Tracking the route of molecular oxygen in O₂-tolerant membrane-bound [NiFe] hydrogenase

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[NiFe] hydrogenases catalyze the reversible splitting of H₂ into protons and electrons at a deeply buried active site. The catalytic center can be accessed by gas molecules through a hydrophobic tunnel network. While most [NiFe] hydrogenases are inactivated by O₂, a small sub-group, including the membrane-bound [NiFe] hydrogenase (MBH) of Ralstonia eutropha, is able to overcome aerobic inactivation by catalytically reduction of O₂ to water. This O₂ tolerance relies on a special [4Fe₃S] cluster that is capable of releasing two electrons upon O₂ attack. Here, the O₂ accessibility of the MBH gas tunnel network has been probed experimentally using a "soak-and-freeze" derivatization method, accompanied by protein X-ray crystallography and computational studies. This combined approach revealed several sites of O₂ molecules within a hydrophobic tunnel network leading, via two tunnel entrances, to the catalytic center of MBH. The corresponding site occupancies were related to the O₂ concentrations used for MBH crystal derivatization. The examination of the O₂-derivatized data furthermore uncovered two unexpected structural alterations at the [4Fe₃S] cluster, which might be related to the O₂ tolerance of the enzyme.

In other enzymes, the molecular tunnels connect the protein exterior with the active site deeply buried within the protein matrix. Such tunnels can serve as permanent or gated substrate pathways as found in DhaA haloalkane dehalogenase or cholesterol oxidase, respectively (7, 8). In lipoxygenases, two permanent hydrophobic tunnels that might be used individually by the two substrates, lipid and dioxygen, merge at the active site (9, 10). The nature of gas-substrate tunnels has also been described in various metalloenzymes by using protein structure and gas diffusion kinetics (11, 12). A recent study on the gas tunnel network of the metalloenzyme, O₂-tolerant [NiFe] hydrogenase and its O₂-sensitive counterparts revealed major structural differences in dimension and complexity (13). Furthermore, tunnel oxygen-tolerant [NiFe] hydrogenase | metalloproteins | iron–sulfur cluster | X-ray crystallography | crystal derivatization

Significance

Tracking the route of substrates, intermediates, and inhibitors in proteins is fundamental in understanding their specific function. However, following the route of gases like molecular oxygen within enzymes has always been challenging. In protein X-ray crystallography, gases can be mimicked using krypton or xenon (with a higher electron count); however, these have a different physical behavior compared to true substrates/inhibitors. In our crystal structure of the O₂-tolerant membrane-bound [NiFe] hydrogenase (MBH) from Ralstonia eutropha, we were able to show the direct path of molecular oxygen between the enzyme exterior and the active site with the "soak-and-freeze" derivatization method. This technique might be useful to detect O₂ traveling routes in many other enzymes.

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 5MDJ, 5MDF, 5MDK, and 4TTT).

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In other enzymes, the molecular tunnels connect the protein exterior with the active site deeply buried within the protein matrix. Such tunnels can serve as permanent or gated substrate pathways as found in DhaA haloalkane dehalogenase or cholesterol oxidase, respectively (7, 8). In lipoxygenases, two permanent hydrophobic tunnels that might be used individually by the two substrates, lipid and dioxygen, merge at the active site (9, 10). The nature of gas-substrate tunnels has also been described in various metalloenzymes by using protein structure and gas diffusion kinetics (11, 12). A recent study on the gas tunnel network of the metalloenzyme, O₂-tolerant [NiFe] hydrogenase and its O₂-sensitive counterparts revealed major structural differences in dimension and complexity (13). Furthermore, tunnel oxygen-tolerant [NiFe] hydrogenase | metalloproteins | iron–sulfur cluster | X-ray crystallography | crystal derivatization

In many enzymes, the catalytic centers are deeply buried within the protein matrix. In these cases, substrate access and the transport of intermediates and product release are often facilitated by dedicated molecular tunnels (1). A sheltered active site in combination with molecular tunnels can have several advantages for the enzyme: (i) an increase of the catalytic efficiency compared with free diffusion, (ii) protection of the active site from reactive and inhibitory compounds, (iii) protection of unstable intermediates, (iv) improved transport time between active sites, (v) increased substrate selectivity, and (vi) regio- and/or stereospecific control (1, 2). In multifunctional enzymes, molecular tunnels connect the different sites of intermediate generation and utilization. A very well-studied enzyme with multiple active sites is the class I amidotransferase, carbamoyl phosphate synthetase (CPS). The intermediates ammonia and carbamate migrate through a permanent tunnel with a length of almost 100 Å. The CPS structure also revealed a difference in tunnel diameter adjusting to the dimensions of the intermediate (3). The intramolecular tunnel of glutamine phosphoribosylpyrophosphate amidotransferase (GPA-Tase) shows a different architectural feature. GPATase catalyzes the conversion of phosphoribosylpyrophosphate into phosphoribosylamine. Here, the intermediate tunnel between the two active sites only exists temporarily upon binding of the correct ligand to the active site, which further induces consecutive conformational changes of the protein backbone (4). In enzymes like glutamate synthase and glucosamine 6-phosphate synthase, the substrate tunnel is obstructed and switches between an open state and a closed state through substrate-induced conformational changes of amino acids, which act as a gate (5, 6).
Here, we present the crystal structure of MBH prepared by a “soak-and-freeze” derivatization method that reveals defined O$_2$ sites in two hydrophobic tunnels leading to the active site and a “dead-end” cavity. These observations are in good agreement with molecular dynamics (MD) simulations suggesting an O$_2$ “hopping” process through these tunnels.

