Purification, Characterization, and Preliminary Crystallographic Study of Copper-Containing Nitrous Oxide Reductase from *Pseudomonas nautica* 617

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ABSTRACT: The aerobic purification of *Pseudomonas nautica* 617 nitrous oxide reductase yielded two forms of the enzyme exhibiting different chromatographic behaviors. The protein contains six copper atoms per monomer, arranged in two centers named Cu<sub>A</sub> and Cu<sub>Z</sub>. Cu<sub>Z</sub> could be neither oxidized nor further reduced under our experimental conditions, and exhibits a 4-line EPR spectrum (<i<g<sub>x</sub>= 2.015, A<sub>x</sub>= 1.5 mT, g<sub>y</sub>= 2.071, A<sub>y</sub>= 2 mT, g<sub>z</sub>= 2.138, A<sub>z</sub>= 7 mT>) and a strong absorption at ~640 nm. Cu<sub>A</sub> can be stabilized in a reduced EPR-silent state and in an oxidized state with a typical 7-line EPR spectrum (<i<g<sub>x</sub>= 2.021, A<sub>x</sub>= 0 mT, g<sub>y</sub>= 2.178, A<sub>y</sub>= 4 mT>) and absorption bands at 480, 540, and ~800 nm. The difference between the two purified forms of nitrous oxide reductase is interpreted as a difference in the oxidation state of the Cu<sub>A</sub> center. In form A, Cu<sub>A</sub> is predominantly oxidized (<i>S</i> = ½, Cu<sup>1.5+</sup>–Cu<sup>1.5+</sup>), while in form B it is mostly in the one-electron reduced state (<i>S</i> = 0, Cu<sup>1+</sup>–Cu<sup>1+</sup>). In both forms, Cu<sub>Z</sub> remains reduced (<i>S</i> = ½). Complete crystallographic data at 2.4 Å indicate that Cu<sub>A</sub> is a binuclear site (similar to the site found in cytochrome c oxidase) and Cu<sub>Z</sub> is a novel tetracopper cluster [Brown, K., et al. (2000) Nat. Struct. Biol. (in press)]. The complete amino acid sequence of the enzyme was determined and comparisons made with sequences of other nitrous oxide reductases, emphasizing the coordination of the centers. A 10.3 kDa peptide copurified with both forms of nitrous oxide reductase shows strong homology with proteins of the heat-shock GroES chaperonin family.

Denitrification constitutes one of the main branches of the global nitrogen cycle. In this process, nitrate is reduced to dinitrogen through the series of reactions outlined below.

\[
2\text{NO}_3^- \rightarrow 2\text{NO}_2^- \rightarrow 2\text{NO} \rightarrow \text{N}_2 \text{O} \rightarrow \text{N}_2
\]

This respiratory process is coupled to ATP generation. The enzymes involved in denitrification are usually synthesized under anaerobic conditions in the presence of nitrogen oxides, although denitrification can also occur in the presence of oxygen (1, 2). Many denitrifying bacteria grow at the expense of N2O as the sole electron acceptor for the oxidation of organic compounds (3). Nitrous oxide reductase (N2OR)<sup>1</sup> catalyzes the conversion of N2O to N2, the last step of the complete denitrification pathway in denitrifying bacteria. The reaction catalyzed by N2OR is

\[
\text{N}_2\text{O} + 2\text{H}^+ + 2\text{e}^- \rightarrow \text{N}_2 + \text{H}_2\text{O}
\]

N2OR has been purified from various microorganisms: *Pseudomonas (Ps.) stutzeri* (formerly *Ps. perfectomarina*) (4), *Rhodobacter capsulatus* (formerly *Rhodopseudomonas capsulata*) (5), *Rhodobacter sphaeroides* f. sp. *denitrificans* (6), *Paracoccus denitrificans* (7), *Wolinella succinogenes* (8), *Achromobacter cycloclastes* (9), *Ps. aeruginosa* (10), *Thiosphaera pantotropha* (11), *Thiobacillus denitrificans* (12), and *Achromobacter xylosidans* (13). While these N2O-ORs are soluble proteins, a membrane-bound enzyme was detected in the gliding soil bacterium *Flexibacter canadensis*.

