as its mutant (36) were recently claimed to be partially reversible upon dialysis. The results from the studies of these modified heme systems greatly enhanced the understanding of the active site environment and catalytic mechanism of these hemoproteins.

It is difficult to establish the chemical identity of the modified heme because of the auto-recovery behavior of green CPO. Q-band continuous wave (CW), pulsed electron nuclear double resonance spectroscopy (ENDOR), Mössbauer and electron paramagnetic resonance (EPR) studies of allylbenzene-inactivated CPO had proposed the green heme species as an allylbenzene N-alkylated heme. A nitrogen in one of the heme pyrrole rings is alkylated by the terminal carbon of allylbenzene and the compound I oxygen is ligated with the internal double-bond carbon, forming a five-membered heterocyclic chelate ring (17, 54). However, none of the above mentioned techniques was able to establish to which pyrrole nitrogen the allylbenzene moiety is attached.

To better understanding the catalytic mechanism of CPO, the topology of substrate access orientation to the heme active center, the mechanism of spontaneous heme dealkylation and restoration of catalytic activity in this system, I report a novel process
for establishing the exact structure of the N-alkylated heme center of CPO. Being aware of the significantly increased stability of the isolated N-alkylated heme (16), the modified heme was extracted out of the green CPO followed by NMR spectroscopic and LC-MS studies to define the structure of the isolated “green heme”.

2.2 Materials and Methods

2.2.1 Materials

*Caldariomyces fumago* was purchased from ATCC (Manassas, VA). Deuterated solvents were from Cambridge Isotope Laboratories (Andover, MA). Deionized water with 18.2 megohm·cm resistivity were freshly produced from a Milli-Q BioCel Water Purification System (Millipore, Billerica, MA). All other reagents were of the highest purity available from Fisher Scientific (Pittsburg, PA) or Sigma-Aldrich (St. Louis, MO).

2.2.2 Growth of Mold and Preparation of CPO

Stock cultures of *C. fumago* were inoculated on Potato Dextrose Agar (PDA) plates and grew for 7 days at 28 °C as a surface mat. The surface mat from three plates of inoculum was used to inoculate 300 mL of Fructose Peptone Salts (FPS) medium supplemented with 50 mg/L carbenicillin and homogenized for 30 sec with a sterile tissue homogenizer (Omni TH115). The homogenate was equally divided into six 125 mL Erlenmeyer flasks plugged with autoclaved cotton balls. After 7 days of growth at 22 °C in a rotary shaker (Bruker), the mold mycelium was homogenized again and inoculated into six 4 L baffled culture flasks, each containing 2 L of FPS medium. The inoculum was grown at 19 °C in a shaker with a rotation speed of 200 rpm. Chloroperoxidase production was monitored kinetically via the standard MCD assay (55) three times a week. The *C. fumago* medium containing secreted CPO was harvested when the MCD
chlorination activity stop increasing, typically 12-14 days after incubation, followed by isolation and purification of CPO from the growth medium according to the method established by Morris et al. (56) with minor modifications, using acetone instead of ethanol in the solvent fractionation step.

The Reinheitzahl value (Rz, A398 nm/A280 nm) was used to check the purity of the final sample. Protein preparations with Rz values of 1.44 or higher were concentrated to ~3 mM with ultrafiltration and stored at 4 °C for future use. Concentrations of native CPO were determined using an extinction coefficient of 91 200 M⁻¹ cm⁻¹ at 400 nm.

2.2.3 Preparation, Extraction and Purification of Allylbenzene-Modified Heme

Green CPO was prepared as described by Debrunner et al. (17). Modified green heme was extracted by the conventional acid-butanolone method (57). Since the extracted green heme can automatically form a μ-oxo dimer (54), 0.1 μL of 12 N HCl was applied to the extracted fraction to depolymerize the dimer. To eliminate the native heme auto-recovered during green heme preparation and other possible contaminant, the extracted green heme was vacuum dried and then subjected to further purification with high-performance liquid chromatography (HPLC) on aPhera C18 column (Waters) with a mobile phase of 70:30 (v/v) methanol/H₂O. The purified green heme fraction was collected, vacuum dried and then stored at -20 °C for structure characterization. Unless otherwise specified, the entire procedure for the green heme preparation was performed at 4 °C.

2.2.4 Spectroscopic Characterization of Green Heme

UV-Visible spectrophotometric characterization of the green heme and its cyanide adduct was obtained in methanol at room temperature on a Cary 300 Bio spectrophotometer (Varian, Palo Alto, CA).
Mass spectra were performed on Finnigan LCQ Deca XP MAX system (ThermoFinnigan, San Jose, CA) with Surveyor autosampler and Surveyor LC Pump. The mass spectrometer was equipped with an ESI source operated in the positive ion mode. Secondary mass spectrometry (MS/MS) experiments were performed to analyze the targeted ions. The operating conditions of the MS detector were optimized with a solution of 90:10 (v/v) acetonitrile/water at a flow rate of 20 µL/min. The signal abundance of major peak was maximized as follows: sheath gas flow, 30 arbitrary units; spray voltage, 5 kV; heated capillary temperature, 260 °C; capillary voltage, 45 V; tube lens offset, 55 V, a relative collision energy of 30% was used for all MS/MS experiments with an isolation width of 3.0 μ.

The NMR sample was prepared by dissolving green heme in 500 µL deuterated methanol-D4. The cyanide adducts of the green heme were prepared by addition of twenty equivalence of cyanide from a freshly made 1 M stock solution of KCN in 99.9% D₂O. Demetallation of the green heme was achieved by treatment with 5% (v/v) DCl/methanol-s4 (18).

Proton NMR spectra of both cyanide-bound and demetallated green heme were recorded on a Bruker Avance 600 (Ultrasound) NMR spectrometer operating at a proton frequency of 599.97 MHz. The most intense solvent signal was suppressed with presaturation during relaxation delay. Chemical shift values were referenced to the residual HDO signal at 4.76 ppm. All NMR spectra were acquired at 298 K unless otherwise specified.

Phase-sensitive NOESY spectra for both cyanide-bound and demetallated green heme dissolved in DMSO-D6 were acquired with a mixing time of 1.5 to 300 ms and a
repetition time of 0.02 to 3 seconds. Typical NOESY spectra were collected with 256 experiments in the F1 dimension using the hypercomplex method (58). Total number of 64-4000 scans were accumulated for each F1 experiment, which was acquired with 2048 complex points in the F2 dimension over a spectral width of 24 to 200 ppm. A 200 ms presaturation with a weak decoupler power was used in all NOESY experiments to suppress the residual solvent signal. Magnitude COSY spectra were acquired with the same parameters in NOESY, using standard Bruker pulse sequence (cosyprqf).

Clean TOCSY (59) spectra of demetallated green heme were recorded over different spectral windows using 2048 F2 points and 256 complex F1 points of 48 scans. Solvent suppression was achieved by a 200-ms direct saturation during the relaxation delay period. A mixing time of 60 ms was used to allow effective spin lock for protons with different relaxation properties.

All two-dimensional data were processed on a HP Workstation with a Pentium 4 processor using Topspin 1.3 (Bruker). Various apodization functions were employed to emphasize protons with different relaxation properties. For example, apodization over 256, 512, and 1024 points was used to emphasize fast relaxing broad cross-peaks at the expense of resolution, whereas apodization over 2048 points is necessary to emphasize slowly relaxing cross-peaks. All two-dimensional data for the green heme were zero-filled to obtain 2048 x 2048 matrices as required by the large hyperfine shift dispersion exhibited by the paramagnetic nature of this heme adduct.

2.3 Results

The modified heme species with a bright green color was isolated from the allylbenzene inactivated CPO. Through purification, a red colored species was separated
from the dominant green fraction, which should be the protoheme \( b \) derived from the trace amount of auto-recovered CPO. Small amount of other green species (<1%) was also isolated from the main product, which suggested the presence of isomers of the allylbenzene \( N \)-alkylated heme. The purified green heme species is very stable at room temperature and shows no noticeable optical change for weeks.

### 2.3.1 Mass Analysis

The Mass spectrum of purified green heme showed a dominant peak at \( m/z \) 759.4 (Figure 2.1A), which can be assigned to a doubly charged \( \mu \)-oxo dimer as shown in the inset of Figure 2.1A with a total MW of 1517. The secondary MS/MS spectrum of this peak clearly showed the largest fraction at \( m/z \) 750.0 (Figure 2.1B), which is consistent with the MW of allylbenzene \( N \)-alkylated heme plus an oxygen atom (inset of Figure 2.1B).

Upon addition of 2 equivalents of HCl, the major component of green heme in mass spectrum is observed at \( m/z \) 750.2, which is consistent with result in the absence of HCl and previously reported electrospray mass spectrometric analysis of the green AB-CPO (16). This \( m/z =750.2 \) species is therefore assigned to the green heme monomer resulted from the depolymerization of the \( \mu \)-oxo dimer. The secondary MS/MS spectrum of the \( m/z \) 750.2 peak (Fig. 2.2) clearly shows the cleavage of one of the propionic acid side chain, removal of the allylbenzene adduct together with the oxygen bridge, and other fragments as shown in the insets of Fig. 2.2. Therefore, our MS studies confirm the presence of a modified heme with an allylbenzene moiety added to one of the pyrrole nitrogens as reported in literature. However, the exact position of the modification cannot be established by our MS analysis. To elucidate the chemical nature of allylbenzene modified heme, detailed NMR spectroscopic studies were carried out as described in the next section of this chapter.
Figure 2.1 MS (A) and MS/MS (B) spectra of purified green species extracted from green CPO. The insets are the schematic structures of allylbenzene modified heme μ-oxo dimer (A) and monomer (B), respectively.
Figure 2.2 MS/MS spectra of depolymerized green heme. The insets are the hypothetic structures of the green heme monomer and subsequent fragments.

Figure 2.3 MS/MS spectra of demetallated green heme. The insets are the hypothetic structures of the demetallated green heme and subsequent fragments.
The demetallation of the green heme was achieved by addition of excess amount of HCl to green heme solution. Mass spectrum of the demetallated green heme confirms the success of iron removal with a major component at m/z 697.4 and no peak for the green heme monomer or dimer was observed (Figure 2.3). From the mass spectrum, the demetallated green heme is shown to maintain the allylbenzene N-alkylated porphyrin structure with the iron-bound oxygen protonated to form a hydroxyl group.

2.3.2 UV-Visible Spectroscopy Results

The UV-visible spectrum of the as-isolated green heme in methanol showed broad electronic absorption bands at 384 and 612 nm, which is inconsistent with previously reported 408, 504, and 596 nm for μ-oxo green heme (54). The difference may be attributed to solvent effect since the solvent used in the literature was not specified. No observable changes in the UV-vis spectrum of the as-isolated green heme solution was noticed upon addition of CN suggesting the isolated green heme exists as a μ-oxo dimer. The UV-visible spectrum of the green heme monomer has a Soret band at 404 nm and blue shifted to 395 nm upon cyanide binding (Figure 2.4).

2.3.3 Proton NMR Results

The proton NMR spectrum of the as-isolated green heme (Figure 2.5A) displays signals within the normal spectral window typical of standard diamagnetic compounds suggesting the presence of a magnetically coupled system, in complete agreement with the μ-oxo dimer nature postulated previously (54). After depolymerization, with addition of concentrated DCl, a typical five-coordinated high-spin iron-porphyrin proton NMR spectrum (Figure 2.5B) is observed. Because of the high-spin nature of the green heme monomer, considerably broad resonances are observed in the NMR spectra that provide
only limited information about the structural properties of the green heme. Therefore, cyanide was added to the sample to convert the green heme monomer to a six-coordinated low-spin derivative that gives much sharper and better resolved signals in the proton NMR spectrum (Figure 2.5C).

However, although the resonances of green heme-CN$^-$ are well resolved in the proton NMR spectrum, no further structural information can be extracted because no Nuclear Overhauser Effect (NOE) can be obtained even with a combination of irradiation power and duration in 1D experiments. Similarly, no cross peaks were detected in all routine 2D experiments including NOESY, COSY, and TOCSY. The lack of NOE information could be resulted from the sizable paramagnetic leakage quenching (60), the molecular mass unfortunately falls into the intermediate motion region, or other unknown reasons. Methods involving changing the temperature, using different ligands, such as azide or imidazole, and using different deuterated solvents, such as chloroform, DMSO and the
viscous solvent ethylene glycol as suggested by Dr. La Mar (41, 61), all failed to solve this frustrating problem. Therefore, the iron had to be removed from the green heme by the addition of an excess amount of DCI to enable further NMR study of the green heme structure. The proton NMR spectrum of the demetallated green heme supports the presence of an N-allylbenzene-protoporphyrin adduct (N-ABPP) as shown in Figure 2.5D (43, 62).

Complete assignment of the signals for the N-ABPP was achieved through two-dimensional (2D) NOESY measurements as well as through bond connectivities (COSY and TOCSY) with confirmation through comparison with the assignments made for other N-alkylprotoporphyrins (Table 2.1) (43, 62). Figure 2.6 shows the NOESY spectrum of N-ABPP collected in DMSO-D6 buffer with a mixing time of 35 ms. The COSY and TOCSY spectra as shown in Figure 2.7 and Figure 2.8 clearly shows the correlation between coupled protons and helped us unambiguously confirm the resonance assignments.

2.4 Discussion
2.4.1 Assignment of the Porphyrin Resonances of N-ABPP

The proton NMR spectrum of N-ABPP shows a high similarity to the NMR spectra of N-Methylprotoporphyrin IX (N-MePP) isomers and N-griseofulvin-protoporphyrin (N-GfPP) (43, 62). The successful assignments of these N-alkylprotoporphyrins demonstrated that resonance positions of different proton types in N-alkylporphyrins display the following pattern: (a) meso-H at 9.9-10.4 ppm, (b) vinyl-Hα at 7.9-8.4 ppm, (c) vinyl-Hβ at 6.0-6.5 ppm, (d) propionate-Hα at 4.0-4.4 ppm, (e) methyl groups at 3.4-3.8 ppm, (f) propionate-Hβ at approximately 2.9-3.2 ppm, and (g) N-methyl or methylene protons at -3.0 to -4.6 ppm. Therefore, the four most down-field shifted single-proton resonances located at 10.78, 10.72, 10.60 and 10.35 ppm, must arise from the four
Figure 2.5 The proton NMR spectra of (A) the green heme dimer in deuterated methanol-D4, (B) the green heme monomer in deuterated methanol-D4, (C) the green heme-CN adduct in deuterated methanol-D4, and (D) the demetallated green heme (N-ABPP) in deuterated DMSO-D6. All spectra were collected at 298 K with solvent presaturation.
Table 2.1 Proton NMR chemical shifts (ppm) of the resonances for the N-ABPP and the N$_{III}$ regioisomer of N-methylprotoporphyrin (N-MePP) (43, 62).