**Results and Discussion**

In our protein crystallography experiments, we applied the soak-and-freeze method, recently established at the European Synchrotron Radiation Facility (ESRF) (24), to investigate whether O$_2$ reaches the [NiFe] active site through a dedicated hydrophobic tunnel network. This technique differs fundamentally from previous crystal derivatization methods as the crystals are flash-frozen in the corresponding gas in its liquid form while still under pressure. This facilitates crystal derivatization at high pressure using difficult-to-handle, highly mobile gases like krypton and molecular oxygen. The specially designed sample supports allow for higher throughput and direct use at synchrotron beamlines. MBH crystals were grown and subsequently exposed to molecular oxygen in a gas pressure cell at 56 bar and 70 bar, which is equivalent to dissolved O$_2$ concentrations of 72.8 mM and 91 mM, respectively. After incubation for time periods ranging from 15 to 70 min, the derivatized crystals were flash-frozen in liquefied O$_2$ at 77 K (Table 1). Subsequently, the gas pressure cell was depressurized and the crystals were transferred to liquid nitrogen (LN2) for analysis at the synchrotron beamline. For all O$_2$-derivatized crystals, a nonderivatized reference (REF) dataset was collected using MBH crystals from the same crystallization batch (Table 2). The O$_2$ positions were determined by examining the Fo$^\text{O_2}$–Fo$^\text{Reference}$ electron density map calculated using structure factors of the O$_2$-derivatized (OxD) and REF datasets (Fig. 2 and Fig. S1, including additional final 2mFo – Dfc, Fo – De difference and simulated-annealing omit electron density maps for all four O$_2$ molecules). The dioxygen positions were discriminated from water molecules by (i) their residence in hydrophobic tunnel cavities (Table 3), (ii) comparison with the REF dataset showing no difference in electron density at the same positions, (iii) inspection of geometry and distances to surrounding amino acids and (iv) correlation with the krypton sites of Kr-derivatized MBH (Fig. S2). Using the soak-and-freeze technique, we identified seven O$_2$ locations (Oxy1–7) in the best dataset (Fig. 1), which were located within the hydrophobic gas tunnel network. Oxy1–4 were modeled into the protein structure because of an occupancy of at least 50%. The real-space correlation coefficients for all four are above 91% (Table 3). Three of the O$_2$ molecules (Oxy5–7) showed occupancies of less than 40%, and were therefore not modeled in the structure [Protein Data Bank (PDB) ID code 5MDL]. The majority of the O$_2$ sites found in the structure describe a continuous pathway between the protein exterior and the [NiFe] active site. One O$_2$ molecule (Oxy2) resides in a hydrophobic pocket surrounded by amino acids M301, V461, S471, F569, E570, L573, and L591 (subscript indicates subunit: L, large subunit; S, small subunit) (Figs. 1 and 2 and Table 3). Owing to the 2D representation of Figs. 1 and 2, Oxy2 appears to be close to the [NiFe] active site, but the actual distance is 16.3 Å. A maximum occupancy of 100% was observed for Oxy1 at 70 bar after 15 min of soaking in pressurized O$_2$ before freezing. Using Oxy1 as a benchmark, we can clearly deduce a correlation of O$_2$ occupancy with pressure and incubation time (Table 1). To inspect the positions of the O$_2$ molecules with respect to the gas tunnel network, the latter was calculated with the program CAVER 3.0 (25), using the highly conserved residue R530L as the initial starting point (Fig. 1). The program predicts two subunit-crossing tunnels, which merge before reaching the [NiFe] active site. Tunnel A, with a minimal diameter ($d_{min}$)
of ~1.4 Å and a length of ~33 Å, is categorized as the primary gas tunnel, whereas tunnel B (d_{min} of ~1.3 Å and length of ~52 Å) may serve as a secondary pathway. Approximately 56% of the amino acids forming the tunnel surface are of hydrophobic character (Fig. S3). The hydrophobic tunnel entrances presumably facilitate access of the gaseous substrates. The highest occupied O₂ site (Oxy1), located in branch 1, has no obvious connection to the protein exterior and might serve as a substrate storage cavity. A similar storage cavity has been proposed by Montet et al. (26) for the O₂-sensitive hydrogenase from Desulfovibrio gigas, assuming that some of the hydrophobic cavities serve as gas reservoirs. They have suggested that the concentration of the gaseous substrate in the enzyme affects the enzyme activity. Comparable conclusions have been drawn from the MD simulation results from [NiFeSe] hydrogenase (27). Baltazar et al. (27) deduce from cavities with trapped H₂ molecules that they either hinder H₂ transport to the active site or promote access by storing gas molecules. Considering the low solubility of H₂ in water, the hydrophobic tunnel system might aid in elevating the H₂ concentration above the surrounding medium level. To further inspect a gas reservoir feature, MD simulations of simultaneous H₂ and O₂ transport in the corresponding ratio through the tunnel network, as well as amino acid exchanges within the proposed tunnel wall, might shed further light on this topic. Oxy3 is located within tunnel B near the bottleneck (smallest tunnel diameter) consisting of the amino acids A127L, V131L, and A202L. Oxy4 is positioned close to the bottleneck of tunnel A and the catalytic center (Figs. 2 and 3). The bottleneck in tunnel A has a distance of <3 Å from the [NiFe] active site and consists of amino acids E27L, R530L, P596L, and C597L.

