

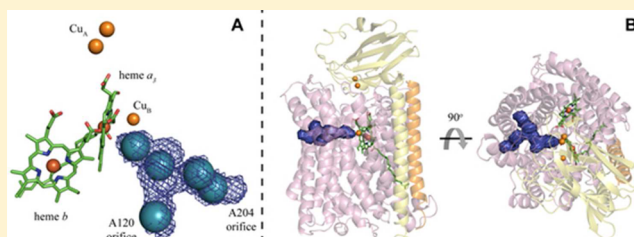
# Mobility of Xe Atoms within the Oxygen Diffusion Channel of Cytochrome *ba*<sub>3</sub> Oxidase

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## Supporting Information

**ABSTRACT:** We use a form of “freeze-trap, kinetic crystallography” to explore the migration of Xe atoms away from the dinuclear heme *a*<sub>3</sub>/Cu<sub>B</sub> center in *Thermus thermophilus* cytochrome *ba*<sub>3</sub> oxidase. This enzyme is a member of the heme–copper oxidase superfamily and is thus crucial for dioxygen-dependent life. The mechanisms involved in the migration of oxygen, water, electrons, and protons into and/or out of the specialized channels of the heme–copper oxidases are generally not well understood. Pressurization of crystals with Xe gas previously revealed a O<sub>2</sub> diffusion channel in cytochrome *ba*<sub>3</sub> oxidase that is continuous, Y-shaped, 18–20 Å in length and comprised of hydrophobic residues, connecting the protein surface within the bilayer to the *a*<sub>3</sub>–Cu<sub>B</sub> center in the active site. To understand movement of gas molecules within the O<sub>2</sub> channel, we performed crystallographic analysis of 19 Xe laden crystals freeze-trapped in liquid nitrogen at selected times between 0 and 480 s while undergoing outgassing at room temperature. Variation in Xe crystallographic occupancy at five discrete sites as a function of time leads to a kinetic model revealing relative degrees of mobility of Xe atoms within the channel. Xe egress occurs primarily through the channel formed by the Xe1 → Xe5 → Xe3 → Xe4 sites, suggesting that ingress of O<sub>2</sub> is likely to occur by the reverse of this process. The channel itself appears not to undergo significant structural changes during Xe migration, thereby indicating a passive role in this important physiological function.



Cytochrome *c* oxidases are terminal enzymes of respiration that comprise the heme–copper oxidase superfamily.<sup>1–3</sup> They are found in the plasma membrane of prokaryotes and the mitochondrial inner membrane of eukaryotes, wherein they catalyze the reduction of molecular oxygen to water with concomitant oxidation of cytochrome *c* and generation of a proton gradient across the membrane. Accordingly, these enzymes contribute to the energy needs of a cell and are crucial for aerobic life.

Cytochrome *ba*<sub>3</sub> oxidase (*ba*<sub>3</sub>) is a B-type cytochrome *c* oxidase from the plasma membrane of *Thermus thermophilus* and is the preferred terminal enzyme of cellular respiration at low oxygen tensions.<sup>4</sup> The enzyme contains the four redox cofactors: heme *a*<sub>3</sub>, heme *b*, binuclear Cu<sub>A</sub>, and Cu<sub>B</sub> as illustrated in Figure 1A.<sup>4–6</sup> Electrons enter the enzyme at the Cu<sub>A</sub> site and tunnel from there to the low-spin heme *b* and move to the heme *a*<sub>3</sub>–Cu<sub>B</sub> dinuclear center (dnc) depending on its chemical state.<sup>3,7,8</sup> Protons enter from the cytoplasm and exit to the periplasm via specially adapted pathways.<sup>9</sup>

Previously, we described a channel in *ba*<sub>3</sub> using X-ray crystallography<sup>10</sup> using static Xe and Kr pressurization.<sup>11</sup> This Y-shaped channel was defined by five major and two minor discrete gaseous atom binding sites that extend from Xe1 near the buried dnc to two possible entry/exit points at the membrane-facing surface of the protein (Figure 1A,B). To the extent that the Xe atom serves as a mimic for the O<sub>2</sub> molecule (see below and ref 10), the previous work demonstrated the

existence of a structurally defined, bifurcated channel within *ba*<sub>3</sub> whose function is to attract O<sub>2</sub> from the lipid bilayer into the dnc. The relative affinity for Xe and Kr at discrete sites within the channel could be determined in terms of relative occupancy under different pressurization conditions,<sup>10</sup> but no information concerning movements of Xe atoms within the channel was obtained from the static structures (Protein Data Bank code 3BVD).

