Specific Function of the Met-Tyr-Trp Adduct Radical and Residues Arg-418 and Asp-137 in the Atypical Catalase Reaction of Catalase-Peroxidase KatG*

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**Background:** Catalase activity in catalase-peroxidases occurs through a poorly defined mechanism requiring residues Arg-418, Asp-137, and the amino acid adduct (M₂₅₅Y₂₂₉W₁₀₇). Results: Along with an adduct radical, Arg-418 stimulates O₂ release from dioxyheme, the rapid formation of which requires Asp-137. Conclusion: The electronics of the Arg-418:MYW-radical interaction enables discharge of O₂ during H₂O₂ turnover. Significance: An atypical catalase mechanism operates in KatG.

Catalase activity of the dual-function heme enzyme catalase-peroxidase (KatG) depends on several structural elements, including a unique adduct formed from covalently linked side chains of three conserved amino acids (Met-255, Tyr-229, and Trp-107, Mycobacterium tuberculosis KatG numbering) (MYW). Mutagenesis, electron paramagnetic resonance, and optical stopped-flow experiments, along with calculations using density functional theory (DFT) methods revealed the basis of the requirement for a radical on the MYW-adduct, for oxyferrous heme, and for conserved residues Arg-418 and Asp-137 in the rapid catalase reaction. The participation of an oxyferrous heme intermediate (dioxyheme) throughout the pH range of catalase activity is suggested from our finding that carbon monoxide inhibits the activity at both acidic and alkaline pH. In the presence of H₂O₂, the MYW-adduct radical is formed normally in KatG[D137S] but this mutant is defective in forming dioxyheme and lacks catalase activity. KatG[R418L] is also catalase deficient but exhibits normal formation of the adduct radical and dioxyheme. Both mutants exhibit a coincidence between MYW-adduct radical persistence and H₂O₂ consumption as a function of time, and enhanced subunit oligomerization during turnover, suggesting that the two mutations disrupting catalase turnover allow increased migration of the MYW-adduct radical to protein surface residues. DFT calculations showed that an interaction between the side chain of residue Arg-418 and Tyr-229 in the MYW-adduct radical favors reaction of the radical with the adjacent dioxyheme intermediate present throughout turnover in WT KatG. Release of molecular oxygen and regeneration of resting enzyme are thereby catalyzed in the last step of a proposed catalase reaction.

The catalase-peroxidase enzyme, KatG, is found in many microorganisms but in the human pathogen *Mycobacterium tuberculosis* (*M. tb*) it is the only “catalase” (2 H₂O₂ → O₂ + 2 H₂O) and plays an important role in protecting this organism against oxidative stress (1–2). The structural origins of its high catalase activity are therefore of special interest since KatG may fulfill a role different from the ubiquitous monofunctional catalases, especially in organisms that contain genes for both the monofunctional and dual function enzymes (3). KatG in *M. tb* is also responsible for peroxidative activation of the pro-drug isoniazid (isonicotinic acid hydrazide, INH) used to treat tuberculosis infection, though a genuine peroxidase function for the enzyme in normal bacterial physiology is not known. Mutations in *M. tb* KatG that confer resistance to INH create ongoing challenges worldwide in treating TB.

The catalase activity of KatG depends upon three conserved amino acids (Met-255, Tyr-229, and Trp-107, *M. tb* numbering, MYW) with side chains post-translationally linked into a unique adduct on the distal side of the heme pocket (Fig. 1) (4–7). Mutation of any of these residues reduces catalase activity to less than one percent of the rate in wild-type (WT) KatG, but without compromising peroxidase activity (8–12). These findings demonstrate that the MYW-adduct enables formation of a unique intermediate required for efficient turnover of H₂O₂. Using rapid freeze-quench (RFQ) EPR, we reported the characterization of a narrow doublet radical signal (17-Gauss linewidth, aH1,H2 = 11 and 2.5 Gauss for β-methylene hydrogens) from a modified tyrosine in *M. tb* KatG (13–14). More recent isotope labeling and mutagenesis experiments (15), allowed assignment of the radical found during catalase turnover to the MYW-adduct and elucidated a function for this unique cofactor first suggested several years ago (16). Explaining the operation of this radical in the catalase reaction particular to KatG is among our present goals.