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(14). With the exception of N₂ORs from *Rhodobacter sphaeroides*, which contains zinc and nickel (6), and from *Wolinella succinogenes*, which is a heme–copper protein (8), all other purified N₂ORs contain only copper cofactors. Copper N₂ORs exist in several forms, distinguished by their redox and spectroscopic properties and by their enzymatic activity, for which numbers and trivial names are commonly used as reviewed in (3).

The most widely used model regards the copper sites of N₂OR as two binuclear centers: Cuₐ and Cu₂Z. Cuₐ, responsible for the electron transfer from an external donor to the catalytic center (15), is spectroscopically and structurally similar to the Cuₐ center of cytochrome c oxidase (COX) (16–20). The structure of this latter center was determined (21–25), and its characteristics are reported in a recent review by Beinert (26). Cu₂Z is believed to be the catalytic center where N₂O reduction occurs, and its binuclear nature was assumed from chemical analysis and spectroscopic studies [see (3) for a review]. Another, more recent, model considers Cuₐ and Cu₂Z as variants of the same binuclear copper center, the catalytic center being undetectable by visible or EPR spectroscopies (27). This model, based on results from UV–visible, MCD, and EPR spectroscopies, makes way for speculation as to the possible nature of the catalytic center.

In this manuscript we report the purification, crystallization, and biochemical and spectroscopic characterization of two forms of N₂OR isolated from the halophilic denitrifier *P. nautica* 617. These two forms, called A and B, differ in their spectroscopic and redox properties. The gene sequence coding for N₂OR and the deduced amino acid sequence are also presented. The X-ray data have demonstrated that the two copper centers of N₂OR have different features: Cuₐ is a binuclear copper center similar to the one found in COX, and Cu₂Z, the putative catalytic site, is a novel tetranuclear copper center never before found in biological systems (28). Based on the data presented in this manuscript, we propose a model in which the two purified forms A and B correspond to two different oxidation states of Cuₐ, with Cu₂Z remaining redox-inactive under our experimental conditions.

**EXPERIMENTAL PROCEDURES**

**Growth of *P. nautica* and N₂OR Purification.** The reclassification of the type strain of *Pseudomonas nautica* as *Marinobacter hydrocarbonoclasticus* has been proposed (29). However, strain 617 awaits classification, and we will continue to refer to it as *P. nautica* 617 to avoid confusion. Three 200 L batches of *P. nautica* 617 (Pasteur Institute Collection, ref. *P. nautica* no. 617/1.85) were grown under denitrifying conditions in artificial seawater at 30 °C with 10 mM nitrate as electron acceptor (30). Yeast extract (0.1%) and lactate (1%) were used as carbon and energy sources. The medium was supplemented with a separately sterilized Starkey oligoelement solution (31) (0.2 mL/L of culture). Cells were harvested in the late exponential phase of growth by centrifugation. Approximately 600 g of cells (wet paste) was resuspended in 100 mM Tris-HCl, pH 7.0, and broken in a French press. The crude extract was centrifuged at 8000g for 20 min and twice at 125000g for 1 h to remove cell debris and membranes. The resulting supernatant was used for N₂OR purification. To avoid freeze–thaw cycles in intermediate purification fractions, all steps were carried out nonstop at 4 °C until the protein was considered to be pure. The soluble extract was applied to an anion exchange DEAE-Biogel column (4.5 × 40 cm, Bio-Rad) equilibrated with 10 mM Tris-HCl, pH 7.6. A 10–400 mM Tris-HCl, pH 7.6, linear gradient was applied, and N₂OR-containing fraction eluted at 100–125 mM Tris-HCl. This fraction was loaded onto a Superdex 75 column for gel filtration (2.6 × 56 cm, Pharmacia) run with 300 mM Tris-HCl, pH 7.6. After dialysis, the N₂OR-containing fraction was applied to a Source 15Q anion exchange column (1.6 × 25 cm, Pharmacia) equilibrated with 10 mM Tris-HCl, pH 7.6. A 10–500 mM Tris-HCl, pH 7.6, gradient was applied, and forms A and B of N₂OR were eluted successively between 160 and 200 mM Tris-HCl, pH 7.6. The final yield was approximately 135 mg of total enzyme (A + B).

