Characterization of Anammox Hydrazine Dehydrogenase, a Key N₂-producing Enzyme in the Global Nitrogen Cycle

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Anaerobic ammonium-oxidizing (anammox) bacteria derive their energy for growth from the oxidation of ammonium with nitrite as the electron acceptor. N₂, the end product of this metabolism, is produced from the oxidation of the intermediate, hydrazine (N₂H₄). Previously, we identified N₂-producing metabolism, is produced from the oxidation of the intermediate, c-type hemes in each subunit. These nuances make HDH a powerful reductant in nature, to N₂ (Reaction 3) (5–7).

An estimated 30–70% of all N₂ that is released into the atmosphere is produced by anaerobic ammonium-oxidizing (anammox) bacteria (1, 2), which represent one of the latest scientific discoveries in the biogeochemical nitrogen cycle. These organisms gain their energy for growth from the oxidation of ammonium, with nitrite as the electron acceptor, to produce N₂. With the employment of advanced molecular tools, these bacteria have been detected in nearly every anoxic environment where fixed nitrogen compounds are degraded (3). Besides its biogeochemical and ecological relevance, the anammox process has found worldwide application in ammonium removal from wastewater as an environment-friendly and cost-effective alternative to conventional systems (4).

In our current understanding, anammox catabolism is composed of three consecutive, coupled reactions with two intermediates, nitric oxide (NO) and hydrazine (N₂H₄): 1) the one-electron reduction of the substrate nitrite to NO (Reaction 1); 2) the activation of the second substrate ammonium with NO and the concomitant input of three electrons to synthesize N₂H₄ (Reaction 2); and 3) the oxidation of hydrazine, the most powerful reductant in nature, to N₂ (Reaction 3) (5–7).

**REACTIONS 1–3**

\[
\begin{align*}
\text{NO}_2^- + 2\text{H}^+ + e^- & \rightarrow \text{NO} + \text{H}_2\text{O} \\
(E_0 = + 0.38 \text{ V}) \\
\text{NO} + \text{NH}_4^+ + 2\text{H}^+ + 3e^- & \rightarrow \text{N}_2\text{H}_4 + \text{H}_2\text{O} \\
(E_0 = + 0.06 \text{ V}) \\
\text{N}_2\text{H}_4 & \rightarrow \text{N}_2 + 4\text{H}^+ + 4e^- \\
(E_0 = -0.75 \text{ V})
\end{align*}
\]

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Hydrazine Dehydrogenase

The four electrons that are released during hydrazine oxidation then drive the reduction reactions (Reactions 1 and 2).

The enzyme that catalyzes hydrazine oxidation and thus effectively produces about half of all N₂ emitted into the atmosphere is called hydrazine dehydrogenase (HDH) or hydrazine-oxidizing enzyme. In our previous work, we identified HDH from *Kuenenia stuttgartiensis* (KsHDH) as the gene product of kustc0694 and determined some of its catalytic properties (5). A hydrazine-oxidizing enzyme highly related to kustc0694 was isolated before from the anammox enrichment culture KSU-1, but the exact reaction catalyzed by it has so far remained unknown (8). In the genome of *K. stuttgartiensis*, kustc0694 is one of the 10 paralogs of hydroxylamine oxidoreductase (HAO)-like proteins (3, 6). HAO-like proteins have two structurally well-characterized representatives, NeHAO from *Nitrosomonas europaea* (9–11) and hydroxylamine oxidase from *K. stuttgartiensis* (kustc1061; here denoted as KsHOX) (12). NeHAO is a key enzyme in aerobic ammonium-oxidizing bacteria, which catalyzes the four-electron oxidation of hydroxylamine to nitrite (Reaction 4), whereas KsHOX is a dominant protein in the anammox bacterium catalyzing the three-electron oxidation of hydroxylamine to NO (Reaction 5).

\[
\begin{align*}
\text{NH}_2\text{OH} + \text{H}_2\text{O} & \rightarrow \text{NO}_2^- + 5\text{H}^+ + 4e^- \\
(E_0' &= +0.065 \text{ V})
\end{align*}
\]

\[
\begin{align*}
\text{NH}_2\text{OH} & \rightarrow \text{NO} + 3\text{H}^+ + 3e^- \\
(E_0 &= -0.030 \text{ V})
\end{align*}
\]

**REACTIONS 4 and 5**

Despite a limited sequence identity (~30% at the amino acid level), NeHAO and KsHOX are structurally highly similar (10–12). Both are homotrimeric proteins in which each monomer binds eight c-type hemes (Fig. 1). The arrangement of 24 hemes of both proteins is fully superimposable. Within a subunit, seven His/His-ligated hemes constitute an electron-wiring circuit toward an external electron acceptor (12, 13). To facilitate efficient electron transport, these hemes may be electronically coupled, resulting in highly convoluted EPR spectra of as-isolated, fully oxidized (all ferric) NeHAO (14, 15). One heme (heme 4) forms part of a structurally conserved catalytic center (Fig. 1). This heme 4 is covalently bound to a tyrosine residue from a neighboring subunit. This unusual cross-link induces a characteristic absorption band at about 460 nm in the UV-visible spectrum of the reduced protein (9, 10, 12, 14). After this absorption band, catalytic heme 4 is termed the P₄₆₀ cofactor. Due to small structural changes around the P₄₆₀ catalytic site (Fig. 1C), KsHOX oxidizes hydroxylamine to NO rather than to nitrite as NeHAO does (12). Interestingly, both HAO and KsHOX can use hydrazine as a substrate too (Reaction 3), although with lower catalytic efficiency (kcat/Km) than their physiological substrate, hydroxylamine (9, 12).

Here, we characterized KsHDH in detail and identified it as a homotrimeric octaheme protein in which the three subunits are covalently bound to form a P₄₆₀-like prosthetic group as found in NeHAO and KsHOX. In contrast to the two latter enzymes, the homotrimerics of KsHDH themselves were found to form octamers in solution. KsHDH specifically oxidized hydrazine to N₂ in accordance with Equation 3 and was inhibited by NH₂OH and NO. The genome of *K. stuttgartiensis* harbors a close paralog of kustc0694, namely kustd1340, whereas close orthologs of kustc0694 and kustc1340 were detected in all anammox genomes sequenced thus far. Considering a close structural relationship with other HAO-like proteins, we addressed which features determine the specificity of HDH from anammox bacteria.

**Results**

**HDH Is a Homotrimeric Protein with Covalently Bound Subunits Forming Octamers in Solution**—Hydrazine dehydrogenase from *K. stuttgartiensis* was purified as a bright red protein that, when resolved by native PAGE, displayed one prominent band with an estimated molecular mass of ~200–220 kDa (Fig. 2A). When resolved by SDS-PAGE, the protein remained at the top of the gel. Occasionally, a smaller band was observed that migrated with a size of ~67 kDa (Fig. 2B, arrow). MALDI-TOF MS verified that all visible bands were derived from the same protein that had been annotated as the octaheme protein kustc0694. Notably, 18 of 83 predicted peptides in the m/z 500–4,000 range were detected after tryptic digestion, including two peptides (molecular masses 2,173.9 and 2,615.2 Da) that distinguished kustc0694 from its close paralog kustd1340 (96% sequence identity at the amino acid level; supplemental Fig. S1). Peptides comprising a CX₃CH motif for covalent heme c binding remained beyond detection. Linear MALDI-TOF MS established a molecular mass of 201,800 ± 300 Da of the holoenzyme. This mass matched the theoretical value (201,610.5 Da) of a homotrimer with each monomer (62,271.5 Da) possessing eight bound heme c (Mₚ = 616.5) molecules. Thus, the high molecular mass of native KsHDH indicated that the three subunits were covalently bound to each other. Calculation of the subunit theoretical mass took into account N-terminal cleavage (after Val-107) following an alternative translation start site at Met-75 (see below; supplemental Fig. S1). Cleavage of the N-terminal leader sequence (32 amino acids) would facilitate export of HDH into the anammoxosome, an anammox-specific cell organelle where catabolism resides and where processed HDH is specifically localized, as recently shown by immunogold labeling (16).

