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Perchlorate Reductase Is Distinguished by Active Site Aromatic Gate Residues^{*[5]}

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Perchlorate is an important ion on both Earth and Mars. Perchlorate reductase (PcrAB), a specialized member of the dimethylsulfoxide reductase superfamily, catalyzes the first step of microbial perchlorate respiration, but little is known about the biochemistry, specificity, structure, and mechanism of PcrAB. Here we characterize the biophysics and phylogeny of this enzyme and report the 1.86-Å resolution PcrAB complex crystal structure. Biochemical analysis revealed a relatively high perchlorate affinity ($K_m = 6 \mu\text{M}$) and a characteristic substrate inhibition compared with the highly similar respiratory nitrate reductase NarGHI, which has a relatively much lower affinity for perchlorate ($K_m = 1.1 \text{ mM}$) and no substrate inhibition. Structural analysis of oxidized and reduced PcrAB with and without the substrate analog SeO_3^{2-} bound to the active site identified key residues in the positively charged and funnel-shaped substrate access tunnel that gated substrate entrance and product release while trapping transiently produced chlorate. The structures suggest gating was associated with shifts of a Phe residue between open and closed conformations plus an Asp residue carboxylate shift between monodentate and bidentate coordination to the active site molybdenum atom. Taken together, structural and mutational analyses of gate residues suggest key roles of these gate residues for substrate entrance and product release. Our combined results provide the first detailed structural insight into the mechanism of biological perchlorate reduction, a critical component of the chlorine redox cycle on Earth.

Perchlorate has recently gained notoriety because of the identified environmental prevalence of this exotic ion (1). Its

recently documented widespread distribution in both terrestrial and extraterrestrial planetary and non-planetary samples suggests that perchlorate is prevalent throughout our solar system (1, 2). To date, perchlorate is primarily recognized as a groundwater contaminant resulting from Cold War munition manufacturing (3). However, recent studies have additionally determined a natural atmospheric source (1, 4). Widespread perchlorate abundance has also been identified on Mars (5–7), on our moon, and in meteorites (2). On Mars, the unique perchlorate physical chemistry is responsible for maintaining water in the liquid state regardless of the environmental extremes (8). Furthermore, Martian perchlorate may represent a significant energy and oxygen resource for human exploration (9). Although the exact abiotic mechanisms of natural perchlorate neogenesis are not completely understood, there are no known biogenic mechanisms. In contrast, terrestrial perchlorate diagenesis primarily occurs biologically through microbial respiration (10).

Dissimilatory (per)chlorate-reducing bacteria can use perchlorate or chlorate (collectively (per)chlorate) as a terminal electron acceptor during anaerobic respiration (11). Canonical dissimilatory (per)chlorate-reducing bacteria utilize a highly conserved pathway that is phylogenetically distributed through horizontal gene transfer (3, 12, 13). Perchlorate is sequentially reduced to chlorite by the perchlorate reductase (PcrAB), which is then dismutated into chloride and oxygen by the chlorite dismutase (Cld). The biogenic oxygen is respired by the same microorganism, generally through the use of a high affinity cytochrome *cbb*₃ oxidase (3). Due to oxygen biogenesis, these unique chemotrophs represent a distinctive metabolic juxtaposition between aerobic and anaerobic metabolisms. Although an impressive body of literature focuses on the biochemistry, genetics, and evolution of the Cld, relatively little is known about the PcrAB. We have shown that PcrAB is an oxygen-sensitive heterodimeric periplasmic enzyme that catalyzes the $2e^-$ reduction of perchlorate to chlorate, which is further reduced by $2e^-$ to form chlorite (3, 14). The catalytic subunit PcrA is phylogenetically closely related to the NarG subunit of nitrate reductase (15). Both proteins are in the DMSO⁴ reductase superfamily and are strongly associated with intermediate

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[5] This article contains supplemental Fig. S1 and Table S1.

The atomic coordinates and structure factors (codes 4YDD, 5CH7, 5CHC, and 5E7O) have been deposited in the Protein Data Bank (<http://www.pdb.org/>).

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⁴ The abbreviations used are: DMSO, dimethylsulfoxide; [Fe-S], iron-sulfur cluster; [4Fe-4S], four-iron/four-sulfur cluster; [3Fe-4S], three-iron/four-sulfur cluster; MGD, molybdopterin guanine dinucleotide; *Ec*, *E. coli*.

electron-shuttling protein subunits (PcrB and NarH, respectively). Given the important and ancient roles of [Fe-S] clusters in biology and electron transfer, it is furthermore important to consider the detailed structural relationships of the five [Fe-S] clusters in PcrAB and NarGH.

Here we perform an unprecedented structure and function characterization of PcrAB. The crystal structures revealed all metal ion cofactors along with a novel gating mechanism at the active site that evidently endow the enzyme with unique biophysical characteristics. These, in addition to comparative phylogenetic and mutational studies, identified the evolutionary basis for this enzyme and revealed the unique and distinguishing features between PcrAB and the closely related NarGH respiratory complex.

Experimental Procedures

Expression and Purification of PcrAB—Cells of wild-type or mutant *Azospira suillum* PS were initially grown aerobically from single colonies in acetate, lactate, pyruvate growth medium (12) in 5-ml cultures and incubated overnight at 37 °C at 250 rpm. This culture was then used to inoculate 1 liter of the same growth medium and incubated at the previously listed conditions. Anaerobic cultures were prepared by adding 100 ml of the overnight aerobic culture to sterile 1-liter glass bottles, and fresh sterile acetate, lactate, pyruvate growth medium containing 5 mM NaNO₃, 10 mM NaClO₃, and 10 mM NaClO₄ was added until the bottles were completely filled with no headspace. The sealed bottles were then incubated without shaking at 37 °C for 18–24 h. Cultures changed from yellow to pink when all of the electron acceptors were consumed. This typically yielded 5–6 g of wet cell paste/ liter of anaerobic culture. Cells were harvested anaerobically by filling centrifuge tubes with the anaerobic cultures in an anaerobic chamber (Coy Laboratory Products), and the tubes were sealed, centrifuged at 10,000 × g for 10 min, and returned to the anaerobic chamber where the supernatant was decanted. The pink cell pellets were resuspended in anaerobic buffer A (50 mM Tris, pH 8.0, 10% (v/v) glycerol) to a density of 0.4 g of cells/ml, supplemented with 1 mg/ml lysozyme and 0.1 mg/ml DNase I (Sigma), sealed in glass bottles, and frozen at –80 °C until needed.

PcrAB was purified anaerobically using techniques described previously with some modifications (16). Frozen cells were thawed, lysed under a nitrogen stream with an EmulsiFlex-C3 cell homogenizer (Avestin), and centrifuged at 150,000 × g for 45 min. The clarified lysate was kept anaerobic until it was loaded onto three columns connected in tandem and equilibrated with anaerobic buffer A using an ÄKTA Explorer FPLC (GE Healthcare). The tandem series of columns consisted of a 1 × 20-cm S-Sepharose HP column (GE Healthcare), a 2.5 × 40-cm Q-Sepharose FF column (GE Healthcare), and finally a 2.5 × 5-cm CaPure-HA[®] hydroxyapatite column (Tosoh Bioscience). Once the sample was loaded and the UV/visible traces on the FPLC detector returned to baseline, the first two columns were removed, and PcrAB was eluted from the hydroxyapatite column using a 300-ml linear gradient to buffer B (buffer A with 400 mM NaH₂PO₄). PcrAB typically eluted at 200–250 mM NaH₂PO₄. Fractions containing activity as described below were pooled and concentrated in sealed cen-

trifugal concentrators with 100-kDa-molecular mass-cutoff filters (Amicon) to <800 μl. The concentrated sample was loaded onto a HiPrep Sephacryl S-300 HR 16/60 size exclusion column (GE Healthcare) that was equilibrated with anaerobic buffer C (buffer A with 150 mM NaCl) at a flow rate of 0.25 ml/min. PcrAB typically eluted at 50–60 ml. Fractions shown to be pure by SDS-PAGE were pooled and concentrated as described previously to 7 mg/ml, snap frozen in gasketed cryovials with liquid N₂, and stored at –80 °C until use. Protein concentrations were determined via the Bradford assay (Sigma). PcrAB containing the PcrA W461E mutation was purified in the same manner as wild-type PcrAB.

Steady-state Kinetics—The cNarGHI from *Escherichia coli* was a generous gift from the Weiner laboratory (University of Alberta). Activity assays of PcrAB and *EccNarGHI* were performed in an anaerobic chamber by monitoring the substrate-dependent oxidation of NADH (Sigma) with phenazine methosulfate (Sigma) as an electron shuttle (14). 450 μM NADH was incubated with 100 μM phenazine methosulfate and protein sample in 50 mM MOPS, pH 7.2, and reactions were initiated by the addition of substrate. For reactions containing *EccNarGHI*, the buffer was supplemented with 0.01% (v/v) Thesit detergent (Sigma) to prevent the enzyme from precipitating. NADH oxidation rates were determined by monitoring the changes in absorbance at 340 nm (*A*₃₄₀) using an Infinite F200 Pro plate reader (Tecan) at 25 °C. Reaction rates were determined by fitting the data in Origin 6.0 (MicroCal). Although the *EccNarGHI* *K_m* value for nitrate established with our NADH/phenazine methosulfate assay was consistent with previous reports, the *k_{cat}* values were lower (17). This suggests that phenazine methosulfate is not as efficient as an electron donor to *EccNarGHI* as other electron donors, such as methyl viologen, benzyl viologen, and plumbagin (17, 18), but it does not affect the affinity of the enzyme for its primary substrate.