<table>
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<th>Shift</th>
<th>Assignment</th>
<th>Shift</th>
<th>Assignment</th>
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meso protons on the porphyrin ring. The relatively more downfield shifted position of meso protons in N-ABPP compared to those in N-MePP and N-GfPP suggests a bigger ring-current effect of the macrocycle due to the out-of-plane distortion of the N-alkylated pyrrole (62-64). Among the four meso protons, the meso-H$_\gamma$ and meso-H$_\delta$ shall reveal distinctive NOE correlations with only the 6- and 7-propionate protons, and with only two methyl groups (1- and 8-CH$_3$), respectively. The meso-H$\alpha$ and meso-H$\beta$ shall both show NOEs with one methyl and one vinyl-H$\alpha$. In the NOESY spectrum (Figure 2.6), the signal
Figure 2.6 The 600-MHz phase-sensitive $^1$H NOESY spectra of 1 mM N-ABPP in DMSO-D6 taken at 298 K with a mixing time of 35 ms. Cross-peak assignments are as follows: 1, H-$\delta_{
uc{meso}}$:1-CH$_3$; 2, H-$\delta_{
uc{meso}}$:8-CH$_3$; 3, H-$\gamma_{
uc{meso}}$:6-Ha; 4, H-$\gamma_{
uc{meso}}$:6-Ha'/7-Ha; 5, H-$\gamma_{
uc{meso}}$:7-Ha'; 6, H-$\gamma_{
uc{meso}}$:6-Hfb; 7, H-$\gamma_{
uc{meso}}$:7-Hfb; 8, H-$\gamma_{
uc{meso}}$:AB propyl-2H; 9, H-$\gamma_{
uc{meso}}$:AB propyl-3H; 10, H-$\gamma_{
uc{meso}}$:AB propyl-3H'; 11, H-$\alpha_{
uc{meso}}$:2-Ha; 12, H-$\alpha_{
uc{meso}}$:3-CH$_3$; 13, H-$\beta_{
uc{meso}}$:4-Ha; 14, H-$\beta_{
uc{meso}}$:5-CH$_3$; 15, 2-Ha:2-Hf; 16, 4-Ha:3-CH$_3$; 17, AB benzene $m$-H:AB benzene $p$-H; 18, AB benzene $m$-H:AB benzene $o$-H; 19, AB benzene $o$-H:AB propyl-1H; 20, AB benzene $o$-H:AB propyl-2H; 21, AB benzene $o$-H:AB propyl-1H'; 22, AB benzene $o$-H:AB propyl-3H; 23, AB benzene $o$-H:AB propyl-3H'; 24, 6-Ha:6-Ha'; 25, 6-Ha:5-CH$_3$; 26, 6-Ha:6-Hf; 27, 6-Ha':5-CH$_3$; 28, 6-Ha':6-Hf; 29, 7-Ha:8-CH$_3$; 30, 7-Ha:7-Hf; 31, 7-Ha':8-CH$_3$; 32, 7-Ha':7-Hf; 33, AB propyl-1H:AB propyl-2H; 34, AB propyl-1H:AB propyl-1H'; 35, AB propyl-1H:AB propyl-3H; 36, AB propyl-1H:AB propyl-3H'; 37, AB propyl-2H:6-Hf; 38, AB propyl-2H:AB propyl-1H'; 39, AB propyl-2H:AB propyl-3H; 40, AB propyl-2H:AB propyl-3H'; 41, AB propyl-1H':AB propyl-3H; 42, AB propyl-1H':AB propyl-3H'; 43, AB propyl-3H:AB propyl-3H'.
Figure 2.7 The 600-MHz phase-sensitive $^1$H COSY spectra of 1 mM N-ABPP in DMSO-D6 taken at 298 K. Cross-peak assignments are as follows: 1, 2-Hα:2-Hβ; 2, 2-Hα:2-Hβ'; 3, 4-Hα:4-Hβ; 4, 4-Hα:4-Hβ'; 5, AB benzene m-H:AB benzene p-H; 6, AB benzene m-H:AB benzene o-H; 7, 6-Hα:6-Hβ; 8, 6-Hα':6-Hβ; 9, 7-Hα:7-Hβ; 10, 7-Hα':7-Hβ; 11, AB propyl-1H:AB propyl-2H; 12, AB propyl-1H:AB propyl-1H'; 13, AB propyl-2H:AB propyl-1H'; 14, AB propyl-2H:AB propyl-3H; 15, AB propyl-2H:AB propyl-3H'; 16, AB propyl-3H:AB propyl-3H'. Cross-peak 15 can only be seen with lower contour levels.
Figure 2.8 The 600-MHz phase-sensitive $^1$H TOCSY spectra of 1 mM N-ABPP in DMSO-D6 taken at 298 K. Cross-peak assignments are as follows: 1, 2-Hα:2-Hβ; 2, 2-Hα:2-Hβ'; 3, 4-Hα:4-Hβ; 4, 4-Hα:4-Hβ'; 5, AB benzene p-H:AB benzene m-H; 6, AB benzene p-H:AB benzene o-H; 7, AB benzene m-H:AB benzene o-H; 8, 6-Hα:6-Hβ; 9, 6-Hα':6-Hβ; 10, 7-Hα:7-Hβ; 11, 7-Hα':7-Hβ; 12, AB propyl-1H:AB propyl-2H; 13, AB propyl-1H:AB propyl-1H'; 14, AB propyl-1H:AB propyl-3H; 15, AB propyl-1H:AB propyl-3H'; 16, AB propyl-2H:AB propyl-1H'; 17, AB propyl-2H:AB propyl-3H; 18, AB propyl-2H:AB propyl-3H'; 19, AB propyl-1H':AB propyl-3H; 20, AB propyl-1H':AB propyl-3H'; 21, AB propyl-3H:AB propyl-3H'.
at the lowest field (10.78 ppm) shows NOEs to two resonances at 3.89 and 3.36 ppm that correspond to resonance positions of methyl groups. Therefore the 10.78 ppm peak must arise from the meso-Hδ as it is the only proton that could give such an NOE pattern. Consequently, peaks at 3.89 and 3.36 ppm must arise from either 1- or 8-CH3. The meso-H at 10.72 ppm displays several NOE correlations with peaks within the propionate α-proton and β-proton regions and can be thus assigned to the meso-Hγ. The other two meso-Hs at 10.60 and 10.35 ppm, so far can be putatively assigned to the meso-Hα and Hβ, respectively, because in other N-alkylprotoporphyrins, meso-Hα is always lower shifted than meso-Hβ. Accordingly, the methyl resonance at 3.66 ppm and the vinyl-Hα resonance at 8.65 ppm both showing NOE to the 10.60 ppm meso-H, can be assigned to the 3-CH3 and 2-Hα, respectively. Similarly, the 10.35 ppm meso-H NOE correlated 3.61 and 8.46 ppm resonances can be assigned to the 5-CH3 and 4-Hα, respectively.

The NOE correlation between the assigned 3-CH3 at 3.66 ppm and the 4-Hα at 8.46 ppm is in complete agreement with the above proposed assignments. Additionally, the observation of NOE correlations between the 5-CH3 at 3.61 ppm and two propionate α-protons (4.41 and 4.26 ppm) also solidly confirmed the previous assignments and leads the assignment of the 4.41 and 4.26 ppm peaks to the 6-propionate α-protons. As a result, the other two propionate α-protons at 4.21 and 4.03 ppm must derive from the 7-propionate group. The assignments of the vinyl β-protons and propionate β-protons are archived through COSY and TOCSY by the spin correlations with corresponding vinyl and propionate α-protons and are listed in Table 2.1. Therefore, all the 22 nonexchangeable protoporphyrin IX proton resonances are unambiguously assigned through a combination of 2D NMR experiments.
2.4.2 Assignment of the Linking Group Resonances of N-ABPP

The spectrum of N-ABPP displays two very high-field resonances at -5.88 and -6.49 ppm, which are the result of strong shielding from the ring-current of porphyrin. Similar high-field resonances are assigned to the N-linking group protons in N-MePP and N-GfPP (43, 62), and thus the -5.88 and -6.49 ppm peaks must arise from the propyl 3-Hs where the allylbenzene backbone is linked with the heme pyrrole nitrogen. The strong correlations revealed in NOESY, COSY and TOCSY spectra firmly establish the geminal partnerships between these two protons. The inequivalence of the two propyl 3-protons indicates a relatively fixed confirmation of the N-ABPP.

Shown in the COSY spectrum (Figure 2.7), the cross-peaks between the -0.42 ppm and both the propyl 3-protons direct the assignment of the -0.42 ppm resonance to the propyl 2-H. Subsequently, the 0.32 and -0.74 ppm peaks correlated both with the propyl 2-H and with each other in COSY must arise from the propyl 1-protons.

The two-proton resonance at 4.46 ppm displays NOE correlations to all of the assigned propyl protons and therefore must be derived from the two ortho-Hs of the benzene ring. Again through the COSY and TOCSY spectra, the two meta-Hs and the para-H on the benzene ring are assigned to the 6.04 and 6.56 ppm peaks, respectively.

The multiplicity of the assigned peaks shown in the 1D spectrum and the complementary correlation information revealed by all the 2D experiments are in total agreement with the above assignments. Therefore, the unequivocal assignments for all the allylbenzene protons are successfully accomplished.
2.4.3 Identification of the Predominant Regioisomer of N-ABPP

In the NOESY spectrum, the assigned propyl-2H, propyl-3H, and propyl-3H’ from allylbenzene moiety, all show indubitable space correlations with the meso-H$_y$, which indicates that the N-alkylation must occur at either pyrrole ring III or IV. The conclusive evidence for the N-alkylation position is the NOESY correlation between the propyl-2H of allylbenzene moiety and the 6-propionate H$\beta$ of the porphyrin side chain, which indicates that the nitrogen of the pyrrole ring III is alkylated by allylbenzene in green CPO.

The exact same conclusion for the identification of the N-ABPP regioisomer can also be obtained through analyzing the porphyrin resonances pattern by comparing with the reported results for N-MePP regioisomers (62). It is concluded that, in the N$_{III}$ and N$_{IV}$ regioisomers of N-MePP, the propionate $\beta$-proton resonance of the ring that bears the alkyl group is shifted upfield by approximately 0.5 ppm, whereas in the N$_I$ and N$_{II}$ regioisomers, the $\beta$-protons derived from the two propionate groups are essentially equivalent. For the N-ABPP from green CPO, the 6-propionate H$\beta$ (2.76 ppm) is upfield shifted by 0.48 ppm comparing to the 7-propionate H$\beta$ (3.24 ppm), indicating the N$_{III}$-alkylation of the allylbenzene modification. Moreover, it is also concluded that in the N$_{III}$- and N$_{IV}$-MePP regioisomers, the alkylation results in an approximate 0.15 ppm upfield shift for the vinyl H$\alpha$ resonance on the neighboring ring (4-vinyl for N$_{III}$, 2-vinyl for N$_{IV}$) comparing to that on the ring opposite to the alkylated ring (2-vinyl for N$_{III}$, 4-vinyl for N$_{IV}$). While for the N$_I$ and N$_{II}$ regioisomers, the alkylated ring vinyl H$\alpha$ resonance is approximate 0.4 ppm upfield shifted to the corresponding resonance on the other ring. As shown in Table 2.1, the 4-vinyl H$\alpha$ (8.46 ppm) of N-ABPP is upfield...
shifted by 0.19 ppm compared with the 2-vinyl Ha (8.65 ppm), which therefore also indicates that the N-alkylation is on the nitrogen of pyrrole ring III.

With the above definitive signal assignment, the structure of the N-ABPP is clearly defined as N_{III}-(3-phenyl-2-hydroxyalkyl) protoporphyrin IX (Figure 2.9A) and the green heme extracted from green CPO is consequently established as shown in Figure 2.9B. By comparing with the N-alkylated porphyrin models that has no rigid regiospecificity (40, 46, 48, 52) or only small regiospecificity (29), and the N-alkylation of cytochrome P-450 heme that favors the alkylation of pyrrole ring I (23, 38), and IV (22, 38), or shows no regiospecificity (21, 24-25, 43), CPO exhibits a high regio- and stereospecificity with a different N-alkylation position on the prosthetic heme. The unique regioselectivity of the alkylated heme could be related to the specific exposure of the heme group in CPO and the mechanism of CPO catalyzed epoxidation in general and allylbenzene initiated inactivation of CPO in particular as discussed in the next section of this Chapter.

Figure 2.9 The Structure of N-ABPP (A) and green heme (B) from CPO calculated by MM2 energy minimization. Note the distance between the AB propyl 2-H and the porphyrin 6-Hβ is consistent with the observed NOE in NOESY.
2.4.4 Mechanism of Epoxidation and Substrate Binding of CPO Revealed by the Regiospecific N_{III}-alkylation

A well-accepted none-concerted epoxidation versus \( N \)-alkylation competition mechanism was proposed for cytochromes P450 as depicted in Figure 2.10 (28, 30). When a terminal olefin substrate encounters the compound I oxygen, the activated oxygen can oxidize either the terminal (pathway A) or the internal (Pathway B) carbon of the double bond. It is postulated that only pathway B can lead to heme \( N \)-alkylation by the terminal carbon of the substrate, which causes enzyme inactivation (28).

The determination of the high regio- and stereospecific \( N_{III} \)-alkylation of the heme provides a good viewpoint to the \( N \)-alkylation and active site topology of CPO. Additional constraints rather than electronic factors must exist to control the site of pyrrole \( N \)-alkylation at the CPO active center and the orientation of alkene addition to the heme face. Although currently there is no allylbenzene or styrene bound CPO crystal structure available, useful information can be suggested from other crystal structures of CPO, because the enzyme active site has only minor conformational change upon substrates binding (65). As shown in the crystal structures of CPO (Figure 2.11A) (14, 65), amino acid residues Val67, Ile68, Leu70, Ala71, Asn74, Phe103, Ile179, Val182, Glu183, Phe186 and Ala267 are located at the distal side close to the heme center. These residues together with the heme group, form the substrate binding pocket at the active site.

The regiospecificity of allylbenzene \( N_{III} \)-alkylation strongly suggests that when the internal carbon of the double bond of allylbenzene is oxidized by CPO compound I, the terminal carbon of the double bond is located on top of the pyrrole ring III nitrogen of heme. Therefore, the benzene ring of allylbenzene must head to the opposite side (above
Figure 2.10 Postulated catalytic pathways of (A) epoxidation only or (B) possible for both epoxidation and N-alkylation. The pathway (C) is the recovery of the N-alkylation and is only observed in CPO and two P450 mutants as stated in the introduction.

Figure 2.11 CPO distal side binding pocket top view (A), and side view with incorporated substrate allylbenzene (B).

The pyrrole I, Figure 2.11B), which is reasonably the end of the substrate access channel where allylbenzene gets into the active site. Orientations of allylbenzene different from that shown in Figure 2.11B are very unlikely because of the steric constrains caused by the distal residues. Noticeably, the benzene ring of allylbenzene is “coincidentally” sandwiched by the phenyl side chains of Phe103 and Phe186, which are located at the bottom of the substrate access channel and are almost parallel to the heme plane. The
sandwich structure is also revealed in the crystal structure of CPO complexes with
cyclopentanondione, nitrate and acetate (65), indicating that the phenyl side chains of
Phe103 and Phe186 may work together as a clamp to hold substrates at fixed orientation,
leading to the formation of one enantiomer. The benzene ring of allylbenzene is therefore
oriented perpendicular to the heme plane when allylbenzene enters the active site, placing
the double bond carbons on top of pyrrole III that facilitates the N-alkylation pathway
and affords the N\text{III}-alkylated product. The flexibility of Phe103 (14) makes the double-
phenyl clamp capable of holding substrates using steric and edge-to-face $\pi-\pi$ interactions.
Meanwhile, Asp74 and Glu183, with their hydrophilic side chains, may also help to guide
allylbenzene. Finally, other distal side residues could provide steric hindrances to further
control substrate orientation (Figure 2.11B).

Similar assumption was made concerning the function of Phe103 and Phe186 in the
oxidation of benzyl alcohol catalyzed by CPO (3), although no direct or indirect evidence
was provided. Therefore, it is the first time that the Phe103-Phe186 double phenyl clamp
function is demonstrated with experimental support. To the best of my knowledge, this
double phenyl clamp motif has never been reported in literature. Although more evidence
is required to corroborate this novel motif, it extends and inspires our understanding of
enzyme substrate binding strategy.

The proposed CPO active site topology together with the above mentioned
epoxidation versus $N$-alkylation mechanism can flawlessly explain the spontaneous
restoration of the green CPO and the enantioselectivity of CPO catalyzed epoxidations.
With the double-phenyl clamp holding the benzene ring of allylbenzene, a small
distortion force from the steric clashes with distal residues is applied to the $N$-alkylation
bond, which will decrease its stability and lead to the epoxidation reaction. This is why the extracted green heme, without the interference of the protein backbone, becomes much more stable in either dimer or monomer forms. For CPO catalyzed enantioselective epoxidations, using the styrene epoxidation as an example, the double-phenyl clamp holds the benzene ring of styrene at approximately the same position as that of allylbenzene. The terminal carbon of the double bond will be placed close to the Compound I oxygen because it is one carbon shorter than styrene, and therefore undertakes the epoxidation pathway, which strongly supports the proposed mechanism shown in Figure 2.10. The highly enantioselective epoxidation activity can be attributed to the presence of the double-phenyl clamp that holds the benzene ring at a orientation that favors the formation of (R)-enantiomer. This hypothesis is supported by the fact that (R)-styrene epoxide (~50% ee) is the main product from CPO catalyzed oxidation of styrene (66).