The oligomeric state of KsHDH was corroborated by the results from sedimentation velocity analytical ultracentrifugation (AUC SV) and analytical size exclusion chromatography coupled to multiangle light scattering (SEC/MALS) analyses (Table 1 and Fig. 3, A and B). Both methods showed a prominent species corresponding to the homotrimer (α₃, calculated mass of 201.6 kDa) and a monomeric species (M₁ = 61,000–75,000) in minor amounts. Strikingly, both methods also revealed aggregates with molecular masses of 1,760.7 and 1,731.7 kDa as determined by SEC/MALS and AUC SV, respectively (Table 1). In the AUC sedimentation profile, these aggregates represented the major peak (Fig. 3A). Amino acid analysis of both the monomeric and octameric species confirmed the theoretical mass (162,271 Da) of the holoenzyme. This result is in accordance with the MALDI-TOF MS data (201,800 Da), suggesting that the octameric species was a native superstructure. The octameric state was retained in both SDS-PAGE and native PAGE. The mass distribution in both techniques indicated that the octamer was a multimer of three subunits. The subunit sedimentation coefficient S₂₀,W was determined as 5.9 S, corresponding to an approximate molecular mass of 62,000 Da, in agreement with SEC/MALS data (62,271.5 Da). High-resolution SEC/MALS chromatograms featuring the octameric state of KsHDH were obtained when the column was processed with the N-terminal leader sequence (see below). This method resulted in chromatograms featuring the octameric state of KsHDH. The high molecular mass of native KsHDH indicated that the three subunits were covalently bound to each other. Calculation of the subunit theoretical mass took into account N-terminal cleavage (after Val-107) following an alternative translation start site at Met-75 (see below; supplemental Fig. S1). Cleavage of the N-terminal leader sequence (32 amino acids) would facilitate export of HDH into the anammoxosome, an anammox-specific cell organelle where catabolism resides and where processed HDH is specifically localized, as recently shown by immunogold labeling (16).
two major peaks with elution volumes as observed for the original sample. This indicated a dynamic equilibrium between the \( \alpha_3 \) and \( \alpha_{24} \) species in solution. Transmission electron micros-

copy (TEM) of negatively stained KsHDH samples supported the AUC SV and SEC/MALS results and revealed assemblies with an approximate size of 16 nm, next to smaller (8 nm) globular particles, accounting for the homotrimers (Fig. 4A). After glutaraldehyde cross-linking and repurification by SEC, these 16-nm particles having a 4-fold symmetry were the most abundant species observed besides particles with a 5-fold symmetry in minor amounts (Fig. 4, B and D–F). The 16 × 16-nm particles were visible as tetrameric configurations and as two parallel dumbbells, probably representing orthogonal views of the same particle (Fig. 4, D and E). These large particles would then be composed of eight egg-like protein molecules, their tops pointing to each other. One may note that one such assembly would carry a calculated 192 heme c molecules. KsHOX did not migrate as higher molecular weight aggregates, neither on SEC/MALS nor on AUC SV chromatograms. TEM showed a homogeneous distribution of 8 × 8-nm particles only (Fig. 4C). Their size and shape fitted the dimensions (9 × 9 × 9 nm) determined for the tulip-shaped KsHOX homotrimeric structures (Fig. 1A) (12).
Hydrazine Dehydrogenase

TABLE 1
Characterization of KsHDH oligomers in solution

<table>
<thead>
<tr>
<th>Stoichiometry</th>
<th>Calculated mass (kDa)</th>
<th>AUC S20,w</th>
<th>smax</th>
<th>smax/s20,w</th>
<th>Mass from AUC SV (kDa)</th>
<th>Mass from MALS (kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>α</td>
<td>67.2</td>
<td>5</td>
<td>6.0</td>
<td>1.20</td>
<td>68</td>
<td>75.0 ± 2.0</td>
</tr>
<tr>
<td>α3</td>
<td>201.6</td>
<td>10.97</td>
<td>12.4</td>
<td>1.13</td>
<td>220.7</td>
<td>182.1 ± 1.8</td>
</tr>
<tr>
<td>α24 = (α3)8</td>
<td>1612.9</td>
<td>43.33</td>
<td>49.7</td>
<td>1.15</td>
<td>1731.7</td>
<td>1760.2 ± 7.0</td>
</tr>
</tbody>
</table>

AUC SV coefficients are given in Svedberg units (1 S = 10^{-13} s). 
smax: maximal sedimentation coefficients at which smax = 0.00361 ml^3.

MALS analyses accounted for any shape.

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kustd1340 were readily detected in other sequenced anammox genomes (79–96% sequence identities of the N-terminally cleaved gene products; supplemental Fig. S1). However, kustc0694 differed from all others by the presence of a 75-amino acid N-terminal extension. This extension was most likely an artifact caused by an erroneously annotated start codon, the actual translation start being Met-75. In agreement with this, 21 nucleotides upstream of Met-75, a Shine-Dalgarno sequence (AGGAGG) was present. Met-75 was followed by an N-terminal leader sequence (32 amino acids) seen at this position in all other HDH homologs. Moreover, the first peptide (M_r = 2,691.2 Da) that we could detect in tryptic digests was the one immediately following Val-107 in the KsHDH sequence.

Multiple-protein sequence alignment affiliated the HDH-like proteins with octaheme proteins with known crystal structures and functions (NeHAO and KsHOX) (supplemental Fig. S1). This affiliation was supported by sequence conservation of the eight heme c binding motifs, including the positions of the seven histidines that act as distal ligands to the seven His/His-ligated heme c moieties involved in electron transfer. An unusual feature of the HDH proteins was the presence of a CX_4CH, rather than a CX_2CH, binding motif for heme 3. Taking into account the conserved position of this motif and of the distal histidine (H3'), this heme 3 would be His/His-ligated, making it an electron-transferring one, but possibly with a different spatial orientation as compared with heme 3 in NeHAO and KsHOX.

Notwithstanding their limited amino acid sequence identity (~30%), protein folds and the arrangement of heme cofactors of NeHAO and KsHOX are virtually identical (Fig. 1) (10–12). This similarity particularly concerns the structure of the catalytic heme 4 and its surroundings. Sequence comparison suggested that the structural similarity might include the HDH members. Notably, key amino acids involved in the covalent binding of an adjacent subunit to heme 4 (a tyrosine) and in proton withdrawal from the substrates (an aspartate-histidine pair) could be detected in the HDH sequences, next to the amino acids forming part of a hydrogen-bonding network that transports protons to the water-filled cavity at the substrate entrance site (Fig. 1, A and C). A critical difference between NeHAO and KsHOX is the presence a tyrosine (Tyr-358) that assists in the addition of a water molecule to a nitrosyl intermediate during nitrite formation in the former protein (see Reaction 4) (12). In KsHOX, this tyrosine is moved away from the catalytic site by a two-amino acid contraction, and the position of the NeHAO Tyr-358 is occupied by a methionine (Met-323). This very same contraction and methionine replacement were also found in the HDH members. One may note that neither

Sequence Analysis of KsHDH and Its Homologs Related to HAO-like Octaheme Proteins from Other Anammox Species—MALDI-TOF MS indicated KsHDH to be a homotrimmeric protein possessing eight c-type hemes per subunit and identified the protein as the gene product of kustc0694 rather than of the close paralog kustd1340. Close homologs of kustc0694 and
hydroxylamine oxidation to NO (Reaction 5) nor hydrazine oxidation to N$_2$ (Reaction 3) includes a water addition. Opposite to the catalytic site near the proximal histidine of heme 4 (P$_{460}$), there is a distinction between the three proteins (Fig. 1, B and C) that might be of relevance in determining substrate specificity. Here, the carboxylate oxygen of Asp-231 in NeHAO is at hydrogen-bonding distance to the proximal histidine of heme 4 (P$_{460}$), whereas the carbonyl oxygen of Ala-201 occupies this position in KsHOX. Remarkably, in the HDH members, a cysteine was observed at this site, possibly affecting the heme 4 redox potential (see below).