Crystallization, Data Collection, and Structure Determination of PcrAB—PcrAB (7 mg/ml) was mixed with an equal volume of the crystallization solution (0.1 M Tris, pH 8.25, 20% (w/v) PEG 6000) using the hanging drop vapor diffusion method in the anaerobic chamber. All crystallization buffers were degassed before use, and crystallization trays were stored in the anaerobic chamber for at least 3 days before use. The brown rod-shaped crystals grew over 6 weeks at ~20 °C. For W461E PcrA of PcrAB mutant, crystals were grown in 23% (w/v) PEG 3350 and 0.1 M Tris, pH 8.5, with an equal volume of protein and crystallization solution. The hexagonal plate crystals appeared after 1 month at ~20 °C in the anaerobic chamber. A single crystal of wild-type or mutant PcrAB was transferred into crystallization solution supplemented with cryoprotectant of 20% (v/v) and subsequently 30% (v/v) ethylene glycol for wild-type PcrAB or 25% (v/v) ethylene glycol for mutant PcrAB. The crystal was mounted on a loop on a B3S base (MiTeGen) and inserted in a MicroRT capillary tube (MiTeGen) with 50 μl of reservoir solution in the sealed end of the tube to prevent dehydration. The mounted crystal was then transported outside of the anaerobic chamber and immediately snap frozen in liquid N₂.

The reduced PcrAB crystal (Protein Data Bank code 5CH7) was obtained by soaking 4 mM Na₂SO₃ in the presence of 1.2

PcrAB Is Specialized for Scarce Perchlorate

mM reduced methyl viologen and 2.8 mM sodium dithionite. This solution was added to a drop containing a single oxidized PcrAB crystal (crystallized in 20% (w/v) PEG 6000, 0.1 M Tris, pH 8.5) in the anaerobic chamber for 18 h at $\sim 20^\circ\text{C}$. Reduced methyl viologen was prepared as a 10 mM stock solution using the reservoir solution and a small amount of zinc powder as a reductant, which was subsequently removed via centrifugation at $20,000 \times g$ for 1 min in the anaerobic chamber. For selenite-bound PcrAB (Protein Data Bank code 5CHC), a PcrAB crystal was soaked with 4 mM Na_2SeO_3 in the presence of reduced methyl viologen and sodium dithionite for 18 h at $\sim 20^\circ\text{C}$ as described above. The reduced PcrAB crystals were then transferred into crystallization solution supplemented with the aforementioned reducing agents and cryoprotectants. The mounting procedure for the reduced crystals was the same as described above.

The x-ray diffraction data of PcrAB crystals were collected at BL12.3.1 SIBYLS beamline (sibyls.als.lbl.gov) at the Advanced Light Source (19). For oxidized PcrAB, two data sets from different spots of the same crystal were collected at 100 K and integrated using the XDS package (20). The data were then merged and scaled using AIMLESS in the CCP4 suite (21). POINTLESS indicates a space group of $P2_12_12_1$ for three molecules of PcrAB per asymmetric unit for oxidized and reduced PcrAB.

The oxidized PcrAB structure was determined via molecular replacement using Phaser in the Phenix suite (22) using ethylbenzene dehydrogenase (Protein Data Bank code 2IVF) α - and β -subunits as a search model (23). The molecular replacement solution was then processed with AutoBuild in Phenix (22). The resulting model was manually fit by Coot (24) and refined by Phenix.refine with non-crystallographic symmetry restraints.

The reduced PcrAB structures and W461E PcrA of PcrAB mutant structure were determined by molecular replacement using oxidized PcrAB as a search model. The molecular replacement solutions of reduced PcrAB were processed the same way as oxidized PcrAB except the twin law ($-h-l,k,h$) was applied for the mutant PcrAB (estimated twin fraction, 0.16) during refinement. The x-ray data collection and refinement statistics are listed in Table 2.

The protein data bank codes are 4YDD (oxidized PcrAB), 5CH7 (reduced PcrAB-Phe¹⁶⁴ gate switch), 5CHC (reduced PcrAB- SeO_3^{2-} bound), and 5E7O (PcrAB mutant-W461E PcrA). All the structure figures are depicted using Chimera (25). The hydrophobicity scale is based on Kyte and Doolittle (26).

PcrAB Site-directed Mutagenesis—The *pcrA* gene was amplified from *A. suillum* PS genomic DNA using PcrA_AD-F and PcrA_AD-R primers (supplemental Table S1). This PCR product was purified and combined with the pCR[®]-Blunt II-TOPO[®] vector using the Zero Blunt[®] TOPO PCR cloning kit (Life Technologies) and transformed into XL1 Blue chemically competent *E. coli* (Stratagene). Positive transformants were grown in LB supplemented with 50 $\mu\text{g}/\text{ml}$ kanamycin, and plasmid DNA was isolated using the QIAprep Spin Miniprep kit (Qiagen). This plasmid was used as a template for PCR-based site-directed mutagenesis with primers specific to particular point mutations (supplemental Table S1) (27). These PCRs were

digested with DpnI (New England Biolabs) and transformed again into XL1 Blue cells. Positive transformants were grown, plasmid DNA was isolated as done previously, and positive mutations were screened using Sanger sequencing. Plasmids containing the correct mutations were digested with ApaI and BamHI (New England Biolabs) along with the pNPTS138 suicide vector (12), purified, ligated using T4 DNA ligase (New England Biolabs), and transformed into XL1 Blue cells. Transformants were screened via colony PCR, and pNPTS138 plasmids containing correctly mutated *pcrA* were isolated and confirmed by Sanger sequencing. Allelic replacement of *pcrA* with the point mutants was performed as described previously (12). Growth curves of all strains were performed anaerobically using a Sunrise 96-well plate reader (Tecan) according to previously published protocols (12). Chlorate accumulation experiments were performed in anaerobic bottles with 10 mM perchlorate and 10 mM acetate and monitored via OD_{600} and ion chromatography using an ICS1500 (Dionex) outfitted with an AS25 column (Thermo Scientific).

Phylogenetic Analysis—Phylogenetic analysis of the catalytic subunit of the perchlorate reductase (PcrA) was performed using the computer cluster at the University of California Berkeley QB3 Computational Genomics Resource Laboratory. The molybdopterin oxidoreductase protein family (Pfam PF00384) formed the starting alignment for a phylogenetic analysis with FastTree2 (version 2.1.7) (28). From the resulting tree, a clade consisting of DmsA and DorA was chosen as an appropriate outgroup to known PcrA proteins. Non-basal sequences were combined with the top 100 blastp hits from both NCBI and Integrated Microbial Genomes (IMG) databases. CD-HIT (29) was used to cluster sequences at a threshold of 0.8. The remaining sequences were aligned using MUSCLE (30) and trimmed with trimAL (31), and these informative positions were used to reconstruct the phylogeny with FastTree2. Salient features of clades, including known function of members, number of sequences containing a predicted twin arginine translocation signal (32), and sequence motifs (33) of proposed active site residues, were visualized on the tree.

Results

Comparative Steady-state Kinetics of PcrAB Reveals Unique Features—PcrAB kinetic analysis revealed nearly identical steady-state parameters for (per)chlorate at substrate concentrations below 200 μM (Table 1A). Although chlorate followed typical Michaelis-Menten kinetics (Fig. 1A), perchlorate showed significant substrate inhibition above 200 μM with an estimated inhibition constant (K_i) of ~ 7.5 mM (Fig. 1B). Interestingly, a critical inhibitory concentration for perchlorate was not noted over the range analyzed (0–200 mM). PcrAB also mediated turnover of NO_3^- , BrO_3^- , and IO_3^- (Table 1A), but no activity was observed with SO_4^{2-} , SO_3^{2-} , SeO_4^{2-} , SeO_3^{2-} , AsO_4^{3-} , AsO_3^{3-} , NO_2^- , WO_4^{2-} , or MoO_4^{2-} . Neither BrO_4^- nor IO_4^- could be investigated due to their reactive nature. The catalytic efficiencies (k_{cat}/K_m) for all substrates of PcrAB were within the same order of magnitude (Table 1A), suggesting no evolutionary specificity for any of these oxyanions.

E. coli cytoplasm-facing nitrate reductase (*EccNarGHI*) also had activity toward (per)chlorate (Table 1B) albeit with

TABLE 1**Steady-state kinetic parameters and Bond lengths and molecular dipoles of substrates**

Steady-state kinetic parameters show relatively high substrate affinity for PcrAB in comparison with *EccNarGHI*. A, wild-type PcrAB; B, *EccNarGHI*; C, PcrA W461E. The steady-state kinetics were determined using the NADH/PMS assay. Error represent one standard deviation of triplicate assays. D, Bond lengths and molecular dipoles of substrates (52). N.S., not a substrate. *, Iodate was also previously determined to not be a substrate of *EccNarGHI* (17). †, Estimated values due to the solubility limitations of NaIO_3 .