The observations based on the crystallographic structure were compared with MD simulations of MBH in aqueous solution, as described in further detail in Materials and Methods. The resulting 300-ns-long trajectories showing the diffusion of O₂ and H₂ molecules suggested a hopping-like movement of O₂ and H₂ molecules (examples for diffusion pathways of four individual O₂ molecules are shown in Movies S1–S4) through the main tunnels (Figs. 2 and 3 and Figs. S3, S5, and S6). Notably, diffusion outside the tunnel system and no permanent tunnel opening was detected in the simulations, which is in agreement with previous calculations performed on a structurally similar O₂-sensitive “standard” [NiFe] hydrogenase (21). Diffusion of gas molecules through the previously proposed water channel (19) was predicted to be unfavorable compared with the gas diffusion through tunnels A and B. O₂ diffusion (hopping process) through the gas tunnel network resulted in high O₂ densities at the Oxy1 and Oxy3 sites (molecule occupancies of >75%) (Fig. 3), which is in excellent agreement with the results of the soak-and-freeze experiment. Furthermore, O₂ molecules were detected in the vicinity of all other experimentally proposed

### Table 1. Characteristics of O₂ derivatization experiments

| Dataset | Pressure, bar | Time, min | Resolution, Å | Occupancy Oxy1, % | Fe4-E76 distance, Å | OH⁻ ligand, % |
|---------|--------------|-----------|---------------|-------------------|-------------------|--------------|
| 1       | 56           | 15        | 1.35          | 63                | 2.8               | 31           |
| 2       | 56           | 15        | 1.30          | 72                | 2.8               | 27           |
| 3       | 56           | 15        | 1.42          | 71                | 3.1               | 36           |
| 4       | 56           | 15        | 1.44          | 80                | 2.8               | 28           |
| 5       | 56           | 30        | 1.31          | 79                | 2.5               | 22           |
| 6       | 56           | 30        | 1.45          | 73                | 2.8               | 39           |
| 7       | 56           | 30        | 1.27          | 92                | 2.4               | 35           |
| 8       | 56           | 30        | 1.32          | 91                | 2.5               | 41           |
| 9       | 56           | 60        | 1.55          | 91                | 2.6               | 38           |
| 10      | 56           | 77        | 1.37          | 62                | 2.9               | 36           |
| 11      | 70           | 15        | 1.41          | 100               | 2.6               | 45           |
| 12      | 70           | 15        | 1.45          | 91                | 2.7               | 32           |

### Table 2. REF dataset experiments

| Dataset | Resolution, Å | Fe4-E76 distance, Å |
|---------|---------------|---------------------|
| 1       | 1.40          | 3.37                |
| 2       | 1.41          | 3.30                |
| 3       | 1.38          | 3.50                |
| 4       | 1.36          | 3.39                |
| 5       | 1.50          | 3.43                |
locations, as well as within a 3-Å radius from the active site. Due to the side-chain flexibility at room temperature, the positions of the O₂ sites change during the course of the simulations, particularly in tunnel A. This impedes a direct comparison with the experimentally determined values. The assignment of these sites, however, was computationally predicted on the basis of residence times and site occupancies by the O₂ molecules (Fig. 3).

Analysis of transition probabilities between the preferred O₂ locations showed that, except for Oxy2, all O₂ positions in the MBH were connected by the tunnel network, which was also predicted by CAVER 3.0 (Fig. 2). In addition, Oxy5 and Oxy6, located close to the proposed tunnel entries, showed the highest transition probabilities (28.6% and 23.2%, respectively) to the “bulk region” (solvent and undefined regions within the protein), making these positions potential entry sites for O₂ (Fig. 3B and Table 4). The probability for a transition from the bulk region to branch 1 (Oxy1 and Oxy7) was much lower compared with transitions from the bulk region to positions Oxy5 and Oxy6 at the potential entry points (Table 4). Deeply buried sites, such as Oxy3 and the branch 1 region, showed higher self-transition probabilities (Table 4) and occupancies (Fig. 3B) than regions close to the protein surface (Oxy5 and Oxy6), indicating longer residence times for O₂ at these sites. The only difference between MD and CAVER 3.0 predictions was that in the MD simulations, the observed O₂ densities in tunnel A were lower than in tunnel B (Fig. 3), while CAVER 3.0 predicted tunnel A more densely populated.