**Analytical Methods.** SDS–PAGE electrophoresis was performed with the discontinuous buffer system of Laemmli at 12.5% (32). Native molecular weight determination by gel filtration was performed on a prepacked Superdex 200 HR 10/30 column (Pharmacia). Elution buffer was 50 mM potassium phosphate buffer, pH 7.0, containing 150 mM NaCl. The flow rate was 0.5 mL/min, and 200 µL of protein or standard was injected. The molecular mass standards were ferritin (440 kDa, Pharmacia), catalase (232 kDa, Pharmacia), aldolase (158 kDa, Pharmacia), bovine serum albumin (67 kDa, Sigma), ovalbumin (43 kDa, Sigma), chymotrypsinogen A (25 kDa, Pharmacia), and ribonuclease A (13.7 kDa, Pharmacia). Protein concentration was determined by the method of Lowry et al. (33). Copper content was routinely estimated using the method described by Poillon and Dawson (34). Inductively coupled plasma emission analysis (ICP), performed at the University of Georgia, was also used for copper content determination as well as to investigate the presence of other metals in the enzyme. N₂OR activity was determined spectrophotometrically, at 25 °C, by following the oxidation of reduced methyl viologen at 600 nm under anaerobic conditions (35, 13). The stock solution of enzyme was 125–180 µM. The reaction was performed in 50 mM phosphate buffer, pH 7.1, and was initiated by addition of N₂O-saturated water. N₂OR specific activity is expressed as micromoles of N₂O reduced per minute per milligram of enzyme. Poly(vinylidene difluoride) protein sequencing membranes were used for electroblotting, and N-terminal sequencing was performed on a 476 Sequenator (Perkin-Elmer, Applied Biosystems Division, Foster City, CA). C-terminal sequence analysis was performed on a Procise 494C sequencer (Perkin-Elmer, Applied Biosystems Division) using a chemical procedure slightly modified from the one described in (36). The alkylated thiohydantoins were identified on-line by reversed-phase analysis on a 140C microgradient system, from the same firm, using a linear gradient made from 3.5% tetrahydrofuran/35 mM sodium acetate, pH 3.8, as solvent A, and 100% acetonitrile as solvent B. Prior to sequence analysis, the protein was adsorbed on a ProSorb sample preparation cartridge and, after several subsequent washes with water, treated with phenylisothiocyanate to modify the lysine chains (37). Electrospray mass spectrometry was performed on a Bio-Q quadrupole mass spectrometer equipped with an electrospray ionization source (Micromass, Altrincham, U.K.) as used in (38).
Molecular Biology. *Ps. nautica* cells were grown as previously described. Genomic DNA was isolated using a standard phenol–chloroform extraction protocol (39, 40). The full N2OR encoding gene was amplified by PCR using two sets of oligonucleotide primers. In the first reaction, a degenerate primer, 5’-GARTACTAGTGCTTCGG-3’, representing the gene sequence of the first residues of the N-terminus was used as the forward primer, while an analogous degenerate primer was designed according to homologies with other N2OR amino acid sequences (5’-GATGTCGATCARYTGGTCRTTC-3’) and used as the reverse primer, resulting in the amplification of a DNA fragment of approximately 1300 bp. To amplify the remaining 550 bp of the N2OR gene, a second PCR was performed with the following oligonucleotide primers: 5’-GTCCAAGTT-TTCGAAAGACCG-3’ (forward) and 5’-CATSCGRCCSAC-CATYTCCAT-3’ (reverse). In both PCR reactions, 50 ng of genomic DNA, 50 pmol of each primer, 2.5 mM of each dNTP, and 5 units of *taq* DNA polymerase in 1x PCR buffer (Amersham Pharmacia Biotech) were used in a total volume of 50 μL. The reactions were performed in a thermal cycler (Biometra) using the following program: 1 cycle at 94 °C for 5 min, followed by 35 cycles of 0.5 min at 94 °C, 1.5 min at 56 °C, 2 min at 72 °C, and a final extension step for 10 min at 72 °C. The PCR products were analyzed by electrophoresis in a 0.8% agarose gel in the presence of ethidium bromide using 1x TAE buffer. The fragments of interest were purified from contaminants by gel extraction using the QiAquick Gel Extraction Kit (QIAGEN) and subsequently ligated into the commercial pGEM-Teasy vector from Promega according to the manufacturer’s instructions. The resulting DNA plasmids were transformed into Epicurian Coli XL1-Blue competent cells (Stratagene) which were grown overnight in Luria Broth medium with 100 μg/mL ampicillin at 37 °C. Positive recombinant clones were screened after their white color and confirmed by restriction digestion analysis of plasmidic DNA with NotI. Plasmidic DNA from single transformants was isolated using the Wizard Plus Minipreps DNA Purification System (Promega) and sequenced at the Mayo Clinic Molecular Biology Facility.

