

Characterization of Cytochrome 579, an Unusual Cytochrome Isolated from an Iron-Oxidizing Microbial Community[∇]

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A novel, soluble cytochrome with an unusual visible spectral signature at 579 nm (Cyt₅₇₉) has been characterized after isolation from several different microbial biofilms collected in an extremely acidic ecosystem. Previous proteogenomic studies of an Fe(II)-oxidizing community indicated that this abundant red cytochrome could be extracted from the biofilms with dilute sulfuric acid. Here, we found that the Fe(II)-dependent reduction of Cyt₅₇₉ was thermodynamically favorable at a pH of >3, raising the possibility that Cyt₅₇₉ acts as an accessory protein for electron transfer. The results of transmission electron microscopy of immunogold-labeled biofilm indicated that Cyt₅₇₉ is localized near the bacterial cell surface, consistent with periplasmic localization. The results of further protein analysis of Cyt₅₇₉, using preparative chromatofocusing and sodium dodecyl sulfate-polyacrylamide gel electrophoresis, revealed three forms of the protein that correspond to different N-terminal truncations of the amino acid sequence. The results of intact-protein analysis corroborated the posttranslational modifications of these forms and identified a genomically uncharacterized Cyt₅₇₉ variant. Homology modeling was used to predict the overall cytochrome structure and heme binding site; the positions of nine amino acid substitutions found in three Cyt₅₇₉ variants all map to the surface of the protein and away from the heme group. Based on this detailed characterization of Cyt₅₇₉, we propose that Cyt₅₇₉ acts as an electron transfer protein, shuttling electrons derived from Fe(II) oxidation to support critical metabolic functions in the acidophilic microbial community.

Biological oxidation of Fe(II) by acidophilic microbial communities found in mines with exposed pyrite ore accelerates the dissolution of FeS₂ and acidification of the mine water, resulting in acid mine drainage (AMD), a global environmental problem (8). One of the most-intensively studied AMD sites is the Richmond Mine at Iron Mountain, CA, where copious biofilm communities are found in extremely low-pH (0.5 to 1.0) solutions (2). Most of these communities are pink biofilms dominated by *Leptospirillum* group II bacteria, with lower abundances of *Leptospirillum* group III bacteria and several archaeal species (4). A *Leptospirillum* group II bacterium-dominated biofilm was collected at the “5-way” site at the Richmond Mine (11) and analyzed by metagenomic sequencing (5-way community genomics data set [24]). Proteomic characterization by mass spectrometry (MS) of a similar biofilm isolated from the “AB end” site of the Richmond Mine identified an abundant extracellular protein from *Leptospirillum* group II bacteria, encoded by gene 20 on sequencing scaffold 20 (gene 14-20), that has a CXXCH heme binding motif common to *c*-type cytochromes but otherwise insignificant sequence similarity to known proteins (17). The results of gel electrophoresis and N-terminal sequencing confirmed that this protein contained heme and was abundant in the extracellular

fraction. The first 40 amino acids deduced from the environmental genomic sequence were nearly identical to the N-terminal sequence deduced for the Fe(II)-oxidizing cytochrome 579 (Cyt₅₇₉) purified from an isolate of *Leptospirillum ferriphilum*. The reduction potential of *L. ferriphilum* Cyt₅₇₉ was estimated to be ≥660 mV, and the cytochrome was fully reduced in the presence of excess Fe(II) at pH 2.0 (17). A cytochrome with very similar spectral and pH-dependent-redox properties had also been isolated from *Leptospirillum ferrooxidans* (10). The ability of *L. ferriphilum* and *L. ferrooxidans* Cyt₅₇₉ to oxidize Fe(II) at low pH led to the hypothesis that this novel cytochrome identified in the biofilm acted as the primary Fe(II) oxidant for *Leptospirillum* group II bacteria.

Here we report the purification and characterization of Cyt₅₇₉ from a *Leptospirillum* group II bacterium-dominated biofilm collected at Richmond Mine. The results of detailed biochemical and MS studies of Cyt₅₇₉ from the biofilm suggest that it functions as a periplasmic electron transfer protein.

MATERIALS AND METHODS

Isolation of extracellular proteins. Richmond Mine biofilm samples were collected in 50-ml conical Falcon tubes (BD Biosciences, San Jose, CA), frozen at the site on dry ice, and later stored at –80°C. Biofilm samples were collected from the AB end site (near the junction of the “A drift” and B drift) in January 2004; from the C drift site (15 m beyond the AMD dam) in November 2005; and from the “UBA” site (in the A drift) in November 2005. A map describing the field site can be found in online supplementary information of reference 11. To obtain the extracellular fraction, the biofilm was thawed, suspended in 110 ml 0.2 M H₂SO₄ (pH 1.1), and homogenized in a glass tube by using several vigorous strokes of a tight-fitting, round, glass pestle. The resulting homogeneous cell suspension was stirred for 2 h at 4°C and then centrifuged at 24,000 × g for 12

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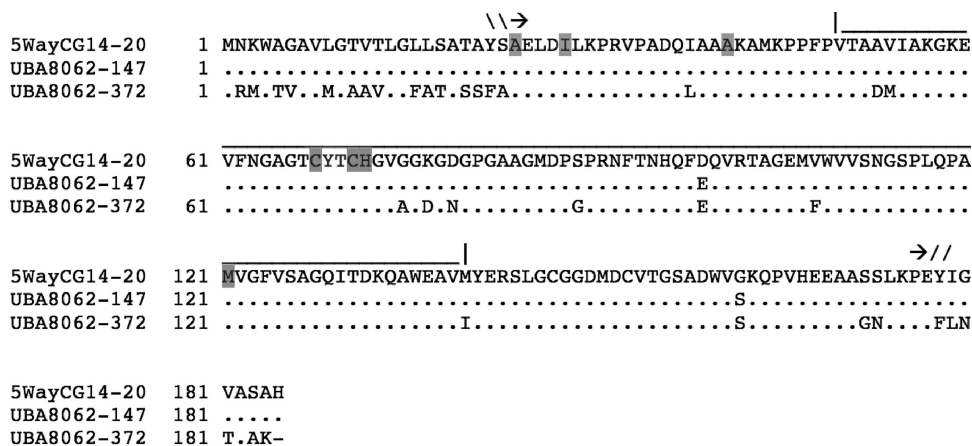


FIG. 1. Alignment of amino acid sequences of Cyt₅₇₉ from the 5-way and UBA genomic datasets. One gene from the 5-way site (5WayCG14-20) and two from the UBA site (UBA8062-147 and UBA8062-372) encode variants of Cyt₅₇₉. Three N-terminal start sites observed by sequencing isolated proteins are indicated in gray, as are the predicted heme binding residues Cys68, Cys71, His72, and Met121 (see model in Fig. 8B). The line above the alignment indicates that portion of Cyt₅₇₉ used for structural modeling. Arrows indicate the N-terminal signal cleavage site and the observed C terminus.

min. The supernatant is the extracellular fraction used for cytochrome purification. For proteomic analysis, proteins from a 10-ml sample of the extracellular fraction of the biofilm from the C drift were precipitated with 10% trichloroacetic acid and the precipitate was collected by centrifugation, rinsed twice with cold methanol, and air dried.