Table 1A. PcrAB – Wild-type					
Substrate	Perchlorate	Chlorate	Bromate	Iodate	Nitrate
K_M (μM)	6.0 ± 2.1	7.4 ± 2.1	4.4 ± 3.8	11 ± 5.4	23 ± 5.2
k_{cat} ($\text{e}^- \text{min}^{-1}$)	27.1 ± 1.9	27.1 ± 1.4	30.5 ± 4.5	35.9 ± 3.3	51.1 ± 2.4
k_{cat}/K_M ($\text{M}^{-1} \text{s}^{-1}$)	7.5×10^4	6.1×10^4	1.2×10^5	5.4×10^4	3.7×10^4

Table 1B. <i>Ec</i> NarGHI					
Substrate	Perchlorate	Chlorate	Bromate	Iodate	Nitrate
K_M (μM)	1060 ± 154	113 ± 35	2690 ± 502	N.S.*	202 ± 65
k_{cat} ($\text{e}^- \text{min}^{-1}$)	344 ± 12	928 ± 78	462 ± 39	N.S.*	1440 ± 14
k_{cat}/K_M ($\text{M}^{-1} \text{s}^{-1}$)	5.4×10^3	1.4×10^5	2.9×10^3	N.S.*	1.2×10^5

Table 1C. PcrAB – PcrA W461E					
Substrate	Perchlorate	Chlorate	Bromate	Iodate [†]	Nitrate
K_M (μM)	301 ± 14.1	20.5 ± 3.2	4.8 ± 2.1	$1.27 \times 10^2 \pm 3.5 \times 10^4$	$1.59 \times 10^4 \pm 4.5 \times 10^3$
k_{cat} ($\text{e}^- \text{min}^{-1}$)	54.1 ± 4.1	72.1 ± 2.3	31.0 ± 1.7	85.1 ± 15	43.2 ± 3.5
k_{cat}/K_M ($\text{M}^{-1} \text{s}^{-1}$)	3.0×10^3	5.9×10^4	1.1×10^5	1.1×10^1	4.5×10^1

Table 1D. Substrate Bond Lengths and Molecular Dipoles					
Substrate	Perchlorate	Chlorate	Bromate	Iodate	Nitrate
Bond Length (\AA)	1.50	1.53	1.69	1.82	1.26
Molecular Dipole (D)	0	2.02	3.06	4.46	0

very different kinetics. *EccNarGHI* had a much lower affinity for all substrates analyzed relative to PcrAB and showed ~ 200 -fold lower affinity toward perchlorate (PcrAB $K_m = 6 \mu\text{M}$; *EccNarGHI* $K_m = 1.1 \text{ mM}$). *EccNarGHI* catalytic efficiencies were inversely proportional to substrate size, being lower for the larger oxyhaloanions (ClO_4^- and BrO_3^-) relative to the smaller NO_3^- and ClO_3^- . No activity was observed toward IO_3^- (Table 1B). In contrast to PcrAB, *EccNarGHI* was uninhibited by increasing concentrations of perchlorate and displayed typical Michaelis-Menten kinetics throughout the range tested (Fig. 1B). Surprisingly, although not considered its natural substrate, *EccNarGHI* had a much higher maximum perchlorate turnover rate than PcrAB with a k_{cat} nearly 10-fold higher (Table 1, A and B). At $\sim 125 \mu\text{M}$ perchlorate, both PcrAB and *EccNarGHI* had nearly identical turnover rates, whereas below that PcrAB had a turnover rate up to 5-fold greater than *EccNarGHI* (Fig. 1C), suggesting that it evolved to take advantage of the low concentrations of (per)chlorate naturally found in the environment (4).

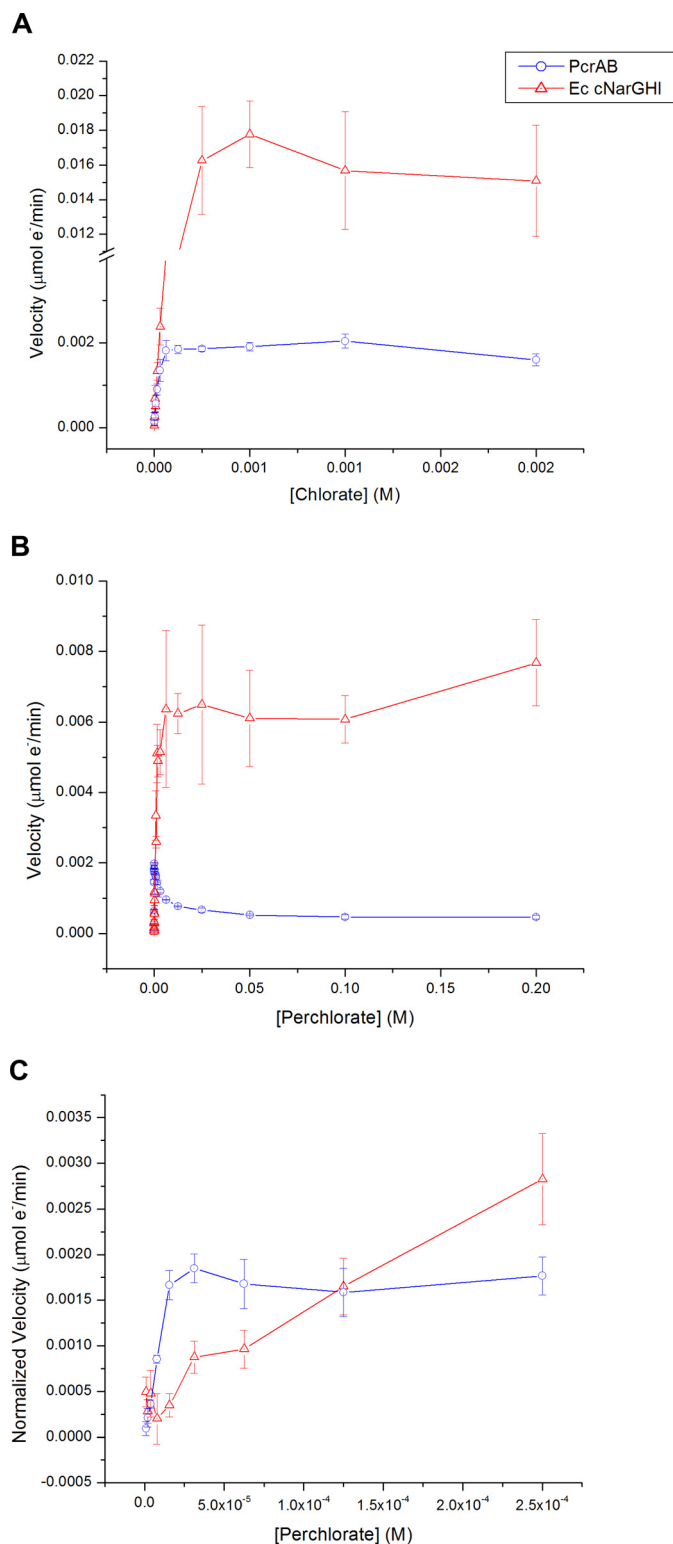


FIGURE 1. Steady-state kinetic curves for PcrAB and *EccNarGHI* for chlorate (A), perchlorate (B), and perchlorate (C) at low substrate concentrations ($<200 \mu\text{M}$). Error bars represent one standard deviation of triplicate assays.

Crystal Structure of Oxidized PcrAB Reveals a Unique Gated Active Site—To better understand why PcrAB is more efficient at low perchlorate concentrations, we crystallized the purified enzyme anaerobically and determined the crystal structure of

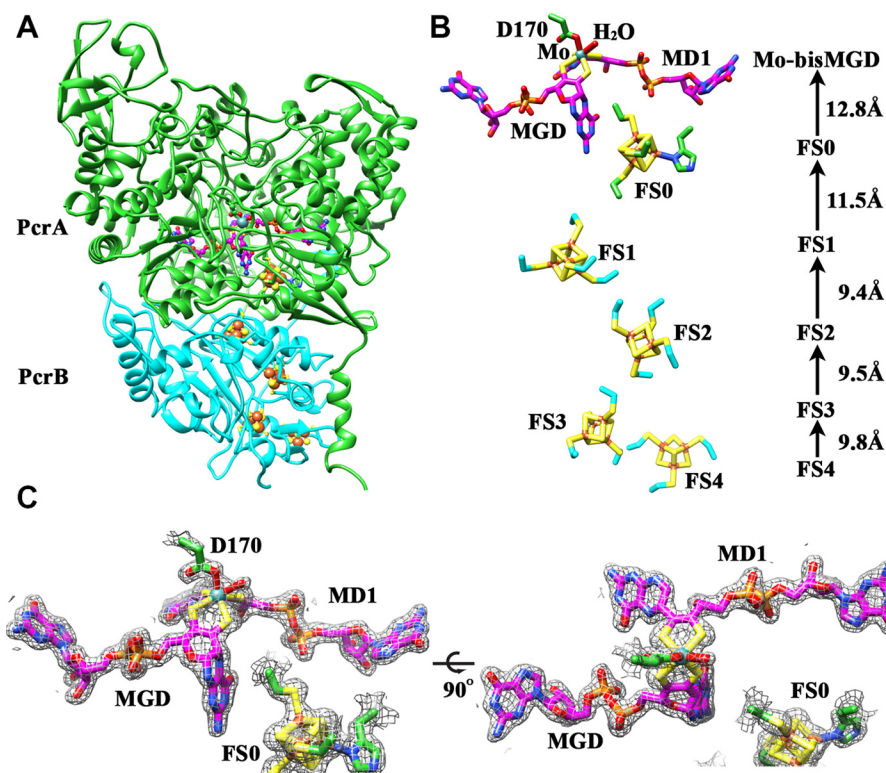


FIGURE 2. Crystal structure of oxidized PcrAB heterodimer reveals [Fe-S] clusters are 10–12 Å apart from the adjacent cluster and the Mo-bis-MGD cofactor. *A*, overall crystal structure of oxidized PcrAB. PcrA and PcrB are shown in green and cyan ribbon, respectively. [Fe-S] clusters are shown in sphere (iron, brown; sulfur, yellow), bis-MGD cofactor is shown in magenta ball-and-stick, and molybdenum is shown in turquoise sphere. *B*, the relative position of cofactors in PcrAB along with Asp¹⁷⁰ and a water molecule that coordinate to the active site molybdenum atom. Edge-to-edge distances between cofactors are shown. Labeling of the [Fe-S] clusters is consistent with previous publications (35, 36). The tricyclic pyranopterin of the molybdopterin cofactor is labeled as MGD, and the bicyclic dihydropterin is labeled as MD1. *C*, electron density $2|F_o| - |F_c|$ simulated annealing omit maps (gray) at 1.5 σ contour level of PcrA cofactors and active site for oxidized PcrAB. The tricyclic pyranopterin of the molybdopterin cofactor is labeled as MGD, and the bicyclic dihydropterin is labeled as MD1.