### Table 3. Characteristics of O₂ molecules in O₂-derivatized MBH

| O₂ position | Close environment (distance < 4 Å) | \(F_{O} - F_{O}^{\text{reference}}\) peak [σ (contour level)] | Occupancy, % | Atomic displacement factor of O₂ atoms, Å² | Real space correlation coefficient |
|-------------|-----------------------------------|----------------------------------------------------------|-------------|---------------------------------|----------------------------------|
| 1           | W130L, V280L, I283L, G284L, T473L | 14.94                                                   | 100         | O1: 29.61                       | 0.98                             |
| 2           | M30L, C32L, V46L, F569L, E570L, L573L, L591L | 8.41                                                   | 50          | O2: 27.98                       | 0.93                             |
| 3           | A127L, L128L, V131L, A202L, Y206L | 11.42                                                   | 62          | O1: 29.99                       | 0.96                             |
| 4           | S21L, E22L, I25L, T475L | 3.69                                                   | 56          | O2: 28.47                       | 0.91                             |
| 5           | K33L, V36L, L37L, L207L | <4                                                   | <40         | O1: 31.35                       | 0.91                             |
| 6           | L150L, I449L | <5                                                   | <40         | O1: 37.63                       | 0.91                             |
| 7           | L438L, S441L, A442L, L462L, L465L, L474L | <3                                                   | <40         | O1: 21.90                       | 0.91                             |

![Fig. 3.](image-url) Comparison of the experimentally determined O₂ positions in the predicted gas tunnels with the O₂ probability densities derived from MD simulations. (A) O₂ probability densities obtained from MD simulations (pink surface) are superimposed on the structure of O₂-derivatized MBH. The MBH tunnels calculated in PyMOL (29) are shown as a gray surface consisting of tunnel A, tunnel B, and branch 1. O₂ positions determined by the soak-and-freeze approach (24) are indicated as cyan and yellow spheres. The O₂ molecules 5–7 are shown only here, as they are not modeled in the actual MBH structure (PDB ID code 5MDL) due to their low occupancy. The backbone of the MBH is drawn in C-alpha representation, and the active site is drawn as a ball/stick model. (B) Calculated transition network consists of six sites (nodes) illustrated as red circles. The percental occupancy of each site is indicated within the circles. Transitions between nodes are highlighted as arrows, weighted according to their probabilities. Blue arrows mark regions with the highest transition probability coming from the bulk region “state 0” (as a percentage).
as the main access pathway (Fig. 1). This finding may result from the increased flexibility of the amino acid side chains narrowing tunnel A in the simulations.

The spatial probability density of $\text{H}_2$ in the gas tunnel network was similar to that predicted for $\text{O}_2$ (Fig. S6). However, distinct differences in the diffusion properties of $\text{H}_2$ and $\text{O}_2$ were encountered. For the $\text{H}_2$ diffusion, tunnel A was strongly favored over tunnel B, as reflected by (i) the high transition probabilities between Oxy5 and Oxy4 (Table S1) and (ii) the increased occupancies of these sites (Fig. S6). Concomitantly, the low occupancy of Oxy3 (45.7% vs. 76.9% for $\text{H}_2$ and $\text{O}_2$, respectively) reflects a reduced diffusion of $\text{H}_2$ through tunnel B. Because of the smaller size and lower molecular weight of $\text{H}_2$ compared with $\text{O}_2$, higher $\text{H}_2$ densities were found in close vicinity (2.7-Å distance) to the [NiFe] center, as well as outside the MBH tunnel system in the protein matrix (Fig. S5).

To investigate whether the pressure treatment, which comes along with the soak-and-freeze method, had an impact on the overall structure of MBH and its gas tunnels, we compared the $\text{O}_2$-derivatized structure and a structure obtained after high-pressure (2,000 bar of helium) treatment of MBH crystals with the nonderivatized reference structure of MBH at ambient pressure (28). The crystal structures were aligned using the program PyMOL (29), and we obtained root mean square deviations (rmsds) of 0.043 Å and 0.195 Å when comparing the $\text{O}_2$-derivatized structure and the high-pressure structure with the reference MBH structure, respectively (Fig. S7). The greater differences with the high-pressure structure are also reflected by the B-factors (temperature factors). The B-factor comparison between the three structures was visualized using PyMOL in a range between 10 Å$^2$ and 50 Å$^2$ (Fig. 4). All $\text{O}_2$-derivatized MBH structures revealed no significant changes in the overall temperature factors upon pressurization compared with the REF datasets. Nevertheless, a further increase of the pressure to 2,000 bar led to a marked rise in the B-factors. The pressure-induced higher flexibility of amino acids even led to an unresolved disordered loop structure close to the entrance of tunnel A and in an enlargement of the surrounding area (Fig. 4). Furthermore, branch 1 of the hydrophobic tunnel is capped by a surface-located $\alpha$-helix, which does not show an increase in flexibility through pressurization either at 70 bar or 2,000 bar. This supports the proposal of branch 1 being mainly a dead end and not a transient gas tunnel with a lockable lid. Notably, the [NiFe] active site, as well as the medial [3Fe4S] cluster and the distal [4Fe4S] cluster of the electron relay, maintained their spatial structure upon pressurization. In all three MBH structures,