**KsHDH Is an N$_2$-producing Enzyme**—KsHDH catalyzed the four-electron oxidation of hydrazine to N$_2$ according to Reaction 3. This four-electron stoichiometry was apparent when following the reaction with excess ferric cytochrome c as the electron acceptor (Fig. 5A). Experiments with double-labeled hydrazine (H$_2$N$^{15}$N-$^{15}$NH$_2$) confirmed that double-labeled N$_2$ was the end product of this reaction, fitting the expected 1:1 stoichiometry (Fig. 5B). Hydrazine oxidation followed Michaelis-Menten kinetics with a $V_{\text{max}} = 11 \pm 1.2$ $\mu$mol min$^{-1}$ mg$^{-1}$ of protein and a $K_m = 10 \pm 2.2$ $\mu$M for hydrazine (Table 2). Hydroxylamine did not serve as a substrate, neither in oxidative nor in reductive directions, when assays were performed in the presence of oxidized or reduced cytochrome c, respectively. In the absence of cytochrome c, hydroxylamine disproportionation into ammonium and nitrogen species of higher oxidation states (NO$_2^+$, NO, N$_2$O, N$_2$) did not occur. Other nitrogenous compounds tested (NH$_4^+$, NO, NO$_2^-$, NO$_3^-$) were not converted in assays with oxidized or reduced cytochrome c as cosubstrate either. Instead of being substrates, hydroxylamine ($K_i$$_{NH_2OH} = 7.9 \pm 1.8$ $\mu$M) and NO ($K_i$$_{NO} = 2.5 \pm 0.9$ $\mu$M) were strong competitive inhibitors of hydrazine oxidation activity. These findings demonstrated that KsHDH indeed was a dedicated hydrazine dehydrogenase that makes N$_2$.

The catalytic properties of KsHDH were similar to those reported for its homolog (HDH/hydrazine-oxidizing enzyme) purified from the anammox enrichment culture KSU-1 (Table 2). Octaheme KsHOX from *K. stuttgartiensis* (kustc1061) and from other anammox bacteria oxidizes hydrazine as well, albeit with lower catalytic efficiency ($k_{\text{cat}}/K_m$). Hydrazine also served as a substrate for NeHAO, surprisingly with $k_{\text{cat}}$ and $K_m$ values that were superior to genuine HDH, although hydrazine does not play a role in *N. europaea* metabolism; this organism and other aerobic ammonium-oxidizing microorganisms are not able to synthesize hydrazine, which is an exclusive property of anammox bacteria (3, 5–7).
Hydrazine Dehydrogenase

KsHDH Harbors a P460 Prosthetic Group—As expected from its bright red color, linear MALDI-TOF MS, and protein sequence analyses, KsHDH was a multiheme protein. The UV-visible electronic absorbance spectrum of as-isolated, fully oxidized KsHDH displayed features typical for ferric heme c proteins, namely a dominant absorption peak having a maximum at 408 nm and a broad absorption band in the 550 nm region (Fig. 6). After reduction by dithionite, pronounced absorption peaks emerged with maxima at 420, 524, and 554 nm, corresponding to the Soret, β, and α bands, respectively, that are typically for ferrous c-type hemes. However, an absorption band at 473 nm revealed the presence of a P460-type ferrous heme group in KsHDH as is typical for NeHAO and in KsHOX and in agreement with the presence of a conserved Tyr in the sequence alignments (9, 10, 12, 14). The sequential reduction of KsHDH by the addition of small aliquots of dithionite demonstrated that the His/His-ligated hemes were reduced first (Fig. 7, A and B). Only after their reduction, the P460 band emerged, implying that its redox potential would be lowest. Equilibration of KsHDH at a series of defined potentials supported this conclusion (Fig. 7C) and suggested an Em of about −420 mV for P460 in KsHDH, whereas the His/His-ligated hemes reduced at

![Graph showing kinetic values for mammalian cytochrome c](image)

FIGURE 5. Hydrazine oxidation to N2 by KsHDH. A, hydrazine oxidation (initial concentration, 10 μM) by KsHDH (0.65 μg) was followed by recording the reduction of 50 μM cytochrome c at 550 nm. No reaction was observed when followed in the presence of 50 μM hydroxylamine (dashed line). The concentration of reduced cytochrome c was calculated on the basis of the increase of absorbance at 550 nm (Δε550 = 19,600 M⁻¹ cm⁻¹). B, formation of labeled 30N2 from double-labeled hydrazine (H215N-15NH2, 16 nmol) was followed over time in the presence of 50 μM cytochrome c and 63 ng of KsHDH.

**TABLE 2**

Catalytic and structural properties of hydrazine dehydrogenase from *K. stuttgartiensis* and other hydrazine-oxidizing enzymes

Kinetic values are given for mammalian cytochrome c as electron acceptor. KSU-1, anammox strain KSU-1. NA, not applicable; NR, not reported.

<table>
<thead>
<tr>
<th></th>
<th><em>K. stuttgartiensis</em> HDH</th>
<th><em>K. stuttgartiensis</em> HOX</th>
<th>KSU-1 HZO/HDH</th>
<th>KSU-1 HOX</th>
<th>Brocadia anammoxidans HOX</th>
<th>Nitrosomonas europaea HAO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vmax N,NH2 (μmol min⁻¹ mg⁻¹)</td>
<td>11 ± 1.2</td>
<td>1.6 ± 0.0</td>
<td>6.2</td>
<td>0.54</td>
<td>1.1</td>
<td>14</td>
</tr>
<tr>
<td>kcat N,NH2 (s⁻¹)</td>
<td>36</td>
<td>4.9</td>
<td>13.4</td>
<td>1.1</td>
<td>3.4</td>
<td>47</td>
</tr>
<tr>
<td>kcat K₃,N₂H₄ (μM⁻¹ s⁻¹)</td>
<td>10 ± 2.2</td>
<td>54 ± 3.3</td>
<td>5.5</td>
<td>25</td>
<td>18</td>
<td>4</td>
</tr>
<tr>
<td>kcat K₃,N₂H₄ (μM⁻¹ s⁻¹⁻¹)</td>
<td>3.5</td>
<td>0.1</td>
<td>2.4</td>
<td>0.942</td>
<td>0.19</td>
<td>12</td>
</tr>
<tr>
<td>kcat NH₃OH (μM⁻¹ s⁻¹)</td>
<td>NA</td>
<td>4.8 ± 0.2</td>
<td>NA</td>
<td>9.6</td>
<td>21</td>
<td>28.5</td>
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<tr>
<td>kcat NH₃OH (μM⁻¹ s⁻¹)</td>
<td>NA</td>
<td>15</td>
<td>NA</td>
<td>19</td>
<td>64</td>
<td>95</td>
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<tr>
<td>kcat NH₃OH (μM⁻¹ s⁻¹⁻¹)</td>
<td>NA</td>
<td>4.4 ± 0.9</td>
<td>NA</td>
<td>33</td>
<td>26</td>
<td>3.6</td>
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<tr>
<td>kcat NH₃OH (μM⁻¹ s⁻¹⁻¹)</td>
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<td>3.4</td>
<td>NA</td>
<td>0.58</td>
<td>2.5</td>
<td>26</td>
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<tr>
<td>kcat NH₃OH (μM⁻¹ s⁻¹⁻¹)</td>
<td>NA</td>
<td>7.9 ± 1.8</td>
<td>NA</td>
<td>2.4</td>
<td>NA</td>
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<tr>
<td>kcat NO (μM)</td>
<td>2.5 ± 0.9</td>
<td>NA</td>
<td>NA</td>
<td>NR</td>
<td>NR</td>
<td>NA</td>
</tr>
<tr>
<td>Total size (kDa)</td>
<td>210.5 (~1,600)</td>
<td>184</td>
<td>130</td>
<td>118</td>
<td>183</td>
<td>200</td>
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<tr>
<td>Subunit composition</td>
<td>α₃/(α₂)α₄</td>
<td>67.2</td>
<td>61.3</td>
<td>62</td>
<td>53</td>
<td>58</td>
</tr>
<tr>
<td>Catalytic heme optical maximum (nm)</td>
<td>473</td>
<td>468</td>
<td>472</td>
<td>468</td>
<td>468</td>
<td>463</td>
</tr>
<tr>
<td>Catalytic heme midpoint potential (mV)</td>
<td>−420</td>
<td>−300</td>
<td>NR</td>
<td>NR</td>
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<td>19–21</td>
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midpoint redox potential ($E_m$) values ranging between $-70$ and $-410$ mV. The $P_{460}$ reduction potentials of NeHAO and KsHOX ($E_m = -260$ and $-300$ mV versus NHE, respectively) are substantially higher (12–14).