The double-phenyl clamp theory is strongly supported by the CPO F103A mutant (Wang, et al. unpublished results), in which the phenyl side chain of Phe103 is removed and the double-phenyl clamp function is disabled. This active site change shall not stop the N-alkylation of heme, but may affect the stability of the N-alkylated heme product because the deletion of the phenyl side chain can remove or reduce the distortion force. As predicted, this mutant can be inactivated by allylbenzene but is not able to reverse to the resting state upon standing. Additionally, the activity and enantioselectivity of the epoxidation will also be affected by this mutation. For small substrates such as styrene, the ee value of the product should be decreased because the removal of the phenyl ring will allow the substrate rotate more freely. Indeed, the styrene epoxidation product of
CPO F103A mutant only has an ee of ~3% (Chen, et al. unpublished results), a dramatic decrease compared to that of wide-type CPO.

In conclusion, the modified heme in allylbenzene inactivated CPO was extracted, demetallated and characterized by UV-vis, ES-MS/MS, and extensive 1- and 2-D NMR methods. The results indisputably demonstrated that allylbenzene induced alkylation of heme occurs at the nitrogen atom of heme pyrrole ring III in CPO. Based on this discovery, a new mechanism of CPO catalysis and topology of heme active site is proposed, in which a Phe103-Phe186 double-phenyl clamp together with other distal residues are suggested to play an important role in adjusting the position of phenyl substrates, making CPO a highly stereospecific catalyst. This mechanism could be further explored by determining the crystal structure of CPO complexed with allylbenzene or styrene, site-specific mutagenesis, specially designed substrate and theoretical calculations, and could be expanded in exploring the inhibition and epoxidation of P450 enzymes.

2.5 References


CHAPTER III. THE ACTIVE SITE STRUCTURE OF SOYBEAN PEROXIDASE AS PROBED BY 1D AND 2D NUCLEAR OVERHAUSER ENHANCEMENT EFFECT

3.1. Introduction

Plant peroxidases are a ubiquitous class of enzymes that catalyze the oxidation of a wide variety of organic and inorganic substrate molecules at the expense of hydrogen peroxide or other organic hydroperoxides. Plant peroxidases have found important and widespread medical and environmental applications, such as antigen conjugates for color visualization in immunoassays, catalysts in the treatment of various industrial wastes, and key component of biosensors in analytical diagnosis (1-3). The most widely studied plant peroxidase, horseradish peroxidase (HRP), has been historically used for such purposes because of its favorable stability and ease of isolation. However, the limited availability of the horseradish root extract combined with the shortage of alternative sources of the enzyme has made the use of HRP a relatively expensive practice (4-5). Therefore, an abundant and moderately inexpensive source of peroxidases for commercial utilization, especially for large scale environmental applications in the treatment of various industrial wastes to remove hazardous and/or toxic materials is highly desired.

Soybean peroxidase (SBP) is a very attractive candidate for such purposes as the enzyme shows a very similar reactivity profile with that of HRP (6). Meanwhile, SBP is readily obtained in reasonable yield from soybean seed hulls that are plentiful and inexpensive (7). Moreover, the conformational, thermal and kinetic stabilities, together with the catalytic efficiency of SBP are substantially higher than HRP (8-11). As a result, SBP has received an extensive amount of attention as a less expensive, more active, and more stable replacement of HRP for immunological, biocatalytical, biosensing
applications during recent years (12-21).

Most of the previous investigations have been concerned with the physiological and biological functions of SBP. Less attention has been paid to the structural and chemical properties of the enzyme. The lack of structural information is a result of the difficulty encountered in obtaining suitable single crystals for X-ray diffraction, a common problem for most glycoprotein peroxidases (22). The presence of more than one closely-similar isozyme which leads to the microheterogeneity of the enzyme preparations is an additional problem for crystallization (23). Alternatively, the non-glycosylated recombinant SBP (rSBP) was expressed in *Escherichia coli* and its structure in the tris-bound form was solved at a relative low resolution (2.8 Å, 88.2% overall completeness) (6). Other structure studies of this enzyme include electronic absorption and resonance Raman (24), tryptophan fluorescence (25), Edman sequencing, and MALDI-TOF MS studies (26). Unfortunately, none of these studies were able to unequivocally reveal the structure of the active site cavity in native SBP. Despite this fact, structural information about many peroxidases from various sources has accumulated over the years through use of other indirect methods, such as computer modeling (27), sequence homology comparison to structurally characterized CcP (28-32), and especially, NMR spectroscopy (33-34). The successful application of NMR spectroscopy to the elucidation of the heme active-site structural features of numerous peroxidases encouraged my interest in the project.

The primary objective of this study is to characterize the heme active-site structural properties of SBP through use of proton NMR spectroscopy, since information about the active-site structure of heme peroxidases is pivotal to the understanding of the structure-
function relationship of this class of enzymes (35). The hyperfine-shifted signals of the heme protons were assigned unambiguously, and the NMR parameters and peak patterns of SBP are compared with those of HRP and CcP. The effects of the secondary metal ions, Ca$^{2+}$, on the NMR features of SBP are discussed. Paramagnetically shifted signals are fully assigned through use of the 1D and 2D nuclear Overhauser enhancement (NOE) experiments and comparison with the spectral properties of HRP and CcP.

3.2. Materials and Methods

Crude SBP was purchased from Bio-Research Products, Inc. (North Liberty, IA) and was further purified by passage through a DEAE Sephadex A-50 anion exchange column (2.6×20 cm) followed by gel filtration (sephadex G-75, 2.6×100 cm). Proteins were eluted from the anion exchange column with a linear salt gradient consisting of 0 to 0.5 M NaCl in 10 mM phosphate buffer at pH 6.8. The procedure gives typical protein preparations with Rz values of 2.8 or even higher.

The concentration of the enzyme samples was determined spectrophotometrically using $\varepsilon_{403} = 94.6$ mM$^{-1}$cm$^{-1}$ for SBP (9). The molecular weight of SBP was determined to be 42 kD through SDS electrophoresis. On the basis of the Soret band absorption, it was found that commercially purchased SBP preparations were about 60% pure by weight.

Enzymatic activity of SBP preparations (both native and Ca$^{2+}$ depleted) was measured by both ABTS and guaiacol assays with each assay triplicated and the results averaged. ABTS is the registered trade name for 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) diammonium salt, which has been suggested to be the most ideal substrate for peroxidase assays (36). Typically, the aqueous reaction mixture consisted of 1.5 mM ABTS, 1.0 mM H$_2$O$_2$, and 0.1 μg/ml SBP. Hydrogen peroxide concentration was
determined by its molar absorptivity of 39.4 M⁻¹cm⁻¹ at 240 nm (37). The reaction mixture was buffered at pH 6.0 with 10 mM phosphate and the reaction was started by the addition of H₂O₂. The progress of the reaction was monitored at 406 nm (ε₄₀₆ = 18.6 mM⁻¹cm⁻¹) with the rate constant calculated from the first 15 seconds of the reaction. For the guaiacol assay, the reaction mixture consisted of 10 mM sodium phosphate, 6.5 mM guaiacol, 0.1 μg SBP/ml, and 0.06 mM H₂O₂. The reaction was carried out at pH 7.0 and was monitored at 470 nm (ε = 26.6 mM⁻¹cm⁻¹) after addition of H₂O₂. The rate constant was calculated from the first 60 seconds of the reaction (38). All assays were carried out at room temperature.

The removal of Ca²⁺ from SBP was accomplished by the same procedure as described for HRP (39). Native SBP was incubated with 60 ml of 6 M guanidine hydrochloride containing 10 mM EDTA at pH 7 for 8 hours. The incubation mixture was then dialyzed three times against 4 L of 10 mM phosphate buffer (pH 7) containing 10 mM EDTA at 4 °C over a period of 24 hours, followed by dialysis at the same condition without EDTA. The reconstitution of Ca²⁺ was achieved by addition of a 100-fold excess of CaCl₂ to the calcium-depleted SBP (0.7 mM) in 10 mM Tris buffer at pH 6.8. Incubation of the native SBP with excess Ca²⁺ showed no uptake of extra Ca²⁺ into the protein as determined by atomic absorption analysis, indicating that all calcium-binding sites were occupied. Tris buffers were used in calcium reconstitution experiments in order to minimize interference from formation of insoluble or refractory calcium phosphate which might otherwise be experienced with phosphate buffers.

The heme group of SBP was easily extracted from the native enzyme by the reported 2-butanone procedure (40). It was found that the prosthetic group of SBP is indeed an
iron protoporphyrin IX (Figure 1.1B) as was earlier suggested, since the NMR spectrum of the extracted prosthetic group is identical to that of the standard protoheme. Therefore, no efforts were made to reconstitute the standard iron protoporphyrin IX into the apoprotein of SBP.

Proton NMR spectra of both native and calcium-depleted SBP samples were recorded at 20, 25, 30, 35, 40 °C on a Bruker Avance 600 (Ultrashield) NMR spectrometer operation at a proton frequency of 599.93 MHz. Protein concentrations in NMR samples varied from 2.5 to 2.8 mM in either H2O or D2O buffers (10 mM phosphate, pH 6.8). The NMR samples in D2O were prepared by at least five times of isotope exchange of the protein solution with D2O buffer at pH 6.8 (uncorrected for any isotope effect). The isotope exchanges were carried out in a 2.0 ml Millipore Centricon YM-30 centrifugal filter unit at 4 °C. The 1H NMR spectra of the high-spin ferric native, calcium-depleted, and calcium-reconstituted SBP were collected with a 125 kHz spectral width, 16384 data points, and a repetition rate of approximately 15 per second. Generally, 20,000 to 60,000 scans were needed to obtain a decent spectrum due to the high spin nature of the protein. Free induction decays were exponentially multiplied by a line broadening factor of 100 Hz before Fourier transformation to improve the signal-to-noise ratio of the spectra.

The low-spin cyanide-adducts of the protein were prepared by adding a 20 fold excess of freshly prepared KCN to the high-spin enzyme samples. The NMR spectra of the ferric low-spin forms were obtained by collecting 1000-2000 scans with 4 K data points over a 62.5 kHz bandwidth, and a repetition time of 0.2 second with solvent presaturation during relaxation delay. The spectra were obtained by multiplication of the free induction decays with a 20 to 40 Hz apodization. Fast repetition spectra of the low-
spin protein-cyanide complexes were also recorded with a 125 kHz sweep width, 2048 data points, and a repetition rate of 10 ms to detect the fast relaxing proton signals which are intrinsically broad. Chemical shift values at different temperatures for both the ferric high- and low-spin forms of SBP are referenced to the residual HDO signal according to equation 11 (41):

$$\delta_{(HDO)} = 7.835 - \frac{T(K)}{96.9} \quad (11)$$

The spin-lattice relaxation times ($T_1$) of the hyperfine-shifted protons for the cyanide complex of SBP were measure by the standard inversion-recovery pulse sequence. A repetition rate of 3.3 per second and a 90° transmitter pulse of 7.6 $\mu$s was used. The $T_1$ values were estimated from the null point with the relationship of $T_1 = \tau/\ln2$.

The NOE experiments on the cyanide complex of the protein were performed by irradiation of the resonance of interest prior to the 90° pulse. Typically, the peak of interest is irradiated for 100 ms with a weak decoupler power of 65 dB and a repetition time of 0.2 second to obtain steady state NOEs. The buildup of NOE is time dependent and is expressed by equation 12 for an isolated two-spin system (42):

$$\eta_i(t) = \frac{\sigma_{ij}}{\rho_i} (1 - e^{-\rho_i t}) \quad (12)$$

where, $\rho_i^{-1} = T_{1i}$, the selective spin-lattice relaxation time of $H_i$, $t$ is the duration of the saturating pulse on $H_j$, and $\sigma_{ij}$ is the cross-relaxation between $H_i$ and $H_j$. In the high-field case ($\omega^2 r_{ij}^2 \gg 1$), the cross-relaxation is given by equation 13 (42):

$$\sigma_{ij} = \frac{-h^2 \gamma^4}{10 r_{ij}^6} \tau_c \quad (13)$$

where $\tau_c$ is the reorientation time of the vector defined by $H_i$ and $H_j$, and $r_{ij}$ is the length of this vector. For short irradiation times ($<30$ ms) $\eta_i$ takes the form as in equation 14:
\[ \eta_i(t) = \sigma_i t \] (14)

and is independent of the spin-lattice relaxation time of the nucleus.

2D phase-sensitive NOESY and magnitude COSY spectra for the cyanide-bound derivative of SBP were recorded at 600 MHz by using presaturation to eliminate the residual water signal. All pulse sequences utilized here employed single 90° hard pulses to maximize excitation bandwidth. NOESY spectra with a spectral width of about 60 ppm were recorded at 35 and 40 °C with a repetition rate of 5 s\(^{-1}\) and a mixing time of 20 ms to avoid spin diffusion. A total of 2000 experiments were collected with 1024 data points in the F2 direction. The COSY spectra were collected at 25 °C with other parameters similar to NOESY. All 2D spectra data consist of 4096 data points in the acquisition dimension and of 1024 in the indirect dimension. The data were multiplied in both dimensions and were zero-filled and Fourier-transformed to obtain 1024 × 1024 real data points.

3.3. Results

3.3.1. Proton NMR of the Native Ferric High-Spin SBP

The proton NMR spectra of the native ferric high-spin SBP in D_2O solution are shown in Figure 3.1. No additional hyperfine shifted signals were detected when the experiments were carried out in H_2O solution. A very broad signal “A” at 98 ppm, which was observed for other hemoproteins (34, 43-45), was also detected for SBP. Because this broad, far downfield shifted signal has been assigned to the N-δ1-H of the proximal histidine (Figure 3.2) in several hemoproteins, the presence of this peak is a strong indication that the proximal ligand in SBP is the same as that in most traditional hemoproteins, which is consistent with the crystal structure of rSBP. Signal “A” is exchangeable and its exchange behavior varies with different proteins. For manganese
and lignin peroxidase this signal disappear soon after the sample is placed in D$_2$O (34, 45), while for HRP the signal maintains its intensity even after the protein is dissolved in D$_2$O for several weeks (43). The exchange rate of this signal in SBP is very similar to that of the corresponding signal in HRP, as the signal remains detectable even after 3 weeks in D$_2$O buffer.

Figure 3.1 The 600 MHz proton NMR spectra of 2.5 mM native high-spin SBP in 10 mM phosphate buffer at pH 6.8 and 298K.

3.3.2. Proton NMR of the Cyanide Adduct of Native SBP

The proton NMR spectrum of the cyanide-ligated SBP complex in D$_2$O solution is shown in Figure 3.3A. Sharp lines for the paramagnetically shifted signals were observed since the CN$^-$ adducts of all heme peroxidases contain low-spin ferric iron (S = 1/2). Additional peaks were detected when the spectrum was recorded in H$_2$O solution (Figure 3.3B). The spectral features of SBP-CN$^-$ display a high degree of similarity to those of HRP-CN$^-$ and CcP-CN$^-$ shown in Figures 3.3C and 3.3D. The close similarity of the proton NMR spectra of these peroxidases is an indication that they share the same protoheme IX prosthetic group.
Figure 3.2 The schematic representation of the active site structure of the cyanide-ligated form of most heme peroxidases showing the presence of both the proximal and distal histidine. The protoporphyrin plane is represented by the two bold horizontal bars.