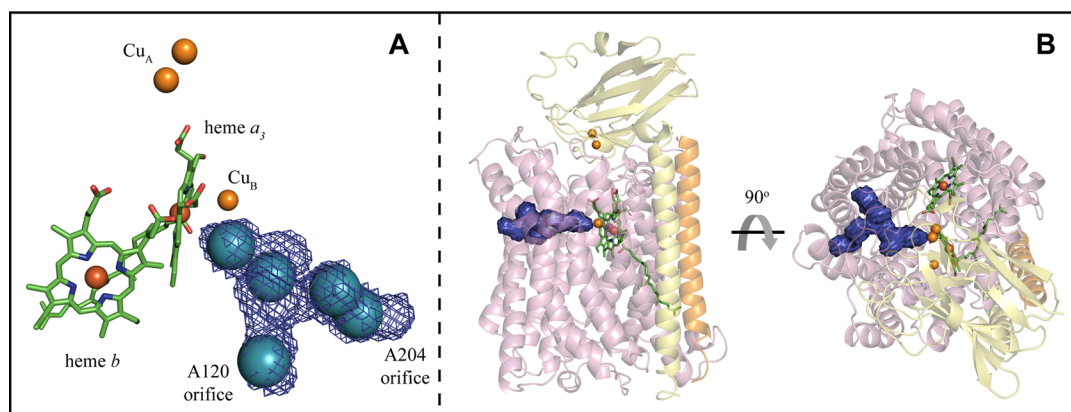
Xe pressurization of crystals has been used to identify a putative O<sub>2</sub> diffusion channel in *Rhodobacter sphaeroides* cytochrome *c* oxidase,<sup>12</sup> similar to that in *ba*<sub>3</sub>, and to define hydrophobic sites in other non-heme-containing proteins.<sup>13</sup> For integral membrane proteins, Xe has been used to probe O<sub>2</sub> channels in photosystem II,<sup>14</sup> the NH<sub>3</sub> channel in the Amt-1 transporter,<sup>15</sup> and N<sub>2</sub>O sites in the NMDA receptor.<sup>16</sup> Xe pressurization has also been used to perturb the conformation and activity in bacteriorhodopsin<sup>17</sup> and to delimit detergent micelles in crystals.<sup>18</sup> In soluble enzymes, Xe has been used to define the pathways for O<sub>2</sub>,<sup>19</sup> CH<sub>4</sub>,<sup>20</sup> CO,<sup>21</sup> and NO<sup>22</sup> into active sites. In heme nitric oxide/oxygen binding (H-NOX) domains, redistribution of Xe atoms within a hydrophobic channel is implied by anisotropic electron density and

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**Figure 1.** (A) Metal centers of *ba*<sub>3</sub> with Xe sites (blue spheres). The Y-shaped hydrophobic channel of *ba*<sub>3</sub> (blue mesh) is overlaid on the Xe sites; the orifice to each fork of the channel is labeled by an adjacent residue. (B) Transmembrane (left) and overhead (right) views of *ba*<sub>3</sub> illustrating the shape and location of the hydrophobic channel (blue surface). Subunit I is lavender, subunit II, which contains the soluble Cu<sub>A</sub> domain, is yellow, and subunit IIa is orange; heme *a*<sub>3</sub> and heme *b* are green.

increasing occupancy closer to the heme.<sup>22</sup> Aside from the latter observation, no crystallographic study of Xe binding sites has extracted direct experimental evidence concerning the relative movements of Xe atoms within hydrophobic channels or cavities.

Ligand migration, however, as inferred from locations of static sites, has been extensively studied in the globins using Xe pressurization crystallography.<sup>13</sup> The interpretation of data from these studies has benefitted from being carried out with crystals of relatively small proteins that diffract to high resolution, enabling detailed description of the binding sites. Petsko and co-workers<sup>23</sup> first reported crystal structures of myoglobin pressurized under several atmospheres of Xe gas, revealing four unique binding sites. Xe has been used, both experimentally and theoretically, to examine proteinaceous CO and O<sub>2</sub> binding sites within sperm whale myoglobin,<sup>24</sup> human hemoglobins,<sup>25</sup> the hemoglobin, HbI, from the blood clam, *Scapharca inequalis*,<sup>26</sup> and human cytoglobin.<sup>27</sup> Theoretical work<sup>28</sup> suggested that the Xe binding sites in Mb may function as transient binding sites for ligands migrating from the solvent to the active site. However, as pointed out by Olson and colleagues,<sup>25</sup> most experimental data, including Laue X-ray crystallography,<sup>29</sup> do not support the theoretical conclusions but suggest, instead, that ligands escape and enter myoglobin by the His(E7) gate and are trapped in the distal pocket; this holds for the human hemoglobins as well (ref 25 and references therein). In the case of bacterial truncated hemoglobins and the neuronal mini-globin from the sea worm, ligands also enter the protein through an apolar tunnel between the C-terminal ends of the E and H helices<sup>27</sup> and are subsequently trapped in the distal pocket. Blocking this apolar tunnel with a Trp mutation or filling the tunnel with Xe markedly decreases both the fraction of ligands that escape to solvent and the bimolecular rate constant for entry of ligands to the active site.<sup>30</sup> As noted, this behavior is to be distinguished from the other globins, in which ligands appear to enter and escape through the distal ligand gate.

Xe lacks several features of another oxygen mimic, carbon monoxide, whose spectroscopic properties have been widely used to examine its photoinduced release from and short ranged recombination with ferrous hemes within a variety of heme-containing proteins. Transient IR spectroscopies have also been used to follow recombination of CO in the heme—

