Enzyme-Mechanism of Isoniazid Activation in Class I and Class III Peroxidases*

Received for publication, March 3, 2004, and in revised form, June 29, 2004
Published, JBC Papers in Press, July 1, 2004, DOI 10.1074/jbc.M402384200

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There is an urgent need to understand the mechanism of activation of the frontline anti-tuberculosis drug isoniazid by the Mycobacterium tuberculosis catalase-peroxidase. To address this, a combination of NMR spectroscopic, biochemical, and computational methods have been used to obtain a model of the frontline anti-tuberculosis drug isoniazid bound to the active site of the class III peroxidase, horseradish peroxidase C. This information has been used in combination with the new crystal structure of the M. tuberculosis catalase-peroxidase to predict the mode of INH binding across the class I heme peroxidase family. An enzyme-catalyzed mechanism for INH activation is proposed that brings together structural, functional, and spectroscopic data from a variety of sources. Collectively, the information not only provides a molecular basis for understanding INH activation by the M. tuberculosis catalase-peroxidase but also establishes a new conceptual framework for testing hypotheses regarding the enzyme-catalyzed turnover of this compound in a number of heme peroxidases.

Isoniazid (isonicotinic acid hydrazide, INH) is a prodrug that forms a key part of the frontline chemotherapy used to treat tuberculosis in both developed and developing countries. More than 8 million people per year are diagnosed with tuberculosis, and nearly 2 million of those people die (1). The continued rise in drug-resistant and multidrug-resistant strains of tuberculosis and the scale of the tuberculosis epidemic has stimulated fundamental research to elucidate the molecular mechanisms of anti-tuberculosis drugs, including INH, with a view that this information can initiate the discovery of new antimicrobial targets and alternative treatment regimes. Although INH has been in clinical use for the treatment of tuberculosis since the 1950s, the mechanism of activation of this prodrug and its mode of action are still not fully established. However, numerous in vitro and in vivo studies have shown that a catalase-peroxidase (CP), endogenous to Mycobacterium tuberculosis (the principal causative agent of the disease), is required for activation of INH (discussed in Ref. 2). This dimeric, heme-containing enzyme of ~160-kDa molecular mass is a member of class I of the superfamily of plant, fungal, and bacterial peroxidases and displays characteristic catalatic and peroxidatic activities (3–5). The latter activity using hydrogen peroxide has been invoked as a means of oxidizing INH (6–8). An additional manganese dependence has also been suggested based on studies using the related Mycobacterium smegmatis CP (9, 10), although this activity may not play an essential role in INH oxidation by the M. tuberculosis CP (8, 11). However, M. tuberculosis CP has been shown to display the ability to oxidize Mn2+ as well as halogenated compounds using a chloroperoxidatic activity (12).

The determination of the crystal structure for a recombinant form of M. tuberculosis CP (mtCP) (2) has been a significant step toward understanding enzyme-catalyzed activation of INH. This structure provides a three-dimensional framework for rationalizing how structural elements within the active site of this enzyme can contribute to drug activation assuming a binding site for INH can be identified. Computational analyses of the mtCP structure identified an energetically favorable binding site for INH near the δ-meso edge of the heme (2). This location is similar to that observed for other small aromatic compounds in the crystal structures of class I, II, and III peroxidases (13–16). An alternative binding site has also been suggested based upon crystallographic studies of the Burkholderia pseudomallei CP, which places INH in a surface loop structure (17) ~10 Å away from the binding site of small aromatic compounds located near the δ-meso edge of the heme. The local environment of INH in this surface loop is considerably different from that near the δ-meso edge of the heme, and activation of the INH in this case is proposed to involve an electron transfer pathway that would be unique to CP enzymes (17). Presently there is no reported structure of an INH-peroxidase complex, and it is clear that identification of an INH binding site is essential for defining key residues and other structural elements that catalyze the activation of the drug. Interestingly, it has been shown that the class III peroxidase, horseradish peroxidase (HRP), can oxidize INH using hydrogen peroxide in a classic peroxidatic mechanism involving the for-
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mation of Compound I to produce reactive species capable of reducing nitro blue tetrazolium (18). Later work from the same group demonstrated similar activity using partially purified *M. tuberculosis* CP (6). Studies have shown that Compound II can be stably generated in the reaction of HRP Compound I with INH, further supporting the operation of a classic peroxidatic mechanism of INH oxidation for this enzyme (19). The same authors also identified Compound I in the reaction of *M. tuberculosis* CP with INH and speculated that a similar peroxidatic pathway may be utilized (19). Based upon these data and the amenability of HRP to characterization of enzyme-enzyme interactions, a number of enzyme-enzyme complex systems have been developed to study the nature of the active site of the enzyme. The complexes are formed by exchanging INH with substrate, inhibitors, or other small molecules in solution. The complexes are then characterized by solution techniques such as relaxation, two-dimensional NMR spectroscopy or EPR. A number of specific enzyme-enzyme complex systems have been developed to study the nature of the active site of the enzyme.