Protein Crystallization. Purified N2OR (form B) from *Ps. nautica* was concentrated to 5 mg/mL in a buffer solution of 100 mM Tris-HCl, pH 7.6. Crystals were obtained by the technique of vapor diffusion at 20 °C with hanging drops of 3 μL of protein solution, mixed with 3 μL of reservoir solution containing typically 18% PEG 4000, 0.1 M BICINE, 0.6 M NaCl, 15% 2-propanol, 10 mM spermine-4HCl at a final pH of 9.5. X-ray diffraction data were collected on BM14 (ESRF). The three-dimensional structure solved by the MAD (multiple wavelength anomalous dispersion) technique at 2.4 Å will be described elsewhere (28).

Spectroscopic Measurements. UV–visible spectra of enzyme solutions were recorded on either a Shimadzu UV-160A or a Shimadzu UV-265FS spectrophotometer. The crystal visible spectra were recorded with the ORIEL microspectrophotometer on ID09 beamline (ESRF), using unpolarized light. The crystals were taken from the mother liquor, transferred for a few seconds to the cryo-cooling solution (10% ethylene glycol), mounted on a cryo-loop, and flash-frozen to 100 K in a cold stream of nitrogen gas (Oxford Cryosystems). Care was taken to record optical densities in the linearity range (OD < 2.5). The dark current corresponding to electronic noise was measured, and the baseline correction was performed with the buffer in which the crystals were soaked. EPR measurements were performed on an X-band Bruker EMX spectrometer equipped with a dual-mode cavity (model ER4116DM) and an Oxford Instruments continuous-flow cryostat. EPR spectra were simulated using the program WINEPR–SimFonia (Bruker). Spin quantitations were made by double integration of a signal obtained under nonsaturating conditions and comparison to a Cu(II)–EDTA standard.

Redox Titrations. Redox titrations of N2OR were performed anaerobically in 100 mM Tris-HCl, pH 7.6, at 20 °C. A mixture of 16 mediators covering redox potentials ranging from −550 to +430 mV, each at the final concentration of 2 μM, was used to facilitate electron transfer during the oxidation and reduction cycles. The mediators used were as follows: triquat, methyl viologen, neutral red, safranin, phensafarin, antraquinone-2-sulfonic acid, 2-hydroxy-1,4-naphthoquinone, phenazine, indigo carmine, indigo tetrasulfonate, duroquinone, 5-hydroxy-1,4-naphthoquinone, 1,4-naphthoquinone, 1,2-naphthoquinone, 2,6-dichlorophenol indophenol, and potassium ferricyanide. The protein solution (174 μM) was reduced by addition of aliquots of sodium dithionite (in 100 mM Tris-HCl, pH 9.0) or oxidized with potassium ferricyanide (in 100 mM Tris-HCl, pH 7.6). The potential was monitored by using a platinum electrode as a measuring electrode and a Ag/AgCl electrode as a reference electrode. Calibration was performed with a saturated solution of quinhydrone at pH 7.0. Samples were collected under anaerobic conditions at different potentials, and the spin concentration of each spectrum was plotted against the measured redox potential.