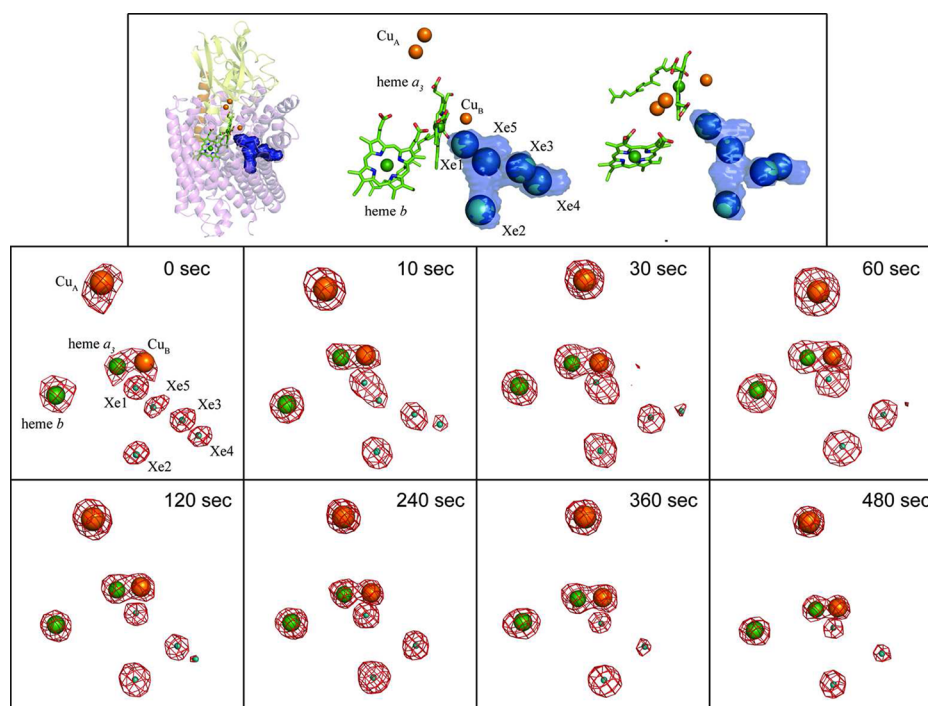
copper oxides.<sup>31</sup> Where CO falls short as a mimic of O<sub>2</sub> is that its 14 X-ray scattering electrons require the availability of highly diffracting protein crystals to identify its position. A useful feature of Xe as a dioxygen mimic, however, in addition to its similar size and hydrophobicity, is that its 54 electrons, combined with significant anomalous X-ray scattering effects, allow facile identification of atomic positions and quantification of occupations even in crystals exhibiting only modest diffraction.

In this paper, we utilize an adaptation of traditional methods to trap intermediates using the apparatus developed by Soltis et al.<sup>11,32</sup> During this operation, pressurized Xe binds to hydrophobic pockets within the protein matrix.<sup>23</sup> For heavy atom derivatization, crystals are frozen immediately after depressurization to capture as much Xe as possible within the protein. In our approach, the crystals are pressurized for several minutes, quickly depressurized to 1 atm, and frozen by plunging into liquid nitrogen, but the time interval at 1 atm prior to freezing is varied from 0 to 480 s, allowing adsorbed Xe to diffuse out of the crystals. By crystallographically quantifying the positions and occupancies of Xe atoms as a function of time, a kinetic model can be derived describing their motions. To our knowledge, this is the first data to quantitate movement of gas molecules within a large integral membrane protein.

## EXPERIMENTAL PROCEDURES

**Protein Purification and Crystallization.** Protein preparation and purification was done following the methods described by Chen et al.<sup>33</sup> The modified crystallization procedure is detailed in the Supporting Information.

**Xe Derivatization and Trapping.** Xe derivatives of *ba*<sub>3</sub> were produced following the procedure previously employed<sup>10</sup> using a single crystal pressurization cell developed at the Stanford Synchrotron Radiation Lightsource (SSRL).<sup>11,32</sup> Crystals were pressurized to 100 psi (689.48 kPa) for 5 min followed by release of the pressure in the chamber. To capture the different stages of Xe diffusion out of the protein under the reduced Xe pressure, the crystals were removed from the chamber at select time intervals (0, 10, 30, 60, 120, 240, 360, and 480 s) and flash frozen into liquid nitrogen. Approximately 20 s is required to remove the crystal from the pressurization vessel and plunge it into liquid nitrogen.



**Figure 2.** Location and occupancies of Xe sites at eight time points during diffusion of Xe out of pressurized crystals. The top panel illustrates the location of the Xe sites lining the hydrophobic channel. The left and center figures in the top panel are in the same orientation as the other panels. The figure on the top right is an  $\sim 40^\circ$  rotation toward the reader. The  $|F^+| - |F^-|$  anomalous difference Fourier map contoured at  $6\sigma$  (red mesh) for the Cu, heme Fe, and Xe atoms of a Xe-pressurized crystal is shown for each time point. The 0 s panel shows the relative location and naming convention of the Xe and metal sites (the binuclear  $\text{Cu}_A$  center was refined as a single atom given the resolution of the data). The average occupancy of Xe at each site at each time point (0–480 s) was determined by occupancy normalization using a total of 19 data sets (Experimental Procedures).

**Data Collection and Structure Determination.** X-ray diffraction experiments were conducted on SSRL beamlines 7-1 and 11-1. All data sets were collected using  $\lambda = 1.127 \text{ \AA}$  to simultaneously maximize the anomalous signal from the Fe, Cu, and Xe ( $f'' = 1.91, 2.74, \text{ and } 4.37$  electrons, respectively, at this wavelength) in the Xe-derivatized  $ba_3$  crystals. Each data set consisted of 360 frames collected with an oscillation angle of  $1^\circ$  with 10 s exposure per frame. Diffraction data were recorded at 100 K on an ADSC Q315 CCD detector, and no significant decay was observed at the resolution used in refinement (Table S1). The data sets were integrated and scaled using MOSFLM and SCALA.<sup>34</sup> Detailed data collection and refinement statistics are presented in Table S1.