Purification of Cyt₅₇₉. Proteins in the extracellular fraction (150 ml) were precipitated with (NH₄)₂SO₄ and redissolved in ~5 ml sample buffer (SB) containing 20 mM H₂SO₄ and 100 mM NH₄(SO₄)₂ at pH 2.2. A light red precipitate at 45% NH₄(SO₄)₂ saturation was gelatinous, indicating the presence of exopolysaccharides. A deeper red precipitate at 95% NH₄(SO₄)₂ contained 75 to 80% of the protein found in the extracellular fraction. This precipitate was dialyzed for 16 h at 4°C against 1 liter SB. The dialysate was loaded onto an SP-Sepharose FF column (5 ml) preequilibrated in SB. The column was washed with 2 column volumes of SB, and the red fraction (9 ml; 4 mg total protein) eluted with 100 mM sodium acetate (NaOAc), pH 5.0. Visible spectroscopy indicated that the pH 5.0 fraction was highly enriched in Cyt₅₇₉. The remaining protein was removed with a 0 to 2 M NaCl gradient (30 ml) in pH 5.0 buffer. Between 1.2 M and 2.0 M NaCl, light yellow fractions (3 ml each; 2 mg total protein) eluted that had visible spectra consistent with the presence of *c*-type cytochromes (α -band at 552 nm for reduced samples).

Immunogold labeling of Cyt₅₇₉ and transmission electron microscopy (TEM) of biofilms. Polyclonal antibodies were produced in rabbits (Covance, Denver, PA) by using the cation-exchange fraction of Cyt₅₇₉ as the antigen. Prior to immunization, the antigen was concentrated by using MicroCon spin filters (10-kDa-molecular-mass cutoff; Millipore, Billerica, MA) and resuspended in phosphate-buffered saline. Immunoblotting of a biofilm lysate indicated a high specificity of the antibody preparation for Cyt₅₇₉ (data not shown). Antibodies were purified from serum by using a Melon gel antibody purification kit (Pierce, Rockville, IL).

A biofilm sample was frozen under high pressure (Bal-tec HPM 010) and freeze substituted in 0.2% glutaraldehyde and 0.1% uranyl acetate in acetone. The sample was then rinsed in acetone and embedded in LR White resin. Microtomed sections (~70 nm thick) were mounted on carbon-coated, Formvar film-covered nickel grids and blocked with bovine serum albumin and cold-water fish gelatin (Sigma Aldrich, St. Louis, MO). The anti-Cyt₅₇₉ antibody was used as the primary antibody, and goat anti-rabbit antibody conjugated with 10-nm gold particles was used as the secondary antibody. After being labeled, samples were fixed in 0.5% glutaraldehyde. Prior to analysis, all samples were stained with uranyl acetate and lead citrate. Samples were observed with an FEI Tecnai 12 TEM operated at 120 kV. Images were recorded on film, and the negatives were scanned and digitally processed to optimize contrast by using Adobe Photoshop.

Separation of forms of Cyt₅₇₉. The fraction enriched in Cyt₅₇₉ (3 mg) from the C drift biofilm in 100 mM NaOAc, pH 5.0, was concentrated (as described above) to ~1 ml and dialyzed for 16 h against 1 liter of 25 mM L-histidine-HCl, pH 6.2. The dialyzed Cyt₅₇₉ fraction was loaded onto a 1- by 30-cm chromatofocusing column (PBE 94 Polybuffer exchange; Amersham Biosciences, Piscataway, NJ) preequilibrated with 2 column volumes of pH 6.2 buffer and eluted with PBE 74

Polybuffer, pH 5.0. Two red fractions eluted from the column at pH 5.5 (0.3 mg) and pH 5.1 (1.0 mg), and a third fraction was eluted with 1 M NaCl in 100 mM NaOAc, pH 5.0 (1.5 mg).

pH-dependent Fe(II) oxidation of Cyt₅₇₉. The C drift site Cyt₅₇₉ fraction (1.5 mg/ml) was diluted 1:10 in 100 mM glycine-200 mM SO₄²⁻, pH 2.0, and oxidized with a small amount of Fe₂(SO₄)₃ in pH 2.0 buffer. The oxidized Cyt₅₇₉ was then diluted 1:10 further in buffer that contained 30 mM FeSO₄-200 mM SO₄²⁻ in a 1.5-ml quartz cuvette, and the visible spectrum was obtained after 1 min. Low-pH buffers (pH 1.2 to 4.0) were prepared according to the method described by Schnaitman et al. using glycine and β -alanine (19). The spectrum was retaken after 10 min to ensure that the reaction had reached equilibrium. In all cases, the reaction was >95% equilibrated after 1 min.

Protein MS. Intact-protein characterization was performed by high-resolution Fourier transform ion cyclotron resonance (FTICR)-MS analysis. All FTICR mass spectra were acquired with a Varian 9.4-Tesla HiRes electro-spray FTICR-MS. Micromolar solutions of the purified Cyt₅₇₉ proteins were prepared in 50:50 water-acetonitrile (with ~0.1% acetic acid added). Using a syringe pump (flow rate of 1.75 μ l/min), the analyte was directly infused into a Z-type electrospray ion source. After generation, ions were accumulated in an external hexapole for 1 s and then transferred into the high-vacuum region with a quadrupole lens system. Detection then followed in the cylindrical analyzer cell of the MS. Calibration of the MS was accomplished externally with the various charge states of the protein ubiquitin, resulting in a mass accuracy of plus or minus 3 to 5 ppm and mass resolutions of 50,000 to 160,000 Da (full width at half maximum), as previously described (7). Ion dissociation was accomplished by infrared multiphoton dissociation (IRMPD) with a Synrad carbon dioxide laser (75-W maximum power and 10.2- μ m wavelength). For this experiment, the desired parent ion was isolated by ejecting all other ions from the analyzer cell and dissociated with infrared laser irradiation (30% maximum laser power for 1.5 s), and the resulting fragment ions were measured at high resolution in the FTICR analyzer cell.