![Figure 1: UV–visible spectra of *Ps. nautica* N2OR. (A) Form A; (B) form B; and (C) dithionite-reduced enzyme.](image-url)
RESULTS

N\textsubscript{2} OR Purification. After the first purification step, N\textsubscript{2}-OR exhibited a purple color, with an intense absorption peak at \~540 nm. However, after the second anion exchange step, a separation was achieved between a predominantly purple form (called A) and a blue-colored form (called B), with different visible spectra (Figure 1A,B). Form B eluted at a slightly higher ionic strength than form A. Both forms were active and exhibited similar specific activities: 55 ± 6 and 23 ± 14 \textmu{}mol of N\textsubscript{2}O reduced min\textsuperscript{-1} (mg of enzyme)\textsuperscript{-1} for forms A and B, respectively. However, maximum specific activity was only achieved with a preincubation of the enzyme with dithionite-reduced methyl viologen. N\textsubscript{2} OR was found to be sensitive to dilution. Upon dilution of N\textsubscript{2} OR stock solution from 10 to 1 mg/mL, the specific activity decreased with time in an exponential manner to reach the value of approximately 1 \textmu{}mol of N\textsubscript{2}O reduced min\textsuperscript{-1} (mg of enzyme)\textsuperscript{-1} 3 h after dilution. The two forms were

F IGURE 2: DNA and deduced amino acid sequences of Ps. nautica N\textsubscript{2} OR. PCR oligonucleotide primers are indicated in boxes, and pairs of primers are identified by the type of arrow above the box.
**FIGURE 3:** Amino acid sequence comparison of N₂ORs. Comparison was performed using the ClustalX program (43). Labels are as follows: Ps.n., *Pseudomonas nautica* (this work); Ps.s., *Pseudomonas stutzeri* (NCBI sequence number AAA25907); Ps.a., *Pseudomonas aeruginosa* (CAA46381); A.c., *Achromobacter cycloclastes* (CAA75425); Pa.d., *Paracoccus denitrificans* (CAA52798); S.m., *Sinorhizobium meliloti* (ACC44023); and R.e., *Ralstonia eutropha* (CAA46383).

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[Figure showing amino acid sequence comparison]

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indistinguishable by SDS–PAGE, yielding a band at ~65 kDa and one at ~16 kDa (see below). Mass spectrometric analysis gave a mass of 65 382 ± 34 Da for N2OR. The molecular mass of the monomer calculated from the amino acid sequence is 65 373 Da. Native molecular mass determination by gel filtration yielded a value of 120 kDa, indicative of a dimer.

Copurification of a 10.3 kDa Peptide with N2OR. Both forms of N2OR consistently copurified with a peptide. Its apparent molecular mass, as estimated by SDS–PAGE, was 16 kDa. Electrospray ionization mass spectrometry provided a value of 10 289.9 ± 0.4 kDa. This peptide could not be separated from N2OR by anion exchange, hydrophobic interaction, adsorption, or gel filtration chromatographies, indicating a strong association between the two proteins, but precipitated upon N2OR crystallization. Therefore, the spectroscopic and activity data presented in this article were obtained with samples containing this peptide. Amino-terminal sequencing of the 10.3 kDa peptide, following electroblotting, yielded the sequence MKIRPLHDGV-VIKKKNFVIKDASIIGLP. Sequence comparison revealed high homology with proteins of the GroES heat shock chaperonin family (41, 42).

Metal Content of N2OR. Total copper determination was performed on various batches of both forms of N2OR. The number of copper atoms found was 10.7 ± 1.7 per dimer, a value slightly higher than those previously reported for this enzyme (3). No significant amounts of other metals were found by plasma emission analysis.

Gene and Protein Sequence. In Figure 2, the partial DNA sequence and the deduced amino acid sequence of Ps. nautica N2OR are presented. Figure 3 shows the comparison of the complete N2OR amino acid sequence (obtained by N-terminal sequencing, C-terminal sequencing, and translation of the PCR amplification product) with other known N2OR sequences. Ligands of CuZ and CuA centers, as determined by X-ray analysis (28), are marked in black and gray, respectively. These amino acids are conserved in all sequences shown. While the coordination sphere of CuZ was not predicted or observed before, the ligands of CuA were successfully predicted by Zumft (3). Trp563 (identified by an arrow in Figure 3) binds CuA through the carbonyl group, which is in accordance with the fact that this residue is not totally conserved (one sequence out of seven shows a histidine instead of a tryptophan).