Using the wild-type recombinant structure of  $ba_3$  as a search model (PDB code 1XME), molecular replacement was done using MOLREP.<sup>35</sup> All data sets were refined in the same fashion by rigid body and restrained refinement with REFMAC5<sup>36</sup> using overall  $B$  factor refinement and a geometry weighting term of 0.3.  $2|F_o| - |F_c|$  and  $|F_o| - |F_c|$  electron density maps, and anomalous difference Fourier maps, clearly revealed the positions of the Xe atoms as well as Cu and Fe atoms (Figure 2) consistent with previous results.<sup>10</sup> Unbiased coordinates and occupancies of the Fe, Cu, and Xe sites were refined using SHARP<sup>37</sup> against the highly redundant anomalous data sets (Table S1) and expected anomalous scattering factors (Table S2). Because all neighboring Xe–Xe distances are greater than the van der Waals contact distance (4.4 Å) (Table S3 and Figure S2), the Xe sites were atomically resolved at the resolution of the data sets. Structures representing the highest resolution data set at each time point have been deposited with the RCSB (PDB codes 3S33, 3S38, 3S39, 3S3A, 3S3B, 3S3C, and 3S3D, corresponding to time points 10, 30, 60, 120, 240,

360, and 480 s, respectively). The 0 s time point structures are very similar to the previously deposited, 3.37 Å resolution structure (PDB code 3BVD, ref 10). However, because the previously deposited structure is not in the same space group, the newer but lower resolution data sets were used for the 0 s time point in the analysis (Tables S1 and S2).

**Occupancy Normalization.** The accuracy of the anomalous signal in a given data set depends upon diffraction quality, crystal orientation and absorption effects, site-specific radiation damage, and other systematic errors,<sup>38,39</sup> which can lead to apparent differences in the occupancies of anomalous scattering centers. To compare Xe occupancies across multiple data sets, we used biochemical information to define an internal reference for normalization. Cytochrome  $ba_3$  oxidase contains four redox cofactors: heme  $b$ , heme  $a_3$ , a single  $\text{Cu}_B$  site, and the binuclear  $\text{Cu}_A$  center. Analytical data and previous structures of  $ba_3$  showed the consistent full occupancy of these four metal centers, i.e., the occupancies of  $\text{Fe}_b$ ,  $\text{Fe}_{a_3}$ , and  $\text{Cu}_B$  are reasonably taken to be unity. However, as shown in Table S2, individual metal site occupancies vary significantly from 1.0. Utilizing the biochemical constraint, we normalized the occupancies of the metal sites as the mean of  $\text{Fe}_b$ ,  $\text{Fe}_{a_3}$ , and  $\text{Cu}_B$  relative to 1.0. For example, for data set 13 at 240 s (Table S2) the refined occupancies of the  $\text{Fe}_b$ ,  $\text{Fe}_{a_3}$ , and  $\text{Cu}_B$  sites are 0.978, 0.960, and 1.034, respectively, with mean 0.991. Multiplication by  $1/\text{mean}$  normalizes the occupancies of all the metal and Xe sites for a given data set (FeFeCuB columns in Table S2). Normalized values per site are then averaged over the redundant data sets at each time point (AVG, DEV in Table S2).

The  $\text{Cu}_A$  center provides an internal control for this normalization procedure. For example, the normalized



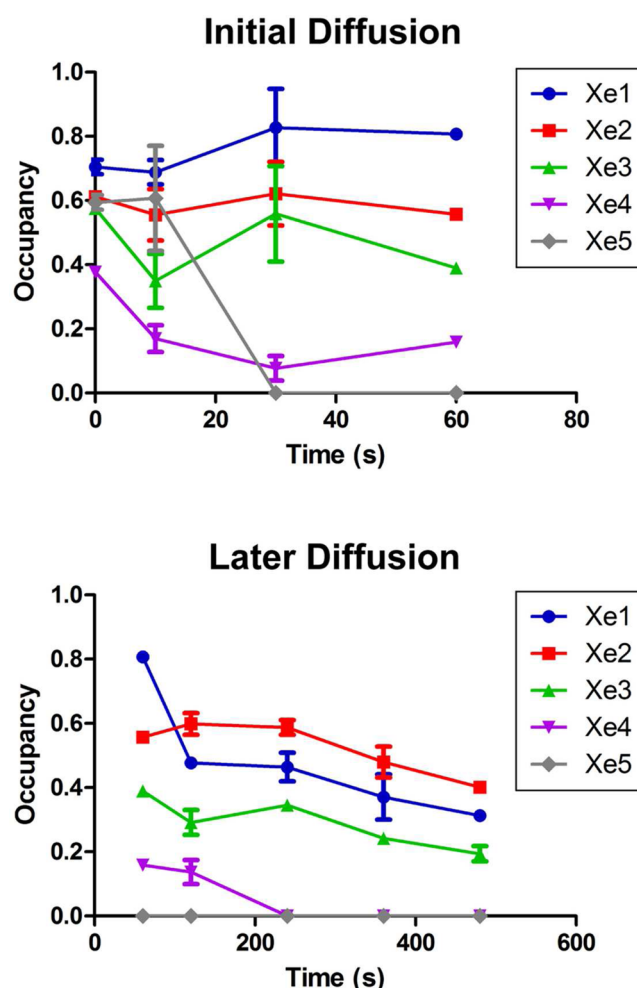
occupancy of Cu<sub>A</sub> for data set 13 is 1.609/0.991 = 1.624 and the average for three data sets at 240 s is 1.704 (Table S2). The normalized average occupancy for Cu<sub>A</sub> for all 19 data sets is  $1.662 \pm 0.048$  or an error of less than 3%. The apparent occupancy of the Cu<sub>A</sub> center, refined as a single site, is a noninteger value because the binuclear center (Cu–Cu distance 2.7 Å) is not resolved at the  $\sim 4$  Å resolution of the data sets (Table S1). Hence, normalization based on the expected occupancy of the Fe<sub>b</sub>, Fe<sub>a3</sub>, and Cu<sub>B</sub> sites in *ba*<sub>3</sub> is validated by results for the Cu<sub>A</sub> center, allowing for internal correction of Xe occupancies within each data set, and comparison of average Xe site occupancies at different time points.