To verify amino acid differences in Cyt₅₇₉ variants, purified samples were denatured, reduced, and digested with trypsin (sequencing grade; Promega, Madison, WI). Peptides were analyzed by using one-dimensional liquid chromatography-tandem MS (LC-MS-MS) on a Thermo Fisher linear-trapping quadrupole instrument. All MS-MS spectra were searched with DBDigger (22) against a database of all proteins predicted by genomic sequencing of biofilm samples (15, 24), as well as all potential amino acid variants of Cyt₅₇₉. The output data files were then filtered and sorted with the DTA Select algorithm (21) using the following parameters: fully tryptic peptides only; delta correlation value of at least 0.08; cross-correlation scores of at least 25 (+1 ions), 30 (+2 ions), and 45 (+3 ions); and at least two unique peptides per protein.

Amino acid variants were also verified from crude extracellular fractions of biofilms from the A bend, C drift, and UBA sites. Extracellular proteins were denatured, reduced, trypsin digested, and analyzed by using two-dimensional LC-MS-MS on a linear-trapping quadrupole instrument as previously described

(11, 15, 17). The MS-MS spectra were searched and filtered by using the same method as described above for the purified protein.

Structural modeling of Cyt₅₇₉. For the best possible results of homology modeling, several different techniques were combined (9) with our high-throughput computational system, AS2TS (29). Pairwise sequence alignments using both Smith-Waterman (20) and FASTA (16) and multiple sequence alignments using PSI-BLAST (1) and CLUSTALW (23) were carried out. PSI-BLAST analyses were performed on the nonredundant set of protein sequences in the NCBI database, with an E-value threshold of 0.001. After five iterations on NR sequences, the final PSI-BLAST run was restricted to sequences corresponding to PDB structures.

Secondary structure predictions were tested by using PSIPRED (12) and PHD (18). Structural alignments between all identified templates and preliminary models were calculated by LGA (28), and these results were used to further guide the process of three-dimensional (3D) model construction. Regions of insertion-deletion and uncertain sequence-structure alignments were built as loops. These regions were modeled using LGA (28) by "grafting" in suitable fragments from related structures in PDB. Finally, SCWRL (5) was used to add coordinates for missing side chain atoms.

General methods. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed according to the method of Laemmli (14). The protein concentration was estimated according to the method of Bradford (6). Trypsin digestion and N-terminal sequencing of proteins were performed as described previously (17). Gel filtration was performed on a 1- by 30-cm Superdex 75 column (Amersham Biosciences, Piscataway, NJ) equilibrated with 100 mM NaOAc, pH 5.0, containing 150 mM NaCl. Bovine serum albumin (67 kDa), ovalbumin (43 kDa), chymotrypsin (25 kDa), and RNase A (13 kDa) were used as molecular-mass standards.

RESULTS

Environmental genomic data indicate three distinct Cyt₅₇₉ genes. In addition to the previous metagenomics data for the 5-way site, we examined a second genomic data set obtained from a biofilm collected at the UBA site, which was dominated by a *Leptospirillum* group II species closely related to the characterized species from the 5-way site (15). Two homologs of Cyt₅₇₉ were identified. One is encoded by gene 8062-147, with an amino acid sequence 99% identical to the amino acid sequence encoded by gene 14-20 from the AB end site; the amino acid sequence encoded by a paralog of this gene, 8062-372, is 83% identical to that encoded by 14-20 (Fig. 1).

Cyt₅₇₉ purification from biofilms. Cyt₅₇₉ was purified from the acidic wash of the C drift biofilm by using ammonium sulfate precipitation and cation-exchange chromatography at low pH. Visible spectroscopy of the deep red band that eluted at pH 5.0 confirmed the characteristic absorption peak at 579 nm, consistent with the assignment of this cytochrome as Cyt₅₇₉. Examination of the purified protein by circular dichroism (CD) spectroscopy indicated a structure that is 70% α -helical, 3% β -strand, 8% turn, and 20% disordered when compared with the structures indicated by reference CD spectra (data not shown). These results were distinctly different from those of similar analyses of a purified membrane cytochrome, Cyt₅₇₂, which consists largely of β -strands (11).

The visible spectrum of purified Cyt₅₇₉ oxidized with Fe(III) at pH 2.0 exhibited a Soret band at 427 nm. In addition, a weak absorption band at 695 nm characteristic of an axial methionine ligand was observed in concentrated solutions (>0.2 mM) of oxidized Cyt₅₇₉ (data not shown). Upon reduction of isolated Cyt₅₇₉ with 500 μ M sodium ascorbate, the Soret band shifted to 441 nm and β (539 nm) and α (579 nm) bands were observed (Fig. 2A). The Soret band of the reduced spectrum also had a distinct shoulder at 419 nm, a feature absent in the spectrum of reduced Cyt₅₇₉ isolated from *L. ferriphilum* (17).