EXPERIMENTAL PROCEDURES

**Materials**—Deuterium oxide, 3-nitrobenzoic acid and hydrogen peroxide (30% w/v) were obtained from Sigma-Aldrich. HRP was also obtained from Sigma Aldrich and was used without any further purification. Protein concentration of HRP was determined spectrophotometrically by measuring absorbance at 403 nm (29). Recombinant soybean APX (rsAPX) and recombinant pea APX (rpAPX) were prepared as described previously (32). Hydroxylamine HCl was obtained from Sigma Aldrich and was used without any further purification. KCN was from Carlo Erba (Milan, Italy), and INH was obtained from Caterina Ghio (ICQEM-CNR, Pisa, Italy) and were obtained from Merck.

**NMR Spectroscopy**—Samples were prepared by exchanging HRPC into 50 mM NaH2PO4 buffer in either H2O (buffer A) or deuterium oxide (buffer B) by allowing the excess solid KCN followed by re-adjusting the pH to 7.0. Protein concentration of INH into both samples was performed by the stepwise addition of excess solid KCN followed by re-adjusting the pH to 7.0. Protein concentration of samples prepared in buffer B ranged from 0.4 to 0.8 mM, whereas the protein concentration of samples prepared in buffer B was 2.5 mM.

One-dimensional 1H NMR and titration experiments were performed using a standard water-suppression sequence at 318 K using either a Bruker DRX 500 or a Bruker AVANCE 800 spectrometer. Spectra for HRPC in buffer A were collected at 500 MHz, whereas spectra for the HRPC-cyano adduct in buffer B were collected at 900 MHz. Titration of INH into both samples was performed by the stepwise addition using 1.0, 0.5, 0.25, 0.1, and 0.01 mM stock solutions of INH (prepared in buffer A or B) until no further change in the spectra were observed (>250 mM).

Two-dimensional NMR experiments including NOESY (35) and clean total correlation spectroscopy (36) spectra were collected using a Bruker AVANCE 800 spectrometer at both 318 and 900 K. T1 values of the INH model was obtained using distances derived from relaxation data (37). All two-dimensional NMR experiments were phase-sensitive. For NOESY spectra, spectra widths of 50,000 and 20,000 Hz were used with mixing times of 20 and 50 ms and recycle delays ranging from 300 to 500 ms. For scalar experiments only the smallest spectral width was used with a recycle delay of 500 ms. Spectra were acquired using 800–1024 free induction decays with 4096 complex data points. Data were processed using a sine-squared bell filter with shifted 45° or 30°. INH spectra were zero-filled to give the final 4096 × 2048-processed maps.

Data were analyzed using the XEASY program (ETH, Zurich, Switzerland) (39) on an IBM RISC 6000. NOESY cross-peaks were integrated with the standard routines of the XEASY program and translated in proton-proton distances using cross-peaks between protons at fixed distances for calibration. For NOESY correlation maps, spectra were measured in the low spin state and, therefore, have much faster electronic relaxation rates compared with unliganded HRP. It follows that nuclear relaxation rates for the cyanide adduct are much less affected by the paramagnetic center.