Crystallographic Data. Blue crystals (Figure 4) grew in about 30 days in the space group P61, with unit cell dimensions a = b = 211 Å, c = 166 Å, and 3 dimers per asymmetric unit (55% solvent content, Vm = 2.73). The refined three-dimensional model has an R-factor of 21.6% (Rfree = 25.0%) and contains a dimer of 580 × 2 residues, 12 copper atoms, and 728 water molecules. The CuA center is composed of two copper atoms and is similar to the CuA site in subunit II of COX (22–25). The remaining four copper ions are assembled in the CuZ center, a novel cluster shaped as a distorted tetrahedron with seven His and three hydroxide ion ligands (Figure 5). A more complete description of the structure is presented elsewhere (28).

Spectroscopic Characterization of Forms A and B of N2OR. The absorbance spectra of forms A and B of Ps. nautica N2OR are shown in Figure 1. Both forms of the enzyme show absorbance bands in the ranges 540–560 and 620–640 nm, but the peak around 540 nm is more intense in form A. In addition, the spectrum of form A shows absorption features at ~480 and ~800 nm. UV/Visible spectroscopy showed that reduction of both forms of the enzyme by sodium dithionite resulted in a decrease in absorption at 540 nm and the bleaching of the 480 and 800 nm features, with a single remaining peak around 640 nm (e640 nm = 7.1 ± 0.3 mM−1 cm−1 for the protein dimer) (Figure 1C). This corresponds to the so-called form III of the enzyme, obtained by the addition of excess reductant to native N2OR (4, 44, 45). Oxidation of either form of N2OR with potassium ferricyanide resulted in a significant increase in absorption at 480, 540, and 800 nm (data not shown). The UV–visible spectrum of the crystals was characteristic of an almost fully reduced enzyme.

Both enzyme forms were also studied by EPR spectroscopy in the as-purified (Figure 6), dithionite-reduced, and ferricyanide-oxidized states (Figure 7A,B). The spectrum of the fully reduced enzyme could be simulated by assuming a single species with a 4-line hyperfine splitting in the g region (g∥ = 2.015, A∥ = 1.5 mT, g⊥ = 2.071, A⊥ = 2 mT, g∥ = 2.138, A∥ = 7 mT) and line width values of 5.5, 5.0, and 5.5 mT for g∥, g⊥, and A∥, respectively. As demonstrated by the oxidized minus reduced difference spectrum (Figure 7C), the fully oxidized spectrum could be simulated by adding, in equal proportions, the simulated signal of the fully reduced enzyme and a CuA-like signal exhibiting a 7-line hyperfine splitting in the g region (g∥ = 2.021, A∥ = 4 mT, g∥ = 2.178, A∥ = 4 mT) with line width values of 8.0, 8.0, and 3.0 mT for g∥, g⊥, and A∥, respectively. The 4-line signal was assigned to CuZ by spectral deconvolution of N2OR spectra at different oxidation states (Figure 7). The spectra of forms A and B, as-purified, were simulated in the same manner assuming that both forms are intermediate redox states between the fully oxidized and the fully reduced form (Figure 6). The proportions of CuA (oxidized) and CuZ signals used to simulate the various spectra are summarized in Table 1.

The amount of EPR-detectable copper spins was also determined for both forms of the enzyme in different oxidation states (Table 1). For both forms, the EPR spectrum of oxidized N2OR quantified approximately 2 spins per monomer, whereas the fully reduced form yielded ap-
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proximately 1 spin per monomer. The as-purified form A contains about 90% of the spin intensity when compared to the ferricyanide-oxidized state. In the case of form B, only 60% of the spin intensity is present.