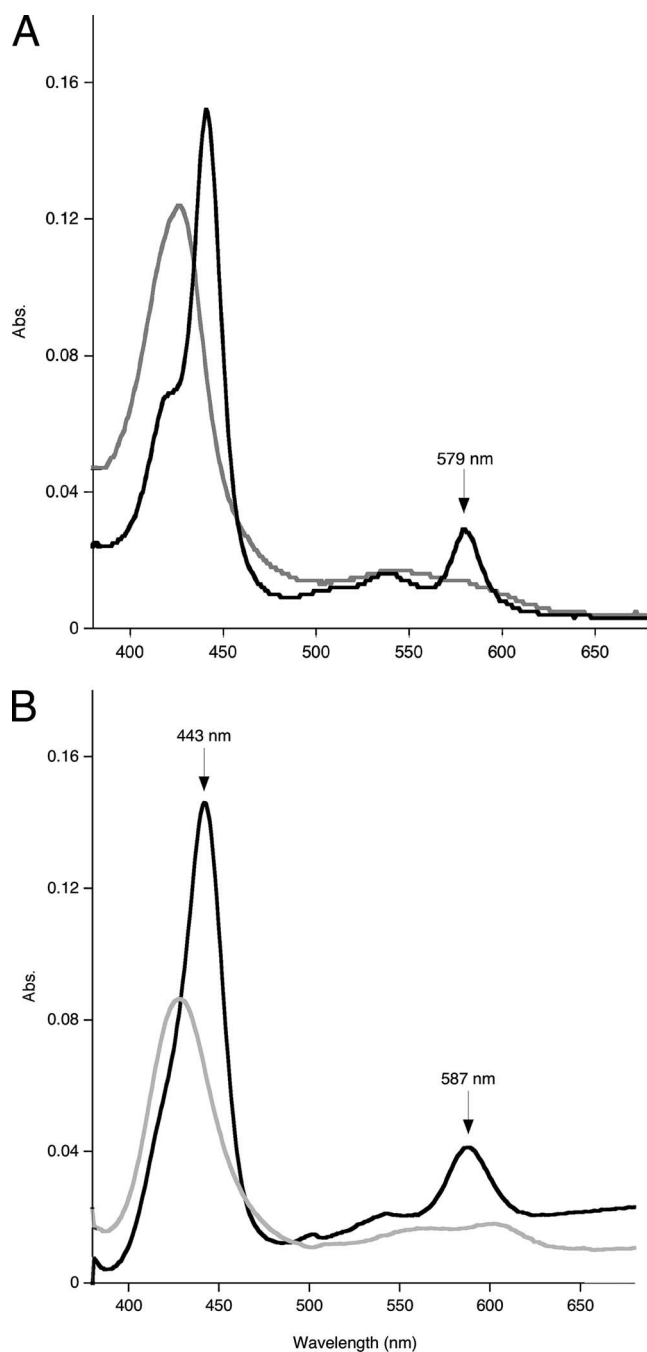


FIG. 2. Visible spectroscopy of Cyt₅₇₉. (A) Cyt₅₇₉ (0.015 mg/ml) isolated from the C drift biofilm in 100 mM glycine–200 mM SO₄²⁻, pH 2.0, was treated separately with 5 μ l of 10% Fe₂(SO₄)₃ [23% Fe(III)] (gray line) and 5 μ l of 1 mM sodium ascorbate (black line) in quartz cuvettes. The spectra were compared to those of the same solutions lacking Cyt₅₇₉. (B) Cyt₅₇₉ (1.5 mg/ml) was diluted by adding 50 μ l into 450 μ l of 0.2 M NaOH, 500 μ M sodium ferricyanide (gray line) or 2 mM sodium dithionite (black line), and 500 μ l of pyridine was added. Abs., absorbance.

The alkaline pyridine hemochrome spectrum had a Soret band at 443 nm and an α band at 587 nm (Fig. 2B). The results of SDS-PAGE of this fraction revealed two closely spaced protein bands at \sim 16 kDa (Fig. 3). Since MS proteomics of this frac-

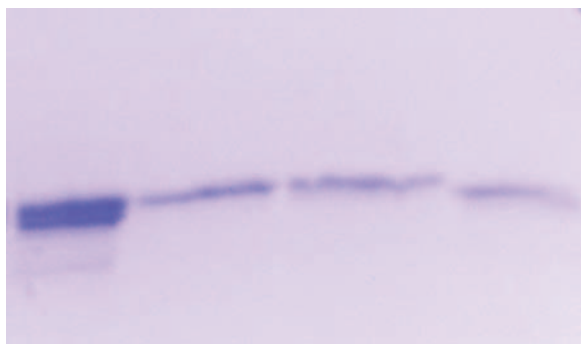


FIG. 3. Separation of different forms of Cyt₅₇₉. Chromatofocusing was used to fractionate a Cyt₅₇₉ sample, and proteins were analyzed on a 10 to 20% acrylamide gel using SDS-PAGE. First lane, C drift biofilm Cyt₅₇₉ fraction; second lane, C1 fraction; third lane, C2 fraction; and fourth lane, C3 fraction.

tion digested with trypsin indicated that >98% of the peptides were from Cyt₅₇₉, we concluded that two protein species represented different forms of Cyt₅₇₉ (data not shown). The results of Edman degradation identified two N-terminal sequences of Cyt₅₇₉ from the C drift biofilm (AELDILKPRV and ILKPRVPAD) that corresponded to the predicted amino acid sequence for all the Cyt₅₇₉ variants. Identical N-terminal sequences were obtained for a Cyt₅₇₉ preparation from the AB end site, the original proteomic sample (data not shown). The predicted N-terminal cleavage site to give the N-terminal sequence AELDILKPRV of signal peptidase I is between residues 23 and 24 for the variant sequences of Cyt₅₇₉. The Cyt₅₇₉ fraction eluted as a single band at an apparent molecular mass of 20 kDa from a Superdex 75 gel filtration column, consistent with the assignment of Cyt₅₇₉ as a monomer.

Cyt₅₇₉ was localized in *Leptospirillum* group II cells by TEM imaging of a thin section of the C drift biofilm that had been treated with polyclonal antibodies raised against Cyt₅₇₉ and a secondary gold-labeled antibody. Visualization of the antibody-treated thin section by TEM indicated that Cyt₅₇₉ was localized on the exterior of the *Leptospirillum* group II cells and was not distributed throughout the biofilm (Fig. 4). Since Cyt₅₇₉ contains a signal peptide and has no other hydrophobic regions in its amino acid sequence, we hypothesize that it is located in the periplasm of *Leptospirillum* group II cells.

Multiple forms of Cyt₅₇₉, separated by chromatofocusing. As mentioned above, the results of SDS-PAGE indicated that multiple forms of Cyt₅₇₉ were present in the purified fraction. The forms were too close in molecular weight to separate successfully by gel filtration. However, the forms of Cyt₅₇₉ were separated by using a preparative chromatofocusing column. Two red bands were eluted at pH 5.5 (C1) and pH 5.1 (C2) in a pH gradient of 6.2 to 5.0. The red fraction remaining on the column was eluted with pH 5.0 1 M NaCl buffer (C3). All three red fractions had nearly identical visible spectra; however, C1 had a Soret band for the oxidized Cyt₅₇₉ that was shifted to 425 nm, compared to 428 nm for C2 and C3. The results of SDS-PAGE of the separated Cyt₅₇₉ fractions confirmed that the pH 5.5 and pH 5.1 fractions represented the higher band in the crude Cyt₅₇₉ fraction, while the pH 5.0 1 M NaCl fraction represented the lower band (Fig. 3). N-terminal sequencing of

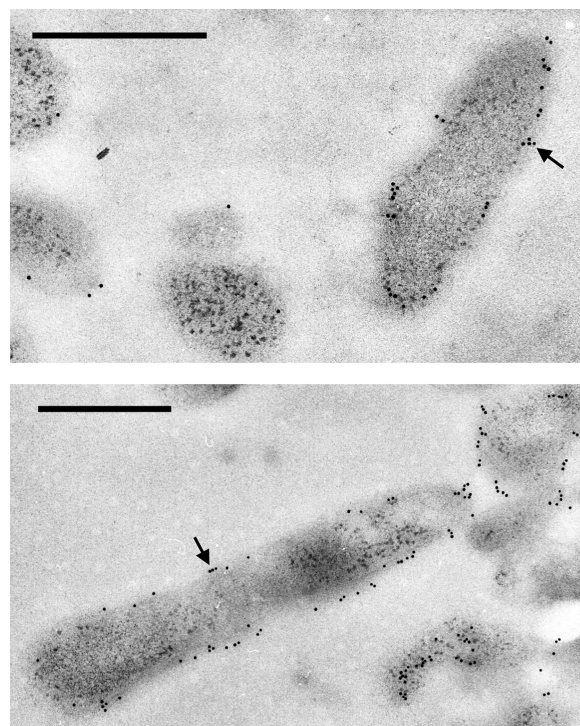


FIG. 4. TEM images of immunogold-labeled biofilm. Ultrathin section of biofilm showing Cyt₅₇₉ distribution on the edges of cells, possibly in the periplasm, and along the exterior of cells. Two representative fields are shown. Black arrows show gold particles; scale bars show 500 nm.

the individual bands revealed different start sites for each (Table 1). Cyt₅₇₉-specific polyclonal antibodies detected all three forms of the protein.