**Molecular Dynamics Simulations**—Before molecular dynamics simulations, models of HRPC, the HRPC-cyano adduct, HRPC-INH, and HRPC-cyano-INH were constructed using the modeling program Sybyl (Tripos Associates, Inc.) on an Evans and Sutherland model PS390 graphics work station. The starting model for HRPC was the x-ray structure (42) (Protein Data Bank code 1ATJ). Residues reported in the x-ray structure were used in the modeling studies. Cyanide was docked onto the HRPC model by coordinating its carbon atom to the HRPC iron at an orientation essentially perpendicular to the heme plane based upon coordinates of the x-ray structure for the CcP-cyanide complex (43). The orientation of INH in HRPC-INH and HRPC-cyano-INH models was obtained using distances derived from relaxation data and NOEs as described above. All models also contained the two calcium ions present in the HRPC crystal structure.

Molecular dynamics simulations were carried out on the four models listed above with the AMBER 4.1.2 software package (44) using an IBM/S2 parallel computer. Throughout the calculations the SHAKE algorithm was used (45), and geometrical restraints were applied to maintain the four disulfide bridges in the HRPC crystal structure (Cys-11-Cys-91, Cys-44-Cys-49, Cys-97-Cys-301, and Cys-177–Cys-209). All force field parameters (except those for the heme, the proximal histidine residue, calcium ions, and cyanide) were taken from the AMBER 4.1 force field data base (46). For NOESY spectra, NOEs were easily measured at various concentrations of enzyme. Because the affinity constant for the adduct or complex is known, from the value of T1obs it is possible to obtain a distance limit between the metal ion and the nucleus according to the Solomon-Bloembergen equation (40, 41). For INH binding to HRPC, T1obs of interest are those of the ring proton resonances of INH that can easily be measured at various concentrations of enzyme. Because the fluid ring is expelled but does flip, INH INH complexes are defined as the ring carbons in the ortho position relative to pyridyl nitrogen, whereas INH C3H protons are in the meta position (Fig. 1A). A reasonable estimate of the diamagnetic contribution T1d can be obtained from the T1obs of the same resonances in the presence of the HRPC-cyano adduct, provided that INH binds to this adduct in a manner similar to that which it binds HRP. This assumption is reasonable because the iron in the HRPC-cyano adduct is in a low spin state and, therefore, has much faster electronic relaxation rates compared with unliganded HRP. This relaxes INH activation rates for the cyano adduct are much less affected by the paramagnetic center.
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RESULTS AND DISCUSSION

Affinity of INH for HRPC—Numerous studies of HRP have shown that this enzyme is able to catalyze the oxidation of a variety of aromatic donor compounds using the classic peroxidatic cycle involving the formation of Compounds I and II (for review, see Dunford et al. (21)). This mechanism was also invoked to describe the oxidation of INH (Fig. 1A) by HRP (19). 1H NMR spectra of HRPC and the HRPC-cyanide adduct in the absence and presence of INH are shown in Figs. 2 and 3, respectively. Titration of HRPC with INH yielded an apparent equilibrium affinity constant \( K_D \) of 84.0 ± 0.5 \( M^{-1} \) corresponding to a calculated apparent \( K_D \) value (1/\( K_D \)) of 11.9 \( M^{-1} \), which is slightly higher compared with reported values obtained using isothermal titration methods, 1.1–3.4 \( M^{-1} \) (57), and may simply reflect subtle differences between the two techniques. Titration of the HRPC-cyanide complex with INH yielded a similar \( K_D \) of 70.9 ± 0.4 \( M^{-1} \), corresponding to a \( K_D \) of 1.44 \( M^{-1} \), suggesting that the hydrazinyl moiety of INH has a similar binding site in the presence and absence of cyanide and that INH is not directly coordinated to the heme.