**Modeling Kinetics.** Using the Simbiology module of MATLAB (Natick, MA) and the normalized occupancies of Xe from our structures, we modeled the kinetics of Xe diffusion out of *ba*<sub>3</sub> and obtained rate constants for each of the processes. Details of the model and results of the simulation are provided in the Results and Discussion, and the corresponding coupled differential equations are given in the Supporting Information.

## RESULTS AND DISCUSSION

Xe-pressurized crystals of recombinant, wild-type *ba*<sub>3</sub> were flash frozen after release of pressure at times 0, 10, 30, 60, 120, 240, 360, and 480 s; 19 data sets having between 3.3 and 5.5 Å resolution were collected (Table S1). Following structure solution and refinement (Table S1), we monitored the presence of Xe at the five major binding sites using isomorphous and anomalous difference Fourier maps (Figure 2). The refinement protocol uses the anomalous scattering from three metals sites in the native enzyme, Cu<sub>B</sub>, Fe<sub>b</sub>, and Fe<sub>a3</sub>, to obtain internally normalized occupancies of the Xe atoms for each data set (Table S2). The normalized occupancy for the binuclear Cu<sub>A</sub> center serves as an independent, internal indicator of error (see Experimental Procedures). The average occupancy, along with the calculated error bars, for each Xe site at each diffusion time point is plotted in Figure 3. The estimated error of the quench times is 1–2 s.<sup>40</sup>

**Protein Structure and Xe Locations.** As shown in Table S1, the diffraction resolution of the 19 data sets ranges from 3.3 to 5.0 Å. Within this range, no side-chain movements or backbone displacements relative to native structures of *ba*<sub>3</sub> were observed upon Xe binding. This is also true for comparisons with *ba*<sub>3</sub> structures at different redox states.<sup>41</sup> Using isomorphous and anomalous difference maps, we located between three and five Xe atoms in each data set; early time points revealed the most Xe atoms while later time points showed fewer. The locations for each of the Xe sites in all of the structures are similar to the corresponding sites in the previously reported structures,<sup>10</sup> and the same nomenclature is used to identify the individual sites (Figure 2, top panel). Variations in position within the five discrete binding sites are small (Figure S2); distances between neighboring Xe atoms, which are greater than the van der Waals contact distance of  $\sim 4.4$  Å, are given in Table S3. The amino acid residues surrounding each Xe site are primarily hydrophobic without propensity for a specific amino acid type, e.g., aromatics, to dominate (see Supporting Information of ref 10). The only noticeable change in a Xe position is a shift of Xe4 by  $\sim 2$  Å away from the Xe3 site toward Val201 and Ala204, which occurs only at diffusion time points 30 and 120 s. The Xe4 site is the least occupied of these time points and is closest to the outside of the protein near Ala204.<sup>a</sup>



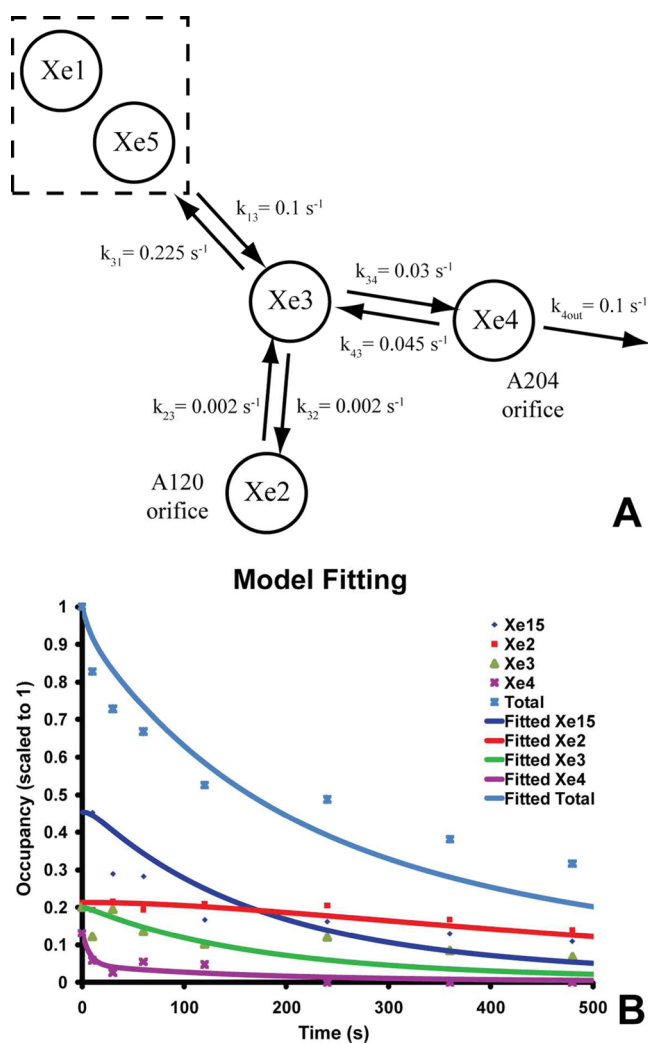
**Figure 3.** Normalized occupancies at each Xe site plotted vs time. Error bars represent standard error,  $s/\sqrt{n}$ , where  $s$  is the standard deviation and  $n$  is the number of observations, i.e., data sets at that time point. The top panel shows 0–60 s data points, and the bottom panel shows 60–480 s data points.