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3 The abbreviations used are: KatG, catalase-peroxidase; *M. tb*, *M. tuberculosis*; *Bp*, *B. pseudomallei*; KatG[R418L], KatG[R418L] mutant of KatG; KatG[D137S], D137S mutant of KatG; BLC, bovine liver catalase; EPR, electron paramagnetic resonance; RFQ, rapid freeze-quench; MYW, Met Tyr Trp.
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Residues beyond the MYW-adduct but near the heme are also required for high catalase activity in KatG. For example, replacement of Arg-418 with aliphatic amino acids reduces catalase turnover to less than 2% that of the WT rate (17, 18). This residue, despite its location remote from the heme pocket, was proposed to induce electronic steering effects on the heme through its guanidinium group, which interacts with the phenolate oxygen of Tyr-229 (Fig. 1) at alkaline pH. In the low pH range, the Arg side chain is found in a non-interacting conformation (17). The pH dependence of the conformational change was shown crystallographically for Burkholderia pseudomallei KatG, and alternate conformations of the Arg-418 side chain are also seen in the two subunits of the M. tb KatG structure. Another residue, Asp-137, located in a substrate access channel is also essential for KatG catalase activity for reasons that are not completely clear (19).

Investigating the role of Arg-418 (and briefly that of Asp-137) in the context of a potentially novel catalase reaction mechanism seemed warranted and was pursued for this work. According to current evidence, this mechanism in KatG has certain features in common with, and others quite distinct from, those known for the monofunctional catalases. In all cases, the reaction is assumed to proceed through a species at the oxidation level of peroxidase Compound I (ferryl heme:porphyrin) known for the monofunctional catalases. In all cases, the reaction radical assuming Compound I formation (the latter not seen experimentally); 2) conversion of ferryl heme to dioxygen by the second peroxide, in enzyme bearing the adduct radical; 3) recombination of unpaired electrons restoring the resting enzyme and releasing O2.

This mechanism or similar ones have been proposed before (16) but detailed verification of the steps is of ongoing interest. Here, we focus on three issues: 1) formation and discharge of dioxygen in KatG; 2) the role of an MYW-adduct radical specific to the KatG reaction; and 3) the role of key remote residues in the mechanism. Optical stopped-flow spectrophotometric and rapid freeze-quench EPR experiments were used to compare reactions in wild-type (WT) M. tb KatG and in catalase-deficient mutants. Supporting the experimental work is a theoretical analysis using Density Functional Theory (DFT) to model the interaction of Arg-418 with the reactant in Rxn. 3, a dioxygen:MYW-adduct radical structure. The latter became of obvious importance based on results collected for the catalase-deficient mutant KatG[R418L]. A new perspective emerges on how the rapid KatG catalase reaction depends directly on the interactions between the MYW-adduct radical, dioxygen, and the conserved residue Arg-418 and on the role of Asp-137 in the reaction pathway.

EXPERIMENTAL PROCEDURES

Mutagenesis, Overexpression, and Purification of M. tuberculosis KatG Enzymes—The WT M. tb katG gene in the pKAT II plasmid was a gift from Dr. Stewart Cole. KatG[R418L] and KatG[D137S] mutants were prepared using the Quickchange (Stratagene) kit for site-directed mutagenesis. Mutagenesis primers used to introduce the desired replacements (mutated codons are in bold) were as follows: (R418L) 5’-CAAGCTTGA-TCCACCTAGACATGGGTCCCG-3’ and 5’-GGGACCCAT-TGCTAGGTGGATCAGCTTG-3’; (D137S) 5’-CAAGCTTGCCCCTCCAAAGCCCAAGTGGTG-3’ and 5’-CAAGCTTGCGGTTGGGCGG-3’. Successful mutations were confirmed by sequencing of mutated plasmids (GENEWIZ, South Plainfield, NJ). Overexpression of recombinant WT KatG and its mutant enzymes, and their purification by FPLC were achieved as described in previous reports (23); the heme precursor δ-aminolevulinic acid was added to culture media to

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\begin{align*}
\text{[Fe}^\text{III}]\text{Por(MYW)} & \rightarrow \text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)} \\
\text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)} & \rightarrow \text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)}^* \\
\text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)}^* & \rightarrow \text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)}^* \\
\text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)}^* & \rightarrow \text{[Fe}^\text{IV}=\text{O}]=\text{Por}^*\text{(MYW)} + \text{O}_2
\end{align*}
\]