The INH-HRPC Complex—Previous assignments of signals for the HRPC-cyanide adduct (28) enabled comparative analysis of spectra obtained from two-dimensional-NOESY experiments of this complex in the absence and presence of INH. A selective perturbation of the 1H NMR spectrum for the HRPC-cyanide adduct is observed upon the addition of INH (Fig. 3 and Table I) but does not indicate the presence of any large conformational changes associated with INH binding to the enzyme. Residues most affected include Arg-38, Phe-41, His-42, and Gly-169 with smaller perturbations observed for Phe-152 and Phe-172. Notably, signals of Phe-179 are somewhat obscured by overlapping INH C3H ring resonances. Dipole-dipole couplings between the INH C2H protons and the C18H3 and C17\(^\text{15}\)H protons of the heme (Fig. 4) are clearly observed as well as a weak NOE connectivity between the INH C3H and the His42 Cε1H protons (Fig. 5). A few other dipole-dipole couplings between INH and the HRPC-cyanide adduct can also be observed, but resonances associated with the latter are unassigned. Additionally, titration of 20 \( M \) INH with increasing

mass spectrometer equipped with a 15 m × 0.250-mm column in the positive electronic impact mode.

**Fig. 1. Structures of aromatic and pyridyl compounds.** A, INH, B, BHA, C, salicylhydroxamic acid. Shown are oxidation products of INH isonicotinic acid (D), isonicotinamide (E), pyridine-4-carboxaldehyde (F).
amounts of HRPC (up to a concentration of 0.8 mM) led to a reduction of the $T_1$ obs values for C2H and C3H ring protons of INH, from 8.7 and 7.1 s to 3.4 and 1.2 s, respectively. The low spin HRPC-cyanide adduct was used as a reference value for the diamagnetic species (as the paramagnetic contribution to $T_1$ obs of INH resonances produced by the HRPC-cyanide adduct is small compared with that of the high spin HRPC). This resulted in lower distance limits of separation for contacts between the INH C2H or C3H protons and the heme iron of 7.1 and 5.4 Å, respectively.

The crystallographic coordinates of unliganded HRPC (42), in the presence and absence of a docked cyanide ligand, were used as a starting point for molecular dynamic calculations. These calculations yielded solution equilibrated models of HRPC and the HRPC-cyanide adduct, which showed only minor deviations from the crystal structure (root mean square deviation of 0.9–1.0 Å for backbone atoms and 1.2–1.3 Å for side chains). All secondary structure elements as well as other key structural features, including the four disulfide bridges, two calcium ion binding sites, and the heme binding site, remained intact. In addition, water molecules placed in a 10-Å shell surrounding HRPC were observed to minimize at "conserved" positions in the active site of the x-ray structure, near the heme C20 and iron in the distal pocket. Models of the HRPC-INH and HRPC-cyanide-INH complexes, generated without explicit constraints on the position of INH, yielded positions of INH in both these complexes that were consistent with experimental data obtained from the NOEs.

The proposed modes of binding for INH to the HRPC-cyanide adduct and to HRPC are shown in Fig. 6, A and B. INH occupies the same region of the active site in a similar orientation to that seen in the crystal structure of HRPC with the structurally similar benzhydroxamic acid (BHA; Fig. 1B and Fig. 6C) (16). Comparison of the crystal structure of HRPC-BHA (16) and the NMR-based model of HRPC-INH demonstrates that both BHA and INH can access a similar but not identical set of interactions that may relate to their differing affinities (Fig. 6D). The pyridinyl moiety of INH is located in a hydrophobic “aromatic binding pocket” (25), near the heme-18 methyl and makes a number of non-bonded contacts to residues including Gly-69, Ala-140, Pro-141, and Phe-179 as well as to the heme itself. In the HRPC-INH and HRPC-cyanide-INH models the hydrazide moiety of INH forms hydrogen bonds to a backbone carbonyl oxygen, INH N3-Pro-139 O (2.7–2.8 Å), and to the distal cata-
lytic histidine residue, INH O-His-42 Nε2 (3.0 Å). In the HRPC-INH model (Fig. 6B) the INH carbonyl oxygen also forms a hydrogen bond (2.7 Å) to the conserved water located directly above heme iron in the distal pocket but not to the side chain of Arg-38, as would be predicted from the HRPC-BHA crystal structure (16). This is probably due to the high mobility observed for this group in the molecular dynamic simulations. The lack of an interaction may be an artifact of this calculation but may also suggest that this interaction is somewhat weaker compared with others in this particular complex. Flexibility of this residue has been observed in the active site of CcP (43). In addition, in comparative thermodynamic studies of HRPC the flexibility of Arg-38 has been implicated to play a role in reducing affinities for certain aromatic hydroxamic acid analogues compared with BHA (57). In contrast, the INH carbonyl oxygen in the HRPC-cyanide-INH model (Fig. 6A) does appear to form a hydrogen bond with the Arg-38 Nε2 group (2.9 Å) and may also interact with the nitrile group of the cyanide ligand (3.1 Å); however, the significance of these differences may only be small given the similarities in the affinities for INH of the HRPC and HRPC-cyanide complexes.