Assignment of the hyperfine shifted signals for the SBP-CN⁻ complex was achieved through comparison with the assignments made for HRP-CN⁻ and CcP-CN⁻ (Table 3.1) (34, 46-52) and examination of the active site structure revealed by the rSBP crystal structure (6) with confirmation through one- and two-dimensional NOE measurements as well as through bond connectivities (COSY). The 1D NOE experiments were carried out
by irradiation of the hyperfine shifted signals with a weak decoupler power for a short period of time, typically 100 ms. Some NOE difference spectra are shown in Figures 3.4(B to E), and Figures 3.5(B to F).

**Figure 3.3** The proton NMR spectra of the low-spin protein-cyanide complexes. (A) SBP in 10 mM phosphate D₂O buffer at pH 6.8, 298 K.; (B) SBP in the same buffer with 90% H₂O and 10%D₂O at 298 K.; (C) HRP in the same buffer with 90% H₂O and 10%D₂O at pH 6.3, 301 K; and (D) CcP in the same buffer with 90% H₂O and 10%D₂O at pH 6.3, 301 K. Spectra (C) and (D) are from ref (34).
The spectrum of the low-spin SBP cyanide complex collected with a rapid repetition rate at 298 K is shown in Figure 3.6. Two broad resonances (peaks “z” and “y” each of single proton intensity) were observed in the downfield and upfield region, respectively. Peak “y” is poorly resolved from the methyl peak “b” and shows only as a shoulder of peak “b” at 298 K. These two fast relaxing signals were also detected earlier for several other low-spin heme peroxidases, such as CMP-CN (43, 52).

Shown in Figure 3.7 and 3.8 are the NOESY spectra of SBP-CN in D$_2$O buffer with a mixing time of 20 ms collected at 308 K and 313 K, respectively. The NOESY spectra of SBP-CN in 90% H$_2$O and 10% D$_2$O buffer with a mixing time of 20 ms collected at 323 K is shown in Figure 3.9. The NOESY spectra provide a greatly improved resolution, especially for overlapped peaks which are hard to differentiate from 1D NOE results. The clear NOESY connectivities and the results from scalar (COSY; Figure 3.10) connectivities prove the validity of the NOE connectivities observed in 1D NOE experiments and lead to the proposed assignments for all of the hyperfine shifted protons from the heme active site in SBP-CN. Furthermore, resonances from protons that are near to the heme active center but are buried in the diamagnetic envelope were also revealed from NOESY spectra as shown in Table 3.2. The suggested assignments for hyperfine shifts are further verified by the Curie intercepts predicted from the temperature dependence of the resonances (Figure 3.11).

**3.3.3. Role of Endogenous Calcium**

Atomic absorption spectroscopic analysis showed that native SBP contains an average of 2.9 Ca$^{2+}$ ions per molecule of SBP based on the Soret absorption of the protein (Wang, unpublished results). This result is in complete agreement with the results
Table 3.1a Proton NMR chemical shift (ppm), T₁ (ms), and Curie plots intercept (ppm) values of the isotropically shifted resonances for the low-spin cyanide derivatives of SBP, HRP and CcP.

<table>
<thead>
<tr>
<th>Peak</th>
<th>SBP-CN⁻</th>
<th>HRP-CN⁻</th>
<th>CcP-CN⁻</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Shift</td>
<td>T₁</td>
<td>Int</td>
<td>Shift</td>
</tr>
<tr>
<td></td>
<td>ppm</td>
<td>ms</td>
<td>ppm</td>
<td>ppm</td>
</tr>
<tr>
<td>aᵇ</td>
<td>26.7</td>
<td>---</td>
<td>---</td>
<td>31.0</td>
</tr>
<tr>
<td>b</td>
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<td>108</td>
<td>-3.9</td>
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<td>---</td>
<td>20.8ᶜ</td>
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<td>e</td>
<td>21.4</td>
<td>94</td>
<td>7.0</td>
<td>19.6</td>
</tr>
<tr>
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<td>20.7</td>
<td>129</td>
<td>1.9</td>
<td>20.1</td>
</tr>
<tr>
<td>g</td>
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<td>94</td>
<td>-5.4</td>
<td>15.6</td>
</tr>
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<td>---</td>
<td>11.3</td>
<td>16.3</td>
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<td>---</td>
<td>10.7</td>
<td>9.9ᶜ</td>
</tr>
<tr>
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</tr>
<tr>
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<td>o</td>
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</tr>
<tr>
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<td>-2.8ᶜ</td>
</tr>
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<td>80</td>
<td>3.5</td>
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</tr>
<tr>
<td>t</td>
<td>-7.7</td>
<td>50</td>
<td>2.4</td>
<td>-6.5ᶜ</td>
</tr>
<tr>
<td>y</td>
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<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>z</td>
<td>-28.3</td>
<td>---</td>
<td>---</td>
<td>-29.9</td>
</tr>
</tbody>
</table>

ᵃ Results for HRP-CN⁻ and CcP-CN⁻ are from ref (34) at 301K. Results of SBP are from NMR spectra collected at 313K.
ᵇ Indicates exchangeable protons.
ᶜ From ref (47-48) measured at 323 K.
Figure 3.4 The 600-MHz proton NMR spectrum of (A) 2.8 mM low-spin cyanide-ligated form of SBP in 10 mM phosphate D$_2$O solution at 313 K, pH 6.8; note peak “d” and “e” overlap at 21.4 ppm. (B) through (E) are the NOE difference spectra obtained upon irradiation of the downfield shifted signals “b”, “c”, “d+e”, and “f”, respectively. The irradiated peaks are indicated by a downward arrow in each difference spectrum.
Figure 3.5 The 600-MHz proton NMR spectrum of (A) 2.8 mM low-spin cyanide-ligated form of SBP in 10mM phosphate D$_2$O solution at 313 K, pH 6.8; (B) through (F) are the NOE difference spectra obtained upon irradiation of the downfield shifted signals “g”, “i”, “s”, “t” and “q”, respectively. The irradiated peaks are indicated by a downward arrow in each difference spectrum.
obtained for *Coprinus cinereus* peroxidase (53) and *Coprinus macrorhizus* peroxidase. However, different results have been reported for an average of 2 Ca$^{2+}$ per molecule content of rSBP(6) and HRP (39, 54). Nevertheless, results of as high as 4 Ca$^{2+}$ per molecule of HRP have also been reported (55). Despite the controversial results reported for the calcium content in HRP, all the above mentioned references have presented the complete removal of the calcium ion from the protein by incubating the enzyme with guanidine hydrochloride and EDTA.

Enzymatic activity measurements indicated that elimination of the guanidine-EDTA removable calcium ion from SBP decreased the initial protein activity by 20% for the ABTS assay (from 338 to 272 μmole/min·mg SBP) and 31% for the guaiacol assay (from 565 to 390 μmole/min·mg SBP). Incubation of the calcium “depleted” SBP with excess of CaCl$_2$ regained the full activity of the enzyme. However, in the case of HRP-C, removal of the bound calcium resulted in approximately 60% drop of the initial activity of the protein (39).
Figure 3.7 The 600-MHz phase-sensitive $^1$H NOESY spectra of 2.8 mM SBP-CN$^-$ in D$_2$O taken at 308 K with a mixing time of 20 ms. Cross-peak assignments are as follows: 1, 8-CH$_3$:7-H$\alpha$; 2, 8-CH$_3$:H$_{\delta_{\text{meso}}}$; 3, 8-CH$_3$:7-H$\beta$; 4, 8-CH$_3$:7-H$\beta''$; 5, 8-CH$_3$:δ$_2$:CH$_3$ Leu243 (4.13 Å); 6, 3-CH$_3$:4-H$\alpha$; 7, 3-CH$_3$:ring-H Phe41 (~4.5 Å); 8, 3-CH$_3$:H$_{\alpha_{\text{meso}}}$; 9, 3-CH$_3$:4-H$\beta_{\text{trans}}$; 10, 3-CH$_3$:4-H$\beta_{\text{cis}}$; 11, H$\beta$ His169:H$\beta'$ His169; 12, H$\beta$ His169:H$\alpha$ His169; 13, 4-H$\alpha$:H$_{\beta_{\text{meso}}}$; 14, 4-H$\alpha$:4-H$\beta_{\text{trans}}$; 15, H$\beta'$ His169:H$\alpha$ His169; The numbers in parenthesis are the distances between protons that give rise to the cross-peak measured in rSBP crystal structure.
Figure 3.8 The 600-MHz phase-sensitive $^1$H NOESY spectra of 2.8 mM SBP-CN$^-$ in D$_2$O taken at 313 K with a mixing time of 20 ms. Cross-peak assignments in addition to those of Figure 3.7 are as follows (numbering continuously from Figure 3.7): 16, 7-H$\alpha$:7-H$\alpha'$; 17, 7-H$\alpha$:H-$\gamma_{\text{meso}}$; 18, H$\beta'$ His169:N$_{\text{ Pep}}$H His169; 19, H$\beta$ Arg38:H$\delta'$ Arg38; 20, H$\beta$ Arg38:H$\beta'$ Arg38.
Figure 3.9 The 600-MHz phase-sensitive $^1$H NOESY spectra of 2.5 mM SBP-CN$^-$ in 90\% H$_2$O and 10\% D$_2$O taken at 323 K with a mixing time of 20 ms. Cross-peak assignments in complete agreement with 1D NOE assignments for distal and proximal histidine protons are as follows (numbering continuously from Figure 3.7 and 3.8): 21, H$\beta$ His169:NH$_p$ His169; 22, H$\beta$ His169:H$\alpha$ His169; 23, NH$_\delta_1$ His42:H$\epsilon_1$ His42; 24, H$\beta''$ His169:NH$_p$ His169; 25, H$\beta'$ His169:H$\alpha$ His169; 26, NH$_p$ His169:H$\alpha$ His169; 27, 6-H$\beta$:H$\beta$ Arg38.
Figure 3.10 The 600-MHz phase-sensitive $^1$H COSY spectra of 2.5 mM SBP-CN$^-$ in 90\% H$_2$O and 10\% D$_2$O taken at 323 K. The spectrum is symmetrized for better resolution. Cross-peak assignments in complete agreement with previous 1D and 2D NOE assignments for hyperfine shifted protons are as follows (numbering differently from NOESY spectra): 1, H$\beta$ His169:H$\beta'$ His169; 2, H$\beta$ His169:H$\alpha$ His169; 3, 4-H$\alpha$:4-H$\beta_{\text{trans}}$; 4, 4-H$\alpha$:4-H$\beta_{\text{cis}}$; 5, NH$\delta_1$ His42: H$\epsilon_1$ His42; 6, H$\beta'$ His169:H$\alpha$ His169; 7, H$\beta'$ Arg38:H$\beta$ Arg38.
Table 3.2 Proton NMR chemical shift (ppm) values of the diamagnetic resonances for the low-spin cyanide derivatives of SBP (assigned through NOESY<sup>a</sup>) and HRP<sup>b</sup>.

<table>
<thead>
<tr>
<th></th>
<th>SBP-CN&lt;sup&gt;−&lt;/sup&gt;</th>
<th>HRP-CN&lt;sup&gt;−&lt;/sup&gt;</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shift (ppm)</td>
<td>308 K 313 K 323 K</td>
<td>308 K 328K</td>
<td></td>
</tr>
<tr>
<td>0.27</td>
<td>0.30</td>
<td>0.39</td>
<td>7-propionate Hβ</td>
</tr>
<tr>
<td>2.37</td>
<td>2.68</td>
<td>2.47</td>
<td>7-propionate Hβ'&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>---</td>
<td>7.01</td>
<td>6.95</td>
<td>7-propionate Hα'</td>
</tr>
<tr>
<td>1.90</td>
<td>---</td>
<td>1.62</td>
<td>α&lt;sub&gt;meso&lt;/sub&gt;-H</td>
</tr>
<tr>
<td>7.35</td>
<td>7.39</td>
<td>7.33</td>
<td>β&lt;sub&gt;meso&lt;/sub&gt;-H</td>
</tr>
<tr>
<td>---</td>
<td>4.34</td>
<td>---</td>
<td>γ&lt;sub&gt;meso&lt;/sub&gt;-H</td>
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<td>6.87</td>
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<td>δ&lt;sub&gt;meso&lt;/sub&gt;-H</td>
</tr>
<tr>
<td>10.0</td>
<td>9.96</td>
<td>9.42</td>
<td>His169 Hα</td>
</tr>
<tr>
<td>5.14</td>
<td>5.22</td>
<td>5.33</td>
<td>Phe 41 ring H</td>
</tr>
<tr>
<td>---</td>
<td>-0.8</td>
<td>-0.54&lt;sup&gt;c&lt;/sup&gt;</td>
<td>-0.61&lt;sup&gt;d&lt;/sup&gt; Arg38 Hβ&lt;sup&gt;′&lt;/sup&gt;</td>
</tr>
<tr>
<td>---</td>
<td>1.7</td>
<td>1.47&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.93&lt;sup&gt;d&lt;/sup&gt; Arg38 H6&lt;sup&gt;′&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> Results for SBP-CN<sup>−</sup> at 308 K and 313 K are obtained in D<sub>2</sub>O buffer; and at 323 K are obtained in 90% H<sub>2</sub>O and 10% D<sub>2</sub>O buffer; note that the isotopic effect may slightly vary the chemical shift values.

<sup>b</sup> Results for HRP-CN<sup>−</sup> are from ref (47-48).

<sup>c</sup> Observed at different contour level. Data not shown.

<sup>d</sup> Results are from ref (56) collected at 323 K.

The proton NMR spectra for both the ferric high-spin and low spin cyanide bound forms of SBP are substantially changed when calcium is partially removed from the protein (Figures 3.12). Similar changes were reported for HRP upon calcium removal (54-55). The spectral changes were interpreted as arising from the substantial alterations in the heme environments, most likely the heme proximal and distal sides. Incubation of the calcium-depleted SBP with excess CaCl<sub>2</sub> resulted in the re-uptake of calcium as is reflected by the proton NMR spectra of the calcium recombined SBP (data not shown) which are essentially identical to those of the native SBP (Figures 3.1 and 3.2). Similar, reversible behavior of the bound calcium ion in both HRP and SBP implies that the ion plays some common function in the two proteins.
Figure 3.11 Curie plots of observed shift vs. reciprocal temperature for the resolved hyperfine shifted resonances of SBP-CN– in the (A) downfield and (B) upfield regions. The letter designation of each peak following the scheme of Figure 3.3A is given to the right of each line, and the extrapolated diamagnetic intercept at \( T^{-1} = 0 \) is given in Table 3.1.

3.4. Discussion

3.4.1. Assignment of the Hyperfine-Shifted Resonances for Native SBP

Assignment of the paramagnetically-shifted signals for high-spin heme proteins was previously thought to be problematic as the signals are invariably broad and application of NOE and two dimensional NMR methods are thwarted by the extremely short relaxation times of the shifted protons. However, the advances in the NMR technology and methodology combined with isotope labeling has allowed the complete assignment of the hyperfine-shifted signals in several high-spin hemoprotein systems such as HRP, CcP and lignin peroxidase(34, 44, 49, 57). On the other hand, the peculiar magnetic property of the high-spin form of heme peroxidases also serves as an important hint for
Figure 3.12 The 600-MHz proton NMR spectra of (A) calcium “depleted” ferric high-spin form and (B) calcium “depleted” ferric low-spin cyanide-ligated form of SBP in 10 mM phosphate buffer in D$_2$O at pH 6.8, 298 K. Proton chemical shift is referenced with respect to the signal of HDO in the sample solution.

the assignment of the isotropically-shifted signals. For high-spin heme-containing proteins, only protons which directly interact with the heme iron through a net of covalent bonds are expected to exhibit substantial hyperfine shifts, as the magnetic anisotropy is negligible in such systems. By analogy with HRP and CcP, it is anticipated that only the resonances of the 1-, 3-, 5-, 8-CH$_3$, of the 2- and 4-H$_\alpha$, of the 6- and 7- H$_\alpha$ protons, and of the H$_\beta$ protons of the proximal His (Figure 3.2) will be well-shifted outside the diamagnetic region in the down field region of the spectrum (44, 49, 51, 58-
The chemical shift values for the paramagnetically-shifted signals for native high-spin SBP are shown in Table 3.3 with the corresponding results from HRP for comparison.