\text{SCHEME 1.}

iron, formally a ferrous heme-dioxygen/ferric heme-superoxy complex analogous to peroxidase Compound III.) In most peroxidases, Compound III is a catalytically inert species slowly formed in the presence of excess H2O2 and it does not directly dissociate oxygen (22).
promote formation of holoenzyme. KatG concentration is expressed as the heme concentration based on the pyridine hemochromogen assay (ε<sub>557 nm</sub> = 34.5 mM<sup>-1</sup>cm<sup>-1</sup>) (24). Catalase and peroxidase activities were determined spectrophotometrically (NT-14 UV/Vis, AVIV Instruments) in replicate analyses using initial rates of reactions, in which the consumption of hydrogen peroxide was monitored at 240 nm for catalase activity (ε<sub>430 nm</sub> = 43.6 M<sup>-1</sup>cm<sup>-1</sup>), or by color production using t-butyl hydroperoxide and o-dianisidine (peroxidase activity) according to previous reports (23).

Kinetic Measurements—RFQ-EPR samples were prepared using an Update Instrument, Inc. model 1000 chemical-freeze quench apparatus, as described previously (25). Briefly, the enzyme (50 μM final) and H<sub>2</sub>O<sub>2</sub> (50 mM final) were mixed at room temperature in a 1:1 volume ratio, and the mixture was allowed to react for the indicated time periods determined by the length of the reactor tubes or, for longer periods by programming a delay interval; it was then sprayed into a funnel attached to an EPR tube submerged in an isopentane bath held at approximately −130 °C. The frozen powder was packed into precision bore EPR tubes, and samples were transferred to liquid nitrogen for storage.

EPR spectra were recorded at 77 K on a Bruker E500 ElexSys EPR spectrometer operating at X-band (9 GHz). Data acquisition and manipulation were performed using XepView and WinEPR software (Bruker). Instrumental parameters were as follows: modulation amplitude, 1 G; microwave power, 0.1 milliwatt; modulation frequency, 100 kHz; microwave frequency, 9.445 (± 0.005) GHz; number of scans averaged, 9. For quantitative EPR analysis, signal intensities were evaluated by double integration of first-derivative EPR signals using TEMPOL and H<sub>2</sub>O<sub>2</sub> as standards. The TEMPOL radical concentration was evaluated optically (ε<sub>128 nm</sub> = 13.2 M<sup>-1</sup>cm<sup>-1</sup>) (26). The spins per heme values are corrected for the effect of dilution of frozen samples by trapped isopentane (packing dilution factor ~0.4).

Errors can be significant for later time points, however, because O<sub>2</sub> gas dilutes the mixed samples in variable amounts during incubations.

The companion experiments for evaluating the kinetics of H<sub>2</sub>O<sub>2</sub> consumption were performed on a stopped-flow spectrophotometer (HiTech Scientific, UK) using the same enzyme and H<sub>2</sub>O<sub>2</sub> concentrations as for EPR samples. The absorbance of H<sub>2</sub>O<sub>2</sub> at 240 nm was monitored for the desired time period. Due to strong absorbance caused by the high concentrations of enzyme and peroxide, the instrument was adapted with a 1.5 mm light path flow-cell.

CO Inhibition of Catalase Activity of KatG—To check the effect of carbon monoxide on catalase activity, KatG and H<sub>2</sub>O<sub>2</sub> solutions were purged with carbon monoxide for 15 min before being loaded into the syringes of the stopped-flow apparatus. The sample reservoirs containing the CO-saturated solutions were sealed immediately after loading, to avoid gas exchange. The consumption of H<sub>2</sub>O<sub>2</sub> was monitored at 240 nm as a function of time after mixing the enzyme with H<sub>2</sub>O<sub>2</sub> in the airtight stopped-flow system. For time courses at pH 6.0 and 8.5, 20 mM potassium phosphate buffer and 20 mM sodium borate buffer were used respectively.