Predicted INH Binding Site in Class I Peroxidases—The NMR-generated model of the HRPC-INH complex is shown in Fig. 7A. A predicted binding mode of the drug is shown in the crystal structures of mtCP (Fig. 7B) (2), rsAPX (Fig. 7C) (29), and CcP (Fig. 7D) (30), based upon superpositions between the

| Residue | Assignment | HRPC-cyanide δ ppm | HRPC-cyanide-INH δ ppm | HRPC-cyanide-INHf δ ppm |
|---------|------------|---------------------|------------------------|-------------------------|
| Heme    | C2H₂      | 2.84                | 2.49                   | 1.96                    |
|         | C3H        | 5.27                | 5.39                   | 5.57                    |
|         | C3H        | -1.33               | -1.43                  | -1.58                   |
|         | C3H        | -2.40               | -2.46                  | -2.55                   |
|         | C7H        | 24.22               | 24.51                  | 24.95                   |
|         | C8H        | 19.24               | 19.24                  | 19.24                   |
|         | C8H        | -2.98               | -3.03                  | -3.11                   |
|         | C8H        | -2.00               | -2.04                  | -2.10                   |
|         | C12H₁      | 6.54                | 6.56                   | 6.59                    |
|         | C13H₂      | -2.52               | -2.43                  | -2.29                   |
|         | C17H₃      | 18.64               | 18.70                  | 18.79                   |
|         | C17H₃      | 9.59                | 9.79                   | 10.09                   |
| Arg-38  | HN         | 5.64                | 5.84                   | 6.14                    |
|         | H₂         | 0.56                | 1.17                   | 2.10                    |
|         | H₂         | -4.96               | -4.41                  | -3.57                   |
|         | H₂         | -0.62               | -0.26                  | 0.29                    |
|         | H₂         | 0.65                | 1.18                   | 1.99                    |
|         | H₂         | 1.41                | Not located            |                        |
|         | H₂         | -6.55               | -5.90                  | -4.91                   |
| Phe-152 | H₆ or H₆₁,₂ | 5.44               | 5.40                   | 5.34                    |
|         | H₆₁,₂      | 6.26                | 6.23                   | 6.18                    |
|         | H₆₁,₂ or H₇ | 7.13               | 7.10                   | 7.05                    |
| Ser-167 | H₆         | 5.70                | 5.86                   | 5.60                    |
|         | H₇         | 3.24                | 3.11                   | 2.91                    |
| Gly-169 | HN         | 10.12               | 9.93                   | 9.64                    |
|         | H₂         | 5.73                | 5.40                   | 4.90                    |
|         | H₂         | 5.01                | 4.80                   | 4.48                    |
|         | H₂         | 12.86               | 12.54                  | 12.33                   |
| His-170 | HN         | 12.21               | 11.86                  | 11.33                   |
|         | H₂         | 9.51                | 8.97                   | 8.15                    |
|         | H₂         | 21.65               | 21.43                  | 21.10                   |
|         | H₂         | 14.31               | 14.26                  | 14.18                   |
|         | H₂         | 20.7                | 17.9                   |                        |
| Thr-171 | HN         | 9.08                | 8.93                   | 8.70                    |
| Phe-172 | HN         | 9.52                | 9.34                   | 9.07                    |
|         | H₆₁,₂      | 7.62                | 7.50                   | 7.32                    |
|         | H₆₁,₂      | 7.16                | 7.08                   | 6.96                    |
| Ile-244 | δ1CH₃      | -2.99               | -2.42                  | -1.55                   |
|         | H₂         | 0.11                | 0.25                   | 0.46                    |
|         | H₂         | 1.10                | 1.29                   | 1.50                    |
| Phe-45  | H₆₁,₂ or H₇ | 4.81               | 4.80                   | 4.78                    |
|         | H₆₁,₂      | 6.37                | 6.35                   | 6.32                    |
|         | H₆ or H₆₁,₂ | 6.99               | 6.97                   | 6.94                    |
coordinates of these class I enzymes and the HRPC-INH complex. Location of this common binding site for the drug, oriented near the δ-meso edge of the heme, is consistent with computational predictions that identify this region as the most favorable binding site in both mtCP and HRPC (2). In rsAPX and CcP, the predicted binding mode of INH in these class I enzymes overlaps with both the GRID interaction maps, which show this site to be the most favored for INH binding as well as the position of salicylhydroxamic acid (Fig. 1C) in the recent APX-salicylhydroxamic acid crystal structure (13). Notably, based upon the crystal structure of B. pseudomallei CP, an alternative binding site has been proposed (17) that, in the mtCP crystal structure, is located in a surface loop of the enzyme (2). However, as discussed in Bertrand et al. (2), the region near the δ-meso edge is suggested to be the most favored site for INH binding in mtCP and is consistent with the location of INH in the HRPC-INH complexes presented here (Fig. 6, A and B).