### Table 4. Transition matrix for $\text{O}_2$ diffusion

| State     | Bulk | Oxy1 | Oxy2 | Oxy3 | Oxy4 | Oxy5 | Oxy6 | Oxy7 |
|-----------|------|------|------|------|------|------|------|------|
| Bulk      | 99.6 | 0.0  | 0.0  | 0.0  | 0.0  | 0.0  | 0.0  | 0.0  |
| Oxy1      | 4.2  | 70.9 | 0.0  | 7.2  | 0.0  | 0.0  | 2.9  | 14.8 |
| Oxy2      | 11.0 | 0.0  | 88.9 | 0.0  | 0.0  | 0.0  | 0.0  | 0.0  |
| Oxy3      | 9.7  | 6.8  | 0.0  | 71.0 | 7.0  | 5.5  | 0.0  | 0.0  |
| Oxy4      | 9.2  | 0.0  | 0.0  | 14.3 | 69.4 | 7.1  | 0.0  | 0.0  |
| Oxy5      | 23.2 | 0.0  | 0.1  | 15.6 | 9.3  | 51.8 | 0.0  | 0.0  |
| Oxy6      | 28.6 | 5.1  | 0.0  | 0.0  | 0.0  | 0.0  | 62.9 | 3.4  |
| Oxy7      | 11.9 | 15.9 | 0.0  | 0.0  | 0.0  | 0.0  | 2.2  | 70.0 |

All directional (column to row) gas hopping probabilities are given as a percentage. The bulk region contains positions in the solvent or undefined areas in the protein.

Fig. 4. Comparison of $\text{O}_2$-derivatized, nonderivatized, and high-pressure MBH structures on the basis of the B (temperature)-factors. All structures are shown in tube representation. Low to high B-factors are set to values from 10 to 50 Å$^2$ and represented in the PyMOL color-bar rainbow ranging from dark blue to red. The tube radius is scaled based on the B-factors. The hydrophobic tunnel is illustrated as a black surface with PyMOL (29). The lower structures are rotated by 110° and provide a view of the entry of tunnel A edged by a white line.
the Ni and Fe atoms have distances of 2.9 Å (Table 5), and they are bridged by a hydroxide ligand, which is entirely in line with the oxidation state of the catalytic center (18, 19). They also showed the typical open conformation of the proximal [4Fe3S] cluster with a bond length of ~2.1 Å between iron 4 and the backbone nitrogen of cysteine 20 (Fig. S8), which is compatible with the as-isolated, superoxidized state of the enzyme (19). Despite the structural conformity, the [4Fe3S] cluster of the O2-derivatized MBH showed two additional structural features compared with the reference structure. First, the carboxylate functional group of E76 is shifted toward Fe4 of the cluster (Fig. 5A), and, second, Fe1 carries a hydroxyl ligand (Fig. 5B). While the distance of E76 and Fe4 amounted to 3.4 Å and 3.2 Å in the reference and high-pressure structure, respectively, it was reduced to 2.6 Å in the O2-derivatized MBH (Fig. 5A and Tables 2 and 5). The E76 shift toward the proximal cluster was observed in all of the O2-derivatized MBH structures (Tables 1 and 2). In the as-isolated structure of the membrane-bound O2-tolerant [NiFe] hydrogenase 1 from Escherichia coli (EcHyd-1), the E76 was found in a double conformation with distances of 4.9 Å and 2.4 Å of the glutamate-derived carboxylate Oε to Fe4 (30). Volbeda et al. (30) proposed that E76 functions as a base for deprotonation of the carboxamido N of cysteine 20, which further enables the Fe4 to bind to the backbone of cysteine 20, and thus stabilizes the high-potential form of the [4Fe3S] cluster. A similar cluster stabilization was proposed for the Fe1-bound hydroxyl ligand in MBH. The presence of this ligand with an Fe-O distance of 1.8 Å between the hydroxyl oxygen and the Fe1 was confirmed in all of the O2-derivatized MBH structures. The hydroxyl ligand has an average occupancy of ~34%, with no obvious relation to the O2 concentration (Table 1). In the REF datasets, however, the Fe1-bound hydroxyl ligand was not visible. In the previously published crystal structures of as-isolated, superoxidized MBH, the hydroxyl ligand occurred with high variation in occupancy, reaching from 0% in PDB ID code 4TTT, to 30% in PDB ID code 4IUC (B-factor: 9 Å²), and up to 88% in PDB ID code 4IUB (B-factor: 11 Å²) (19). At present, it remains unclear whether the Fe1-bound hydroxyl ligand and the E76 shift are related to the 5+/-4+ redox state transition of the [4Fe3S] cluster (19) or represent damage caused by high O2 concentrations. Nevertheless, it is important to note that no remarkable damage due to high O2 pressure and/or radiation was observed for the remaining metal cofactors in the MBH. Moreover, the two structural features might point to a dedi-