**Redox Behavior of N\textsubscript{2}OR.** Redox titrations of N\textsubscript{2}OR were followed by EPR spectroscopy. Both forms of the enzyme gave similar results. The data obtained were fitted to a Nernst equation for a one-electron process and a midpoint redox potential of approximately +260 mV (data not shown). No EPR-silent state was ever detected.

**DISCUSSION**

The unveiling of the tetranuclear nature of Cu\textsubscript{Z} by X-ray crystallography sheds a new light on UV--visible and EPR studies of N\textsubscript{2}OR. Our spectroscopic data on the three oxidation states (oxidized, as-purified, and reduced) of forms A and B of *Ps. nautica* N\textsubscript{2}OR can only be explained if the two copper centers, Cu\textsubscript{A} and Cu\textsubscript{Z}, contribute independently to the spectra. We suggest that the main differences observed between the visible spectra of the two forms correspond to a difference in the oxidation state of Cu\textsubscript{A}. However, we cannot at present exclude more substantial structural varia-

**FIGURE 5:** Schematic representation of the Cu\textsubscript{A} binuclear center and Cu\textsubscript{Z} tetranuclear center of *Ps. nautica* N\textsubscript{2}OR. Distances were determined from crystallographic data.

**FIGURE 6:** EPR spectra of forms A and B of *Ps. nautica* N\textsubscript{2}OR, as-purified. (A) Form A, 215 \mu M enzyme in 160 mM Tris-HCl buffer, pH 7.6, 39 K, microwave power 1.59 mW, modulation 4.05 Gpp, microwave frequency 9.49 GHz. (B) Form B, 267 \mu M enzyme in 160 mM Tris-HCl buffer, pH 7.6, 35 K, microwave power 5.04 mW, modulation 1.05 Gpp, microwave frequency 9.49 GHz. One scan was run for all spectra, and the gain was 7.1 \times 10^4. The simulations combine a 4-line and a 7-line signal according to the proportions reported in Table 1.

**FIGURE 7:** EPR spectra of oxidized and reduced N\textsubscript{2}OR from *Ps. nautica*. (A) Oxidized enzyme; (B) reduced enzyme; (C) difference spectrum “A minus B”. Experimental conditions were 267 \mu M enzyme in 160 mM Tris-HCl buffer, pH 7.6, 35 K, microwave power 5.05 mW, modulation 1.05 Gpp, microwave frequency 9.49 GHz, 1 scan, gain 7.1 \times 10^4.
This page contains a table and text discussing the oxidation states of copper centers in nitrogenase. The table is labeled as Table 1: Proportions of Oxidized CuA Signal (7 Lines) and CuZ Signal (4 Lines) in EPR Spectra. The text continues with a discussion of the spectroscopic characteristics of the CuZ center, including its oxidation states and the implications for the catalytic mechanism of nitrogenase.

The text mentions that the CuZ center, which is responsible for the 7-line signal observed in the EPR spectra of oxidized and as-purified samples, must have access to three different redox states: oxidized (Cu1.5+), reduced (Cu1+), and an intermediate state. The tetranuclear CuZ center, which is responsible for the 4-line signal observed in all spectra, is also discussed.

The text also mentions that the two forms of the enzyme were purified under aerobic conditions, and there was no indication of the existence of a less active “pink” form, that had been described in other organisms. The spectroscopic characteristics of the CuZ center were compared with those of other N2 ORs, and the results suggest that more than two electrons per dimer were necessary to produce an EPR-active form of the catalytic center. This becomes easier to understand, considering the tetranuclearity of the catalytic center, but would suggest that this center may undergo more than one redox transition at this moment. There is no chemical model that would help us to understand the mechanisms taking place in such a structure.

The tetranuclear structure of the catalytic site was quite unexpected. There were earlier reports that more than two electrons per dimer were necessary to produce an EPR-active form of the catalytic center. This becomes easier to understand, considering the tetranuclearity of the catalytic center, but would suggest that this center may undergo more than one redox transition. At this moment, there is no chemical model that would help us to understand the mechanisms taking place in such a structure.

The tetranuclear structure of the catalytic site may be related to the necessity of activating the rather chemically stable and unreactive N2O molecule.

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References