**Table 3.3** Proton NMR chemical shift (ppm) values of the isotropically shifted resonances for the high-spin native SBP and HRP.

<table>
<thead>
<tr>
<th>Signal</th>
<th>SBP</th>
<th>HRP</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>97.7</td>
<td>95.2</td>
<td>His169 N-δ₁-H</td>
</tr>
<tr>
<td>B</td>
<td>79.3</td>
<td>73.7</td>
<td>5-CH₃</td>
</tr>
<tr>
<td>C</td>
<td>69.0</td>
<td>67.7</td>
<td>1-CH₃</td>
</tr>
<tr>
<td>D</td>
<td>59.5</td>
<td>65.8</td>
<td>2-Hₐ</td>
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<td>E</td>
<td>55.8</td>
<td>64.3</td>
<td>8-CH₃</td>
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<td>33.2</td>
<td>49.6</td>
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</tr>
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<td>30.1</td>
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</tr>
<tr>
<td>H</td>
<td>26.7</td>
<td>46.1</td>
<td>6-Hₐ</td>
</tr>
</tbody>
</table>

*a Chemical shifts are relative to the residual HDO signal at 4.76 ppm. Measurement are carried out at 298 K. Results for HRP are from ref (59) collected at 328 K.

*b Assignments are achieved through relative pattern instead of chemical shift value comparison because of the large temperature dependence of the chemical shifts.

### 3.4.2. Hyperfine-Shifted Heme Resonances Assignments for the SBP-CN⁻ Complex

As was stated in Chapter I, the low-spin nature of the cyanide adducts of heme peroxidases invariably gives well-resolved, sharp signals in their proton NMR spectra, providing a wealth of unique information on the electronic, magnetic, and molecular structural properties of the heme pocket. Therefore, the detection, assignment, and interpretation of the isotropically-shifted signals of the low-spin form peroxidase have been one of the most interesting challenges in NMR investigations. Although 2D NMR methods have been applied to several paramagnetic systems, steady-state 1D NOE
experiments remain to be the most sensitive techniques for revealing magnetization transfer and are still the preferred techniques for detecting connectivities between very closely spaced signals \((60)\). Therefore, I have performed a series of 1D NOE experiments, in order to assign the hyperfine-shifted signals for the SBP-CN\(^-\) complex as are shown in Figures 3.4(B to E), and Figures 3.5(B to F).

Because of the relatively longer relaxation time and closer position of the hyperfine-shifted resonances in SBP-CN\(^-\) than those in HRP-CN\(^-\), particular care must be paid when setting up the NOE experiment to prevent power spillage and off-resonance effects which could mask the real NOE effect. As is shown in Figures 3.4B and 3.4C, irradiation of the two heme methyl resonances “b” and “c” gives NOE patterns that can be attributed to a methyl group near either a vinyl or propionate group (Figure 1.1B). The NOE experiments on HRP-CN\(^-\) have shown that the heme methyls which are close to a vinyl group exhibit NOEs to either the \(\alpha\)-CH proton or the \(\beta\)-CH\(_2\) protons depending on the vinyl group orientation as shown in Figure 1.1B \((47-48)\). The \(\alpha\)-CH vinyl proton resonance is normally observed in the down field region because of \(\pi\)-spin delocalization of the unpaired spin density in the vinyl group, whereas the \(\beta\)-CH\(_2\) vinyl proton peaks are usually positioned between 0 to -5.0 ppm. Saturation of peak “b” yields detection of peak “f” at “20.7” ppm and peak “n” at “-1.2” ppm in the difference spectrum. Several other peaks are also observed in the difference spectrum in the diamagnetic aliphatic region, indicating the development of multiple secondary NOEs or spin diffusion \((61)\). The lack of selectivity in NOEs is a result of the relatively longer irradiation times used in setting up the experiment \((48)\). Nevertheless, observation of the NOE to the peak at “20.7” ppm correspond to the result observed for HRP-CN\(^-\) when the 8-CH\(_3\) signal is irradiated \((48)\).
Comparison of the spectral pattern of SBP-CN\textsuperscript{−} with that of HRP-CN\textsuperscript{−} leads to the assignment of resonance “b” to the 5- or 8-CH\textsubscript{3} group (34, 45, 48).

The detection of only two of the four heme methyl proton signals outside the diamagnetic region is consistent with earlier observations for HRP-CN\textsuperscript{−} and other low-spin hemoprotein cyanide complexes. The result has been attributed to the particular orientation of the proximal histidine ring with respect to the heme plane (48, 62). Depending on the position of the proximal histidine, the four heme methyls display a unique isotropic shift pattern as either the 1-CH\textsubscript{3} and 5-CH\textsubscript{3} or the 3-CH\textsubscript{3} and 8-CH\textsubscript{3} pair in the down field region (62-63). Therefore, the methyl peak “c” at 26.6 ppm must arise from either the 1-CH\textsubscript{3} or the 3-CH\textsubscript{3} group of the heme unit. The observed resonance “c” for SBP-CN\textsuperscript{−} is assigned to the 3-CH\textsubscript{3} group, as irradiation of peak “c” gives NOE patterns very similar to those reported for HRP when 3-CH\textsubscript{3} is irradiated (48). By analogy with HRP-CN\textsuperscript{−} (48), the peaks “e” at 21.4 ppm observed in the NOE difference spectrum by saturation peak “c” (Figure 3.4C) can be assigned to 4-H\textsubscript{α}, and the four upfield shifted peaks “m” at -0.9, “o” at -1.7, “p” at -2.1, and “r” at -2.8 ppm should arise from all the four vinyl \(\beta\)-protons of heme. Since peak “o” displays the strongest NOE correlation with the 3-CH\textsubscript{3} group, it can be assigned to the 4-H\textsubscript{β}-t, which is the most proximate proton to the 3-CH\textsubscript{3} group (Figure 1.1B). Saturation of the assigned 4-H\textsubscript{α} (peak “e”) yields the difference trace of Figure 3.4D, in which peak “o” and “p” exhibit clear NOE connections with the 4-H\textsubscript{α}, therefore peaks “p” which has stronger NOE with peak “e” must arise from the other 4-H\textsubscript{β} proton (4-H\textsubscript{β}-c). Consequently, the two remaining peaks “m” and “r” can be assigned to the 2-H\textsubscript{β}-c and 2-H\textsubscript{β}-t, respectively.

The spectral pattern of the four vinyl H\textsubscript{β}’s between SBP and HRP is slightly different,
indicating the existence of minor heme cavity discrepancy in these two peroxidases. The absence of an NOE 4-H$_a$ $\rightarrow$ 3-CH$_3$ (Figure 3.4D) dictate that the 4-vinyl group is close to in-plane and is oriented in the cis position (48, 58). Moreover, upon saturation of 3-CH$_3$, the relative magnitudes of the NOEs, 4-H$_{\beta-t}$ > 4- H$_{\beta-c}$ $\approx$ 2- H$_{\beta-t}$ > 2- H$_{\beta-c}$ are qualitatively consistent with distances estimated for trans-2-vinyl and cis-4-vinyl, therefore unambiguously establishes the orientations of the 2- and 4-vinyl groups in SBP-CN$^-$. 

While peak “c” is now unequivocally assigned to the 3-CH$_3$, peak “b” at 28.0 ppm must arise from the 8-CH$_3$ and the NOE observed for the peak “f” at 20.7 ppm upon saturation of peak “b” must arise from a 7-propionate H$_a$. The assignment of peak “n” at “-1.2” ppm is circuitous, because the equivalent peak of HRP was originally assigned to a leucine methyl group (49) and later on revealed through 2D NMR methods as deriving from an Ile244 methyl, which is in proximity to the 8-CH$_3$ (56). However, in SBP, the corresponding Ile position is replaced with Leu243, whose methyl group is only 4.13 Å from the 8-CH$_3$ (carbon-carbon distance, measured from the rSBP crystal structure). Therefore, peak “n” is assigned to the Leu243 $\delta_2$-CH$_3$.

Figure 3.5F shows the NOE difference spectra obtained upon saturating a far upfield shifted peak “q” at -2.5 ppm. The absence of NOE with any assigned hyperfine heme resonances indicates that it is not in proximity to 2-, 3-, 4-, 7-, or 8-substituents of the heme. The $^1$H NMR results of HRP-CN$^-$/complex (48) reveals that the resonance assigned to the 6-propionate H$_{\beta}$ is almost identical in both chemical shift value (-2.8 ppm) and temperature dependence (Curie plot intercept at 5.4 ppm in comparison with 5.7 ppm for peak “q”), and additionally exhibits similar NOE pattern to peak “q”, leading the final assignment of peak “q” to one of the $\beta$ protons of 6-propionate group.
3.4.3. Distal and Proximal Histidine Assignments for the SBP-CN\(^-\) Complex

The appearance of the exchangeable peak “a” at 26.7 ppm is an indication of the presence of a distal histidine (His42) within the heme pocket of SBP, which is consistent with the rSBP crystal structure. This exchangeable proton has been detected for a number of heme peroxidases, such as HRP, CcP, and LiP(34, 47-48). Support for the existence of the distal histidine is also found in the presence of another labile proton signal “h” at 16.2 ppm which is comparable to the position of 16.3 and 16.5 ppm for the corresponding proton in HRP-CN\(^-\) and CcP-CN\(^-\) (34). No attempt through 1D NOE experiments was made to identify NOE connectivities of the labile protons to and from other signals. The assignment of peak “h” to the His 42 N-\(\delta_1\)-H is somewhat arbitrary, but is confirmed through 2D NOESY experiment as will be later discussed. The absence of NOE to hyperfine peaks when peak “i” at 12.2 ppm is irradiated (Figure 3.5C) suggested that peak “i” is not in contact with any of other hyperfine-shifted signals. The corresponding peak at 13.3 ppm with similar temperature dependence (Curie plot intercept at 3.6 ppm in comparison with 3.8 ppm for peak “i”) in HRP has been assigned to the C-\(\varepsilon_1\)-H of the distal histidine (34). By analogy, peak “i” for SBP can also be assigned to the C-\(\varepsilon_1\)-H of the distal histidine (His42).

The presence of the extremely broad, fast relaxing, far upfield-shifted peak “z” at −28.3 ppm is a characteristic feature of the existence of the proximal histidine within the heme crevice. This peak has been previously detected for the low-spin cyanide adducts of a number of heme peroxidases, such as HRP, CcP, Lip, LPO, and MPO, and is assigned to the C-\(\varepsilon_1\)-H of the proximal histidine imidazole ring (34, 50, 56, 64-65). The line width and chemical shift values of peak “z” in SBP-CN\(^-\) match quit closely to those of the
corresponding peak in HRP-CN\(^{-}\), suggesting the imidazolate character of the proximal histidine for SBP-CN\(^{-}\) as was earlier determined for HRP-CN\(^{-}\) (50, 65). Additional support for the presence of the proximal histidine is found in the observation of another broad, very fast relaxing signal “y” at 30.1 ppm which has been assigned to the C-\(\varepsilon_2\)-H of the proximal histidine in several heme peroxidases (45).

Signals “d” and “g” are assigned to the C-\(\beta\)-Hs of the proximal histidine (His169), as an intense NOE is observed for resonance “g” at 16.3 ppm when peak “d” at 21.4 ppm is irradiated (Figure 3.4D). Support for this assignment is found from the strong reciprocal NOE to peak “d” when signal “g” is saturated (Figure 3.5B), confirming the geminal nature of peak “d” and “g”. Saturation of peak “d” (Figure 3.4D) and “g” (Figure 3.5B) both yields NOEs to peak “i” and several common peaks in the diamagnetic region. Integration of peak “d” reveals that two protons are overlapped at the same position at all temperatures in SBP-CN\(^{-}\), which is the same as in HRP-CN\(^{-}\). Correspondingly the other proton at 21.4 ppm is assigned to the His 169 C-\(\delta_2\)-H.

Another unfortunate two-proton overlap occurs at peak “i” at 12.2 ppm, which shows weak or strong NOE correlation upon irradiation of peak “d” or “g”, respectively. However, as mentioned above, no reciprocal NOE is observed while saturating peak “i”, which has been assigned as the distal His42 C-\(\varepsilon_1\)-H. Therefore the other proton must arise from one of the proximal His170 protons. The corresponding peak in HRP at 11.98 ppm, which is better resolved without overlapping, is assigned to the proximal histidine N\(_{pep}\)H through 2D NMR methods. Therefore, the proton overlaps with peak “i” is tentatively assigned to the His169 N\(_{pep}\)H, which is resistant to isotope exchange in holoprotein, and waiting to be proved by 2D NMR experiments.
The two labile protons “h” and “j” are assigned through comparison with HRP and CcP cyanide complexes to the $N_{61}$ protons of the distal and proximal histidine, respectively. The assignment of peak “h” is relatively straightforward because of the similar chemical shift position as comparing HRP and CcP and has been proved by NOESY experiment as shown later. However, the peak “j” was originally assigned to the $N_{pep}H$ of the proximal histidine by solely comparing the chemical shift value. As just mentioned above, the $N_{pep}H$ is resistant to isotope exchange with exchange lifetimes of several years (66), which is in contrast with the rapid exchangeable behavior of proton “j”. Therefore, the proton “j” is assigned to the most possible candidate, which is the His169 $N_{61}H$ who has a matching Curie plot extrapolated intercept at infinite temperature with “j”.

Although more NOE correlations with proximal histidine protons in the diamagnetic region are revealed through 1D NOE experiments, no further attempt was made because of the relatively poor resolution within the complicated diamagnetic envelope. More confident assignment for these correlations can only be achieved through 2D NMR.

3.4.4. **Distal Arginine Assignments for the SBP-CN$^-$ Complex**

The presence of a distal Arg as catalytically critical residue was suggested to be highly conserved in HRP (47), CcP (30), and other plant heme peroxidases (67). The rSBP crystal structure also revealed a distal Arg38 located right above the prosthetic heme, with its extended, essentially planar trimethylene side chain extends over pyrrole III. The position and T$_1$ of two upfield hyperfine shifted signals “s” at -4.6 ppm and “t” at -7.7 ppm are well comparable with resonances of -4.75 and -6.45 ppm from distal Arg of HRP-CN$^-$ (47). Therefore, signals “s” and “t” are assigned to Arg38 C-$\beta_1$-H and Arg38 C-
δ₁-H, correspondingly.

Both the result from HRP and the crystal structure of rSBP suggest that the trimethylene side chain extends over pyrrole III of the prosthetic heme, making contact with the heme 5-CH₃, 6-β-CH₂, and γ-meso-H. Although resonances from the heme 5-CH₃, and γ-meso-H are not assigned through 1D NOE methods in this study, irradiations of peak “s” and “t” (Figure 3.5D and E) both yield a clear NOE to peak “q” at -2.5 ppm, which has been formerly assigned to 6-propionate Hβ. Reciprocal NOE correlation between peak “q”, “s” and “t” (Figure 3.5F) indicating that Arg38 side chain does extend above pyrrole III, in complete agreement with the rSBP crystal structure.

3.4.5. 2D NOESY and COSY Results for the SBP-CN⁻ Complex

The NOESY experiments were carried out at different conditions: 308 K (Figure 3.7), and 313 K (Figure 3.8) in D₂O, and 323 K (Figure 3.9) in 90% H₂O and 10% D₂O, to minimize the affect of overlaps and to verify previous nonexchangeable and labile proton assignments. Most of the 1D NOE correlations are confirmed in NOESY spectra, especially for correlations between the hyperfine shifts. The assignments of the labile proton peaks “h” and “j”, together with the overlapping peaks “d” and “i”, are all confirmed as shown in Figure 3.9. Moreover, resonances which have been buried in the diamagnetic envelop in the 1D NMR results are now exposed with high resolutions and allow further assignments for the SBP-CN⁻ complex by comparison with HRP-CN⁻ NMR assignments. The assignments and corresponding chemical shift values at different conditions are shown in Table 3.2. All the four meso-protons and the 7-propionate protons of the prosthetic heme, together with the proximal His170 Hα and two distal Arg38 protons are successfully assigned through 2D NMR. The COSY spectrum in 90% H₂O
and 10% D₂O further confirmed the assignments by revealing the spin-spin coupling correlations between proximal and distal histidine protons, 4-vinyl protons and distal arginine protons (Figure 3.10).