oxidized PcrAB at 1.86-Å resolution (Fig. 2A) along with three reduced or mutant crystal structures (Table 2). PcrAB was crystallized in space group $P2_12_12_1$ and formed a trimer of heterodimers. The PISA assembly (34) indicated that PcrAB was stable as a heterodimer with a buried area of 14,000 Å². The oxidized PcrA structure, where the oxidation state of molybdenum is most likely +5 or +6, resembled other DMSO reductase superfamily enzymes (35–37). PcrAB was composed of one Mo-bis-MGD cofactor and one [4Fe-4S] cluster (FS0) for PcrA and three [4Fe-4S] clusters (FS1–3) and one [3Fe-4S] cluster (FS4) for PcrB with all metal atoms and cofactors having an occupancy of 1 (Fig. 2, A and B). All [Fe-S] clusters were coordinated with Cys residues except the FS0 cluster, which had one iron atom coordinated by the His³² residue. A mutation of this His to Cys in *EccNarGHI* resulted in loss of activity toward nitrate (38). The molybdenum atom was coordinated in a trigonal prismatic manner with four proximal thiolates of the bis-MGD cofactor (average Mo–S bond distance of 2.4 Å) along with a Asp¹⁷⁰ residue and an additional oxygen atom (either an oxo, hydroxyl, or associated water molecule) in the two distal positions (Fig. 2B). The Mo–O bond distances were ~2.0–2.1 and 2.4–3.1 Å for Asp¹⁷⁰ and the additional oxygen atom, respectively, in the three PcrA molecules in the asymmetric unit. The cyclization states of the bis-MGD cofactor were in a configuration with one in a bicyclic dihydropterin and the other in a tricyclic pyranopterin structure (Fig. 2C). Most DMSO reductase superfamily members possess both pterins in the tri-

cyclic state; however, the bicyclic and tricyclic pterins have been noted in NarGHI and ethylbenzene dehydrogenase structures (23, 35), suggesting an interconversion mechanism that controls pterin cyclization states (39). These structural features of PcrAB are similar to *EccNarGH* (35, 36), which has a root mean square deviation of 0.974 Å between 710 aligned C α atoms. The positions of cofactors and clusters in PcrAB and *EccNarGH* were nearly identical with respect to one another, suggesting a strikingly similar pathway of electron transfer to the active site despite 66% sequence disparity.

A funnel-shaped substrate access tunnel (volume, ~1089 Å³) with a wider mouth that narrowed near the active site was detected for PcrAB by the CASTp web server (40) (Fig. 3A). Residue Phe¹⁶⁴ in the tunnel near the active site appeared to block substrate access. Two more aromatic residues (Tyr¹⁶⁵ and Trp⁴⁶¹) were adjacent to Phe¹⁶⁴ and together formed a putative hydrophobic gate that controlled active site entry and exit. Calculation of the electrostatic potential of PcrA using Chimera (25) showed the substrate access tunnel was dominated with positive charges (Fig. 3B). This positively charged tunnel creates the relatively high affinity for oxyanions seen in Table 1A. The *EccNarG* tunnel structure at a similar location formed a much wider pocket toward the active site with a tunnel volume of 1682 Å³ (Fig. 3C). The corresponding gate residues in *EccNarG* overlay well with the PcrA structure. However, residue Glu⁵⁸¹ in *EccNarG* that is equivalent to Trp⁴⁶¹ in PcrA changes the polarity near the gate. The electrostatic

TABLE 2
X-ray data collection and refinement statistics for the PcrAB crystal structures

CC, cross-correlation; ASU, asymmetric unit; r.m.s.d., root mean square deviation; ALS, Advanced Light Source.

	PcrAB oxidized	PcrAB reduced-Phe ¹⁶⁴ gate switch	PcrAB reduced-substrate analog bound	PcrAB mutant-W461E PcrA
Data collection	ALS-BL12.3.1	ALS-BL12.3.1	ALS-BL12.3.1	ALS-BL12.3.1
Wavelength (Å)	1.1158	0.9793	0.9793	1.0062
Space group	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁
Cell dimensions				
<i>a</i> , <i>b</i> , <i>c</i> (Å)	132.82, 175.50, 193.16	132.88, 175.67, 193.28	133.93, 176.02, 193.69	135.59, 253.13, 135.80
α , β , γ (°)	90.00, 90.00, 90.00	90.00, 90.00, 90.00	90.00, 90.00, 90.00	90.00, 119.77, 90.00
Resolution (Å)^a	48.35–1.86 (1.89–1.86)	48.38–2.20 (2.24–2.20)	48.66–2.38 (2.42–2.38)	48.32–2.40 (2.44–2.40)
Observations^a	7,305,807 (365,469)	3,407,133 (169,632)	2,724,751 (135,221)	1,141,127 (57,773)
Unique observations^a	372,024 (18,112)	228,466 (11,210)	183,135 (8,943)	306,180 (15,230)
R_{merge}^a	0.178 (1.623)	0.205 (1.619)	0.222 (1.607)	0.145 (0.824)
R_{pim}^a	0.059 (0.528)	0.079 (0.620)	0.086 (0.618)	0.136 (0.758)
Mean I/σ^a	13.2 (2.1)	13.6 (2.1)	13.0 (2.0)	6.8 (2.0)
Completeness (%)^a	99.1 (97.9)	99.9 (99.7)	100 (100)	99.2 (99.8)
Multiplicity^a	19.6 (20.2)	14.9 (15.1)	14.9 (15.1)	3.7 (3.8)
CC(1/2)^a	0.998 (0.759)	0.997 (0.683)	0.996 (0.706)	0.985 (0.521)
Refinement				
R _{work} /R _{free} (%) ^b	20.0/23.6	17.8/23.3	17.0/22.7	19.5/24.8
mol/ASU	6	6	6	12
No. of residues/ASU	3,664	3,661	3,664	7,344
r.m.s.d. bond length (Å)	0.010	0.011	0.010	0.003
r.m.s.d. bond angle (°)	1.436	1.466	1.380	0.657
Average B-factor	29.29	35.0	36.6	38.25
Protein	29.16	35.1	36.8	38.43
Ligands	21.90	29.8	32.0	36.03
Solvent	31.94	34.9	34.9	35.41
Ramachandran (%)				
Favored	96.06	95.05	95.02	93.88
Allowed	3.69	4.70	4.62	5.48
Outliers	0.25	0.25	0.36	0.64
Protein Data Bank code	4YDD	5CH7	5CHC	5E7O

^a Values in parentheses are the statistics for the highest resolution shell of data.^b $R_{\text{work}} = (\sum |F_{\text{obs}}| - |F_{\text{calc}}|) / \sum |F_{\text{obs}}|$ where F_{obs} and F_{calc} are the observed and calculated structure factors, respectively. R_{free} is calculated the same as R_{work} , but 5% of the data were excluded from the refinement.

potential calculation of EccNarG showed the substrate access tunnel was dominated with negative charges (Fig. 3D), which would generate repulsions for oxyanions, especially for larger anionic molecules, and may explain the relatively low EccNarGHI affinity for these molecules (Table 1B).

Phylogenetic Analysis and Mutational Studies of PcrA Reveal Functional Residues—Phylogenetic analysis emphasizing the gate residues (Phe¹⁶⁴, Tyr¹⁶⁵, and Trp⁴⁶¹), an active site residue (Asp¹⁷⁰), and adjacent residues was used to compare closely related DMSO reductase superfamily enzymes. This revealed a monophyletic clade for PcrA closely related to the NarG (Fig. 4). NarG-containing enzymes either face the cytoplasm (cNarG) or periplasm (pNarG) and are differentiated by the presence of twin arginine translocation sequences (32) targeting pNarG to the periplasm. Twin arginine translocation sequences are found on all sequenced perchlorate reductases as well as many other type II DMSO reductases (see supplemental Fig. S1 for the complete phylogenetic tree). Sequence alignments of PcrA, pNarG, and cNarG depict several conserved regions. All sequences contain a hallmark Asp residue (41) at the active site that coordinates the molybdenum atom from a bis-MGD (Fig. 2). Additionally, all PcrA, pNarG, and cNarG sequences contain several conserved residues in the N-terminal portion of the molybdopterin-binding domain. The PcrAB crystal struc-

ture gate residues (Phe¹⁶⁴ and Tyr¹⁶⁵) were also conserved in NarG, whereas the PcrA Trp⁴⁶¹ gate residue was a Glu in NarG.