Closer inspection of the predicted binding modes of INH in the class I peroxidases shows that several common structural features shared with HRPC may play key roles in INH binding. These include the distal histidine residue, which is common to all these enzymes, the ligating water molecule, and the backbone carbonyl groups of Pro-132 and Pro-145 in the rsAPX (29) and CcP (30) structures, respectively (Fig. 7, C and D). These carbonyl groups may interact with N3 of INH as seen in the HRPC-INH NMR models (Fig. 6, A and B). In addition, the conserved distal tryptophan residue, e.g. Trp-107 in mtCP crystal structure; Fig. 7B) also has the potential to form contacts with N3 of INH. Utilization of the main-chain carbonyl group of Pro-132 and the side chain of Trp-41 for binding a related compound, salicylhydroxamic acid (Fig. 1C), have also been reported in the crystal structure of the complex of salicylhydroxamic acid with rsAPX (13). However, the carboxylic acid moiety of Asp-137 in mtCP, which is conserved among CP homologues, has the potential to form interactions with both the N3 and/or the N1 group of INH (Fig. 7B). In HRPC, a water molecule (w313) occupies the equivalent position (Fig. 7A). In rsAPX (Fig. 7C) and CcP (Fig. 7D) this site is occluded somewhat by residues Ala-70 and Ser-81, respectively. It is also interesting to note that the predicted binding mode of INH in the class I peroxidases, based upon superpositions with the NMR-derived INH coordinates in HRPC, does not suggest that the drug will ligate to the heme. In the case of mtCP, this orientation of INH is also consistent with spectroscopic data, which indicate that the enzyme is in the 5-coordinate state when the drug is bound (58–60).

Enzyme-catalyzed Mechanism for INH Activation—Previous studies of mtCP demonstrated that incubation of INH with the enzyme and H2O2 resulted in small but increasing amounts of the acid (Fig. 1D) and amide (Fig. 1E) derivatives of INH as well as the production of the pyridine-4-carboxaldehyde aldehyde derivative (Fig. 1F), all as a function of time (7, 8). As shown in Table II, HRPC as well as the class I peroxidases rsAPX, rpAPX, and CcP are all capable of turning over INH, producing acid and amide reaction product with varying efficiencies. Given the similarities the active sites of these enzymes share with mtCP (discussed in Ref. 2), these data further support the proposal that a common site located near the δ-meso edge of the heme (Fig. 7) is used for INH binding and activation.