Mass spectrometry of separated Cyt₅₇₉ forms. To determine the accurate molecular masses and fragmentation products for the individual forms of C drift biofilm Cyt₅₇₉, the separated proteins were examined by FTICR-MS. The measured average molecular masses of the peaks in each Cyt₅₇₉ fraction are given in Table 1.

The amino acid sequences of each of these proteins were examined by MS-based fragmentation techniques. Isolation and IRMPD fragmentation of the (M + 13H)¹³⁺ ion for the 16,060-Da species revealed a variety of fragment ions, including a sequence tag, MVWVVSNGS, which is representative of the 8062-147-encoded sequence (Fig. 5, upper panel). The

TABLE 1. Forms of Cyt₅₇₉ identified by N-terminal sequence and intact mass

Cyt ₅₇₉ fraction	N-terminal sequence ^a	Avg molecular mass (Da) of species		Sequence start and end
		Major	Minor ^b	
C1	AELDILKPRV	16,058.97	16,045.60	AELD ... LKPE
C2	ILKPRVPAD	15,691.10	15,705.87	ILKP ... LKPE
C3	AKAMPPFV	14,316.57	14,332.22	AKAM ... LKPE
		14,571.63	—	LAAK ... LKPE

^a N-terminal sequences determined in each fraction by Edman degradation. Overlap in sequences is indicated in bold type.

^b —, species not detected by MS.

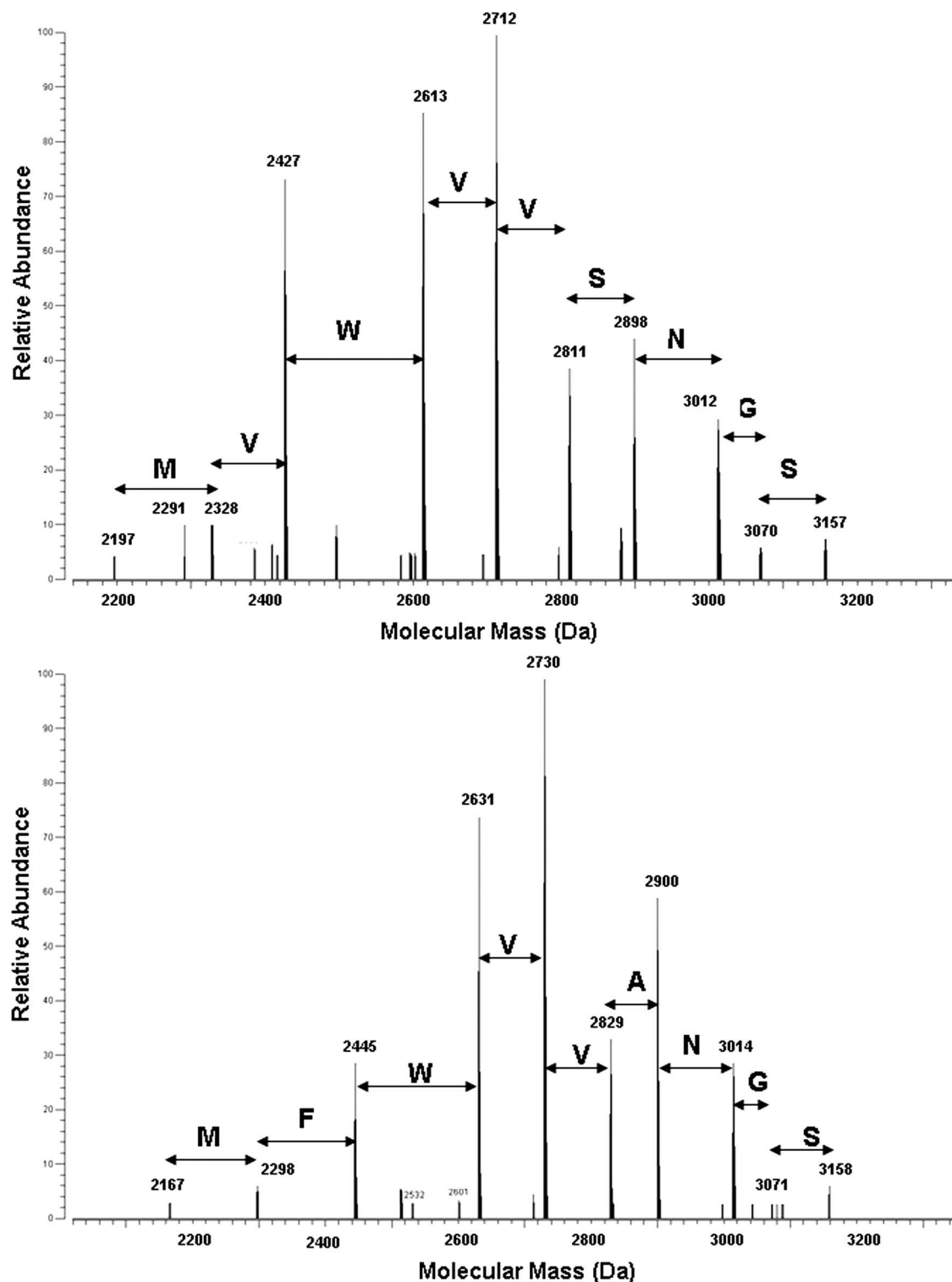


FIG. 5. Sequence tags of Cyt₅₇₉ obtained by IRMPD dissociation of the molecular species. The upper panel shows C1, 16,060 Da, and the lower panel shows C2, 15,690 Da. Amino acids are presented in the single-letter code above the spectra, and these indicate the difference in sequence between the two variants.

larger *b*-type fragment ions verified the presence of a truncated N terminus, supporting the experimentally determined N terminus, AELDILKPRV, and provided sequence information for the first 110 amino acids of the mature protein. Interest-

ingly, some of the smaller *y*-type fragment ions revealed truncation of the C terminus, indicating that this form of Cyt₅₇₉ corresponds to the sequence AELD . . . LKPE of the product of gene 8062-147 lacking the C-terminal eight amino acids. The

TABLE 2. Spectral counts obtained for MXWVVXN sequences of Cyt₅₇₉ from extracellular proteomes

Cyt ₅₇₉ gene	Sequence ^a	Spectral count ^b of sample from:		
		AB end	UBA	C drift
8062-147 (14-20)	MVWVVS N	28	48	57
8062-372	MFWVVS N	85	144	148
8062-372 C drift (S112A)	MFWV VAN	73	7	128

^a Spectral counts are derived from peptide R.TAGEMXWVVXNGSPLQPM VGFVSAGQITDK.Q. Amino acid substitutions are indicated in bold type.