Based on crystal structure and phylogenetic analyses, mutational studies on the gate residues and Asp¹⁷⁰ ligand were performed to determine the impact on cell growth and catalytic efficiency. Replacement of the *A. suillum* wild-type PcrA with PcrA F164A, Y165A, or W461A mutant removed the organism's capacity to grow by (per)chlorate respiration (Fig. 5A), indicating their critical role for PcrAB function. Interestingly, although the PcrA W461A mutant was incapable of growth on (per)chlorate, a W461E mutant retained its capacity to grow with chlorate. Furthermore, the purified W461E mutant PcrAB still displayed activity toward perchlorate albeit with a dramatically increased K_m (Table 1C). The iodate K_m was also greatly increased, whereas the chlorate and bromate values were only slightly impacted (Table 1C). Overall, the steady-state kinetic parameters for this mutant enzyme resembled a midpoint between wild-type PcrAB and EccNarGHI (Tables 1, A–C). The catalytic efficiency of the PcrA W461E mutant for perchlorate ($k_{\text{cat}}/K_m = 3.0 \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$) more resembled that of EccNarGHI ($k_{\text{cat}}/K_m = 5.4 \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$) than wild-type PcrAB ($k_{\text{cat}}/K_m = 7.5 \times 10^4 \text{ M}^{-1} \text{ s}^{-1}$). However, the catalytic efficiency of the PcrA W461E mutant for nitrate decreased 820- and 2670-fold

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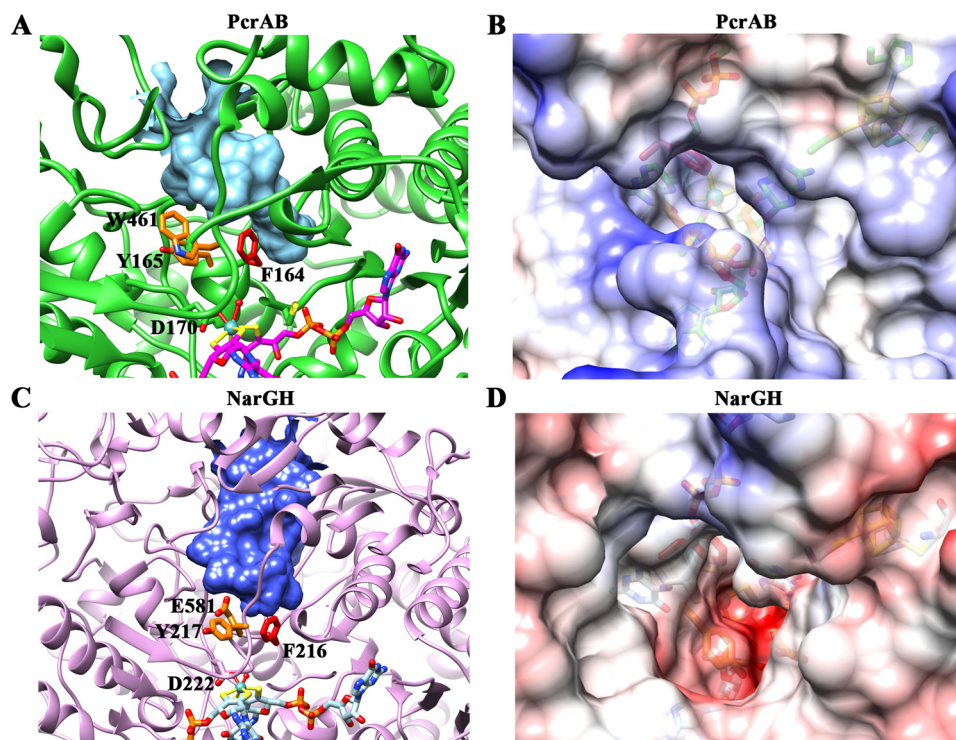


FIGURE 3. Comparisons of substrate accessing tunnels between PcrA and NarG identify the aromatic gate residues and the differences in electrostatic potential. *A*, structure of the closed tunnel in PcrA from the surface of the enzyme to the active site as seen in the oxidized PcrAB crystal structure (Fig. 2*A*) forms a funnel shape (cyan). The surface of the closed tunnel was detected by CASTp (40) and depicted using Chimera (25). The gate residues Phe¹⁶⁴ (red), Tyr¹⁶⁵ (orange), and Trp⁴⁶¹ (orange) are shown in stick. *B*, the electrostatic potential surface of PcrA. The figure is viewed from the mouth of the tunnel down to the active site. The color is shown from negative (red; -10 kcal/mol $\times e$) through white to positive (blue; 10 kcal/mol $\times e$). *C*, structure of closed tunnel (blue) in *EccNarG* (Protein Data Bank code 1Q16) (35) forms a wider tunnel (tunnel volume, 1682 \AA^3) toward the active site than the tunnel in PcrA (tunnel volume, 1089 \AA^3). The corresponding gate residues in NarG are shown in stick with the same color code. *D*, the electrostatic potential surface of NarG by viewing from the same angle as *B*.

compared with wild-type PcrAB and *EccNarGHI*, respectively. Additionally, substrate inhibition for perchlorate was greatly reduced in the W461E mutant, and the kinetics resembled that of *EccNarGHI* (Fig. 5*B*). Surprisingly, this Trp to Glu mutation greatly increased the PcrAB nitrate K_m almost 700-fold (Table 1*C*), which was counterintuitive as this mutation resembles the natural NarG sequence.

The PcrA W461E crystal structure indicated that Glu⁴⁶¹ side chain was highly flexible relative to the corresponding Glu⁵⁸¹ residue in NarG (Fig. 6). This flexibility was in part because the modified active site loses the π -stacking interaction with Tyr¹⁶⁵ and lack of hydrogen bonding with Ser⁵⁹⁴ because the location of Ser⁵⁹⁴ is too far away from Glu⁴⁶¹. In *EccNarG*, Glu⁵⁸¹ is stabilized by hydrogen bonding to Asp⁸⁰¹ (corresponding to PcrA Ser⁵⁹⁴; Fig. 6). The mutation of W461E in PcrA not only introduces a negative electrostatic charge in the tunnel but also puts the Glu⁴⁶¹ side chain in front of the gate between Tyr¹⁶⁵ and Phe¹⁶⁴, which affected the open and closed conformation of Phe¹⁶⁴. This putatively resulted in the decrease in substrate affinity and catalytic efficiency for perchlorate and nitrate (Table 1*C*) as well as the substrate inhibition of perchlorate (Fig. 5*B*).

In contrast to perchlorate and nitrate, the W461E mutation had little impact on the chlorate and bromate kinetics (Table 1, *A* and *C*). These molecules have significantly longer bond lengths than nitrate and perchlorate and thus a greater molecular size (Table 1*D*). Consequently, if steric hindrance was the

major cause of the increased PcrA W461E K_m values, chlorate and bromate would experience a similar shift. This suggests that the mechanism underlying the dramatic kinetic shift (particularly substrate affinities) was not a function of substrate size (Table 1*D*). The defining feature that differentiates the various substrates is the presence (chlorate and bromate) or absence (perchlorate and nitrate) of a molecular dipole and its interaction with the newly introduced polar Glu⁴⁶¹ residue in PcrA. Iodate is the obvious outlier in this proposed trend; however, the significantly larger size of iodate (Table 1*D*) could represent a ceiling at which advantageous dipole interactions with Glu⁴⁶¹ are overcome by unfavorable steric hindrance.

Additional mutations were made in *pcrA* to further elicit the role of the gating residues with respect to trapping transiently produced chlorate generated during perchlorate reduction (Fig. 7). Five *pcrA* mutations were made in accordance with the phylogenetic analysis such that PcrA mutants were created with one point mutation in the PS PcrA relative to another PcrA in the same clade (Fig. 4). PcrA Y165C and F164S/Y165F show significant growth defects, whereas PcrA G169A and G169S grew identically to wild-type PS (Fig. 7), and PcrA Y165F displayed a minor growth defect. The most dramatic result was that PcrA Y165F accumulated $\sim 50\%$ more chlorate than wild-type PS, whereas all other strains accumulated the same or lower amounts relative to the wild-type organism. The PcrA Y165F active site mutation is significant as other dissimulatory