In summary, 16 of all the 22 resonances of the prosthetic heme, all proximal His170 resonances, and all β and δ proton resonances of the distal Arg38 in SBP-CN⁻ complex are identified through the combination of 1D and 2D NOE NMR methods in this work. All hyperfine-shifted signals are unambiguously assigned. Additionally, the cis-, trans-orientation of the two vinyl groups and the orientation of the distal arginine are confirmed as consistent with the recombinant SBP crystal structure.

3.4.6. Attempted Calcium Removal and Reconstitution

Partial removal of calcium from SBP causes substantial changes in the proton NMR spectral features of both the ferric high-spin and cyanide adduct low-spin forms of the protein (Figures 3.12). For the native enzyme, the four heme peripheral methyl proton signals are observed at 79.3, 69.0, 59.5, and 55.8 ppm, while the signals for the Ca²⁺-depleted enzyme are located at 77.8, 66.9, 55.6, and 48.0 ppm. The heme methyl proton peaks of the low-spin cyanide complex of the Ca²⁺-depleted SBP are observed at 29.2 and 28.5 ppm, whereas those of the cyanide complex of the native SBP are located at 29.5 and 28.1 ppm. More minor spectral differences are also noticeable between these spectra. The spectral changes derived from the calcium removal of SBP are similar as reported for HRP (54-55), indicating that calcium ions play the same structural and functional roles in these two peroxidases.

Similar as in HRP, addition of an excess amount of CaCl₂ to the calcium “depleted” SBP regenerates the spectral changes caused upon calcium removal (data not shown),
indicating the reversible binding of calcium within the protein.

The spin state of the heme iron of the Ca\(^{2+}\)-depleted HRP is reported to be much different from that of the heme iron of the native protein (a thermal spin mixing between ferric high and low spin states vs. pure ferric high spin state) as is reflected in the dramatic changes of the chemical shift values of the two forms of the protein (54-55). It is also reported recently that the Ca\(^{2+}\) depletion of HRP does not change the alpha helicity of the protein, but it changes the whole tertiary structure to a loosened, molten globule-like conformation, introduces tertiary structural changes in the immediate vicinity of the heme, and also increase the overall planarity of the heme (68). The high similarity of spectral changes caused by calcium depletion for SBP and HRP suggests that calcium ions play same role in these two closed related enzymes.

3.4.7. Comparison between SBP and HRP

Proton NMR examination of both the high-spin native and the low-spin cyanide ligated forms of SBP suggests the presence of the same prosthetic group, iron protoporphyrin IX, in this protein. Confirmation of this comes from the identical NMR spectral pattern of the extracted prosthetic group from SBP to that of the standard hemin. NMR spectral analysis reveals proximal histidine ligation to the heme iron and the presence of a distal histidine in the active site vicinity. Moreover, the obvious similarity between the proton NMR hyperfine shifts of SBP-CN\(^-\) and HRP-CN\(^-\) indicates that the two enzymes adopt the same heme geometry. The similar structural features of the active cavity of SBP and HRP suggest similar reactivates of these two plant peroxidases, as is reflected by their similar substrate profile. The trans and cis orientations of the 2- and 4-vinyl groups of the heme in SBP and HRP are the same and therefore suggesting identical
heme orientations.

However, differences do exist between SBP and HRP. Major differences include the chemical shift and T_1 values of the two hyperfine-shifted methyl signals and the proton resonance positions of both the proximal and distal histidine residues as were shown in Figures 3.3 and Table 3.1. Additionally, the 2- and 4- vinyl β protons corresponding position are slightly different in SBP (2-H_{β-c} > 4-H_{β-t} > 4-H_{β-c} > 2-H_{β-t}) and HRP (2-H_{β-c} > 4-H_{β-t} > 2-H_{β-t} > 4-H_{β-c}). These dissimilarities can be attributed to the differences in amino acid sequence and/or tertiary structures of the two proteins. The difference of the iron out-of-planarity or heme ruffling between these two closely related peroxidases may also affect their NMR spectral properties significantly.

3.5. References


43. La Mar, G. N., and de Ropp, J. S. (1979) Assignment of exchangeable proximal histidine resonances in high-spin ferric hemoproteins: Substrate binding in horseradish peroxidase, *Biochemical and Biophysical Research Communications* 90, 36-41.


CHAPTER IV. \(^1\)H-NMR STUDY OF PARAMAGNETIC CYTOCHROME P450CAM-CYANIDE COMPLEXES: ASSIGNMENT OF HYPERFINE-SHIFTED HEME RESONANCES

4.1 Introduction

Cytochrome P450 (CYP) enzymes are a widely distributed superfamily of heme-containing, single chain monooxygenases involved in a diversity of vital processes. The P450 enzymes can catalyze the monooxygen insertion of a variety of hydrophobic compounds, such as steroids, fatty acids, and exogenous xenobiotics. All p450s share a common catalytically active species, compound I, which is formed by activating the heme-bound molecular oxygen through a heterolytic O-O bond cleavage (1-2). In contrasting with most other hemoproteins that possess a proximal histidine ligand, the distinctive and highly conserved active site structure in all P450 enzymes utilizes a cysteine thiolate coordinating with the prosthetic heme as the proximal ligand (3).

In spite of the large amount of physicochemical data collected over the years about structure function relationships in P450s, detailed understanding of the effect of substrate binding on the changes of the protein structure are required. Particularly attention shall be put near the dioxygen binding site at the heme iron and in the I-helix groove, because of their suggested important role for oxygen activation (4). Understanding the mechanism and specificity of substrate binding in the cytochrome P450 superfamily is of great importance in explaining its key role in drug metabolism, toxicity, xenobiotic degradation, and several important biosynthetic pathways.

Among the various members of the P450 superfamily, cytochrome P450cam (EC 1.14.15.1; CYP 101; P450cam hereafter), from bacteria *Pseudomonas putida*, is the most
extensively studied and best characterized CYP because of its high solubility, relative stability, and ease of purification. Especially, the mature recombinant enzyme production using *Escherichia coli* expression system makes P450cam more accessible for studies for most purposes. The natural function of P450cam is to catalyze the stereo- and regiospecifically hydroxylation of (1R)-camphor. The heme prosthetic group of P450cam has been shown to adopt a conformation (Figure 4.1) (5) different from the heme in HRP (6) and SBP (7), but similar to that in CPO (8) and CcP (9).

![Figure 4.1 schematic presentations of structure and numbering system for iron protoporphyrin IX in P450cam with both the 2- and 4-vinyls in the cis orientation.](image)

Among the numerous chemical and spectroscopic techniques that are used for the structural investigation of hemoproteins, NMR spectroscopy and x-ray crystallography
have been proved as the most informative and authoritative methods for characterizing the structure of paramagnetic metalloenzymes. NMR spectroscopy, comparing with x-ray crystallography, is more emphasized on analyzing the active site heme cavity environment of hemoproteins in solution, which is more similar to the physiological conditions. Therefore, NMR spectroscopy can provide valuable complementary information to the solid state crystal structures and also enables the possibility of dynamic investigations of hemoprotein-substrates interactions.

Recently, x-ray crystallography has been employed in many studies of P450cam to illustrate enzyme conformation changes and functions of heme propionate groups (10-15). However, only a few controversial NMR studies of paramagnetic P450cam have been reported with only heme methyls assigned even with no reported progress after the year 2000 (16-21), in great contrast with the extensive and in depth NMR studies on heme peroxidases, such as CcP (22-34), HRP (6, 23, 35-51), and CPO (52-57). The stagnation in the paramagnetic P450cam NMR study is mainly caused by the proximal cysteine ligation imparted inefficiency of relaxation and magnetic properties to the heme iron electron, which adversely affect $^1$H NMR spectra resulting short $T_1$, large linewidths and small magnetic anisotropy (58). As a result it is more difficult to extract useful information from protons in proximity to the heme iron. Moreover, the lack of a characteristic structural model such as CcP in peroxidase studies, the relative instability of P450cam compared to heme peroxidases, and the difficulty in the reconstitution of isotopic labeled heme, all exist as obstacles for detailed NMR studies of P450cam.

The urgent need of definitive resonance assignments of P450cam NMR spectrums, together with the successful application of the most powerful two-dimensional (2D)
NMR techniques in characterizing the heme active site structure of chloroperoxidase, which is also a cysteine thiolate hemoprotein, triggered my ambition to use the two-dimensional NMR methods to unambiguously assign the major hyperfine-shifted signals and consequently elucidate the active site structure of P450cam in solution. I present herein the results of a comprehensive NMR study of P450cam-cynaide complexes with and without substrate camphor presence, by utilizing a combination of 2D NMR methods and the Curie intercepts obtained from variable temperature experiments. The results lead to the unequivocal assignment of all heme hyperfine-shifted signals, together with certain correlated diamagnetic resonances. The resulting assigned hyperfine shift pattern yields a valuable insight into the heme cavity of P450cam and will facilitate future assignments of the substrate and amino acid side-chain signals of interest.

4.2 Materials and Methods

4.2.1 Materials

*Escherichia coli* Rosetta II (DE3) cells were from Invitrogen (Carlsbad, CA). Dithiothreitol (DTT), carbon monoxide, hydrogen peroxide, were purchased from Sigma-Aldrich (St. Louis, MO). DEAE sepharose fast flow and Sephadex G-75 were purchased from GE Healthcare (Piscataway, NJ). Water with 18.2 megohm·cm resistivity were freshly produced from a Milli-Q BioCel Water Purification System (Millipore, Billerica, MA). All other reagents were of the highest purity available from Fisher Scientific (Pittsburg, PA). All chemicals were used without further purification.

4.2.2 Preparation of Protein Samples

The wild type (WT) P450cam protein were expressed in *Escherichia coli* strain Rosetta II (DE3) transformed with a plasmid constructed by ligation of pET-30a (+) with
P450cam gene at the restriction sites of NdeI and HindIII. The bacteria were grown in the terrific broth media containing 100mg/L of kanamycin at 37 °C and induced with 1mM IPTG at OD$_{600}$ over 0.8. Induction was allowed to proceed at 30 °C for another 18 hours prior to harvesting by high-speed centrifugation. Cell pellets were resuspended and lysed at 4 °C for 30 minutes in 50 mM Tris-HCl buffer at pH 7.4 containing 50 mM KCl, 0.5 mM DTT, 1 mM camphor, 1 mM EDTA, 200 μM PMSF, 40 U/ml Dnase I, 3 U/ml Rnase A, and 2 mg/ml lysozyme. The sonicated and filtrated cell lysate were then concentrated by ultrafiltration (Amicon cell) with a 30,000 Da cut-off membrane and purified by passage through a fast flow DEAE Sepharose column (50 ml) equilibrated with 50 mM Tris-HCl buffer (pH 7.4). After the initial wash, the protein was eluted with a linear 0 to 0.5 M KCl gradient. The red fractions containing the ~45 kDa proteins were pooled for further purification by size exclusion gel filtration chromatography (Sephadex G-75, 500 ml) with 100 mM potassium phosphate buffer at pH 7.4. All the steps in the protein purification were performed at 4 °C. The purification steps typically yield pure P450cam protein with Rz values ($A_{417}/A_{280}$) higher than 1.4. The concentration of the enzyme sample was determined using the Soret extinction coefficient $\varepsilon_{417} = 115$ mM$^{-1}$cm$^{-1}$ for the native low-spin camphor-free P450cam (59).

### 4.2.3 NMR Spectroscopy

Proton nuclear magnetic resonance ($^1$H NMR) spectra for the native and cyanide-bound proteins with or without excess camphor were recorded at 298 K on a Bruker Avance 600 (Ultrashield) spectrometer operating at a proton frequency of 599.93 MHz. Protein samples for NMR experiments were prepared in D$_2$O buffer (10 mM phosphate, pH 7.4, uncorrected for isotope effect), by at least five times of isotope exchanges.
through a centrifuging microconcentrator (Centricon-30) at 4 °C. All NMR samples contained more than 2 mM purified P450cam and the cyanide-adducts of the protein were prepared by the addition of a 20-fold molar excess of cyanide from a freshly made 1 M stock solution of KCN in 99.9% D₂O. The camphor bound protein samples were prepared by the addition of 1 M camphor stock solution in deuterated methanol-d₄ into the protein solutions until the final camphor concentration reached 7.8 mM.

The ¹H NMR spectra of the ferric low-spin forms (native P450cam without camphor bound, and camphor free and camphor bound P450cam cyanide-adducts) were obtained by collecting 1000-2000 scans with 4 K data points over a 30-40 kHz bandwidth, and a repetition time of 0.2 seconds with solvent presaturation during relaxation delay. A 16-bit digitizer was used and the spectra were obtained by multiplication of the free induction decays with a 20 to 40 Hz apodization. Fast repetition spectra of the low-spin ferric protein forms were recorded with a 125 kHz sweep width, 2048 data points, and a repetition rate of 10 ms to detect the fast relaxing proton signals which are intrinsically broad. The NMR spectrum of the high-spin ferric camphor-bound P450cam was collected with a 125 kHz spectral width, 16384 data points, and a repetition rate of approximately 50 per second. A 12-bit digitizer was used and 20,000 scans were averaged. Free induction decays were exponentially multiplied by a line broadening factor of 100 Hz before Fourier transformation to improve the signal-to-noise ratio of the spectra. The residual solvent signal was suppressed with presaturation during relaxation delay. Chemical shift values were referenced to the residual HDO signal at 4.76 ppm.

The spin-lattice relaxation times (T₁) of the hyperfine-shifted protons for the cyanide complexes of camphor free and camphor-bound P450cam were determined by a variation
of the standard inversion-recovery sequence to include a composite 180° pulse. A repetition rate of 3.3 per second and a 90° transmitter pulse of 7.6 μs was used. The T1 values were estimated from the null point with the relationship of $T_1 = \tau/\ln 2$. The NOE experiments on the cyanide complexes of the protein were performed similar to what is described in Chapter 3, with the irradiation time of 100 ms and a weak decoupler power of 60 dB.

Phase-sensitive NOESY spectra for the cyanide-bound derivatives of P450cam were acquired at 298 K with mixing times ranging from 15 to 20 ms. Typical NOESY spectra were collected with 256 experiments in the F1 dimension. In general, 400 scans were accumulated for each F1 experiment, which was acquired with 4096 complex points in the F2 dimension over a spectral width of 45 or 65 ppm. The residual solvent signal in all NOESY experiments was suppressed using a 200-ms presaturation with a weak decoupler power. COSY spectra were collected with all parameters similar to NOESY.

Pulse sequences utilized in all the experiments employed single 90° hard pulses to maximize the bandwidth of excitation. The Soret band at 440 nm, which is characteristic for the P450cam/cyanide and P450cam/cyanide/camphor complexes, did not shift throughout all the NMR study.

4.3 Results

4.3.1 Proton NMR of the Ferric Forms of the Native Low-Spin Camphor-Free P450cam and the Native High-Spin Camphor-Bound P450cam

The proton NMR spectra of the ferric forms of P450cam (Figure 4.2) are essentially identical to the results reported previously (17, 19, 21). Without the cyanide ligation, the NMR spectrum of the native ferric camphor-free P450cam (Figure 4.2A) exhibits a six-coordinated low-spin feature with a water molecule serving as the distal ligand. The
hyperfine-shifted 19.5 ppm and 14.2 ppm peaks had been assigned previously (19) to the 5- and 1-methyl groups of the prosthetic heme, respectively. A novel far upfield-shifted, fast relaxing broad resonance at -21.7 ppm (Figure 4.2A inset) is noticed for the first time. Based on the extremely short $T_1$ of ~1 ms, the relaxation properties, the large contact shift, the expected distance to the heme iron center, and the similarity of such resonances from CPO (57), this resonance is assigned to one of the $\beta$-CH$_2$ protons of Cys357, the heme iron proximal ligand. Unfortunately, this substrate-free resting state enzyme exhibits a relatively poor stability, which prevents the further application of extensive NMR study.