^b Spectral counts refer to the total number of MS-MS spectra taken for the peptide as an indicator of overall abundance. Each count is the average of the results for three technical replicates.

observed mass is also consistent with removal of the heme group from the protein. However, the predicted average molecular mass of this species at 16,075.26 Da is 16 Da heavier than the measured value stated above. Further studies will determine if the discrepancy between the observed and calculated molecular masses of C1 is due to posttranslational modification or is an artifact of purification and mass spectrometry analysis.

The 15,691 Da (C2) and 14,317 Da (C3) species most closely corresponded to the 8062-372-encoded sequences ILKPR . . . LKPE and AKAMP . . . LKPE, respectively, based on the observed N-terminal sequences (see Table 1). The additional satellite peak at 14,572 Da in C3 was assigned to the sequence LAAAK . . . LKPE, although this N-terminal sequence was not observed by Edman degradation. Based on the measured molecular masses, the C-terminal truncations are identical in the C1 to C3 samples. In each of these cases, the predicted average molecular mass based on the predicted sequence of 8062-372 was 32 Da heavier than the observed mass. To determine if an amino acid variation could account for this difference, the relevant ions from these species were isolated and fragmented by IRMPD as described above. In both C2 and C3, the fragmentation revealed a sequence tag corresponding to the amino acid sequence MFWVVANGS (Fig. 5, lower panel). This sequence was identical to the sequence encoded by gene 8062-372, MFWVVSNGS, except for the Ser to Ala (S112A) variation (in bold), which accounts for a difference of 16 Da. The S112A variation was confirmed by PCR amplification and sequencing of the 8062-372 gene from the C drift biofilm (data not shown). The amino acid variant was also confirmed by the results of two-dimensional LC-MS-MS analyses of the crude extracellular fraction of the C drift biofilm (Table 1). The S112A variation accounts for the observation of the minor species at 15,706 Da (C2) and 14,332 Da (C3). The major species in C2 (15,691 Da) and C3 (14,317 Da and 14,572 Da) may arise from the same posttranslational process as the C1 species.

The S112A variation of the 8062-372 sequence was not found in the genomic data set for the 5-way or UBA genome. However, reexamination of the LC-MS-MS peptide data obtained for the AB end and UBA biofilm extracellular proteomes identified tryptic peptides corresponding to this sequence (Table 2).

Fe(II) oxidation by Cyt₅₇₉ forms. Previous work on Cyt₅₇₉ purified from *L. ferriphilum* and *L. ferrooxidans* demonstrated that

the oxidized form was fully reduced with excess Fe(II) at pH 2.0 (10, 17). When subjected to the same conditions as *L. ferriphilum* Cyt₅₇₉ (30 mM FeSO₄, 0.2 M total SO₄²⁻), the C drift Cyt₅₇₉ fraction before separation by chromatofocusing was ~30% reduced at pH 2.0, as determined by measuring the amplitude of the 579-nm band, in comparison to reduction with sodium ascorbate (data not shown). Studies of the pH dependence of Fe(II) oxidation by Cyt₅₇₉ indicated that minimal oxidation occurred at pH 1 to 2, but the equilibrium shifted to reduced Cyt₅₇₉ at a pH of >3, and Cyt₅₇₉ was almost fully reduced in the presence of 30 mM Fe(II) at pH 4 (Fig. 6). A nearly identical pH dependence of Fe(II) oxidation was observed for the crude Cyt₅₇₉ fraction obtained from the AB end biofilm, as well as the separated Cyt₅₇₉ forms (C1 to C3) obtained by chromatofocusing (data not shown).

Structural model of Cyt₅₇₉. Although no significant homology to Cyt₅₇₉ was found in protein database searches, over 100 candidate structural templates for modeling Cyt₅₇₉ were detected, ranging from 7% to 25% sequence identity. Secondary-structure predictions, along with high levels of structural similarities observed between the analyzed templates, narrowed the candidates to 25. An initial 3D model was constructed based on an alignment of the Cyt₅₇₉ sequence with that of cytochrome *c*₆, 1cyjA (Fig. 7A).

Based on calculated alignments to several structural templates (including RCSB Protein Data Bank accession no. 1cyj, 2dgc, 1w5c, 1ls9, 1h1o, 1jdl, 1kv9, and 1nir), the final 3D model was created, including the position of a *c*-type heme group from cytochrome *c*₆ (13). The heme in Cyt₅₇₉ is likely to be different, as discussed above, due to the unique spectral character of the cytochrome (see also Discussion). This model was

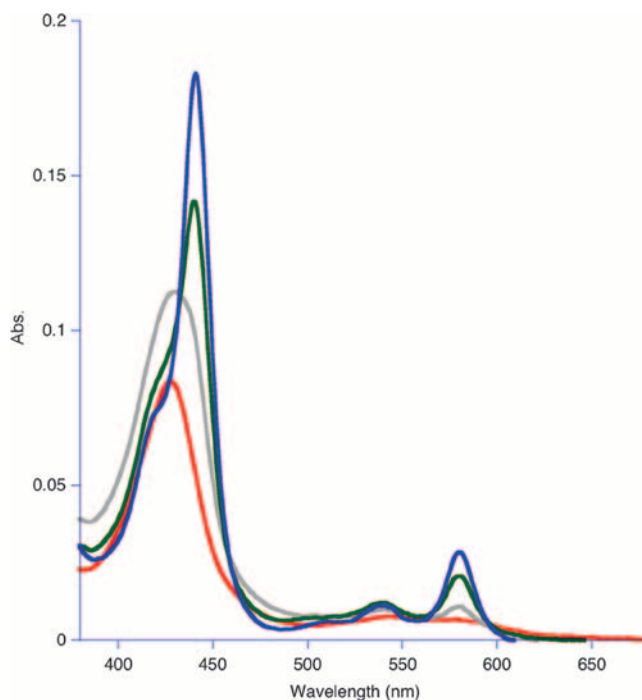


FIG. 6. pH-dependent Fe(II) oxidation by Cyt₅₇₉. The results of redox experiments are shown as follows: pH 1.2 (red), pH 2.0 (gray), pH 3.0 (green), and pH 4.0 (blue). Abs, absorbance.