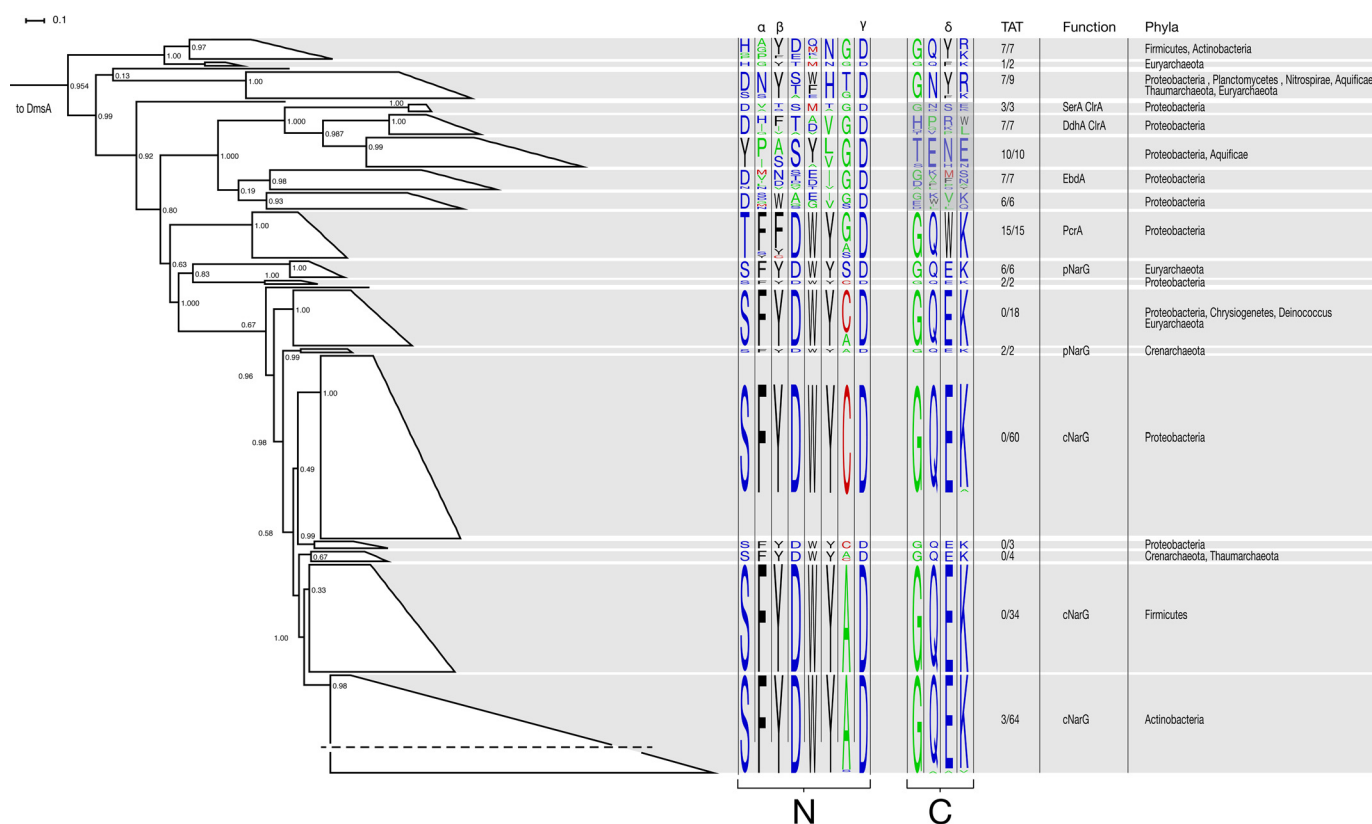


FIGURE 4. Phylogenetic reconstruction of PcrA and closely related DMSO reductase superfamily enzymes using DmsA and DorA as an outgroup. Active site residues identified from the crystal structure of PcrA were visualized for clades with high bootstrap support using sequence motifs. The positions denoted by α , β , and δ in the sequence alignments correspond to the gate residues relative to the *A. suillum* PS PcrA. The positions denoted by γ in the sequence alignments correspond to the Asp residue that has been shown or is predicted to bind to the molybdenum atom at the active site of the corresponding enzyme. N and C represent the N- and C-terminal portions of the MBD in each enzyme. The fraction of predicted twin arginine translocation (TAT) signal sequences, physiological function of known members, and taxonomic affiliation are superimposed on the tree. For the clades containing SerA, ClrA, and EbdA, the darker highlighting in the C-terminal portion of the aligned sequences denotes lower confidence in the alignment. SerA is selenate reductase, ClrA is chlorate reductase, EbdA is ethylbenzene dehydrogenase, PcrA is perchlorate reductase, pNarG is periplasmic nitrate reductase, and cNarG is cytoplasmic nitrate reductase. A horizontal dash in the lowermost cNarG clade denotes the compression of this large group of sequences into a group suitable for visualization.

(per)chlorate-reducing bacteria that have been shown to accumulate chlorate (e.g. *Propionivibrio militaris* MP (42)) contain active sites with this configuration, suggesting that Tyr¹⁶⁵ plays an important role in trapping transiently produced chlorate at the active site of PcrA during perchlorate turnover.

Phe¹⁶⁴ Gating for Substrate Entry and Product Release—To gain insight into the active site gating mechanism, we obtained and characterized two additional novel crystal structures of reduced PcrAB in comparison with the oxidized structures (Table 2). One of the reduced structures contained the substrate analog SeO₃²⁻ bound to the active site molybdenum atom (Fig. 8, A and B). The other structure had the gate residue Phe¹⁶⁴ in a dual occupancy of both open and closed conformations (Fig. 8, C and D). In the SeO₃²⁻-bound structure, one of the selenite oxygen atoms was bound to the molybdenum atom, which was in turn coordinated with Asp¹⁷⁰ in a monodentate fashion (Fig. 8A). The bond distances of Mo–O for selenite and Asp¹⁷⁰ were 2.0–2.1 and 2.2–2.4 Å, respectively, in the three PcrA molecules in the asymmetric unit. The other oxygen atoms of the selenite molecule were orientated such that they did not coordinate to any active site hydrogen bond acceptors. The cyclization states of the bis-MGD rings in the selenite-bound reduced PcrA structure appeared not to change relative

to the oxidized structure (Figs. 2C and 8B). It was proposed in the molybdenum cofactor model complex that the interconversion of bicyclic and tricyclic structures could be solvent-dependent (43), and our results suggest that this interconversion is most likely not redox-dependent.

Soaking with and co-crystallization of substrate analogs have previously been tried with NapA, although nitrate and perchlorate were only found near the outside of the tunnel (44). We successfully obtained crystals with the substrate analog bound to the active site by soaking anaerobically under reducing conditions (see “Experimental Procedures”). Intriguingly, Phe¹⁶⁴ was in the closed conformation in the selenite-bound structure, suggesting that a molecule is trapped at the active site when it is bound to the molybdenum atom.

We obtained a reduced PcrAB structure with the side chain of Phe¹⁶⁴ in dual occupancy of both open (occupancy of 0.45) and closed (occupancy of 0.55) conformations, respectively (Fig. 8, C and D). In this structure, the coordination of Asp¹⁷⁰ to the molybdenum atom was shifted from monodentate to bidentate with bond distances of Mo–O₁ (2.0–2.2 Å) and Mo–O₂ (2.3–2.7 Å) in the three PcrA molecules in the asymmetric unit. This carboxylate shift presumably displaced the water molecule that was originally bound to the molybdenum atom. The water

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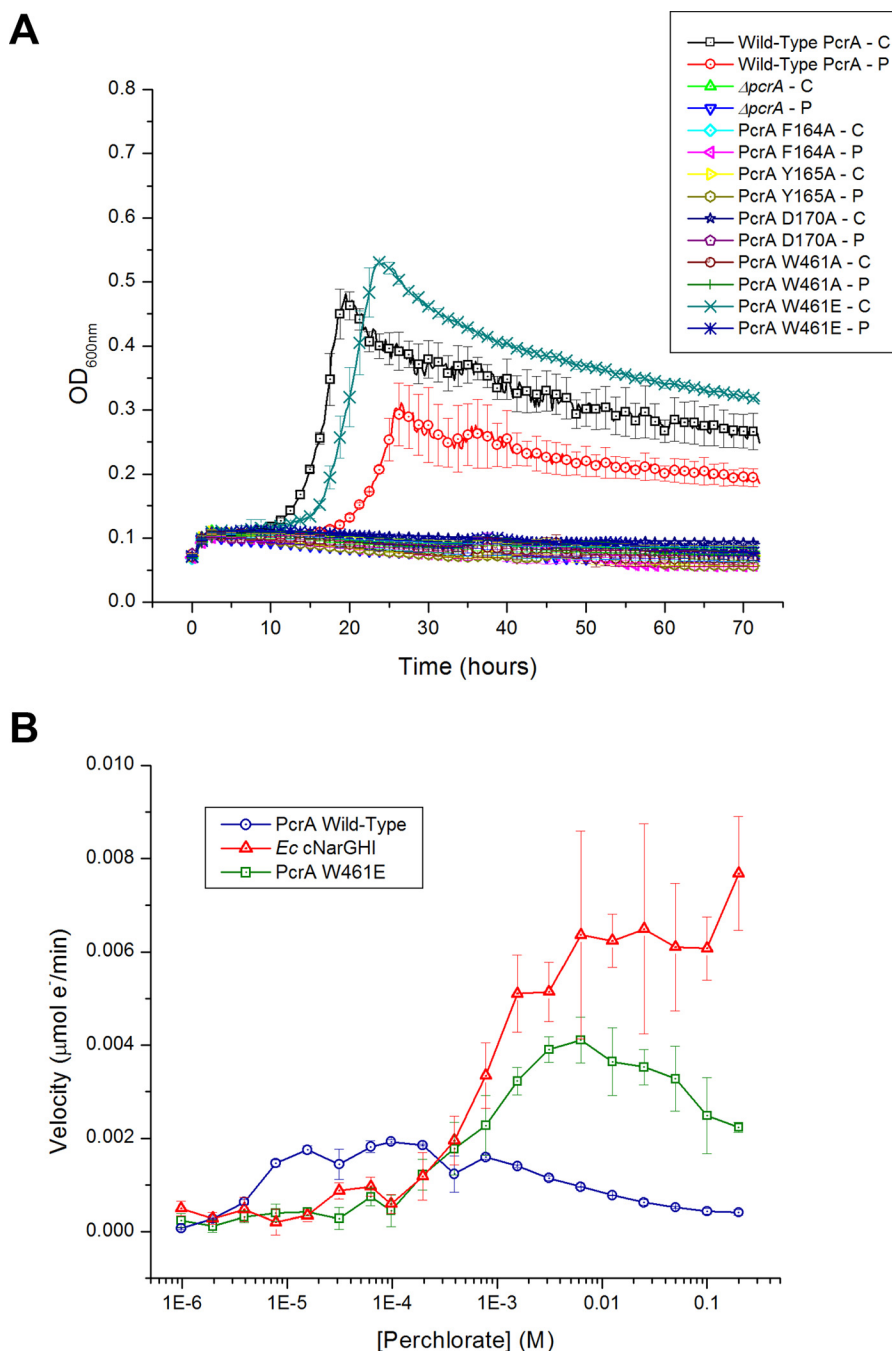