When as-isolated KsHDH was incubated with its substrate, hydrazine, the protein became partially reduced, as concluded from the increase in the absorbances at 420, 524, and 554 nm. The degree of reduction was dependent on the concentration of hydrazine added. Neither hydroxylamine nor NO influenced the absorption spectrum of oxidized KsHDH, but both compounds instantaneously oxidized the dithionite-reduced protein.

**EPR Spectroscopy of HAO-like Octaheme Proteins**—The sequence analysis and biophysical data presented above suggested that the HDH members shared the common architecture of NeHAO and KsHOX enzymes (Fig. 1). This was assessed by inspection of EPR spectra arising from the oxidized forms of all three enzymes.

Previously, Hendrich et al. (15) were able to delineate the complex spectra of fully oxidized NeHAO into four distinct EPR species covering the expected eight Fe$^{3+}$ centers in the eight hemes per subunit (Table 3). Two of the species (species 3 and 4) could be unambiguously assigned to specific hemes. EPR species 4 was the result of spin-coupled heme 4 ($P_{460}$) and heme 6 (see Fig. 1D for the numbering and spatial orientations of the hemes). Importantly, this species was only observed when EPR spectra were recorded in the parallel mode. EPR species 3 was derived from magnetically interacting heme 3 and heme 5. The analyses by the authors (15) left four magnetically isolated low spin hemes unassigned, which were observed as two species (species 1 and 2) covering one and three heme centers, respectively (Table 3). Thus, none of these showed appreciable interaction with each other. A one-electron reduction of NeHAO

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**FIGURE 7. Optical redox determinations of KsHDH.**

**A**. Reduction of as-isolated KsHDH (43 µg ml$^{-1}$; 0.64 µM) by the sequential addition of dithionite aliquots under anaerobic conditions. Spectra were corrected for the effect of dilution by the additions. AU, absorbance units. Arrows, directions of the spectral changes. The absorbance band at 473 nm derived from the $P_{460}$ prosthetic group appeared only after the complete reduction of the His/His-ligated c-type hemes featured in their reduced state by their absorption maxima at 420, 524, and 554 nm. The spectrum of fully reduced KsHDH is highlighted by the red line. B. Reduced minus oxidized absorbance spectra during dithionite reduction. Note that spectral changes occur around distinct isosbestic points at 412, 433, 464, 485, 508, 533, and 562 nm, indicating that the spectral changes as the result of the reduction of the $P_{460}$ catalytic heme do not interfere with those of the His/His-ligated c-type hemes. C. Redox titration of KsHDH ($30$ µM) in an optical transparent thin layer electrochemical cell. Normalized redox potential-dependent spectral changes of the His/His-ligated hemes of KsHDH were followed in oxidative (ox) and reductive (red) directions at 554 nm (black squares, oxidative; blue triangles, reductive) and at 420 nm (red circles, oxidative; green diamonds, reductive). Spectral changes at 473 nm as a result of the reduction of the $P_{460}$ prosthetic group are shown as red triangles. Data were fitted (black line) by a Nernstian curve, assuming a one-electron reduction resulting in an $E_m = -420$ mV.
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TABLE 3
EPR spectroscopic properties of NeHAO, KsHOX, and KsHDH

<table>
<thead>
<tr>
<th>Protein EPR species (iron/monomer)</th>
<th>gxyz values</th>
<th>Comment/Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>NeHAO</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 (1)</td>
<td>1.20; 2.26; 3.03</td>
<td>Uncoupled heme 1</td>
</tr>
<tr>
<td>2 (3)</td>
<td>1.35; 2.19; 3.11 (3.09)</td>
<td>Coupled hemes 7 and 8; includes the easily reducible heme 2</td>
</tr>
<tr>
<td>3 (2)</td>
<td>1.27; 2.28; 3.06</td>
<td>Coupled hemes 3 and 5</td>
</tr>
<tr>
<td>4 (2)</td>
<td>8</td>
<td>Coupled hemes 6 and 4 (Pex); only observed in parallel mode EPR</td>
</tr>
<tr>
<td>KsHOX</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 (1)</td>
<td>1.452; 2.305; 2.955</td>
<td>Heme 1; g values become 1.452, 2.260, and 2.980 after NH2OH incubation</td>
</tr>
<tr>
<td>2a (1)</td>
<td>1.41; 2.145; 3.1412</td>
<td>Reduced after NH2OH incubation; heme 2</td>
</tr>
<tr>
<td>2b (2)</td>
<td>1.41; 2.217; 2.955</td>
<td>Uncoupled hemes 7 and 8; g values become 1.37, 2.217, and 2.980 after NH2OH incubation</td>
</tr>
<tr>
<td>3 (2)</td>
<td>1.40; 2.18; 3.247</td>
<td>Coupled hemes 3 and 5 (Euler angles: 45, –180, and 135°); after NH2OH incubation, each heme reduces by 60%, g changes to 3.265, and the exchange splitting increases from 0.031 to 0.035 cm⁻¹</td>
</tr>
<tr>
<td>4 (2)</td>
<td>NDd</td>
<td>High spin heme, but amount is preparation-dependent (0.03–0.14/monomer; the higher value shown in Figs. 8 (A and B) and 9A)</td>
</tr>
<tr>
<td>5 (minor)</td>
<td>6.07; 5.81; 1.98</td>
<td></td>
</tr>
<tr>
<td>KsHDH</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 (1)</td>
<td>1.6; 2.05; 3.14</td>
<td>Uncoupled heme (heme 1) with broad resonances that sharpen and shift (gxyz = 1.51, 2.256, 3.10) upon reduction by hydrazine</td>
</tr>
<tr>
<td>2a (1) (3.8%) (6.7%) (6.1%)</td>
<td>1.772; 2.06; 2.579</td>
<td>Heme 2; NH4 is reducible; same heme in slightly different environments</td>
</tr>
<tr>
<td>2b (2)</td>
<td>1.772; 2.06; 2.543</td>
<td>Coupled hemes 7 and 8 (Euler angles: 45, 180, and 87°); no reduction by hydrazine, but g changes to 3.05; the exchange splitting is 0.035 cm⁻¹</td>
</tr>
<tr>
<td>3 (2)</td>
<td>1.5; 2.3; 3.0</td>
<td>Coupled hemes 3 and 5 (Euler angles: 45, –180, and 135°); reduction by hydrazine (70%), and g changes to 3.04; the exchange splitting is 0.074 cm⁻¹ and 0.087 cm⁻¹ after reduction</td>
</tr>
<tr>
<td>4 (2)</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>5 (minor)</td>
<td>6.0; 6.0; 2.0</td>
<td>Minor high spin heme (&lt;0.02/monomer)</td>
</tr>
<tr>
<td>6 (minor)</td>
<td>2.099</td>
<td>Possible tyrosine radical coupled to heme 4</td>
</tr>
</tbody>
</table>

a In column 1, numbers in parentheses indicate the integral number of spins, and percentage values represent the percentage of the total of integrated spins.
b Data from Hendrich et al. (15).
c In parentheses, g after 1-electron reduction.
d ND, not determined. Signal accounting for hemes 6 and 4, which only would be observed in parallel mode EPR.