**Temperature at the Crystal after Rapid Freezing in Liquid Nitrogen.** Clearly, passage of the crystal from a pressure of 100 psi (689.48 kPa) Xe at room temperature during the few seconds needed to move the crystal from ambient temperature to 77 K does not result in 100% occupancy of the Xe binding sites, as shown in Table S2. While it is difficult to determine the temperature of each crystal when molecular motion ceases, other studies<sup>40</sup> have shown that the approximate time required to cool a macromolecular crystal from room temperature to 77 K is on the order of 1–2 s. Because the freezing time is  $\sim 1/10$  of the shortest diffusion time, we assume that room temperature equilibrium distributions are trapped upon freezing following the 0–480 s intervals during which outgassing is allowed to occur.

**Xe Egress.** The cumulative occupancy of Xe sites 1, 5, 3, 2, and 4 decreased from  $2.8 \pm 0.38$  Xe atoms to  $0.9 \pm 0.025$  Xe atoms over time (Figure S1), as would be expected as Xe escapes from the crystal. However, the distribution of occupancies at each time point displayed significant variation (Figures 2 and 3), thereby providing an opportunity to probe the mechanism of Xe egress and, by reversal, Xe ( $O_2$ ) access. The Xe1 site, which is nearest to the dinuclear center, is likely the last transient stop prior for  $O_2$  prior to reduction.<sup>42</sup> From 0

to 10 s, all the Xe sites decrease in occupancy and maintain similar occupancy ranking. However, beginning at time point 30 s, the Xe5 site completely disappears and does not reappear at subsequent time points. Concurrently, the occupancies of the Xe1, Xe2, and Xe3 sites increase marginally while Xe4 continues to decrease (Figure 3). Interestingly, before the Xe5 site disappears at time point 30 s, it begins to merge with the Xe1 site at time point 10 s (Figure 2). Such merging of the density was observed only at time point 10 s. Anisotropy of Xe density in a hydrophobic channel has also been noted in H-NOX.<sup>22</sup> By time point 120 s, the occupancy ranking among the Xe sites becomes switched such that the Xe2 site has the highest occupancy (Figure 3). At 240 s and beyond, only the Xe1, Xe2, and Xe3, which are 8–12 Å from each other, sites are occupied.

**A State Model for the Mechanism of Gas Diffusion.** To infer a mechanism for migration of Xe out of the enzyme, we created a simple state model based on the occupancy data. The model (Figure 4A) treats Xe migration as occurring via a series of reactions that move Xe from one site to the next and in



**Figure 4.** (A) A kinetic model for gas diffusion through the hydrophobic channel of *ba3* showing the connectivity of the different Xe sites and their derived rate constants. The sites Xe1 and Xe5 were treated as a single site (see text). (B) Results of the simulation of rate constants after least-squares minimization are plotted vs the experimental occupancies using the model in (A).

which Xe atoms do not interact with each other. The model involves four Xe binding locations in the protein (Xe2, Xe3, Xe4, and a combined site Xe15 = Xe1 + Xe5), plus an external bath. As noted above, the Xe1 and Xe5 sites merge at time point 10 s (Figure 2); therefore, we modeled them as behaving as one by combining their occupancies, i.e.,  $\text{max} = 2$ ,  $\text{min} = 0$ .

Because of spatial constraints within the hydrophobic channel, only those reactions that connect pairs of sites, as shown in Figure 4A, are possible. It is also assumed that there is no reverse flux from Xe in the external bath into the Xe4 site, i.e.,  $k_{4\text{in}}[\text{Xe}]_{\text{bath}} \ll k_{4\text{out}}[\text{Xe4}]$ . Based on the observed changes in Xe2, departure of Xe from this site via the A120 orifice to the external bath<sup>a</sup> does not occur. Hence, in the model, the Xe2 site communicates only with the protein interior (Figure 4A). We then used the Sundials solver in the Simbiology module of MATLAB to numerically solve the set of coupled differential equations corresponding to the reaction model (see Supporting Information for the rate and equilibrium equations). The occupancies under the pressurized condition (equilibrium, time zero) were used to define equilibrium constants for the various reversible reactions. With these constants, four independent rate constants ( $k_{13}$ ,  $k_{23}$ ,  $k_{34}$ , and  $k_{4\text{out}}$ ) were utilized here as input to derive the time-dependent decay of the system and its individual components. The corresponding equilibrium constants,  $K_{13}$ ,  $K_{23}$ , and  $K_{34}$ , permitted calculation of the rate constants of the back reactions:  $k_{31}$ ,  $k_{32}$ , and  $k_{43}$ . To estimate the rate constants in the model, the trust-region-reflective algorithm in the MATLAB “lsqnonlin” function was used to minimize the sum of squares of the residuals between the simulated data and the experimental occupancies at the various sites at different time points.