FIGURE 5. *A*, growth curves for wild-type *A. suillum* PS, $\Delta pcrA$, and PcrA mutant strains under perchlorate or chlorate reducing conditions. C is chlorate, and P is perchlorate. Error bars represent one standard deviation of triplicate growth experiments. *B*, steady-state kinetics of perchlorate reduction by PcrAB containing the PcrA W461E mutation in comparison with wild-type PcrAB and *EcNarGHI*. Error bars represent one standard deviation of triplicate assays.

molecule would then interact with one of the carboxylate oxygen atoms in residue Asp¹⁷⁰ and the side-chain nitrogen atom in residue Asn³⁵ via hydrogen bonding located in the open tunnel (Fig. 8C). Although monodentate and bidentate coordination of Asp to molybdenum atom have been seen in some independent NarGH structures (35, 36, 45), a carboxylate shift has only previously been proposed to protein metals other than molybdenum, such as iron, manganese, cadmium, and zinc (46). In combination with our results, this suggests that reduced PcrAB enables the Phe¹⁶⁴ to switch conformations to open active site access, and this gating may be triggered by the water

release from the active site to the tunnel due to the carboxylate shift of the Asp¹⁷⁰ residue (Fig. 8E).

Discussion

Our results reveal kinetic and structural features of PcrAB that establish it as a specialized member of the DMSO reductase superfamily of molybdopterin enzymes. We found that, although PcrA and NarG are closely related enzymes, they evolved from a common ancestor and diverged to tune their activities to the environments in which their host organisms

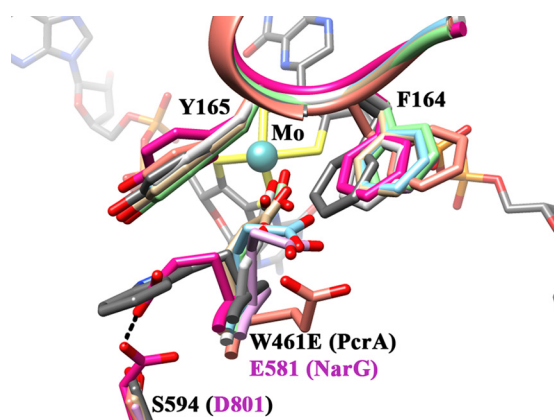


FIGURE 6. The comparison of PcrA W461E and *EccNarG* (Protein Data Bank code 1Q16) in the substrate access tunnel. The overlay of residue W461E PcrA from six chains of PcrA in the asymmetric unit of PcrAB was shown to be highly flexible due to the lack of hydrogen binding with Ser⁵⁹⁴ and the loss of π -stacking with Tyr¹⁶⁵. The corresponding residue Glu⁵⁸¹ in *EccNarG* (magenta) was stabilized by residue Asp⁸⁰¹ via hydrogen bonding. The wild-type PcrAB is shown in dark gray color with Trp⁴⁶¹ stabilized by π -stacking with Tyr¹⁶⁵.

lived. In most environments, perchlorate is present in low micromolar concentrations (47). Thus, a perchlorate reductase would experience selective pressure to evolve a relatively high affinity for perchlorate, whereas a nitrate reductase would experience selective pressure to maintain a relatively low affinity for perchlorate due to the toxicity of chlorite in a non-perchlorate-respiring organism lacking a chlorite dismutase. Consistently, below 200 μM perchlorate, PcrA is a more proficient perchlorate reductase than *EccNarG*. Importantly, the PcrA substrate inhibition by perchlorate does not impact catalytic efficiency at perchlorate concentrations <200 μM .

The unique biophysical parameters (low K_m and substrate inhibition by perchlorate) of the PcrAB are putatively associated with differences in key aromatic residues in the substrate access tunnel of each enzyme. Gate residues (Phe¹⁶⁴, Tyr¹⁶⁵, and Trp⁴⁶¹) were identified from crystal structures of PcrAB. The presence of the Phe¹⁶⁴ residue in PcrA corresponding to Phe²¹⁶ in *EccNarG* suggests this alone cannot explain the low K_m and substrate inhibition. However, the additional aromatic residue Trp⁴⁶¹ in PcrA, which is a Glu in NarG, might serve as a constrictive factor. A constricted active site with more possible contacts between protein and substrate is associated with higher affinity substrate binding (48). As such, the constrictive gate and tunnel positive electrostatic potential could combine to increase PcrA substrate affinities. Consistently, the PcrA W461E mutation partially shifted the PcrAB steady-state kinetics parameters toward those of *EccNarG*. Interestingly, the W461E mutation not only increased the perchlorate K_m by 50-fold but also partially alleviated substrate inhibition. Previously, nitrate inhibition was noted for the periplasmic nitrate reductase NapAB, purportedly due to excess substrate entering the active site tunnel and blocking water release (49). A more constricted active site such as that in PcrA would decrease both the on-rate and off-rate of substrates, products, and intermediates from the active site. Substrate inhibition would therefore be more dramatic when an enzyme must transition through multiple reaction cycles, such as perchlorate reduction, as

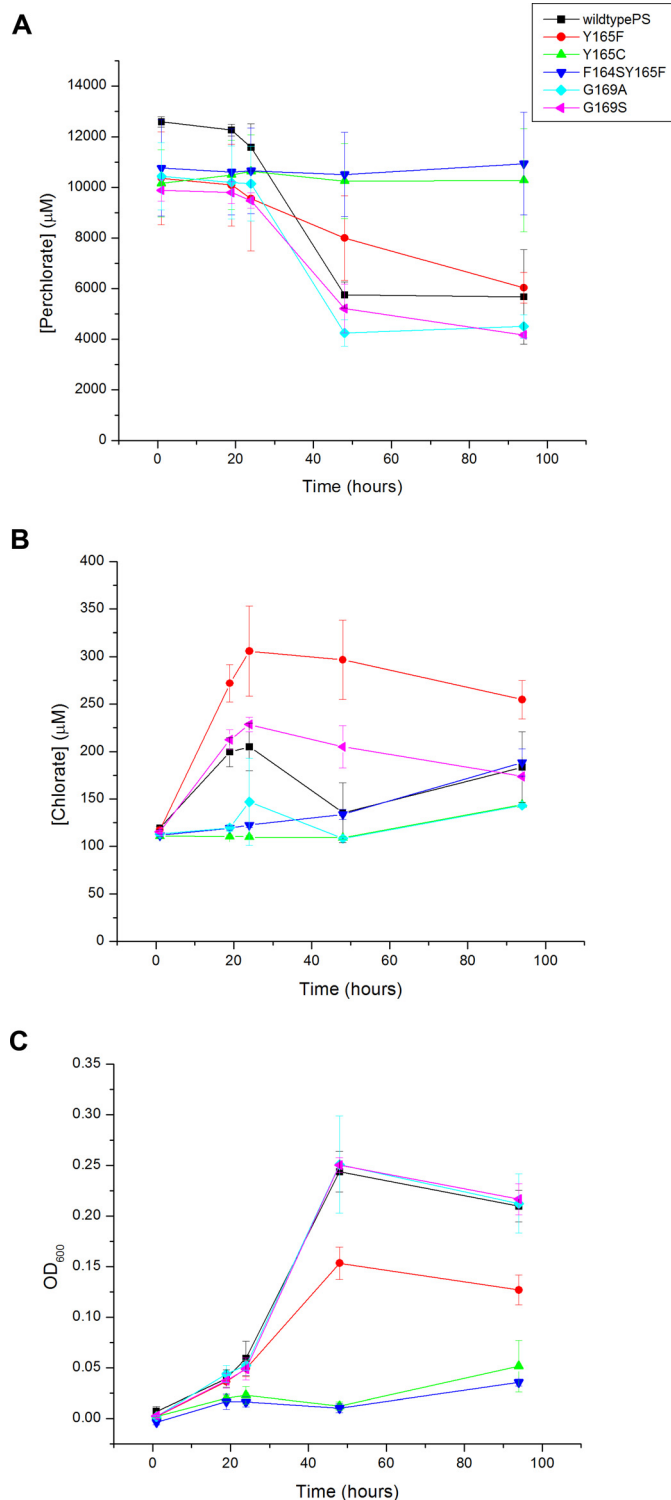


FIGURE 7. Chlorate accumulation experiments of wild-type *A. suillum* PS and *A. suillum* PS cells containing *pcrA* mutations monitoring chlorate (A), perchlorate (B), and OD₆₀₀ (C). Error bars represent the standard deviation of three replicate experiments.

water must leave the active site after the reduction of perchlorate to chlorate before chlorate can bind to the molybdenum atom for subsequent turnover. The lack of substrate inhibition for chlorate and nitrate turnover in wild-type PcrAB, PcrA W461E, and *EccNarG* is consistent with this hypothesis.