Table 5. Selected interatomic distances at the [NiFe] site and the [4Fe3S] cluster

| Interatomic distance | O2-derivatized | High pressure | Reference |
|----------------------|---------------|---------------|-----------|
| [NiFe] active site   | 2.9           | 2.9           | 2.9       |
| Ni-Fe distance, Å    | 2.1           | 2.2           | 2.2       |
| [4Fe3S] cluster      | 2.6           | 3.2           | 3.4       |

Fig. 5. Proximal [4Fe3S] cluster of O2-derivatized MBH. (A) [4Fe3S] cluster of O2-derivatized MBH (PDB ID code 5MDK, blue) and nonderivatized (PDB entry 5MDK, orange) MBH is shown in ballistic representation in its as-isolated state with an open conformation. The 2mFe – DFe electron density map of E76, reference is illustrated as blue mesh and contoured at 1σ. The FgO/Fgreference electron density map is depicted as green mesh and contoured at 3σ. (B) Proximal cluster is shown in its superoxidized (as-isolated) state with an open conformation and the hydroxyl ligand bound to Fe1 with a distance of 1.8 Å. The positive mFO – DFe electron density map of the hydroxyl ligand is illustrated as green mesh and contoured at 3σ.

Fe-OH stretching mode, leading to the conclusion of an O2- derived ligand (19). Therefore, the additional ligand at Fe1 of the [4Fe3S] cluster could be part of an O2-reducing mechanism, where His229, which is in hydrogen-bonding distance to the hydroxyl ligand, might play an important role in delivering protons produced in the H2 oxidation reaction at the active site. A recent publication by Dance (31) describes a theoretical model of the communication between the catalytic center and the proximal cluster via conformational changes of residues terminating in this highly conserved His229. This histidine plays a crucial role in transmitting a proton to the opposite side of the proximal cluster for protonation of sulfur S3, followed by an opening of the Fe4-S3 bond. It has been stated that a close water molecule (4.5 Å away from Fe1) could be involved in an OH-/ OH− turnover to provide the H+ for the separate proximal cluster proton cycle. This result is consistent with our structural findings for the proximal cluster in O2-derivatized MBH. S3 protonation, however, does not lead to the Fe4-N20 bond according to Dance (31). Here, a second protonation/deprotonation cycle is
necessary, which might be facilitated by Glu76. In our structural data, we can confirm an interaction of Glu76 with the proximal cluster under aerobic, oxygen-derivatized conditions. However, a sequence of protonation or different intermediates cannot be assigned in our structures and requires further experiments. It needs to be elucidated whether the O(II) ligand could also originate from the surrounding water network. In this case, the proton released from water dissociation might be transferred via His229 (Fig. 5) to the catalytic center to support O(2) reduction to water. Additionally, the O(II) ligand may stabilize the superoxidized state of the proximal cluster (19).

In summary, the technique of crystal derivatization (soak-and-freeze) enables unprecedented insight into the accessibility of the MBH tunnel network for gas molecules. Our data show that MBH contains several hydrophobic tunnels that can be traced via several O2 molecules residing at defined sites. MD simulations further demonstrate the pathways used by the gas molecules traveling from the protein surface to the deeply buried active site. Furthermore, the MD simulations indicate that O2 and the much smaller H2 molecules likely use the same routes within the gas tunnel network. Additional gas diffusion studies might provide supplementary information on the migration behavior of H2 and O2 within the MBH gas tunnel network under ambient conditions. For example, a previous experimental study on [FeFe] hydrogenases revealed that high H2 levels decreased the inhibition by O2 (32). Our MD simulations already revealed differences in the distribution of O2 and H2, O2 occupies the two transition sites of tunnel A to a much lesser extent than H2 does. O2 might be primarily directed toward branch 1 of gas tunnel B, which is located far away from the vulnerable [NiFe] active site. This indicates a mechanism that, at least to some degree, can separate the two gases H2 and O2 from each other, and would thus be another adaptation of O2-tolerant hydrogenases toward prevention of damage by molecular oxygen. However, our data do not exclude accessibility of O2 to the [NiFe] active site. Understanding the architecture of the tunnel network in MBH is crucial for future modifications of the enzyme functions through engineering of the tunnel characteristics. Site-directed amino acid exchanges allow changing the physical tunnel properties, such as the diameter and hydrophobicity. These modifications may have an influence on both the access of the substrate H2 to the active site and the susceptibility of the enzyme toward O2.

Materials and Methods
Protein Expression and Purification. Native MBH was produced and purified in the as-isolated (air-oxidized) state as described elsewhere (18, 33).