A proposed reaction mechanism for the activation of INH leading to the generation of the activated acyl radical and the production of the amide end product via a diazene intermediate is shown in Fig. 8A (8). Using 15N,15N or 15N,14N isomers of INH, singly labeled at the hydrazide moiety, the level of incorporation of label into the amide end product was determined using HPLC and electrospray mass spectrometry methods. In the case of mtCP, the amide reaction product obtained con-
FIG. 6. Stereodiagrams of compounds bound to the active site of HRPC. A, HRPC-cyanide-INH. B, HRPC-INH. C, HRPC-BHA. D, overlay of HRPC-INH with HRPC-BHA. In all panels the heme (in gray) and protein main chains and side chains are depicted in yellow (for the NMR models of INH complexes) or dark green (for the x-ray coordinates of the HRPC-BHA complex) (16). Water in panels B–D refers to the conserved water molecule located on the distal side of the active site above the heme (listed as HOH 999 in the RCSB Protein Data Bank, code 2ATJ). Carbon atoms of INH are shown in light green (panels A, B, and D), whereas carbon atoms of BHA are shown in dark green (panels C and D). Interactions of INH and/or BHA with cyanide or with possible hydrogen bonding groups in the active site of HRPC are shown by dotted lines (panels A–C). This figure was produced using BOBSCRIPT (63).
tained 50% of the $^{15}$N label, which was interpreted as arising from a splitting of the C-N bond of INH (Fig. 1A) (8) rather than N-N splitting of the hydrazide as previously proposed (7). Identical levels of $^{15}$N label incorporation were obtained here using HRPC, suggesting that HRPC and $mtCP$ share a similar mechanism of activation of INH, including the pathway, which results in formation of the amide reaction product. Based upon these findings and given the similarities in the active sites of HRPC, $mtCP$, rsAPX, and CcP (Fig. 7), an enzyme-catalyzed mechanism for INH activation can be proposed.

Fig. 8B indicates how interactions between structural elements of the active site could stabilize intermediates on the activation pathway of INH in $mtCP$, resulting in the production of the acyl radical. In the first step shown, Compound I, generated after reaction with peroxide, is reduced by INH in a single electron transfer to the heme. In concert, a proton is lost from the hydrazide moiety and could be accepted by His-108. In the second step of the reaction, the C-N bond of the hydrazide is split, yielding diazene. The acyl radical may diffuse from the active site or may remain oriented near the $H_{9254}$-$H_{9254}$ edge of the heme. The diazene reaction intermediate may be stabilized in the active site of the enzyme by interactions with Trp-107.

Table II

| Reaction          | INH | Acid | Amide | Aldehyde |
|-------------------|-----|------|-------|----------|
|                   | 30 min | 60 min | 180 min | 30 min | 60 min | 180 min | 30 min | 60 min | 180 min |
| INH alone         | 200 | 200 | 200 | 0 | 0 | 0 | 0 | 0 | 0 |
| $mtCP$/INH/$t$-BuOOH | 38 | 16 | 1 | 130 | 150 | 159 | 10 | 12 | 9.6 |
| $CcP$/INH/$t$-BuOOH | 135 | 130 | 134 | 42 | 42 | 46 | 5.1 | 3.9 | 5.2 |
| rsAPX/INH/$t$-BuOOH | 118 | 135 | 130 | 35 | 44 | 49 | 2.2 | 3.3 | 3.5 |
| HPRC/INH/$t$-BuOOH | 68 | 44 | 12 | 88 | 122 | 152 | 3.8 | 4.8 | 4.8 |

Reactions were carried out in 50 mM Na$_2$HPO$_4$ buffer, pH 7.5, at 37 °C. In each case, 2 μM enzyme, 200 μM INH, and 10 mM $t$-BuOOH were used.