Oligomerization of KatG during Catalytic Reactions—KatG enzymes (15 μM) were incubated at pH 7.2 for 30 min (37 °C) in the presence of a slow flux of H<sub>2</sub>O<sub>2</sub> generated by aerobic glucose (10 mM) and glucose oxidase. The rate of H<sub>2</sub>O<sub>2</sub> generation was varied from 0–100 μM/min by changing the amount of glucose oxidase. H<sub>2</sub>O<sub>2</sub> flux was separately determined using a horseradish peroxidase-based assay (27). Immediately after incubations, the H<sub>2</sub>O<sub>2</sub>-treated KatG (or mutants) was mixed with an equal volume of 2X SDS-PAGE sample loading buffer and incubated at 100 °C for 5 min. Samples were then loaded onto a precast gradient polyacrylamide gel (PhastGel, 4–15%, GE Healthcare) and electrophoresed using a Pharmacia PhastSystem followed by Coomassie Blue staining (23).

DFT Calculations—Additional support for features of the catalase mechanism was achieved through DFT-based calculations by examining active site model structures and the relative energies of various electronic states. All computations were performed for gas phase models at the level of the well-documented B3LYP hybrid density functional in combination with moderate 6–31G basis sets as implemented in the Gaussian 09 program package (28). A starting model was built based upon the crystal structure of the M. tb KatG enzyme (2cca.pdb) using coordinates for all atoms of the MYW–adduct, including the alpha carbons (formulated as methyl groups) as well as the Arg-418 residue with its side chain in vicinal and remote conformations (as in Fig. 1). Incorporation of dioxyheme (with propinones replaced with methyl groups for simplification) and the proximal histidine completes the structure of the species we focused on for evidence of catalytic properties. The Arg-remote model was optimized with constraints applied to freeze the positions of the α-carbon atoms of each residue and one of the meso carbons (between rings A and B) of the heme. Using the optimized geometry of this starting model, a radical form was produced by removing an H atom from the phenolic oxygen of Tyr-229 of the MYW–adduct as in our prior report (13). Then, relative energies were computed for states of the model having total unpaired spin values from S = 1/2 to S = 7/2 in unit increments (doublet to octet multiplicities). Among the Arg-vicinal models, the sextet (S(total) = 5/2) state was of lowest energy after geometry re-optimization, which led to release of triplet state molecular oxygen. Relative energies for other spin multiplicity states were also computed for the oxygen-dissociated model. The lowest energy states are addressed in detail in “Results.” No negative vibrational frequencies were found in any calculation, showing that all optimized geometries were at true energy minima. Spin contamination values were within acceptable limits, showing that the computed energies and spin distributions are reliable. Spin distributions for each electronic state were evaluated using Natural Orbital Analysis as implemented in NBO 5.0 (29). Additional details are given in the text.

RESULTS

Purified mutant enzymes KatG[R418L] and KatG[D137S] exhibited optical spectra similar to that of WT KatG, with purity ratios, A<sub>407</sub>/A<sub>280</sub> ~0.45–0.5 (not shown), consistent with a normal stoichiometry of one heme per monomer. The spectroscopic features of KatG[R418L] are consistent with a mixture of 5 and 6-coordinate ferric heme in the resting
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enzyme much like that in WT KatG. The KatG[D137S] mutant contains more 5-coordinate heme than WT KatG, with a Soret peak at 405 nm, similar to the analogous mutant D152S of Synchocystis KatG (19). Small differences in the properties of the resting enzymes are not expected to reduce catalase activity (30). For KatG[R418L], the steady-state peroxidase activity assayed with the artificial substrate o-dianisidine was close to wild type (WT) activity (0.7 versus 0.9 units/mg) but its catalase activity (57 units/mg) was about 2% that of WT enzyme. The peroxidase activity of KatG[D137S] was one-third that of WT (0.3 versus 0.9 units/mg), and its catalase activity (24 units/mg) was much lower (0.8% that of WT enzyme).

1) MYW-adduct Radical in WT KatG, KatG[R418L] and KatG[D137S] Mutants—In an earlier study, we presented evidence that a protein-based radical and dioxyheme were simultaneously present in KatG during catalase turnover at neutral pH (13). The radical, which exhibits a narrow doublet EPR signal, was detected using large molar excesses (8000-fold) of peroxide to accumulate it in samples frozen after manual mixing, as well as in samples frozen within milliseconds of mechanical mixing, for which a smaller molar excess of H₂O₂ (1000-fold) was sufficient.