caused the partial reduction and a notable change of spin species 2 (15). Such one-electron reduction would concern the heme with the highest redox potential, which is heme 2 in NeHAO (13). Heme 2 is a key electron sink because it can accept electrons passing via hemes 5 and 3 within the same subunit or passing via hemes 7 and 8 from an adjacent subunit before transferring these to solvent-exposed exit heme 1 (Fig. 1D). In the following, we started from the assumption that heme 2 also has the highest redox potential in KsHOX and KsHDH. Supported by our analyses of KsHOX and KsHDH (see below), we then assigned spin species 2 to non-coupled hemes 7 and 8 together with heme 2, leaving the isolated EPR species 1 to heme 1 (Table 3).

EPR Spectroscopy of KsHOX—Following the same approach as Hendrich et al. (15) and using the same theoretical spin Hamiltonian (Equation 1), we could deconvolute the EPR spectrum of KsHOX into the same spin species and allocate those species to all hemes (Table 3 and Figs. 8A and 9 A and B). Whereas spin species 2 now could be resolved into two sub-species (species 2a (heme 2) and species 2b (hemes 7 and 8); Table 3), the overall EPR spectrum of as-isolated, fully oxidized KsHOX (Fig. 8A) was highly similar to the one described for NeHAO (14, 15). This was expected, because the heme arrangement in both proteins was fully conserved, regarding both the spatial orientations and positions of the hemes and their distances with respect to each other (Fig. 1D). Simulation of the normal mode EPR spectra of KsHOX and the individual heme centers is shown in Fig. 9 (A and B); simulation parameters are listed in Table 3. In the normal mode EPR spectra recorded in this work, the coupled heme pair 6 and 4 remained beyond detection, and because we had no access to parallel mode EPR, these hemes were not determined. Consequently, a total of six of eight hemes per monomer would be observed. In agreement herewith, quantitation of the EPR spectrum versus a copper perchlorate standard (31) of oxidized, as-isolated,
KsHOX (665 μM hemes, as determined optically) yielded a value of 480 μM S = ½ low spin hemes, which is consistent with a total of six EPR-visible hemes. Except for small differences in g values, the main difference between KsHOX and NeHAO was the strength of the magnetic coupling ($D_{zz} - D_{xx}$) between hemes 3 and 5. For KsHOX, a value of 0.031 cm$^{-1}$ was determined for the oxidized enzyme; for NeHAO, the value was 0.08 cm$^{-1}$ (15). For analyses, the same relative orientation of the two heme centers given by Euler angles between the g-tensor axes of 45°, 180°, and 135° were used. These angles differ by only 15° 5° from the angles between the crystallographic axis (55.5°, −173.3, and 115.0°), similar to the orientation in NeHAO. As extensively discussed by Hendrich et al. (15), the lack of precise coincidence of the two sets of axes is, among other reasons, due to the relative orientation of the imidazole planes of the coordinating His residues, being co-planar or not, and/or to the orientation of the imidazole planes relative to the heme meso-position. A peculiar feature of the EPR spectra of KsHOX was the presence of a high spin heme signal at $g_x$ ~ 6 (Table 3 and Figs. 8A and B and 9A). Although prominent, the intensity of the signal varied between different enzyme preparations, never taking integral values. This may suggest that the residual high spin heme signal originated from a heme residue that either was pentacoordinated or was hexacoordinated with water (or OH$^-$), a straightforward candidate being heme 4 lacking the covalent tyrosine binding with the neighboring subunit as is the case for monomeric KsHOX.

From UV-visible light optical changes, it was deduced that incubation of KsHOX with its substrate hydroxylamine caused the two-electron reduction of the c-type heme centers (12). This result was corroborated by EPR spectroscopy. After incubation with NH$_2$OH, hemes 3 and 5 were each reduced by ~60%, whereas one signal had completely disappeared from the EPR spectrum of KsHOX, which is attributed to heme 2 (Figs. 8B and 9A). This observation suggested that hemes 3 and 5 have very similar high midpoint redox potentials. The same conclusion regarding the redox properties of the heme couple 3 and 5 was reached in the study on NeHAO (14). After reduction, the magnetic coupling strength between hemes 3 and 5 increased somewhat to 0.035 cm$^{-1}$, and the g values of the remaining oxidized species differed slightly from those in the oxidized enzyme (Table 3). In addition, some resonances sharpened slightly (e.g. the $g_x$ attributed to hemes 7 and 8) or broadened slightly (e.g. the $g_y$ of heme 1) (Table 3 and Fig. 9B). Appar-
ently, the magnetic properties of hemes 1, 7, and 8 were affected by the reduction of a common neighboring heme, which is heme 2 (Fig. 1D), justifying their assignments. We may note that the same EPR spectral changes were found when KsHOX was incubated with hydrazine, giving further support to our previous finding (12) that binding of hydrazine is associated with the two-electron reduction of His/His-ligated c-type hemes.

Overall, the X-band EPR spectra of KsHOX could be accurately simulated using the parameters listed in Table 3 (Fig. 9, A and B). Using these same parameters, the simulated Q-band spectrum of as-isolated KsHOX agreed well with the experimental spectrum (Fig. 9C), supporting the multicomponent analysis.

EPR Spectroscopy of KsHDH—At first glance, the EPR spectrum of as-isolated KsHDH (Fig. 8C) was quite different from the one of KsHOX (Fig. 8A). In addition, analysis of the EPR spectrum of oxidized KsHDH was complicated by the fact that the majority of the EPR signals appeared to be broadened due to magnetic interactions. This was, for example, evidenced by the absence of separate \( g_x \) resonances (~1.5 < \( g_x < 1.2 \)) or sharp \( g_y \) resonances (at \( g \approx 2 \)–2.2) in oxidized KsHDH. Due to the broad signals obscuring baseline sloping, quantitation of the EPR spectra was less accurate. Consequently, quantitation of the normal mode EPR spectrum of KsHDH (265 \( \mu \)M hemes as determined optically) showed a larger spread (210 ± 50 \( \mu \)M).

Nevertheless, the latter value was in agreement with the presence of six to eight EPR-visible hemes; as before, the coupled hemes 6 and 4 (P460) were not detected. Only after reduction by hydrazine did a magnetically isolated low spin heme center become visible (\( g_x = 1.51, g_y = 2.256, \) and \( g_z = 3.10 \)), which amounted to ~0.45 hemes/monomer as the result of a partial reduction (Table 3 and Figs. 8 (C and D) and 10 (A and B)). The main difference between the spectra of oxidized KsHDH and oxidized KsHOX/NeHAO was that an adequate simulation of the spectrum of KsHDH required that not only hemes 3 and 5 but also the signal allocated to hemes 7 and 8 contribute as a magnetically coupled pair of hemes. Other special features were the presence of a trio of relatively isotropic low spin heme species and of a highly anisotropic low spin heme with \( g_{xx,zz} = 1.6, 2.05, 3.14 \) (Table 3 and Fig. 11 (A and B)). In total, the trio accounted for one heme per monomer at which individual species integrated to approximately one-third of an integral value.