FIG. 7. HRPC-INH complex generated from NMR and INH docked into the active sites of class I peroxidases. A, HRPC-INH. B, $mtCP$/INH. C, rsAPX/INH. D, CcP-INH. In all panels the heme is in gray. Protein main chains and side chains are depicted in dark green for the HRPC NMR model of INH complex, pink for the $mtCP$ crystal structure (2), orange for the APX crystal structure (29), and light green for the CcP crystal structure (30). The conserved water molecule on the distal side of the active site above the heme is shown in red (w1354 for HRPC, w501 for APX, and w595 for CcP). A water molecule in HRPC (w313), which occupies a similar position to one of the oxygen atoms of Asp-137 in $mtCP$, is shown in panel A. Carbon atoms of INH are shown in green. This figure was produced using PyMOL (Ref. 64; www.pymol.org).
Asp-137, and even the oxyferryl group in the same manner as suggested for hydrogen peroxide in the catalytic reaction (61). Transformation of diazene to hydrazine and ammonia (Fig. 8A) may involve deprotonation of the His-108 and Asp-137 side chains in mtCP, although Arg-104 also has the possibility of providing a proton given its proximity, its proposed conformational mobility (43, 62), and its role as an “entropic gate keeper” (57). As discussed previously in the analysis of the INH binding site in other peroxidases, which are capable of activating the drug, the Asp-137 residue is not a conserved feature (2). In HRPC (Fig. 7A) a water molecule, Wat313, occupies a similar position to the carboxylic acid moiety of Asp-137 (Fig. 7B). The ability of Wat313 to occupy this position is due at least in part to the presence of Gly-69, which creates a space large enough to be filled by a well ordered solvent molecule. In APX (Fig. 7C) and CcP (Fig. 7D), Asp-137 is replaced by Ala-70 or Ser-81, respectively, which partially occludes this pocket, preventing a water molecule from occupying a similar position to Wat313 of HRPC. The presence of Asp-137, which is conserved in CPs, may help to explain why INH is turned over most rapidly by mtCP when compared with the other peroxidases under study here. In HRPC, which turns over INH at a reduced rate, the position of the carboxylic acid group of Asp-137 in mtCP is occupied by Wat-313, which may not protonate diazene as efficiently. APX and CcP, which turn over INH most slowly, do not have a water molecule occupying that position and may, thus, rely upon other solvent molecules in the active site or even residues such as the distal arginine (equivalent to Arg-104 in Fig. 8B) to provide protons for the transformation of diazene to ammonia. Ammonia may also be oriented by similar interactions shown for the diazene (Fig. 8B) before the proposed attack by the acyl radical to generate the amide reaction product (Fig. 8A).

To conclude, we have utilized a combination of spectroscopic, biochemical, and computational techniques to obtain a model of INH bound to the active site of HRPC. Furthermore, we have utilized information about the HRPC-INH complex in combination with crystal structures of mtCP (2) and the class I peroxidases APX (13, 29) and CcP (30) to predict the binding mode of INH in these enzymes, which have all been shown to turnover the drug. Using available mechanistic data about the activation pathway involving the formation of the isonicotinamide end product (8), we propose an enzyme-catalyzed mechanism for INH activation that can yield the acyl radical thought to be the activated form of the drug. We, therefore, propose that the models of the enzyme-INH complexes presented here provide a structural framework for further studies designed to test hypotheses regarding in vitro and in vivo activation of INH.

Acknowledgments—We thank Tony Cass, Judit Nagy, Jasmin, and Jeremy Baum for useful discussions in the preparation of this manuscript and Liz Carpenter and Thomas Bertrand for help in the preparation of figures. In addition, we also thank Paolo Carloni for useful discussions regarding the molecular dynamics calculations, Caterina Ghio for allowing us to use the Gaussian 94 program for point charge calculations of isoniazid, Andy Smith for providing coordinates of the x-ray structure of HRPC, and Tom Poulos for providing x-ray coordinates of the CcP-cyanide complex.

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J. Biol. Chem. 2004, 279:39000-39009.
doi: 10.1074/jbc.M402384200 originally published online July 1, 2004

Access the most updated version of this article at doi: 10.1074/jbc.M402384200

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