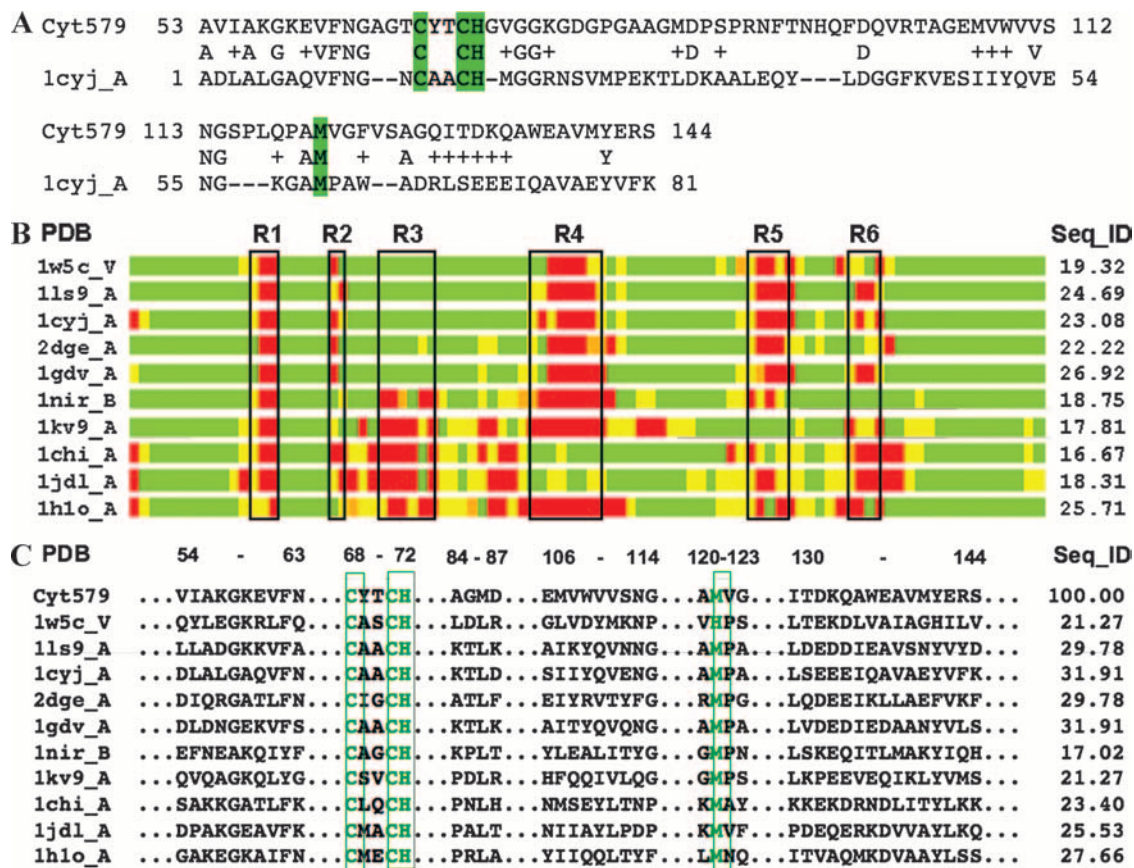


FIG. 7. Modeling of Cyt₅₇₉. (A) The initial structural model of Cyt₅₇₉ was constructed based on sequence alignment with the structure of cytochrome *c*₆, 1cyjA (13). In the alignment, amino acids repeated on the first and second lines are identical, and residues that are chemically similar to those of Cyt₅₇₉ are indicated by plus symbols. Dashes indicate gaps in the alignment. Highlighted residues Cys68, Cys71, His72, and Met121 form the direct interactions with the heme. (B) Regions in the model having structures similar to those of corresponding regions in the structural templates analyzed are aligned in a schematic bar plot. Structural similarity with these templates is indicated as good (green), intermediate (yellow), and nonhomologous (red). Black boxes (R1 to R6) mark regions of structural deviation, or insertions/deletions, observed in structural templates. The region between R1 and R2 corresponds to the conserved CXXCH heme-binding motif. In Cyt₅₇₉, the regions R1 to R6 correspond to the following fragments: R1, 65-AGT-67; R2, 73-GV-74; R3, 78-GDGPGA-83; R4, 93-FTNHQFDQ-100; R5, 115-SPLQPA-120; and R6, 126-SAGQI-130. (C) Residue-to-residue correspondences extracted from structurally conserved regions that were identified within a set of the closest structural templates. The results of the analysis of these regions increased confidence in the calculated sequence alignments used in modeling. The results from calculation of sequence identities between the templates and the model in structurally conserved regions are given in the column labeled "Seq_ID"; in most cases these values are higher than the corresponding Seq_IDs calculated for entire structural alignments shown in panel B.

compared by sequence to structure alignments and in 3D plots with selected structural templates (Fig. 7B and C). The heme orientation and structural elements were compared with the cytochrome *c*₆ structure, 1cyj_A (13) (Fig. 8A). Models for the two major genetic variants of Cyt₅₇₉ were then superimposed to indicate the positions of all nine side chain substitutions, thioether linkages between heme and Cys68 and Cys71, and heme-Fe complex with axial ligands His72 and Met121 (Fig. 8B).

DISCUSSION

In this study, we have purified the abundant, novel bacterial cytochrome first identified by proteogenomic studies in the acidic-wash fraction of biofilms collected at the Richmond Mine in Iron Mountain, CA. We have confirmed the prediction that this protein is Cyt₅₇₉, a modified *c*-type cytochrome that has been implicated as the Fe(II) oxidase in biochemical and physiological studies of *Leptospirillum* isolates (17). In the

initial genomic data set obtained from a biofilm at the Richmond Mine, only one gene was sequenced that coded for Cyt₅₇₉; however, two paralogs of Cyt₅₇₉ were sequenced in a genomic data set from a second biofilm (15, 24). The amino acid substitutions observed in these genetic variants can be predicted in a 3D rendering of the protein structure based on homology modeling (Fig. 8B). It is noteworthy that the predicted variant residues are all located on the surface of the protein and in contact with solvent and thus do not appear to impose any perturbation to structural elements or to the putative interactions with heme. Modeling also predicts a His-Met axial ligation for Cyt₅₇₉ that is consistent with the observation of an absorption band at 695 nm and a mostly helical protein structure that is corroborated by CD spectroscopy.