In KsHDH, the magnetic coupling strength between hemes 3 and 5 (0.074 and 0.087 \( \text{cm}^{-1} \) after reduction) and the Euler angles were found to be nearly the same as in NeHAO (Table 3). For heme pair 7 and 8, the magnetic coupling was weaker (0.035 \( \text{cm}^{-1} \)), and, importantly, the Euler angles were different (45, 180, and 87°), suggesting different relative heme orientations (Table 3). In the KsHOX structure, the difference in orientations of hemes 7 and 8 is evident (Fig. 1D). In the structure of KsHDH, which has yet to be resolved, these Euler angles could...
be different, resulting in the observed spin-spin interaction between hemes 7 and 8. Thus, spin species 2b in KsHDH (Table 3) is derived from two interacting, neighboring hemes, the only candidates being hemes 7 and 8. Consequently, the corresponding spin species in the KsHOX and NeHAO EPR spectra should comprise these hemes 7 and 8 as well, although they are not spin-coupled in these proteins.

Incubation of KsHDH (~35 μM) with excess hydrazine (0.1 mM) resulted in a dramatic change of the EPR spectrum (Figs. 8D and 10 (A and B)). In total, each enzyme monomer became reduced by 2.95 ± 0.4 electrons according to the EPR simulations, whereas a value of 3.2 ± 0.5 electrons was determined from direct double integration of the experimental spectra. More specifically, heme pair 3 and 5 became reduced by ~70%, and the exchange splitting increased to 0.087 cm⁻¹ (Table 3). Reduction of the protein by hydrazine led to the complete disappearance of the trio of relatively isotropic low spin heme species, whereas the highly anisotropic low spin heme became reduced by 55%. Upon reduction, the broad resonances of the latter sharpened and shifted. In contrast, heme pair 7 and 8 remained fully oxidized. Assuming as before that heme 2 has the highest redox potential, this heme 2 was allocated to the trio of relatively isotropic low spin heme species. It is tempting to speculate that within the homotrimeric protein, the three hemes 2 take somewhat different positions, directing electron transfer within and between the subunits and resulting in the three slightly different EPR signals observed. The assignment would leave the isolated highly anisotropic low spin signal to heme 1. It should be noted that treatment of KsHDH with various concentrations of hydrazine resulted in the reduction of 3–4 His/His-ligated hemes as deduced from the UV-visible spectral changes (Fig. 6). The three-electron reduction calculated from the EPR spectra agrees with this.

Except for the signals discussed above, the EPR spectrum of KsHDH showed the presence of two more signals in minor amounts (Table 3). The first one concerned a high spin heme (<0.02 per monomer), highly resembling the one observed in KsHOX. As in the latter protein, this high spin heme could be the result of small contamination of monomeric KsHDH. In the g = 2 region, the EPR spectrum of oxidized KsHDH showed an unusual signal (Fig. 10C) characterized by a peak-to-peak width of 36 Gauss, no resolved hyperfine structure, and an apparent g value of 2.009. Upon reduction by hydrazine, this signal decreased nearly 3-fold in intensity. The signal relaxed rapidly. Its low amount (0.005 spins/monomer) might point to an impurity (but not a high-potential iron-sulfur protein (HIPIP) 4Fe-4S center giving its overall line shape). However, its reduction by hydrazine suggested it to be a component of KsHDH. If so, the signal might derive from a tyrosine radical, more specifically from Tyr-536’ crosslinked to heme 4. This would explain both its rapid relaxation and unusual line shape affected by the magnetism of the S = 5/2 Fe³⁺ of heme 4.

Overall, the EPR spectra of as-isolated KsHDH, both in the absence and presence of its substrate hydrazine (Fig. 8, C and D), could be adequately simulated (Fig. 10, A and B) by the parameters listed in Table 3, as was the case with KsHOX. Although quite different at first appearance, our analyses showed that the EPR spectra of KsHDH were well described by the same framework as those of NeHAO and KsHOX (Table 3). This provided strong evidence that HDH indeed possessed the same architecture and arrangement of His/His-ligated c-type hemes as NeHAO and KsHDH. Again, differences in the EPR spectra were the likely result of only minute differences in the relative orientation of the two imidazole planes of the two histidines that coordinate the low spin hemes and of the precise location of these imidazole planes relative to, for example, the meso-positions of the porphyrin rings. These differences might affect the heme redox potentials, enabling a fine tuning of electron transfer in order to optimize catalytic activities.

Discussion

Here we purified and characterized KsHDH, the gene product of kustc0694. HDH is a key enzyme in anammox metabolism, which oxidizes the intermediate hydrazine to the end product N₂. Biophysical properties, detailed sequence analysis, and UV-visible and EPR spectral data strongly suggested that
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HDH and its close homologs in other anammox genomes were structurally quite similar or nearly identical to two also structurally highly related octaheme proteins, NeHAO and KsHOX. Both NeHAO and KsHOX are capable of hydrazine oxidation, but HDH was specific in this activity (Table 2). This raises the question of how HDH is tuned for its specific action. An answer to this question may be found in differences in the way HDH is structured, in the way hydrazine becomes bound to its catalytic site, and in the way the four electrons and four protons derived from hydrazine oxidation (Equation 3) are abstracted and transferred during catalysis.

A special property of KsHDH was that it assembled into octamers of homotrimers (Table 1 and Figs. 3–5). Such association would require specific amino acid sequences that promote the interaction between the homotrimers. Interestingly, HDH members contained a conserved stretch of 15 amino acids in their C-terminal region, which was absent in HAO and HOX (supplemental Fig. S1). From the crystal structures of the latter two proteins, it is inferred that the stretch could be localized at the tip of each monomer, where it would be well situated to promote the interaction between two egg- or tulip-shaped, covalently bound homotrimer.

As-isolated, catalytically competent KsHDH reacted with hydrazine. This reaction was accompanied with reduction of His/His-ligated c-type hemes. In the case of KsHOX, incubation with hydrazine caused the reduction of two such hemes, whereas a dinitrogen species bound to the P$_{460}$ catalytic center was observed in its crystal structure soaked with hydrazine, which could be a diazene (HN=NH) derivative (Fig. 1B) (12).

Similarly, binding of hydrazine to the catalytic site, immediately followed by a two-electron oxidation would be the first step in the HDH catalytic cycle proposed in Fig. 11. Analogously, hydroxylamine oxidation by KsHOX and NeHAO is initiated by the concomitant binding of this substrate and its two-electron oxidation, yielding a nitroxyln (HN=O) species in a [FeNO]$^+$ configuration as an intermediate (22, 23). An intrinsic problem of such two-electron oxidations is that the first heme that accepts these electrons (heme 6; see Fig. 1D) can carry only one electron at a time. Here, the tyrosine attached to the catalytic heme 4 could function by temporarily storing the second electron as put forward previously (12). This would explain the presence of the P$_{460}$ group in this type of HAO-like proteins that favor oxidative reactions (24).