Cryocryostatization. MBH was cryocryostatized with the sitting-drop vapor diffusion method in 24-well Linbro plates (Jena Bioscience) at a concentration of around 10 mg mL−1 and at 277 K. The reservoir solution contained 20–30% polyethylene glycol 3350 and 100 mM Bis(2-hydroxyethyl)-amino-Tri(hydroxymethyl)-methane buffer at pH 5.5–6.5. Precipitants and protein solutions were mixed in different ratios (7–8 Å) into microbranches (Hampton Research). After 2–4 d, the dark-brown, needle-shaped MBH crystals were directly used for high-pressure cooling and O2 derivatization experiments or were cryo-protected in 15% glycerol and flash-cooled in LN2 as reference data collection.

High-Pressure Cryocooling. The cryoprotectant free cooling of the MBH crystals was performed using a high-pressure cooling system recently developed at the ESRF (28). The needle-shaped crystals were directly harvested from the crystallization drop with specific pluggable sample supports (MiteGen) and transferred into the pressurizing drop tubes at room temperature. The lower parts of the drop tubes were then cooled in LN2, and the crystals were pressurized with 2,000 bar of helium. Thereafter, the crystals were dropped into the bottom of the tube and flash-cooled at 77 K. Finally, the system was depressurized, and the crystals were handled in LN2 and stored under LN2 for further X-ray experiments. Four crystals were high-pressure-cooled at the same time.

O2 Derivatization. The MBH crystals were derivatized with O2 gas in a cryogenic oxygen pressure cell newly developed at the ESRF (24). Each time, one crystal was harvested from the crystallization drop with a specific pluggable sample support (MiteGen) being surrounded by mother liquor, including 15% glycerol. The crystal was then loaded into the pressurizing drop tube at room temperature. The MBH crystals were pressurized at 56 bar (5,600 kPa) and 70 bar (7,000 kPa) in a time frame from 15 to 70 min. Still under pressure, the crystal was dropped into the bottom of the tube and flash-cooled in liquid O2 at 77 K. After depressurizing the system, the crystal was handled in LN2 with a specially designed cryotoolkit and stored in LN2 for further X-ray experiments.

Structure Analysis. Diffraction data were collected at the ESRF. The best high-pressure (HP) dataset was collected at the tunable beamline ID29 (34) with a PILATUS 6M-F detector at a wavelength of 0.976 Å. The best OxD and REF datasets were collected at the tunable beamline ID30B (35) with a PILATUS 6M-F detector at a wavelength of 0.976 Å. All datasets were measured using data collection and the strategy software packages MXCuBE (36) and EDNA (37). The data collection was carried out in the helical rotation mode at 100 K; a rotation increment of 0.05° with an exposure time of 0.037 s for each frame (3,000 images collected); and detector-to-crystal distances of 191 mm, 163 mm, and 256 mm for the HP, OxD, and REF datasets, respectively. The images of the best dataset were indexed, integrated, and scaled using the XDS program package (38) and the CCP4 (39) program SCALA (40). All crystals showed an orthorhombic space group P212121 (unit cell constants HP dataset: a = 73.39 Å, b = 95.70 Å, c = 121.57 Å and α = β = γ = 90.00°; unit cell constants OxD dataset: a = 73.13 Å, b = 95.57 Å, c = 119.83 Å and α = β = γ = 90.00°). Initial phases for MBH (HP, OxD, and REF datasets) were obtained by molecular replacement based on the crystal structure of the as-isolated MBH (PDB ID code 4T7T) as the initial search model using the CCP4 program Phaser (41, 42). Subsequently, different refinement strategies were implemented on the OxD and REF dataset models and electron density interpretation were performed after each refinement cycle using the program Coot (45). The final OxD model has agreement factors Rwork/Rfree of 13.2%/16.3% (46). Structure validations were performed with the programs of the Research Collaboratory for Structural Bioinformatics PDB Validation server (47), MolProbity server (48) and WHAT IF server (49). All molecular graphics representations were created using PyMOL (29). Table 6 summarizes the statistics for crystallographic data collection and structural refinement.

CAVER Tunnel Calculation. The program CAVER 3.0 was used as a PyMOL plugin (25, 29). The settings in the program were manually adjusted. The initial starting point [NH3+ of the guanidine group of the arginine (i.e., R530, for MBH)], shell depth (2 Å), shell radius (3 Å), cluster threshold (3.5 Å), desired radius (5 Å), and maximum distance (3 Å) were maintained as default parameters. The maximum distance specifies the furthest point (from the initial arginine) to start a tunnel calculation. The bottleneck radius is the narrowest part of a given tunnel and corresponds to the minimum probe radius. The tunnel analysis provides a list of calculated tunnels (residues) with their bottleneck residues, which were used to draw Fig. S1. The tunnels are ranked according to a specific cost function (e.g., for two tunnels with an equal radii, the one with the shorter distances has a lower cost) (25). The tunnel with the lowest cost is the primary tunnel.