Detailed biochemical studies of Cyt₅₇₉ isolated from the biofilms have revealed some unexpected features of Cyt₅₇₉. The alkaline pyridine hemochrome spectrum closely resem-

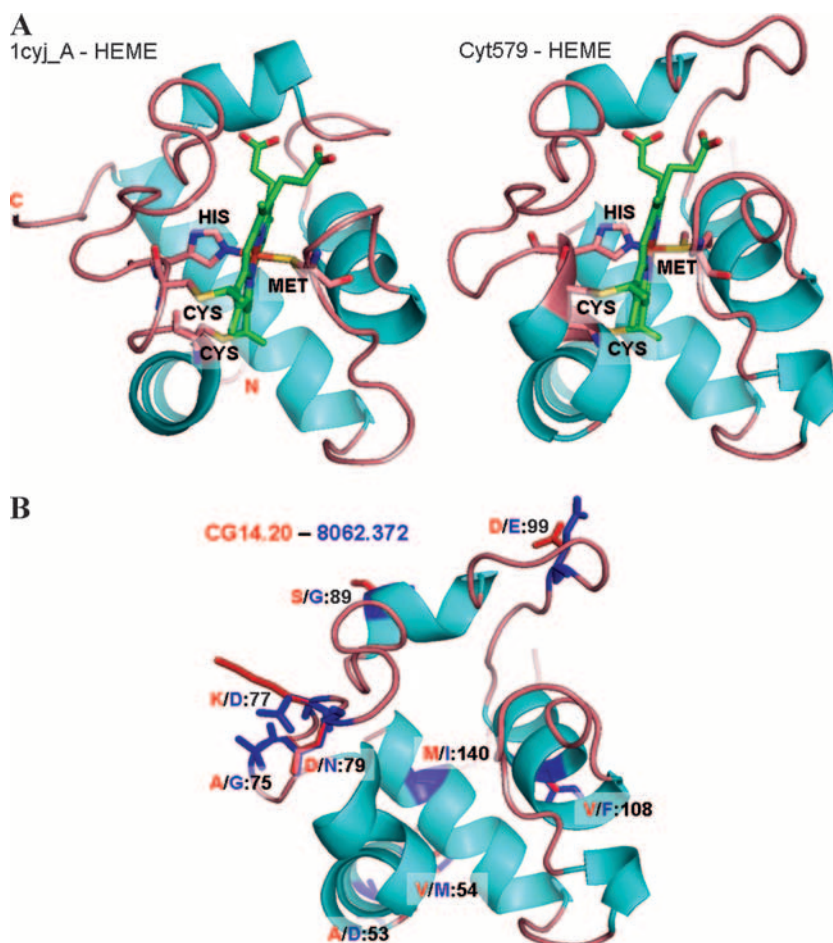


FIG. 8. Structural comparison and variants of Cyt₅₇₉. (A) Structure of cytochrome *c*₆, 1cyjA (13), compared with the final model of Cyt₅₇₉, predicting heme orientation, covalent binding with two Cys residues, and iron coordination complex with axial His and Met residues. (B) Amino acid substitutions are depicted for the two major variants of the Cyt₅₇₉ gene, CG14-20 (red) and 8062-372 (blue) (see Fig. 1 for sequence alignment).

bles the spectrum of heme A (Soret band, 430 nm, and α band, 587 nm), suggesting that the heme in Cyt₅₇₉ may contain a formyl group (3). The spectrum is also consistent with the removal of the heme from the protein, since the α band is red shifted from 579 nm to 587 nm. The presence of a CXXCH amino acid motif and the periplasmic localization of Cyt₅₇₉ are evidence that this unusual heme is covalently bound to the protein, so its removal under alkaline pyridine conditions is unexpected. One interpretation of this result is that the covalent thioether linkages of the modified heme in Cyt₅₇₉ are more sensitive to alkaline pH than those of conventional *c*-type cytochromes.

The second unexpected feature of Cyt₅₇₉ was the isolation of three forms of the protein, truncated at different sites on the N terminus. One of these forms, with a detected N-terminal sequence of AELDILKPRV, was consistent with removal of the predicted signal peptide; however, the other two forms may result from additional proteolysis. Cyt₅₇₉ from *L. ferriphilum* was isolated in one form, corresponding to an N terminus of AELDILKPRV, that is identical to the highest-molecular-weight form of Cyt₅₇₉ from the biofilm (17). These truncations may be due to proteolytic activity during the preparation of

Cyt₅₇₉; however, identical N-terminal sequences were observed for Cyt₅₇₉ preparations from the AB end and C drift biofilms, suggesting that the cleavages are not random and are posttranslational modifications that occur in vivo. N-terminal cleavage sites of Cyt₅₇₉ have been correlated with the different stages of the biofilm life cycle, establishing their ecological relevance (S. W. Singer and M. P. Thelen, unpublished results).

Accurate molecular-mass values for each of the forms of Cyt₅₇₉ were determined by intact-protein analysis using MS. This confirmed the N-terminal cleavage sites observed by Edman degradation and revealed a C-terminal cleavage site. A particularly significant finding was that a sequence variant of Cyt₅₇₉ in the C drift sample was not observed in environmental genomic sequences obtained from Richmond Mine biofilms. The sequence was identified by fragmenting the intact protein and isolating a sequence tag that contained an Ala to Ser variation. The presence of the sequence variant was verified by MS-MS analysis of tryptic peptides. High-resolution intact-protein MS will be invaluable in discriminating between variants of the protein isolated from the environment, allowing the correlation of protein variation with changes in environmental conditions.

The third unexpected feature of Cyt₅₇₉ was that Fe(II) oxidation was not favored thermodynamically at a pH of <3. This result is inconsistent with the results of previous studies with *Leptospirillum* isolates, where complete reduction of Cyt₅₇₉ in the presence of 30 mM Fe(II) was observed at pH 2, and casts doubt on the proposed role of Cyt₅₇₉ as the Fe(II) oxidase for *Leptospirillum* group II bacteria (10, 17).

The properties of Cyt₅₇₉ from *Leptospirillum* group II bacteria are analogous to those of rusticyanin, a periplasmic Cu-containing protein expressed by *Acidithiobacillus ferrooxidans*, an acidophilic Fe(II)-oxidizing bacterium found in environments similar to those where members of *Leptospirillum* group II are found. Biochemical and transcriptomic evidence has implicated rusticyanin as the initial electron acceptor for an outer membrane-bound *c*-type cytochrome, *Cyc2*, which is the proposed Fe(II) oxidase for *A. ferrooxidans* (25–27). In support of this analogy, we have recently purified a novel membrane cytochrome, Cyt₅₇₂, that is expressed by *Leptospirillum* group II in the Richmond biofilms (11). In contrast to Cyt₅₇₉, Cyt₅₇₂ oxidizes Fe(II) at low pH and may donate electrons to Cyt₅₇₉. Efforts to reconstruct the Fe(II)-dependent electron transfer pathway in *Leptospirillum* group II bacteria and clarify the role of Cyt₅₇₉ in this pathway are currently under way.

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