Here we first characterize the behavior of the MYW-adduct radical in WT KatG under various conditions, for comparisons with the catalase-deficient mutants KatG[R418L] and KatG[D137S]. The EPR experiments were performed near the pH optimum (7.2) for WT KatG catalase activity. Fig. 2A shows the narrow doublet EPR signal characteristic of the MYW-adduct radical (13, 15), frozen 8 ms after mixing, corresponding to a concentration of 0.3 spin/heme. Spectra are also shown for samples frozen at two or three later time points as indicated, for WT and mutants respectively. A rise in radical concentration was not seen in the accessible time regimes due to the high reaction rates in each case. The radical signal is detected during a time interval corresponding to the trace for H₂O₂ consumption monitored optically at 240 nm (Fig. 2A, right panel). Increasing peroxide concentration to a 100,000-fold molar excess produced only a small increase in radical yield (up to 0.4 spin/heme), while limiting H₂O₂ concentration to only a 5-fold molar excess still elicited a detectable yield of radical (∼0.05 spins/heme) at 20 ms (not shown). The yields of radical at time points beyond 1 s are not considered to be accurately observed due to the dilution of the samples by oxygen evolution. We note that the decrease in radical signal intensity illustrated in Fig. 2A is not a reflection of its “lifetime” during a catalytic step but rather represents the decreasing steady-state level of the radical intermediate as the peroxide concentration diminishes.

As the loss of catalase activity in certain mutant enzymes might arise from a lack of a required MYW-adduct radical, we examined two catalase-deficient mutant KatGs under the conditions used above. Fig. 2, B and C, show that KatG[R418L] and KatG[D137S] mutants produced the narrow doublet EPR signal in yields similar to WT KatG at the earliest time points. And the persistence of the radical signals coincided with the period required for H₂O₂ consumption (∼30 s for KatG[R418L] and ∼60 s for KatG[D137S]). These findings suggest that the poor catalase activity of these two mutant enzymes arises from some defect in radical utilization. Interestingly, the initial yield of MYW-adduct radical in both mutants was not significantly higher than for WT protein, nor did the radical persist significantly longer than the substrate H₂O₂. These observations suggest that an MYW-adduct radical escape pathway may exist in the mutant enzymes, which limits accumulation despite poor utilization of the radical in a catalase reaction step. We later show evidence for such a pathway from the enhanced formation of surface amino acid radicals capable of forming enzyme cross-links.

2) Dioxyheme Formation—Dioxyheme and the MYW-adduct radical are found concurrently during turnover of excess H₂O₂ by WT KatG (13–14). However, very rapid turnovers prevent evaluation of the kinetics of heme conversion to oxygenated species in the case of wild-type KatG. The KatG[R418L] mutant, which conserves all the catalytically essential heme pocket residues including the intact MYW-ad-
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FIGURE 3. Dioxyheme formation in KatG[R418L] and KatG[D137S]. Spectra were recorded 4 ms after mixing H₂O₂ with KatG[R418L] (A) or KatG[D137S] (B). Enzyme concentrations were 10 μM (final) and the molar equivalents of H₂O₂ were as indicated. All reactions were performed in 20 mM potassium phosphate buffer, pH 7.2 at room temperature using a stopped-flow spectrophotometer. The peaks in the visible wavelength region through which lines are drawn closely match those assigned to typical peroxidase Compound III and indicate the dioxyheme component in both reactions.

FIGURE 4. Carbon monoxide inhibits catalase activity in KatG at low and high pH. H₂O₂ concentration was monitored at 240 nm. A, WT KatG (20 nM) was mixed with 25 mM H₂O₂ in air-saturated (solid lines) or CO-saturated (dashed lines) buffer at pH 6.0 (A) or pH 8.5 (B). C, bovine liver catalase (BLC) activity measured in air (solid line) or CO-saturated (dashed line) phosphate buffer, pH 7.0. The conditions were the same as in A except that BLC concentration was 0.5 nM. D, steady-state peroxidase activity assay of KatG using t-butyl hydroperoxide and o-dianisidine performed aerobically or in CO-saturated acetate buffer.

only traces of dioxyheme are produced with excess H₂O₂ (500 or 5000-fold) (Fig. 3B), with the state of heme in the remainder of the enzyme remaining uncharacterized. We suggest that for the small fraction of mutant enzyme that forms dioxyheme, the WT catalase reaction pathway can operate normally, since the MYW-adduct radical and Arg-418 are both available. The lost catalase activity is then suggested to arise simply from the lack of abundant dioxyheme. These observations indicate the importance of residue Asp-137 in formation of dioxyheme and overall, the results are crucial to formulating key steps in the KatG catalase reaction mechanism.