The native camphor-bound P450cam reveals a five-coordinated high-spin nature (Figure 4.2B) with the distal water expelled upon substrate binding. The resonances at 62.9, 58.6, 40.6, and 36.4 ppm had been tentatively assigned to the four heme methyl groups with the order $8 > 3 > 5 > 1$ (19), which is different from the previous proposed assignments with the order $5 > 1 > 8 > 3$ (21). The considerably broad resonances in the NMR spectrum can provide only limited information about the structural properties of the enzyme. Therefore, no further efforts were made to analyze the spectra of these resting states P450cam in this study.

4.3.2 Proton NMR of the Ferric Low-Spin Cyanide-Adducts of Camphor-Free and Camphor-Bound P450cam

The NMR spectral properties of the cyanide-bound, ferric low spin derivatives of camphor-free and camphor-bound P450cam have been focused on in this study, because their short electronic relaxation times and large magnetic anisotropy yield much sharper
Figure 4.2 600 MHz $^1$H NMR spectra of the ferric forms of (A) low-spin resting state camphor-free P450cam in D$_2$O and 100 mM pH 7.0 phosphate buffer with the inset showing the -21.7 ppm peak, (B) high-spin camphor-bound P450cam in D$_2$O and 100 mM pH 7.0 phosphate buffer containing 4 mM d-camphor, (C) low spin cyanide complex of the camphor-free P450cam in D$_2$O and 100 mM phosphate buffer (pH 7.0) with the inset showing the -21.7 ppm peak, and (D) low spin cyanide complex of the camphor-bound P450cam in D$_2$O and 100 mM pH 7.0 phosphate buffer containing 7.8 mM d-camphor at 298 K with the inset showing the 33.1 ppm peak.
and better resolved signals in their proton NMR spectra. Therefore, valuable information about the electronic and molecular structural properties of the heme pocket can be provided.

The proton NMR spectrum of the substrate-free P450cam-CN\textsuperscript{−} complex in D\textsubscript{2}O solution is shown in Figure 4.2C. The paramagnetically shifted signals were observed with high similarity to that of other heme peroxidase cyanide derivatives (57, 60). The three intense hyperfine-shifted signals “A”, “B”, and “O” with integrated intensities of three protons each are typical of heme methyl groups. Such three methyl peak homologies have been tentatively assigned to the 5-, 1- and 3- heme methyls for the substrate-free P450cam-alkyl isocyanide complexes, respectively (16, 19), which is in contrast to the previously proposed assignments of the pyridine adduct (17) and the cyanide adduct (21). The newly discovered far upfield resonance in the native ferric camphor-free P450cam is also observed after cyanide binding as shown in the inset of Figure 4.2C labeled as peak “Q” at -21.7 ppm with very short T\textsubscript{1} of ~1 ms. Accordingly, this peak “Q” is assigned to the Cys357 \(\beta\)-proton for the same reason as mentioned above.

The proton NMR spectrum of the camphor-bound P450cam-CN\textsuperscript{−} complex in D\textsubscript{2}O solution shown in Figure 4.2D reveals a far downfield-shifted broad peak “a”, which is also strongly relaxed with a short T\textsubscript{1} of ~1 ms. The single proton Peak “a” is previously assigned to a proximal Cys357 C\(\beta\)H in P450cam (21) and CPO (57). The upfield peak at \(-21.7\) ppm, which is observed in either native or cyanide-bound forms of camphor-free P450cam, is not detected with camphor presence, indicating a conformational change around the proximal cysteine upon substrate binding. The binding of camphor to P450cam-CN\textsuperscript{−} complex results in 2-4 ppm low-field bias of the heme methyls (Figure 4.2 C&D). The assignments of methyl peak “b” and “f” to 8-CH\textsubscript{3} and 3-CH\textsubscript{3} was suggested
on the basis of the tentative assignments in the pyridine complex (17). However, disagreement was proposed by Mouro et al., stating that all low-spin P450cam complexes should have the same chemical shift order of the heme methyl groups 5 > 1 > 8 > 3 (19).

The assignment of the hyperfine-shifted signals for the camphor-free and camphor-bound P450cam-CN⁻ complexes was achieved through comparison with the cyanide-adducts of CPO and other heme peroxidases (23, 31, 34, 40-41, 57), in combination with the P450cam crystal structure analysis and the correlation information revealed by the NOESY and COSY spectrum. The chemical shifts and the corresponding diamagnetic shift values predicted from Curie plot intercept at T⁻¹ = 0, as well as the spin-lattice relaxation times for the hyperfine-shifted resonances and their assignments in the ferric cyanide-adducts of camphor-free and camphor-bound P450cam are compiled in Table 4.1, along with the previously reported corresponding parameters in CPO (57).

The NOESY spectrum of the ferric camphor-free and camphor-bound P450cam-CN⁻ in D₂O are shown in Figure 4.3 and 4.4, respectively. For the relatively simple camphor-free P450cam-CN⁻, unambiguous assignment of the hyperfine shifts is achieved based only on the well characterized NOESY spectrum. While for the more complicated camphor-bound P450cam-CN⁻ spectrum, NOESY information combining with the 1D NOE experiments (Figure 4.5B-E), which have higher resolution in the paramagnetic region, as well as the scalar connectivities revealed by COSY (Figure 4.6) lead to the unambiguous assignment for most of the nonexchangeable hyperfine-shifted protons. Some resonances buried in the diamagnetic envelope complex region are also tentatively assigned in the 2D maps. The suggested assignments for hyperfine shifts are further verified by the Curie intercepts predicted diamagnetic shift of the resonances (Figure 4.7).
Table 4.1 Proton NMR chemical shift (ppm), T_1 (ms), and Curie plots intercept (ppm) values of the assigned resonances for the low-spin cyanide derivatives of camphor-free (− cam) and camphor-bound (+ cam) P450cam, and CPO\textsuperscript{a}.

<table>
<thead>
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<tr>
<td>Fe\textsuperscript{III} P450cam-CN\textsuperscript{−} (-cam)</td>
<td>ppm</td>
<td>ms</td>
<td>ppm</td>
</tr>
<tr>
<td>A</td>
<td>24.3</td>
<td>36</td>
<td>b</td>
</tr>
<tr>
<td>B</td>
<td>16.1</td>
<td>34</td>
<td>f</td>
</tr>
<tr>
<td>C</td>
<td>13.3</td>
<td>31</td>
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</tr>
<tr>
<td>D</td>
<td>12.0</td>
<td>30</td>
<td>g</td>
</tr>
<tr>
<td>E</td>
<td>11.5</td>
<td>16</td>
<td>h</td>
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<tr>
<td>F</td>
<td>8.52</td>
<td>---</td>
<td>b</td>
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<td>G</td>
<td>0.5</td>
<td>---</td>
<td>c</td>
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<td>H</td>
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<tr>
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<td>35</td>
<td>q</td>
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<tr>
<td>J</td>
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<td>35</td>
<td>w</td>
</tr>
<tr>
<td>L</td>
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<td>N</td>
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<tr>
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<td>c</td>
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<tr>
<td>P</td>
<td>-5.0</td>
<td>38</td>
<td>u</td>
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<tr>
<td>Q</td>
<td>-21.7</td>
<td>b</td>
<td>w</td>
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\textsuperscript{a} Results for CPO-CN\textsuperscript{−} are from ref (57) and the values from CPO assignment are symmetrically switched along the α-γ meso axis for comparison purpose. For example, the assigned CPO 8-CH₃ results are used to compare with the corresponding values of the 5-CH₃ in P450cam.

\textsuperscript{b} The assignment is not achieved or does not exist in this form.

\textsuperscript{c} The signal is located in the diamagnetic region and is not labeled with letter.
Figure 4.3 The 600 MHz phase-sensitive $^1$H NOESY spectra of 2.0 mM substrate-free P450cam-CN$^-$ in D$_2$O taken at 298 K with a mixing time of 15 ms. Cross-peak assignments are as follows: 1, 5-CH$_3$:6-H$_{\alpha}$; 2, 5-CH$_3$:6-H$_{\alpha}'$; 3, 5-CH$_3$:H$_{\beta\text{meso}}$; 4, 1-CH$_3$:2-H$_{\alpha}$; 5, 1-CH$_3$:H$_{\delta\text{meso}}$; 6, 1-CH$_3$:2-H$_{\beta\text{trans}}$; 7, 1-CH$_3$:2-H$_{\beta\text{cis}}$; 8, 2-H$_{\alpha}$:H$_{\alpha\text{meso}}$; 9, 2-H$_{\alpha}$:2-H$_{\beta\text{trans}}$; 10, 2-H$_{\alpha}$:2-H$_{\beta\text{cis}}$; 11, 2-H$_{\alpha}$:3-CH$_3$; 12, 6-H$_{\alpha}$:6-H$_{\alpha}'$; 13, 2-H$_{\beta\text{cis}}$:2-H$_{\beta\text{trans}}$; 14, 4-H$_{\beta\text{cis}}$:4-H$_{\beta\text{trans}}$; 15, 3-CH$_3$:H$_{\alpha\text{meso}}$; 16, 3-CH$_3$:4-H$_{\alpha}$; 17, 3-CH$_3$:4-H$_{\beta\text{cis}}$; 18, 3-CH$_3$:4-H$_{\beta\text{trans}}$; 19, 4-H$_{\beta\text{trans}}$:4-H$_{\alpha}$. 
Figure 4.4 The 600 MHz phase-sensitive $^1$H NOESY spectra of 2.0 mM camphor-bound P450cam-CN$^-$ in D$_2$O taken at 298 K with a mixing time of 15 ms. Cross-peak assignments are as follows: 1, 5-CH$_3$:6-H$\alpha$; 2, 5-CH$_3$:6-H$\alpha^{'}$; 3, 5-CH$_3$:H$\beta$$_{meso}$; 4, 5-CH$_3$:6-H$\beta$; 5, 5-CH$_3$:6-H$\beta^{'}$; 6, 2-H$\alpha$:1-CH$_3$; 7, 2-H$\alpha$:H$_{meso}$; 8, 2-H$\alpha$:3-CH$_3$; 9, 2-H$\alpha$:2-H$_{trans}$; 10, 2-H$\alpha$:2-H$_{cis}$; 11, 6-H$\alpha$:6-H$\alpha^{'}$; 12, 6-H$\alpha$:6-H$\beta$; 13, 6-H$\alpha$:6-H$\beta^{'}$; 14, 1-CH$_3$:H$_{meso}$; 15, 1-CH$_3$:2-H$_{trans}$; 16, 1-CH$_3$:2-H$_{cis}$; 17, 2-H$_{cis}$:2-H$_{trans}$; 18, 4-H$_{cis}$:4-H$_{trans}$; 19, camphor proton: camphor proton. Cross-peak 6 can only be seen with lower contour levels.
Figure 4.5 The 600-MHz proton NMR spectrum of (A) 2 mM ferric low-spin camphor-bound form of P450cam-CN' in D$_2$O and 100 mM pH 7.0 phosphate buffer containing 7.8 mM d-camphor at 298 K. (B) trough (E) are the NOE difference spectra obtained upon irradiation of the hyperfine-shifted signals “b”, “c”, “f”, and “w”, respectively. The irradiated peaks are indicated by a downward arrow in each difference spectrum.
Figure 4.6 The 600 MHz phase-sensitive $^1$H COSY spectra of 2.0 mM camphor-bound P450cam-CN$^-$ in D$_2$O taken at 298 K after symmetrization for better resolution; note the different scale from that in the NOESY spectrum. Cross-peak assignments in complete agreement with previous NOESY assignments for hyperfine-shifted protons are as follows: 1, 2-Hα:2-H$_{\beta\text{trans}}$; 2, 2-Hα:2-H$_{\beta\text{cis}}$; 3, 6-Hα:6-Hα$'$; 4, 2-H$_{\beta\text{trans}}$:2-H$_{\beta\text{cis}}$; 5, camphor proton: camphor proton.
Figure 4.7 Curie plots of observed shift vs. reciprocal temperature for the resolved hyperfine-shifted resonances of camphor-bound P450cam-CN in the (A) downfield and (B) upfield regions. The letter designation of each peak following the scheme of Figure 4.2D is given to the right of each line, and the extrapolated diamagnetic intercept at $T^{-1} = 0$ is given in Table 4.1.

4.4 Discussion

4.4.1 Assignment of the Hyperfine-Shifted Resonances for Ferric Camphor-Free P450cam-CN- Complex

The ferric camphor-free P450cam-CN- complex shows a typical hemoprotein-cyanide adduct NMR spectrum (Figure 4.2C) with the two well characterized heme methyl groups located at the most downfield region. These two peaks “A” and “B”, as discussed before, shall arise from either 5-CH$_3$ and 1-CH$_3$ or the 8-CH$_3$ and 3-CH$_3$ pair on heme depending on the position and orientation of the proximal ligand (61-62). This is confirmed by the NOESY connections as shown in Figure 4.3, in which peak “A” gives
weak NOEs to peak “F” at 11.1 ppm and peak “G” at 8.5 ppm, typical as the NOE pattern of a heme methyl near a propionate group as shown in many heme peroxidases such as HRP, CcP, and CPO (28, 32, 34, 44, 57). The methyl peak “B”, on the other hand, yields a very weak NOE to peak “C” at 13.3 ppm and two weak NOEs to the upfield peaks “I” at -0.2 ppm and “L” at -1.3 ppm. Clear NOEs are also observed between “C”, “I” and “L”, which is in perfect agreement with the three protons from a same heme vinyl group. Therefore, peak “B” must be either 1- or 3-methyl of the prosthetic heme.

Another methyl peak “O” in the upfield region at -3.3 ppm is observed and has been suggested to be the other heme methyl next to a vinyl group in contrast with the methyl peak “B”. This is confirmed by the weak NOEs between peak “O” and two peaks “M” at -1.6 ppm and “P” at -5.0 ppm, two resonances located in the typical upfield region similar to heme vinyl-β protons in other hemoproteins (23, 40, 57). The last heme methyl resonance in pairing with peak “O” must be buried in the diamagnetic envelope, which is a common feature for most hemoproteins. Interestingly, a weak NOE is observed between peak “C” and peak “O”, which lead to the solid conclusion that peak “O” must arise from the 3-CH₃ because peak “C” has been assigned to the proton from a vinyl group that is near to another heme methyl “B”. Since both vinyl groups in P450cam heme adopt the cis orientation (Figure 4.1), peak “C” can be unambiguously assigned to the 2-vinyl Hα, and consequently, resonance “B” are assigned to the 1-CH₃, with the paring methyl “A” assigned to the 5-CH₃. Resonances “F” and “G” therefore must arise from the 6-propinate α-protons. The assignments of three heme methyl groups results a chemical shift order of 5 > 1 > 8 > 3, in perfect agreement with the previous assignments using magnetization transfer method and isocyanides (19), and in disagreement with the earlier
putative assignment with the order $8 > 3 > 5 > 1$, which was determined solely on homology with HRP ($17, 21$).

The vinyl $\beta$-protons are assigned by comparing their relative NOE strengths to decide the cis- or trans- identity. For example, comparing with “L”, resonance “I” reveals a much stronger NOE with the 1-CH3 peak “B”, and a weaker NOE with peak “C”, yielding the assignment of “L” to 2-vinyl $\text{H}_{\beta}\text{-cis}$ and “I” to 2-vinyl $\text{H}_{\beta}\text{-trans}$. Similarly, the resonances “M” and “P” are assigned to 4-vinyl $\text{H}_{\beta}\text{-cis}$ and 4-vinyl $\text{H}_{\beta}\text{-trans}$, respectively, and a peak at 0.5 ppm showing NOE with both of the 4-$\text{H}_{\beta}$’s is assigned to the 4-$\text{H}_{\alpha}$.