KsHDH was incapable of hydroxylamine oxidation (Table 2), and this compound did not interact with the fully oxidized, as-isolated enzyme. We note that the redox potential of the P$_{460}$ Prosthetic group (actually P$_{473}$) in KsHDH ($E^\circ_m = -420$ mV; Fig. 8C) was 120–150 mV lower than in KsHOX ($E^\circ_m = -300$ mV) (12) and in NeHAO ($E^\circ_m = -260$ mV) (13, 14). The lower $E^\circ_m$ value for the catalytic heme 4 in KsHDH could arise from the cysteine predicted to be in the second coordination sphere of the catalytic heme 4 in KsHDH, but not the HAO and HOX, enzymes (Fig. 1, B and C). If present as a thiolate anion, this cysteine would significantly affect the electronic properties of heme 4, including the bond length with the proximal histidine and, as a result thereof, the redox potential of this heme (25). In the case of KsHDH, the electron-donating (reducing) power of NH$_2$OH might become insufficient for bond formation with the ferric P$_{460}$, preventing turnover of this compound. However, hydroxylamine and also NO were strong competitive inhibitors, and these compounds oxidized reduced KsHDH instantaneously, indicating an interaction with ferrous heme P$_{460}$ as a catalytic cycle intermediate (Fig. 11). In hindsight, the serendipitous finding that hydrazine is an intermediate in the anammox process (26) was apparently due to the inhibition of the HDH reaction by hydroxylamine added to the assays.

In anammox bacteria, hydrazine is the ultimate electron donor in energy conservation, whereas hydroxylamine is probably a side product of the preceding hydrazine synthase reaction (5, 6). Apparently, these bacteria have evolved dedicated enzymes for hydrazine and hydroxylamine metabolism, at which KsHOX recycles hydroxylamine efficiently into hydrazine synthase substrates (NO and three electrons; see Reaction 2). The above oxidation reactions require the well timed transfer of single electrons to an exit heme 1. Branched electron transfer chains in homotrimeric NeHAO and KsHOX enable electron export both to heme 1 of the subunit where substrate oxidation takes place and to heme 1 of a neighboring subunit (Fig. 1D). Possibly, HDH kinetically fine tunes this electron transfer by (reversible) octamerization and by a rearrangement of the electron transfer circuit, including the modified binding of heme 3 by a CX$_5$CH motif.

HDHs from anammox bacteria, which belong to the HAO-like octaheme proteins, are responsible for the production of a significant amount of N$_2$ that is released into the atmosphere. Besides HDH (kustc0694 and kustd1340) and KsHOX (kustd1061), the genome of K. stuttgartiensis harbors seven more HAO paralogs with as yet unknown functions. Most of these are also found in other anammox genomes (6), whereas genes coding for a plethora of related HAO-like proteins can be found in genomic databases (e.g. see Ref. 24). Together, these HAO-like proteins may represent an environmentally important but still poorly understood potential of enzyme activities, all taking advantage of common structural properties but each one tuned for a specific function by subtle modifications of their catalytic sites and multiheme electron transport module.

Experimental Procedures

Protein Purification—HPLC grade chemicals were purchased from Mallinckrodt Baker; all other chemicals were of the highest grade available. All purification steps took place under air and, apart from FPLC, at 4 °C. KsHDH was isolated from K. stuttgartiensis harbors seven more HAO paralogs with as yet unknown functions. Most of these are also found in other anammox genomes (6), whereas genes coding for a plethora of related HAO-like proteins can be found in genomic databases (e.g. see Ref. 24). Together, these HAO-like proteins may represent an environmentally important but still poorly understood potential of enzyme activities, all taking advantage of common structural properties but each one tuned for a specific function by subtle modifications of their catalytic sites and multiheme electron transport module.

Experimental Procedures

Protein Purification—HPLC grade chemicals were purchased from Mallinckrodt Baker; all other chemicals were of the highest grade available. All purification steps took place under air and, apart from FPLC, at 4 °C. KsHDH was isolated from K. stuttgartiensis grown continuously as planktonic cells in a 10-liter membrane bioreactor (5). Routine purification included the following. Biomass (4 liters, $A_{600} = 1.0–1.2$) was harvested by centrifugation at 8,000 × g for 15 min. Harvested cells were resuspended in 5 ml of 20 mM potassium phosphate buffer (pH 7). Cells were broken by three subsequent passages through a French pressure cell operating at 138 megapascals. The cell lysate was incubated with 1% (w/v) sodium deoxycholate on a rotating incubator (20 rpm) for 1 h. After incubation, cell debris was removed by centrifugation (3,000 × g, 15 min), and the obtained supernatant was subjected to ultracentrifugation (150,000 × g, 1 h) in a Discovery 100 ultracentrifuge equipped with a T-1270 rotor (Sorvall, Newtown, CT) to pellet cell membranes. The intense dark red supernatant after ultra-
centrifugation constituted the cell-free extract. An Äkta purifier (GE Healthcare) was used for the subsequent FPLC steps. In both steps, the columns were eluted at a flow rate of 2 ml min⁻¹; the eluate was monitored at 280 nm and collected in 2-ml fractions. To purify HDH, cell-free extract was loaded onto a 30-ml column packed with Q Sepharose XL (GE Healthcare), which was equilibrated with 2 column volumes of 20 mM Tris-HCl buffer (pH 8). After application of the sample, the column was washed with 2 volumes of this Tris-HCl buffer to remove unbound proteins. Hereafter, the sample was eluted isocratically in three steps (200, 400, and 1,000 mM NaCl in the 20 mM Tris-HCl buffer, pH 8). Hydrazide dehydrogenase activity was recovered in the 400 mM NaCl step. Three column fractions (6 ml) that were devoid of hydroxylamine oxidase activity were collected and desalted in the above-mentioned potassium phosphate buffer using spin filters (cut-off 100 kDa; Vivaspin 20, Sartorius Stedim Biotech, Aubagne, France) to prepare the sample for the following FPLC step. In this step, the desalted fractions were applied to a 10-ml ceramic hydroxyapatite (Bio-Rad) column, which was equilibrated with 5 column volumes of 20 mM potassium phosphate buffer, pH 7. Following sample loading, HDH activity was eluted from the column as a broad symmetrical peak upon isocratic elution with 20 mM potassium phosphate buffer (pH 7) for approximately three column volumes. Active fractions were collected and concentrated using spin filters as described above. The purity and identity of the fractions were determined by non-denaturing PAGE and SDS-PAGE as well as MALDI-TOF analysis. Enzyme preparations kept in ice were immediately used for further experiments or rapidly frozen in liquid nitrogen for storage. By this procedure, the enzyme was purified 15-fold with a yield of 21% (Table 4).

Both the recovery and purification factor were likely to be underestimates, because the cell-free extract contained a substantial amount of hydrazine dehydrogenase activity derived from hydroxylamine oxidase (KsHOX, kustc1061). KsHOX was rapidly frozen in liquid nitrogen for storage. By this procedure, the column had been pre-equilibrated with gel filtration buffer (150 mM KCl, 50 mM HEPES-KOH, pH 7.5) at room temperature. 40-µl samples with A₁ cm

280 nm = 5 were injected using an autosampler. Static light scattering analysis was performed in line using a DAWN HELEOS multiangle scattered light photometer (Wyatt Technology Corp., Santa Barbara, CA) (laser wavelength λ = 658 nm). Data were processed using the ASTRA software. MALS analysis accounted for any shape.

**TEM**—Samples of native KsHOX and KsHDH in 25 mM HEPES-KOH, pH 7.5, 25 mM KCl were used directly for negative staining. In addition, a KsHDH sample was subjected to glutaraldehyde cross-linking as follows. KsHDH was diluted to A₁ cm

280 nm ~ 0.5 in gel filtration buffer (150 mM KCl, 50 mM HEPES-KOH, pH 7.5), and 0.1% (v/v; ~10 mM) glutaraldehyde was added. After incubation at 37 °C, the reaction was quenched by the addition of Tris-HCl, pH 8.0, to a final concentration of 167 mM. Afterward, the sample was concentrated and applied to a Superose 6 (10/300 GL) gel filtration column (GE Healthcare, Uppsala, Sweden) at 0.5 ml min⁻¹ and 4 °C. Fractions of the peak eluting at 11±1 ml were pooled, concentrated, and buffer-exchanged to 25 mM HEPES-KOH, pH 7.5, 25 mM KCl using a 100 kDa molecular mass cut-off Amicon ultrafiltration device (Millipore Bioscience, Schwabach, Germany).