3) CO Inhibition and the pH Dependence of the Catalase Reaction—The origin of the pH dependence of the catalase reaction in KatG is not understood in terms of the participation of specific proton transfer events. One unresolved issue is the absence of a dioxyheme optical signature during catalase turnover in alkaline conditions (pH 8.5) (13, 16), raising the question: is there a change in mechanism with pH or merely a change in the identity of the prevailing steady-state intermediate? If dioxyheme is in fact part of a catalytically competent intermediate, it should occur throughout the pH range of the catalase reaction. Furthermore, if this intermediate dissociates oxygen gas, CO could inhibit the catalase reaction by binding tightly to ferrous heme iron, thereby removing enzyme from catalytic competence by trapping it in the stable carbonyl form.

To explore these ideas, the catalase activity of WT KatG was monitored in CO-saturated buffer. The activity at pH 6.0 was inhibited by 95%, but after a lag of several seconds (Fig. 4A). A similar level of inhibition was found at pH 8.5 after a longer lag phase (Fig. 4B). Thus, under both acidic and basic conditions, either free ferrous heme or a dissociable form of dioxyheme must be present during catalase turnover. Formation of free
ferrous heme is difficult to rationalize since there are no reduc-
tants available (except possibly for superoxide anion radical that might be present from side reactions; but this species would tend to combine with ferric heme to give dioxyheme in any case, rather than free ferrous heme), while the formation of dioxyheme is readily rationalized by invoking well-known reac-
tions of heme in peroxidases as given in Scheme 1.

It can be estimated that about 20,000 cycles occur before the maximal inhibition is reached in the CO inhibition process. In contrast to these observations, the activity of the monofunc-
tional enzyme bovine liver catalase (BLC) (Fig. 4C) was not inhibited by CO, confirming that these two enzymes use different catalase mechanisms. The peroxidase activity of KatG assayed with alkyl peroxide is not inhibited by CO (Fig. 4D), which is reasonable as neither ferrous nor dioxyheme is ex-
pected to form under such conditions.

4) Radical Migration and Oligomerization of KatG—Insights into the KatG catalase reaction mechanism and defects in the mutants may also be gained by examining oligomerization of the enzymes. We reasoned that the noted inefficient utilization of the MYW-adduct radical in the mutants might allow for increased electron transfers that ultimately produce surface tyrosyl radicals and then dityrosine cross-links. Such radicals are known to form in WT KatG in the presence of a slow flux of H$_2$O$_2$ (32) under conditions that we propose sustain inefficient catalase turnover; therefore, the oligomerization behavior of the catalase-deficient mutants was explored.

Here, while we do not directly detect radical formation in the presence of enzymatically generated H$_2$O$_2$ used as the sub-
strate, we do show enhanced oligomer formation in the cata-
las-deficient mutants under those conditions. For example, high and moderate yields of oligomers were found for KatG[R418L] and KatG[D137S] respectively, while only traces appeared for WT KatG under the slow-flux conditions (Fig. 5). Oligomerization is an inefficient and/or slow process, as the direct addition of a bolus of H$_2$O$_2$ (1000-fold molar excess) did not produce significant yields (not shown). The enhanced cross-linking of KatG[R418L] is evidence for a higher yield of surface radicals upon decay of the catalytically defective MYW-
adduct radical in this mutant. An enhancement in decay pro-
cesses is more directly evident from the EPR results above, showing that the accumulation of MYW-adduct radical in KatG[R418L] does not exceed that in WT KatG. Decay can be enhanced in this mutant because Arg-418 is required for a rapid Rxn. 3 of Scheme 1 (see DFT results). A slower Rxn. 3 favors escape of the adduct radical through pathways different from its direct reaction with dioxyheme.

KatG[D137S] forms less oligomer than KatG[R418L], but far more than WT. This result is also consistent with enhanced migration of the MYW-adduct radical, as this mutant lacks the dioxyheme species required for a rapid adduct-radical quench-
ing reaction (Rxn. 3, Scheme 1).