MD Simulations. Classical MD simulations were performed to investigate the diffusion of O2 and H2 within MBH. For this, the coordinates of reduced MBH [PDB ID code 3RGW] were inserted randomly, replacing bulk water molecules. The high concentration of oxygen was used to increase the interaction with MBH similar to how it was done for the standard hydrogenase (23). Both gas molecules were simulated as a three-site model as described in literature (23). After stepwise energy minimization and thermal equilibration at 300 K, the system was subjected to a 300-ns-long production run. During the simulation
Table 6. Data collection and refinement statistics

| PDB ID code | O$_2$-derivatized (SMDL) | High pressure (SMDJ) | Nonderivatized (as-isolated) (SMDK) |
|-------------|---------------------------|----------------------|----------------------------------|
| Data collection | ESRF, ID30B | ESRF, ID29 | ESRF, ID30B |
| Space group | P2$_1$2$_1$2$_1$ | P2$_1$2$_1$2$_1$ | P2$_1$2$_1$2$_1$ |
| Cell dimensions a, b, c, Å | 73.13, 95.57, 119.69 | 73.39, 95.70, 121.57 | 73.11, 95.57, 119.83 |
| $\alpha$, $\beta$, $\gamma$ | 90, 90, 90$^\circ$ | 90, 90, 90$^\circ$ | 90, 90, 90$^\circ$ |
| Resolution, Å | 47.78–1.41 | 47.85–1.48 | 47.78–1.50 |
| $R_{merge}$ (58) | (1.49–1.41)$^\dagger$ | (1.56–1.48)$^\dagger$ | (1.58–1.50)$^\dagger$ |
| $R_{merge}$ (58) | 0.064 (0.681)$^\dagger$ | 0.064 (0.839)$^\dagger$ | 0.060 (0.800)$^\dagger$ |
| $I_{(o)}/I_{(e)}$ (59) | 10.7 (2.0)$^\dagger$ | 12.5 (2.0)$^\dagger$ | 12.4 (1.9)$^\dagger$ |
| CC1/2 | 100 (89.6)$^\dagger$ | 100 (82.6)$^\dagger$ | 100 (88)$^\dagger$ |
| Completeness, % | 96.4 (94.4)$^\dagger$ | 99.8 (99.4)$^\dagger$ | 99.8 (98.9)$^\dagger$ |
| Refinement | | | |
| Resolution, Å | 1.41 | 1.48 | 1.50 |
| No. of reflections | 147,390 | 135,294 | 127,184 |
| $R_{merge}$/$R_{merge}$ % | 13.2/16.3 | 14.2/17.2 | 12.5/15.3 |
| No. of atoms/residues | | | |
| Large subunit (HoxG) | 4,774/601 | 4,687/601 | 4,776/601 |
| Small subunit (HoxK) | 2,118/269 | 2,051/268 | 2,142/269 |
| Others | | | |
| [FeS] clusters | 22/3 | 22/3 | 22/3 |
| [NiFe] active site | 9/1 | 9/1 | 9/1 |
| Water | 611/603 | 471/466 | 653/652 |
| Dioxygen | 8/4 | 0 | 0 |
| Cl$^-$ | 1/1 | 1/1 | 1/1 |
| Mg$^{2+}$ | 1/1 | 1/1 | 1/1 |
| Polyethylene glycol | 7/1 | 0 | 0 |
| Mean B-factor (all atoms), Å$^2$ | 19.6 | 27.5 | 25.7 |
| rmsd | | | |
| Bond lengths, Å | 0.008 | 0.010 | 0.007 |
| Bond angles, $^\circ$ | 1.260 | 1.357 | 1.302 |
| Ramachandran plot,$^\S$ (60) | | | |
| Most favored, % | 98 | 98 | 98 |
| Allowed/disallowed | 2/0 | 2/0 | 2/0 |

*One crystal was used.
†Highest resolution shell is shown in parentheses.
‡Half-set correlation coefficient.
§As defined in the program RAMPAGE.

Carried out with Gromacs 4.6.5 (54), a time step of 2 fs was enabled by constraining all bonds containing hydrogen atoms by the LINCS algorithm (51). The NPT ensemble (constant number of particles, pressure (P = 1 atm) and temperature (T = 300 K)) under periodic boundary conditions was realized by the Berendsen pressure (55) and v-rescale temperature (56) methods, applying pressure and temperature coupling constants of $\tau_p = 1.0$ ps and $\tau_T = 0.1$ ps, respectively. Electrostatic interactions beyond a cutoff of 1.4 nm were modeled by the particle mesh Ewald summation (57), and van der Waals interactions were truncated beyond 1.2 nm.

Gas Transition Models. The diffusion of O$_2$ and H$_2$ within the MBH was described by a simple transition model based on the O$_2$ (H$_2$) dynamics extracted from the MD simulations. In total, eight regions were defined. Seven regions correspond to spheres of 5 Å around the amino acids surrounding oxygen molecules (Oxy1–7) in the crystal structure. Bulk solvent (or region 0) contained all gas molecules outside these six defined spheres (solvent and undefined positions in MBH).

Based on this model, transitions between these regions were analyzed for all O$_2$ (H$_2$) molecules using a time resolution of 20 ps. Furthermore, occupancies were calculated taking the last 150 ns of the trajectory into account.

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