For negative staining, 10 µl of a dilute protein solution (A₁ cm

280 nm ~ 1) in 25 mM HEPES/KOH, pH 7.5, 25 mM KCl was pipetted onto a glow-discharged copper grid with carbon-coated Formvar (Plano GmbH, Wetzlar, Germany) and removed after 1 min by gentle blotting from one side with filter paper. The grid was then immediately rinsed with 100 µl of water, blotted again, and treated with 10 µl of 2% (w/v) uranyl acetate solution for 1 min. After removing the staining solution thoroughly by blotting with filter paper, the grid was dried in air for 5–10 min and then inserted in the vacuum port of a FEI Tecnai G2 T20 Twin transmission electron microscope (FEI, Eindhoven, The Netherlands) running at 200 kV accelerating voltage. Electron micrographs were recorded with an FEI Eagle 4K HS, 200-kV CCD camera.

**Enzyme Assays**—Reaction mixtures (1 ml) in 20 mM potassium phosphate buffer (pH 7) were prepared in 2-ml glass cuvettes (1-cm path length) and contained 50 µM bovine heart cytochrome c (Sigma-Aldrich) and enzyme as required. Cuvettes were sealed with rubber stoppers. Reactions were
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monitored at 550 nm in a Cary 50 (Agilent, Santa Clara, CA) spectrophotometer at 37 °C. After recording the absorbance at 550 nm for 1 min, the reaction was started by the addition of a helium-sparged hydrazinium sulfate (Merck, Darmstadt, Germany) stock solution or other substrates to be tested. Inhibitors (1, 10, 50, or 100 μM hydroxylamine; 3, 6, or 9 μM NO) were included before the addition of the enzyme. Hydroxylamine was added from a helium-sparged hydroxylammonium chloride (Merck) stock solution. An NO-containing stock solution (0.9 mM) was prepared by sparging anoxic potassium phosphate buffer with an NO-He gas mixture (1:1, v/v) for 10 min. This stock solution was injected to the assays with a 50-μl gas syringe. To minimize the headspace, the volume of reaction mixtures containing NO was increased to 1.5 ml. Reactions in the reductive direction were assayed by adding partially reduced cytochrome c as an electron donor. Reduction was achieved by mixing cytochrome c (50 μM) with 20 μM ascorbic acid, yielding 40 μM reduced and 10 μM oxidized cytochrome c, respectively. Reaction rates were determined from the initial linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer respectively. Reaction rates were determined from the initial linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package. Quantification was performed on the linear part of the reaction curves using the spectrophotometer (Cary) software package.
Other Analytical Methods—Proteins were identified from polyacrylamide gels by MALDI-TOF mass spectrometry on a Bruker Biflex III mass spectrometer (Bruker Daltonik, Bremen, Germany) operated in reflectron mode. Samples for MALDI-TOF were prepared as described previously (33). Each spectrum (500–4,000 m/z) was analyzed using the Mascot Peptide Mass Fingerprint (Matrix Science, London, UK) against the K. stuttgartiensis database, allowing methionine oxidation as variable modification, 0.2-Da peptide tolerance, and at most one miscleavage. The molecular mass of the KsHDH holoenzyme was determined on the same Bruker Biflex III mass spectrometer, operating in the linear mode.

The formation of gaseous nitrogen compounds and their masses (NO, NO2, N2O, N2) were quantified by gas chromatography (Agilent 6890 equipped with a Poropak Q column at 80°C; Agilent, Santa Clara, CA) combined with a mass spectrometer (Agilent 5975 equipped with a Poropak Q column at 80°C; Agilent, Santa Clara, CA) and its substrate soaks (12) were retrieved from the Protein Data Bank (accession numbers PDB 4N4J–4N4M). The crystal structure of NeHAO (10, 11) was obtained at 408 nm (ε408 = 700 mm–1 cm–1). The value is presented in terms of subunit concentration (67.2 kDa).

Protein Sequence and Structure Analyses—Protein sequences were retrieved from genomic databases (K. stuttgartiensis, PRJNA16685; Jettenia caeni strain KSU-1, PRIDA163683, PRJDB86; Scalindua profunda, taxon object IDs 2017108002 and 2022004002 at the Joint Genome Institute; Brocadia sinica, PRJDB103). Multiple-amino acid sequence alignments were made with the ClustalW2 program at the EMBL-EBI web site. Atomic coordinates and structure factors of KsHOX (kustc1061) and its substrate soaks (12) were retrieved from the Protein Data Bank (PDB) (accession numbers PDB 4N4J–4N4M). The crystal structure of NeHAO (10, 11) was obtained from the PDB under accession numbers 1FGI, 4FAS, 4N4N, and 4N4O.


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References
SUPPLEMENTAL INFORMATION

Characterization of anammox hydrazine dehydrogenase, a key N₂-producing enzyme in the global nitrogen cycle

Wouter Maalcke, Joachim Reimann, Simon de Vries, Julea Butt, Andreas Dietl, Nardy Kip, Ulrike Mersdorf, Thomas Barends, Mike Jetten, Jan Keltjens and Boran Kartal

SUPPLEMENTAL FIGURE LEGEND

FIGURE S1. **Multiple protein sequence alignment of hydrazine dehydrogenase and other HAO-like octaheme proteins.** The Figure shows the alignment of hydrazine dehydrogenase kustc0694 from *Kuenenia stuttgartiensis* (KsHDH), its close homologs and paralogs from other anammox bacteria, hydroxylamine oxidase (KsHOX, kustc1061) from *K. stuttgartiensis*, and hydroxylamine oxidoreductase from *N. europaea* (Neuro_HAO). Predicted N-terminal signal sequences are printed in blue. These sequences are absent in proteins with known crystal structures (KsHOX, NeHAO). The CX₂CH heme binding motifs of the heme c molecules are highlighted red (white letters); primed numbers represent the histidines proximal ligands to the respective heme c molecules as deduced from the KsHOX and NeHAO crystal structures. Heme 3 in KsHDH and its homologs having an unusual CX₂CH binding motif is highlighted pink (white lettering). The tyrosine involved in the covalent binding to the catalytic heme 4 (P₄₆₀) in KsHOX (Tyr-491) and NeHAO is highlighted purple (white letters). The aspartate, histidine and tyrosine residues (Tyr-358 in NeHAO) near the catalytic site are highlighted blue (white letters). Note that the tyrosine is apparently conserved in all proteins. However, in KsHOX the tyrosine is moved away from the catalytic site by several Ångströms by a two-amino acid contraction and it is replaced at that position by a methionine (Met-323) (12). This same contraction is found in the HDH proteins. A 15-amino acid sequence in the C-terminal part, which is specific for KsHDH and its close homologs is printed in bold. Peptide sequences identified for KsHDH by MALDI-TOF analyses are underlined. Protein identifiers and abbreviations represent the following: kust, *K. stuttgartiensis*; KSU-1_HzoB (ZP_10100863) and KSU-1_HzoB (ZP_10098714), hydrazine dehydrogenase/ oxidase (Hzo) from anammox enrichment culture KSU-1; BROSI, anammox bacterium *Brocadia sinica* (PRJDB103); scal, HAOs from anammox bacterium *Scalindua profunda* (Taxon Object IDs 2017108002 and 2022004002 at JGI).
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