5) DFT Calculations and the Function of Arg-418—A crucial role for residue Arg-418 in the catalase reaction mechanism is evident from the results presented above. The observation that replacing Arg-418 with Leu had the effect of extending the pers-

tance of the MYW-adduct radical suggested that the residue is involved in restoring resting enzyme via Rxn. 3 (Scheme 1). DFT calculations were applied to evaluate energies for a series of models of the MYW-adduct radical:dioxyheme species (the reactant in Rxn. 3) to examine whether Arg-418 influences its properties. Models were built to extend earlier calculations for the MYW-adduct radical alone (13) by incorporating dioxy-
heme with its proximal imidazole ligand and Arg-418 in two alternate conformations suggested by the M. tb KatG crystal structure.

The new DFT calculations show that Arg-418 can facilitate conversion of the dioxyheme/MYW-adduct radical unit into triplet (molecular) oxygen, ferric heme, and a closed shell (non-
radical) MYW-adduct. The most relevant results are illustrated by the energy level diagram and the models shown in Fig. 6. The electronic configurations of models include two isoenergetic lower energy states for the Arg-remote model (doublet and quartet) formally consisting of low spin d$^0$ heme iron (S = 0) and a bound oxygen ligand (S = 0) along with the MYW-adduct radical (S = 1/2) for a total spin = 1/2; or low-spin ferric heme iron (d$^5$, S = 1/2) with a bound superoxide anion radical (S = 1/2), along with the MYW-adduct radical (S = 1/2) for S(total) = 3/2. Higher total spin multiplicities were higher in energy for the Arg-remote case. Most notable is that oxygen remained bound in all four electronic states for Arg-remote structures but was dissociated in all Arg-vicinal models.

Panel A shows three energy levels for models selected from the calculated series to illustrate the changes that depend on the Arg-418 configuration. The energy values must be viewed cau-
tiously as the present calculations were performed in vacuo. The overall difference of 44.2 kcal/mol represents the com-
puted energy released upon conversion of the dioxyheme/ MYW-adduct radical unit into the resting enzyme species (S = 3/2 iron, non-radical form of MYW-adduct) plus triplet-state
molecular oxygen when Arg-418 is vicinal to Tyr-229. The intermediate energy level corresponds to the Arg-vicinal/dissociated $^{1}$O$_{2}$ bound state model, with Fe$^{III}$ (S = 1/2) and no MYW-adduct radical, useful to estimate the energy gained solely from the Arg-418–Tyr-229 interaction (17.3 kcal/mol). This value compares well with the 22.3 kcal/mol estimated for the Arg-Tyr hydrogen bonding stabilization calculated for the isolated MYW-adduct model (without heme). Dominating the overall energy gain is the 26.9 kcal/mol contribution (lower part, Fig. 6A) closely corresponding to the singlet-triplet gap for free oxygen.

An important feature of the electronic changes caused by the Arg-Tyr interaction was revealed in DFT calculations on smaller models consisting of the MYW-adduct radical without the dioxyheme component, with and without the Arg interaction (not shown). We found that the guanidinium-phenolate interaction causes a shift in unpaired-electron spin density from the Tyr ring into the Trp indole of the MYW-adduct radical. This shift arises because an increase in negative charge can be stabilized on the phenolic oxygen when the positively charged guanidinium group is nearby. Concomitant with this shift is an increase in net positive character of the Trp indole moiety, the nitrogen of which is adjacent to the oxygen ligand in the models incorporating dioxyheme described above. With dioxyheme present, electron transfer from the “superoxide anion radical” iron ligand into the adduct radical could be favored by the charge and spin redistribution. These electronic changes could not be observed in the extended models described above because of oxygen dissociation and reduction of the adduct radical when Arg was vicinal to Tyr-229.

None of the calculated models indicated covalent changes, such as hydrogen or oxygen transfers to or from the MYW-adduct. Thus, the DFT study predicts that a role for Arg-418 in the catalase reaction is to facilitate conversion of the dioxyheme/MYW-adduct radical species into free triplet-state oxygen, ferric heme iron and the closed shell MYW-adduct to complete the cycle in Scheme 1.