The heme $\alpha$-, $\beta$-, and $\delta$-mesoprotons are tentatively assigned by careful examination of the NOESY correlations with their geminal partners (2-$\text{H}_{\alpha}$ & 3-CH3, 5-CH3, and 1-CH3, respectively) within the proposed common mesoproton resonance range around 1-7 ppm ($40$).

4.4.2 Assignment of the Hyperfine-Shifted Resonances for Ferric Camphor-bound P450cam-CN- Complex

The binding of camphor results a more complicated spectrum (Figure 4.2D), with both the downfield methyl signals shifted by 2-4 ppm to the low-field and the upfield methyl peak overwhelmed by the diamagnetic envelope, indicating an electronic field change in the heme cavity induced by the substrate binding.

The decided chemical shift order of heme methyl groups of $5 > 1 > 8 > 3$ was suggested to be consistent in all ferric low-spin P450cam forms ($19$). Therefore, methyl peaks “b” at 22.0 ppm and “f” at 11.4 ppm are assigned to the 5- and 1-methyl, respectively. The NOE correlations between peak “b” and two downfield-shifted peaks
“d” and “h” are consistent in both 1D and 2D NOE spectra (Figure 4.4 & 4.5B), leading the assignment of “d” and “h” to the two 6-propionate α-protons, which is confirmed by the COSY correlation between “d” and “h” (Figure 4.6). Furthermore, saturation of peak “b” yields detection of a peak at 0.9 ppm and peak “o” at -0.2 ppm in the 1D NOE difference spectrum, which is also revealed in the NOESY spectrum with peak “d” showing the same NOE pattern. Because of the indicated geminal relationships between these resonances and the fact that propionate β-protons are normally located around the upfield region, the “0.9 ppm” peak and peak “o” are assigned as the two 6-propionate β-protons.

The NOE pattern of the assigned 1-CH₃ resonance “f” with a downfield-shifted peak “c” at 14.0 ppm and two upfield-shifted peaks “q” at -0.6 ppm and “s” at -1.6 ppm are very similar to that in the camphor-free P450cam-CN⁻. Correspondingly, peak “c” is assigned to the 2-vinyl Hα and signals “q” and “s” are assigned to 4-vinyl Hβ-trans and 4-vinyl Hβ-cis, respectively. These assignments are solid proved by the COSY result, and therefore, the early proposed putative assignment of peak “S” to an amino acid methyl group (21) must be misled by an overlapping at this position.

The 3-CH₃ is identified from the diamagnetic envelope through careful examination of both 1D and 2D NOE experiments. Similar NOE correlations between 2-Hα and a peak at 0.4 ppm as in the camphor-free P450cam-CN⁻ leads to the assignment of the 3-methyl protons. Assignment of the β-protons from the 4-vinyl group is achieved by revealing the geminal relationships between peaks “p” at -0.3 ppm and “u” at -2.1 ppm from NOESY spectrum without confirmation from the COSY, which is because of the relatively low resolution in that area. The heme α-, β-, and δ-mesoprotons are assigned
again within the 1-7 ppm region with information from NOESY correlations with their geminal heme resonances. The assigned resonances are closely located to the previously assigned mesoproton resonances in the camphor-free P450cam cyanide derivative, suggesting the success of these assignments.

The most upfield shifted resonances “w” displays no correlation with any paramagnetic resonance, indicating it is not derived from any heme proton. Nonetheless, the short T1 (16 ms) reveals that it is in proximity to the heme iron, suggesting its origin from the proton of the substrate camphor, which is located closely to the heme iron in the distal side. The intensive cross peak between the peak “W” and a resonance at 2.3 ppm shown in the COSY map (Figure 4.6) reveals the position of another camphor proton. Further verification of this assignment can be accomplished by using deuterated camphor as substrate.

The unassigned downfield peak “e” and the 1-methyl resonance “f” share several common NOE correlations with peaks in the diamagnetic region, indication that peak “e” may arise from an amino acid near the 1-methyl group.

4.4.3 The Heme thiolate ligand Cys357

The very broad, rapid relaxing, strongly hyperfine-shifted peaks “a” and “Q” at 33.1 and -21.7 ppm share very similar chemical shift value, relaxation time and Curie intercept with the CPO-CN$^-$ hyperfine-shifted resonances at 39.0 ppm and -20.7 ppm, which have been firmly assigned to the $\beta$-CH$_2$ protons of the coordinated proximal cysteine ($57$). However, in contrast to in CPO, the two resonances in P450cam can not coexist at the same time unless in a mixture of camphor-bound and camphor-free forms of P450cam-CN$^-$.

The resonance “a” are only detected in the camphor-bound ferric P450cam-CN$^-$
complex, while resonance “Q” exists in both native and cyanide derivative of camphor-free P450cam, indicating a distance and orientation change upon camphor binding. This camphor induced conformational change is proved by the distances measured from the crystal structure of camphor-free native P450cam and camphor-bound P450cam-CN⁻ (Table 4.2). It is shown that the distances between both the Cys357 𝛽-CH₂ protons and the heme iron increase by ~0.3 Å upon camphor binding. Additionally, the camphor bound induces one of the Cys 𝛽-CH₂ protons shifting toward the 5-methyl group but away from the 𝛾-meso direction. Meanwhile the other Cys 𝛽-CH₂ proton shifts away from both the 5-methyl and the 𝛾-meso directions, indicating an orientation twist of the proximal Cys induced by camphor. The distances of the Cys 𝛽-CH₂ protons to the heme iron in CPO is right in between of the distances in camphor-free and camphor-bound P450cam. This may explain why both of the Cys 𝛽-CH₂ proton resonances can emerge at the same time in CPO-CN⁻.

**Table 4.2** Distances between proximal Cys 𝛽-protons and selected atoms in the low-spin cyanide adducts of camphor-free (- cam) and camphor-bound (+ cam) P450cam, and CPO.

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<th>Proximal Cys</th>
<th>Hβ</th>
<th>Hβ'</th>
<th>Hβ</th>
<th>Hβ'</th>
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<tr>
<td>Heme iron</td>
<td>2.99</td>
<td>4.06</td>
<td>3.29</td>
<td>4.34</td>
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<td>5-CH₃ carbon</td>
<td>4.37</td>
<td>5.58</td>
<td>4.28</td>
<td>5.84</td>
<td>4.27</td>
<td>5.68</td>
</tr>
<tr>
<td>𝛾-meso proton</td>
<td>3.72</td>
<td>5.25</td>
<td>4.42</td>
<td>5.74</td>
<td>4.46</td>
<td>5.95</td>
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<tr>
<th>(- cam) P450cam a</th>
<th>(+ cam) P450cam-CN⁻ b</th>
<th>CPO-CN⁻ c</th>
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<tr>
<td>Hβ</td>
<td>Hβ'</td>
<td>Hβ</td>
</tr>
<tr>
<td>2.99</td>
<td>4.06</td>
<td>3.29</td>
</tr>
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<td>4.37</td>
<td>5.58</td>
<td>4.28</td>
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<tr>
<td>3.72</td>
<td>5.25</td>
<td>4.42</td>
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**a** Distances are measured from the crystal data (PDB code 1PHC).

**b** Distances are measured from the crystal data (PDB code 1O76).

**c** Distances are measured from the crystal data (PDB code 2CIY).
In summary, the presented results provide the first unambiguous assignments for most of the hyperfine-shifted resonances of the ferric low-spin camphor-free and camphor-bound P450cam-CN\(^-\) complexes. These assignments are achieved mainly through 2D NMR methods with the careful analysis of the correlation information in combination with the well established and accepted assignment strategy from earlier hemoprotein NMR studies. The observed methyl order and NOE pattern are consistent with the prosthetic heme structure and previously proposed assignments. Some diamagnetic heme resonances are tentatively assigned according to the NOE correlations and assignments in other hemoproteins. The alternation of the novel proximal cysteine \(\beta\)-CH$_2$ resonances induced by camphor binding is an indication of the conformational change near the proximal side. The presented work solved the trouble that had hindered the $^1$H-NMR study of P450cam for a long time, and can shed more light on future NMR studies of other cytochromes P450 and P450 mutants.

4.5 References


<table>
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CHAPTER V. CONCLUDING REMARKS

In this dissertation I have investigated the native or modified active site structure of three hemoproteins with one- and two-dimensional NMR applications. The fundamental claim I have made in this dissertation is that, NMR techniques are very powerful for the structural elucidation of large organic molecule and hemoproteins.

In Chapter 2, the N-alkylated heme of the allylbenzene inactivated chloroperoxidase was studied, in which the modified heme was extracted, demetallated and characterized by UV-vis, ES-MS/MS, and extensive 1D and 2D NMR methods. The results firmly established the structure of the modified heme as a $\text{N}_{\text{III}}$-(3-phenyl-2-hydroxyalkyl) iron-porphyrin IX, which indicate that the allylbenzene induced CPO inactivation is highly regio- and stereospecific. The result of this study solved a long-lasting mystery and greatly enhanced our understanding about the active site environment and catalytic mechanism of CPO. Based on the combination of our results with the observation in CPO crystal structures, a new mechanism of CPO catalysis and topology of heme active site was proposed. In this mechanism, the phenyl groups of Phe103-Phe186 located at the end of CPO substrate accessing channel function as a clap and play important role in controlling the substrate binding position and orientation. This first-time proposed double-phenyl motif can well explain many catalytic activities displayed by CPO, such as regiospecific epoxidation and the formation of green CPO and its auto-restoration. Although more evidence is required to corroborate this novel motif, it extends and inspires our understanding of enzyme substrate binding strategy.

The biggest unresolved question from this study is that why the green heme monomer-CN\textsuperscript{−} complex does not show any correlation information in all NMR spectra,
such as COSY, NOESY, and ROESY. On the other hand, the green CPO shows no paramagnetically shifted signal, which cannot be explained by the formation of the green heme dimer within the protein. Another task is to further verify the proposed double-phenyl clamp mechanism, which could be explored by determining the crystal structure of CPO complexed with phenyl substrates. Site-specific mutation of the Phe103 and Phe186, and the application of theoretical calculations can also corroborate the elucidation of this mechanism. The future study of the mechanism-based inactivation of CPO can also include the study of allyacetate inactivated green CPO, because allyacetate has no phenyl group and therefore may reveal different binding strategy and N-alkylation preference to the CPO active site.

In Chapter 3, the active site structure of SBP was characterized by one- and two-dimensional NMR spectroscopic studies. All hyperfine shifted resonances from the heme prosthetic group and catalytically important amino acid residues including both proximal histidine that serves as the heme iron ligand and distal histidine that serves as the general acid-base catalyst during formation of compound I of SBP have been unambiguously assigned through (a) measurement of NOE connectivities, (b) prediction of the Curie intercepts from both one- and two-dimensional variable temperature studies, (c) comparison with assignments made for cyanide derivatives of several well characterized heme proteins such as cytochrome c peroxidase and horseradish peroxidase, and (d) examination of the crystal structural parameters of recombinant SBP. The NMR parameters and peak patterns of SBP are compared with those of HRP and CcP. Our study indicates that results from NMR investigations of the protein in solution can complement and verify the results revealed by x-ray diffraction studies of the crystal
form and thus provide a complete and better understanding of the actual structure of the protein. The similar NMR spectral patterns of SBP and HRP suggest very similar overall structural features of the heme cavity and similar activity of these two plant peroxidases. However, differences between the NMR property of the two proteins do exist and suggest structural differences of the two proteins are localized around the 2- and 4- vinyl groups of the heme, which may be responsible for the different activity and stability of the two proteins. No significant spectral changes were observed upon removal of calcium from SBP, indicating that calcium ions play similar structural and functional roles in these two peroxidases.

The next step in this study is to identify signals from all residues in the heme, as well as proximal and distal, environment, and eventually all residues in the heme periphery. Further assignments can be extended beyond those residues in immediate contact with the heme and also include all catalytically relevant residues in the proximal and distal environments. The resulted information from such study can further clarify the high stability and activity of SBP.

In Chapter 4, the first attempt for the assignments of most hyperfine-shifted resonances and some diamagnetic heme resonances of camphor-free and camphor-bound P450cam-CN⁻ complexes are successfully achieved through 2D NMR applications. The observed heme methyl chemical shifts order of $5 > 1 > 8 > 3$ and the NOE pattern are consistent with the prosthetic heme structure and finalized the controversial literature results. Resonances from the two $\beta$-CH$_2$ of the proximal cysteine are observed for the first time. Our results revealed that an alternation of the proximal cysteine $\beta$-CH$_2$ resonances could be induced by camphor binding, which indicated a conformational
change near the proximal side upon camphor binding.

Future work for this study is to assign more resonances from the prosthetic heme and residues around the heme cavity, and especially, from the substrate camphor. The kinetic NMR study of camphor binding is also a challenging but exciting task to solve the frustrating question about how camphor gets into the binding pocket. The NMR study of various P450cam mutants can also shed light on explaining the function of the mutated residues.
APPENDIX A

Sequences of Chloroperoxidase

General information

The wild-type chloroperoxidase protein consists of 299 amino acids as listed in Figure A.

Figure A. Amino acid sequence of full-length of WT Chloroperoxidase.

1  XEPGSGIGYP YDNNTLPYVA PGPTDSRAPC PALNALANHG YIPHDGRAIS
51  RELQNAFLN HMGIANSVIE LALTNAFVVC EYVTGSDCGD SLVNLTLLAE
101 PHAFEHDHSF SRKDYKQGVA NSNDFIDNRN FDAETFQTS SLVNDKTHFD
151 YADMNEIRLQ RESLSNELDF PGWFTESKPI QNVESGFIFA LVSDFNLPDN
201 DENPLVRIDW WKYWFTNESF PYHLGWHPPS PAREIEFVTS ASSAVLAASV
251 TSTPSSLPSG AIGPGAEAVP LSFASTMTPF LLATNAPYYA QDPTLGPN
APPENDIX B

Sequences of Soybean Peroxidase

General information

The wild-type soybean peroxidase protein consists of 304 amino acids as listed in Figure B.

Figure B. Amino acid sequence of full-length of WT Soybean Peroxidase.

1   QLTPTFYRET CPNLFPIVFG VIFDASFTDP RIGASLMRLH FHDCFVQGCD
51  GSVLLNNTDT IESEQDALPN INSIRGLDVV NDIKTA VENS CPDTVSCADI
101 LAIAAEIASV LGGPGWPVP LGRRDSLTAN RTLANQNLPA PFFNLTLQKA
151 SFAVQGLNTL DLVTLSGGHT FGRARCSTFI NRLYNFSNTG NPDPNLNTTY
201 LEVLRARCPQ NATGDNLTNL DLSTPDQFDN RYYSNLLQLN GLLQSDQELF
251 STPGADTIPI VNSFSSNQNT FFSNFRVSMI KMGNIGVLTG DEGEIRLQCN
301 FVNG
APPENDIX C

Sequences of Cytochrome P450cam

General information

The wild-type cytochrome P450cam protein consists of 414 amino acids as listed in Figure C.

Figure C. Amino acid sequence of full-length of WT Cytochrome P450cam.

1  MTTETIQSNA NLAPlppHVP EHLVFDFDMY NPSNLSAGVQ EAWAVLQESN
51  VPDLVWTRCN GGHWIATRGQ LIREAYEDYR HFSSECPFIP REAGEAYDFI
101  PTSMDPPEQR QFRALANQVV GMPVVDKLEN RIQELACSLI ESLRPQGGQCN
151  FTEDYAEFPF IRIFMLLAGL PEEDPHLKY LTDQMTRPDG SMTFAEAKEA
201  LDYLPIIE QRRQKPGTDA ISIVANGQVN GRPIISDAEK RMCGLLVLGG
251  LDTVVNFLSF SMEFLAKSPE HRQELIERPE RIPAACEELL RRFSLVADGR
301  ILTSDYEFHG VQLKKGQIL LPQMSGLDE RENACPMHVD FSRQKVSHTT
351  FGHGSHLCLG QHLARREIIV TLKEWLTRIP DSFIAPGAQI QHKSGIVSGV
401  QALPLVWDPA TTKAV
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