$K_{13} = (k_{13}/k_{31}) = 0.444$ ,  $K_{23} = (k_{23}/k_{32}) = 0.935$ , and  $K_{34} = (k_{34}/k_{43}) = 0.658$  (see Supporting Information). Because of errors in the occupancy data (Table S2) and other potential considerations, the noted value for  $k_{13}$  ( $0.1 \text{ s}^{-1}$ ) is a lower bound on this rate constant. Further increases in this value result in only small changes in the residuals, without reaching convergence; hence,  $k_{13}$  was fixed at this value during minimization.

Interesting relationships emerge from the estimated rate constants. For example,  $k_{23}$ ,  $k_{32}$  ( $0.002 \text{ s}^{-1}$ ) and  $k_{34}$ ,  $k_{43}$  ( $0.03$ – $0.045 \text{ s}^{-1}$ ) are separated by a factor of 15–22.5, meaning that exchange between the Xe3 and Xe4 sites is an order of magnitude faster than between the Xe3 and Xe2 sites. Similarly, exchange between the Xe3 and combined Xe1/Xe5 sites is about 3.5 times faster than between the Xe3 and Xe4 sites. The rate constant for exit from the Xe4 site of the protein,  $k_{4\text{out}}$ , is  $\sim 0.1 \text{ s}^{-1}$  (Figure 4A). However, this is not the rate-limiting step for the overall loss of Xe; rather, slower internal steps significantly impact this global Xe diffusion out of the protein.

The Gibbs free energy for each equilibrium was obtained from  $\Delta G^\circ = -RT \ln K_i$ , where  $R = 1.986 \times 10^{-3} \text{ kcal mol}^{-1} \text{ K}^{-1}$ ,  $T = 298 \text{ K}$ , and  $K_i$  are equilibrium constants  $K_{13}$ ,  $K_{23}$ , and  $K_{34}$ . The corresponding reaction free energies are  $\Delta G^\circ = 0.48$ ,  $0.04$ , and  $0.25 \text{ kcal/mol}$ , respectively, being well below  $RT = 0.59 \text{ kcal mol}^{-1}$  near room temperature. Nevertheless, the general picture indicates that diffusion of Xe away from the active site is thermodynamically unfavorable, though weakly. Future molecular dynamics studies may clarify the meaning of these small free energy differences,<sup>28</sup> but they are beyond the scope of this work.

The nature of the channel may provide qualitative information about the rate at which a hydrophobic ligand

may migrate from site-to-site. Thus, the channel is essentially uniform in diameter and continuous in its hydrophobic nature all the way from the intramembrane protein surface near the Xe4 site into the Xe1 site. This would suggest that the activation energies for passage between the sites are also small compared to  $RT$ , a condition, which suggests that properly defined molecular dynamics simulations of Xe motions in  $ba_3$  may facilitate further understanding of the mechanism shown here. Such studies have been done for  $O_2$ , NO, CO, and Xe diffusion within sperm whale myoglobin<sup>28</sup> and for  $O_2$  and NO in H-NOX domains.<sup>43</sup>

In spite of these approximations and caveats in the  $ba_3$ , Xe occupancy data provide a first experimental view on the nature of gas movement within the protein. Our simple model provides good agreement with most features of the data (Figure 4B), but the rapid partial initial decay of Xe15 (Figure 3) remains to be better understood. While it is possible that other pathways of Xe leakage might exist for the Xe5 site, merging of the anomalous difference electron density between the Xe1 and Xe5 sites (Figure 2) suggests a very low energy barrier between them.

**Reversal of Ligand Diffusion May Suggest a Mechanism for  $O_2$  Ingress.** The results of the modeling suggest that diffusion of oxygen into the dnc of  $ba_3$  may occur via  $Xe4 \rightarrow Xe3 \rightarrow Xe5 \rightarrow Xe1$  sites. Accordingly, it is of interest to compare the hydrophobic channels in other cytochrome  $c$  oxidases of known structure (see Figure 2 of ref 10). In particular, the *Rhodobacter sphaeroides* cytochrome  $aa_3$  oxidase channel, also determined by Xe pressurization,<sup>12</sup> has only one entry point, analogous to the A120<sup>a</sup> orifice (Figure 1A) and roughly overlaps with the  $Xe2 \rightarrow Xe3 \rightarrow Xe5 \rightarrow Xe1$  branch of the  $ba_3$  channel. Because of homology in the structures and hence an expected passive role of the channel in *R. sphaeroides*, diffusion rates may be similar in the two proteins. A recent study of the reaction between fully reduced  $ba_3$  and  $O_2$  indicates a second-order, diffusion-controlled process by which  $O_2$  binds to  $Fe_{a3}$  to form compound A, an oxy complex.<sup>44</sup> Therefore,  $O_2$  likely enters the channel by diffusion, followed by a faster, first-order process in which  $O_2$  moves from the mouth of the channel at A204 to  $Fe_{a3}$ .