DISCUSSION

The new finding, that the MYW-adduct radical is formed in two catalase deficient mutants, demonstrates that residues Asp-137 and Arg-418 are responsible either for formation or discharge of dioxyheme in the proposed catalase reaction mechanism. The very rapid formation of the MYW-adduct radical likely proceeds the same way in all three cases, through a peroxidase Compound I (Scheme 1, Rxn. 1) or possibly in a concerted turnover that bypasses a free Compound I intermediate. Once the initial turnover has occurred, the most likely route to dioxyheme in the proposed scheme (Rxn. 2) is by rapid $H_{2}O_{2}$-reduction of ferryl heme in a reaction common to other peroxidases (33–34) including KatG[Y229F] (10). Residue Asp-137 seems to be crucial for this conversion and may be responsible for an important proton transfer step that enhances the reactivity of ferryl heme (35). This aspartate residue is absent from Class III peroxidases such as HRP, in which Compound III (dioxyheme) is only slowly formed (20 M$^{-1}$s$^{-1}$) in the presence of excess $H_{2}O_{2}$ (34). Thus, facile formation of dioxyheme in KatG distinguishes this enzyme from other heme peroxidases.

A strong argument for the catalytic competence of an intermediate comprised of dioxyheme/MYW-adduct radical is based on the fact that at alkaline pH, the catalase rate of WT KatG is decreased nearly 10-fold compared with pH 7, while the steady-state heme species is different from dioxyheme, and the adduct radical is still present at this pH. Additional suggestions come from the observation of CO inhibition, which indicates that dioxyheme is present over the full pH range of catalase activity despite its lack of accumulation at alkaline pH.

CO binds to Fe(II) heme exclusively. It is not acting like an inhibitor of the substrate $H_{2}O_{2}$, since CO does not bind to the Fe(III) heme or to forms in other oxidation states above Fe(II) heme. The lag period we observed is then a manifestation of
inhibition occurring when CO binds to a small fraction of dioxygenheme enzyme that spontaneously dissociated molecular oxygen to allow replacement by the CO ligand. Such dissociation is apparently rare and many turnover cycles are required to increase the population of molecules that can be siphoned off the catalytic pathway by CO binding.

The observed lag phase is thus consistent with a mechanism-based process and we estimate that if a very small percentage of dioxygenheme (say 1%) were to spontaneously decay by ligand (oxygen) dissociation, several thousand turnover cycles would be needed for inhibition, as was observed. Whether CO is capturing a kinetically competent or some other intermediate species remains to be determined.

The DFT results presented here, combined with results from earlier calculations, provide valuable insights into the behavior of the dioxygenheme/MYW-adduct radical intermediate and Rxn. 3 of the proposed catalase pathway. The calculations show that the side chain of residue Arg-418 promotes discharge of the oxygen ligand and quenching of the adduct radical through electronic effects mediated by the guanidinium salt bridge to the phenolic oxygen of Tyr-229 in the adduct radical. This effect will be favored in WT KatG at pH values near neutral and above that are known to favor the Arg-418/Tyr-229 interaction, and will be absent in the KatG[R418L] mutant.

These phenomena and the pH dependence of the KatG catalase reaction can be integrated to provide insights into the origin of the rate increase occurring from pH 5 to 7 and the decrease from pH 7 to 9 (36). One of the key pH-dependent interactions is very likely the salt bridge between Arg-418 and Tyr-229, which will be favored when the phenolic oxygen of Tyr-229 is deprotonated and the guanidinium group is protonated. Interestingly, dioxygenheme does not accumulate at alkaline pH, while the MYW-adduct radical is present at normal abundance. In the Scheme 1 mechanism, Rxn. 3 should exhibit a higher rate in the higher pH range with the salt bridge in place. In contrast to this, formation of dioxygenheme from ferryl heme is disfavored at high pH, since protonation of the oxo-ligand is required in the typical pathway to the dioxygenheme species (34). Thus, during catalase turnover at alkaline pH, dioxygenheme is not detected because a slow rate of its formation occurs under conditions of efficient quenching: high pH leads to a decreased reaction rate as Rxn. 2 becomes rate-limiting. In the lower pH range, the rate of formation of dioxygenheme will be enhanced but the Arg-418–Tyr-229 interaction is disfavored and quenching of the dioxygenheme/MYW-adduct radical unit becomes slower. Then, dioxygenheme is the observed steady-state species and the turnover rate decreases due to slowing of Rxn. 3.

CONCLUSIONS

Overall, the current results demonstrate that formation of the MYW-adduct radical is not sufficient for efficient catalase turnover by KatG; dioxygenheme and a regulating remote influence (Arg418) on the electronic structure of the MYW-adduct radical must be present for maximal activity. It seems clear that M. tb KatG is structured for very rapid formation of dioxygenheme and robust catalase activity under a wide range of H₂O₂ concentrations, arguably important for the survival of an intracellu-
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