Examination of structures of  $ba_3$  crystallized from detergents suggested that the hydrophobic channel is devoid of crystallographic waters or other small molecules (PDB Codes 1XME and 1EHK). Recently, a 1.8 Å resolution crystal structure of  $ba_3$  in the lipidic cubic phase has identified one water molecule in the hydrophobic channel, that being a water in the Xe1 site hydrogen bonded to a water in the  $a_3$ - $Cu_B$  center;<sup>42</sup> the rest of the channel contains no observable electron density. The higher affinity of the Xe1 site for a hydrophobic ligand, even at longer times (Figure 3), is consistent with the idea that  $O_2$  bound in the Xe1 site has a functional relationship to the active site, being the last “stop” before being reduced to water.

**Xe2 Sites May Function as a Storage Site.** The small changes in occupancy of the Xe2 site over time as Xe diffuses out of  $ba_3$  indicate that very little to no Xe exits the protein through this pathway; hence, the absence of a  $k_{2out}$  in our model. This finding is consistent with studies of the  $ba_3$  I-A120F mutant: even though the phenylalanine replacement physically blocks access to the Xe2 site via the A120 orifice, the mutant retains oxidase activity similar to recombinant, wild-type  $ba_3$ .<sup>42</sup> Hence, the Xe2 arm of the bifurcated channel may not be utilized for primary oxygen input to the enzyme. The much slower rate between the Xe2 to Xe3 sites suggests that

once the Xe2 site is filled by pathway  $Xe4 \rightarrow Xe3 \rightarrow Xe2$ , very little Xe returns to the main input channel, i.e., pathway  $Xe4 \rightarrow Xe3 \rightarrow Xe5 \rightarrow Xe1$ , unless the Xe3 site is empty. Therefore, when  $O_2$  is abundant, the Xe2 site may function as a reservoir for  $O_2$ , which continues to enter through the main input channel during turnover. Under low oxygen tensions, however, the main input channel would become depleted via the Xe4 site, allowing transiently stored  $O_2$  in the Xe2 site to enter the Xe3 site and from there move exergonically to the Xe1 site. This transient supply site would bridge the time when the enzyme consumes all the  $O_2$  in the main input channel during turnover and when the channel gets refilled again. This mechanism is in agreement with the utilization of  $ba_3$  by *T. thermophilus* as the preferred cytochrome  $c$  oxidase when  $O_2$  is scarce.<sup>22</sup> Using internal cavity networks as storage for small molecules in proteins is not without precedence. For example, the truncated hemoglobin family of proteins has been implicated in both  $O_2$  and CO storage,<sup>45</sup> while hydrophobic channels in H-NOX proteins modulate the flux of  $O_2$  and NO, thereby affecting signaling.<sup>22</sup>

## CONCLUSIONS

An extensive Xe-pressurized, crystallographic study supports the presence of a large, hydrophobic channel within which gaseous ligands such as Xe,  $O_2$ , CO, and NO are able to migrate into and out of the active site of the cytochrome  $ba_3$  oxidase. One of the Xe binding sites may also function as a temporary storage site for  $O_2$ . Because the  $ba_3$  structure in Xe pressurized crystals is isomorphous with the native protein [including structures in different redox states<sup>41</sup>] and because the data can be adequately fit with a kinetic model, it appears that the hydrophobic channel functions in a passive manner, and it is unnecessary to consider significant rearrangements within the protein itself. In contrast, in H-NOX proteins dynamics controls  $O_2$  and NO diffusion through “transiently formed pathways”, affecting ligand concentration within an interior cavity proximal to the heme, which in turn affects heme binding and signaling.<sup>22</sup> This behavior is functionally opposite to that in  $ba_3$  where  $O_2$  is a substrate and rapid acquisition from the membrane is necessary for activity and proton pumping.<sup>22</sup> Hence, in  $ba_3$  the populations of Xe in five discrete binding sites, and the rates of exchange among the sites, are determined only by the relative thermodynamic affinity of Xe for the sites and the inherent activation barriers between them. Our observations contribute to a general understanding of how hydrophobic gas molecules gain access to deeply buried active sites while revealing details of their relative mobility within a large integral membrane protein.

## ASSOCIATED CONTENT

### Supporting Information

Protein preparation, purification, and crystallization procedures; Table S1 contains crystallographic statistics for all 19 data sets; the rest of the tables and figures found in this section contain further details on the occupancies of each of the metal sites as well as calculations used in both occupancy normalization and kinetic modeling. This material is available free of charge via the Internet at <http://pubs.acs.org>.

### Accession Codes

The coordinates and observed structure factor amplitudes for the structures in this paper have been deposited in the Protein



Data Bank (PDB codes: 3S33, 3S38, 3S39, 3S3A, 3S3B, 3S3C, 3S3D).

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### Notes

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## ABBREVIATIONS

*ba*<sub>3</sub>, *Thermus thermophilus* cytochrome *ba*<sub>3</sub> oxidase; dnc, dinuclear center.

## ADDITIONAL NOTE

<sup>a</sup>The two arms of the Y-shaped channel are defined as follows: orient the molecule to place the CuA domain at the top (Figure 1A). The fork of the Y facing downward terminates near Ala120; the fork to the right terminates near Ala204. The two orifices of the channel are designated with respect to these residues on the protein surface.

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