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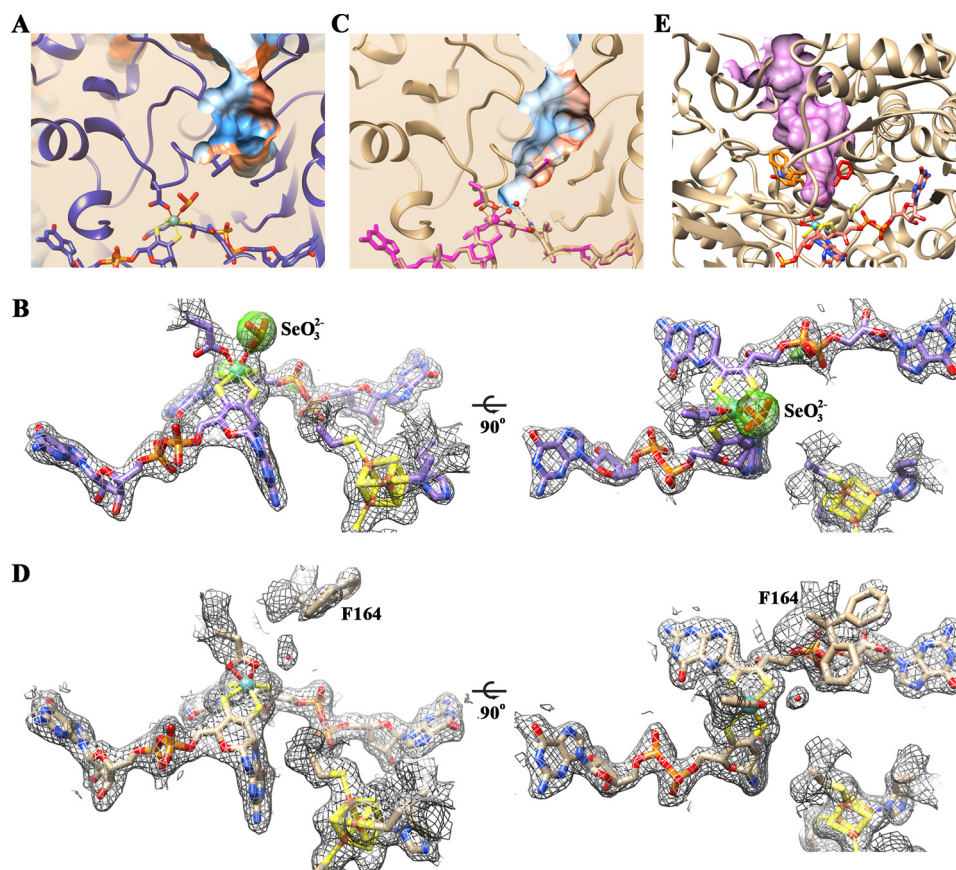


FIGURE 8. Reduced PcrAB structures with substrate analog bound and open and closed conformation of PcrA Phe¹⁶⁴. *A*, active site of PcrA (blue) with selenite (SeO_3^{2-}) coordinated to the molybdenum atom. The occupancy of selenite in the structure is 0.6. The surfaces in *A* and *C* are colored in a hydrophobicity scale (26) where red is the most hydrophobic and blue is the most hydrophilic. *B*, electron density of $2|F_o| - |F_c|$ simulated annealing omit maps at 1.5σ contour level of PcrA cofactors and active site for reduced PcrAB with selenite bound. The anomalous map is shown as a green surface at 3σ contour level. *C*, overlay of the active sites of oxidized (magenta) and reduced (tan) PcrA along with the tunnel to the active site. The reduced PcrA structure reveals dual conformations of the Phe¹⁶⁴ side chain in open and closed states. The tunnel is shown in the open conformation of Phe¹⁶⁴. The water molecule in reduced PcrA (tan) is located inside the open tunnel and hydrogen-bonded with residues Asp¹⁷⁰ and Asn³⁵. *D*, electron density of $2|F_o| - |F_c|$ simulated annealing omit maps at 1.5σ contour level of PcrA cofactors and active site for reduced PcrAB with the side chain of Phe¹⁶⁴ in the open and closed conformations. *E*, structure of the open tunnel in PcrA with Phe¹⁶⁴ in the open tunnel conformation shown as a surface (light purple).

The kinetic observations and unique structural features we identified for PcrAB along with previous publications pertaining to similar molybdenum-containing enzymes (50) support the proposal of a specific mechanism for (per)chlorate reduction by PcrAB (Fig. 9, steps *A–I*). Initially, the Asp residue (Asp¹⁷⁰) is coordinated with an oxygen atom bound to the Mo(VI) atom in a monodentate fashion at the active site with a closed gate (*A*). Previous work with DMSO reductase from *Rhodobacter sphaeroides* has shown that this bound oxygen atom is most likely in a Mo(VI)=O configuration (51). Two sequential proton-coupled electron transfer events reduce the active site through a Mo(V)-OH intermediate (*B*) to Mo(IV) (*C*). Upon $2e^-$ reduction of the active site, the bound water is released from the molybdenum atom and seems to be trapped at the active site by Asp¹⁷⁰ and Asn³⁵ (Fig. 8C). Concomitantly, Asp¹⁷⁰ shifts to a bidentate coordination of the Mo(IV) atom as seen in Fig. 7C. Phe¹⁶⁴ can now swing to open conformation in the reduced enzyme (Fig. 8C). This bidentate coordination of the Asp¹⁷⁰ is a possible trigger that leads to an Phe¹⁶⁴ conformational shift and opening of the tunnel to the active site, allowing water exit and substrate entry. At this point, perchlorate can bind to the active site Mo(IV), which causes the Asp residue to

shift back to monodentate coordination and the gate to close (*D*) as seen in Fig. 8A. Next, electrons are transferred from the reduced active site to the bound perchlorate, resulting in an oxygen atom abstraction and generation of chlorate and oxo-Mo(VI) (*E*). At this point, two additional proton-coupled electron transfer events would occur, forming *F* and subsequently *G*. Between *F* and *G*, we propose that chlorate and water remain trapped in the active site until Asp¹⁷⁰ shifts to a bidentate coordination on the reduced molybdenum and triggers Phe¹⁶⁴ to open the gate. Hypothetically, it is this step that is most likely to account for substrate inhibition during perchlorate turnover. Chlorate is a larger product than that formed in the reduction of other oxyanions, and excess perchlorate in the substrate access tunnel may hinder water exit from such a crowded and constricted region. This situation resembles the proposed substrate inhibition mechanism for NapA (49). The chlorate that has been shown to be accumulated during perchlorate turnover by PS is most likely released from *G* to reform *C* when the gate reopens as the active site becomes reduced for a second time in the catalytic cycle. When chlorate binds to the reduced active site to form *H* as seen in Fig. 8A, an additional oxygen atom abstraction occurs, reducing chlorate to chlorite (*I*), which will

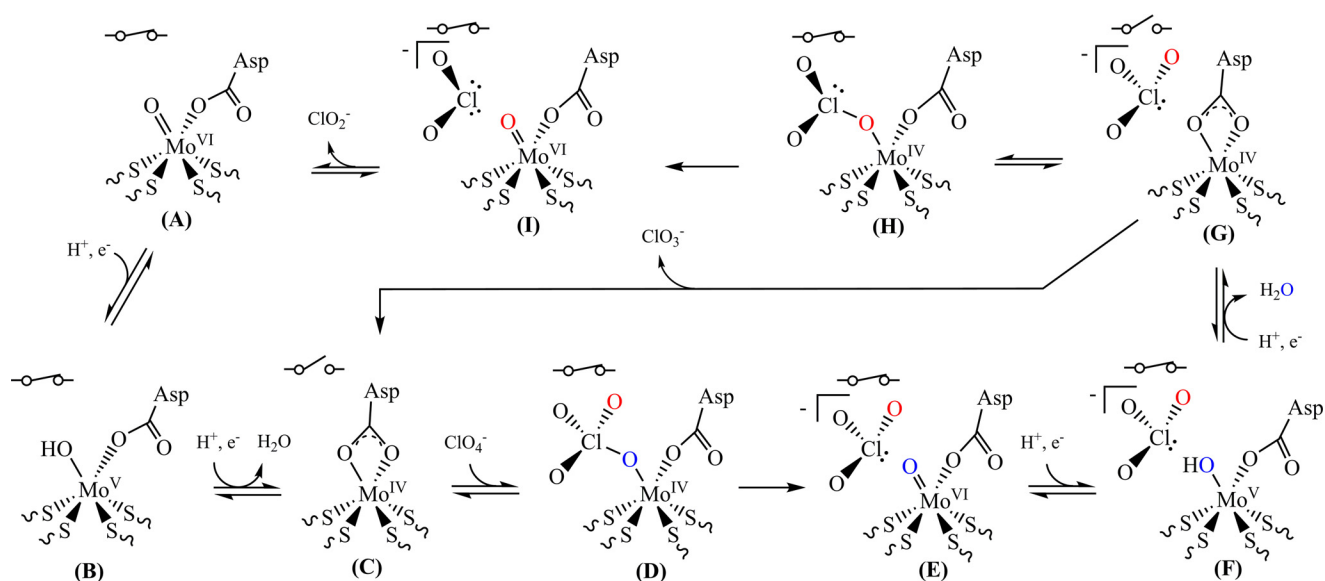


FIGURE 9. **Proposed mechanism for (per)chlorate reduction by PcrAB (A–I).** The Asp residue corresponds to Asp¹⁷⁰ as shown in the oxidized and reduced crystal structures of PcrAB. The symbols for a closed or an open switch represent a closed or open gate, respectively.

diffuse out of the active site to reform A. Notably, this proposed mechanism maintains the molybdenum atom in a six-coordinate geometry throughout the entire catalytic cycle. Although the mechanism is consistent with our results, one main question remains to be answered. Because the open conformation of Phe¹⁶⁴ in PcrA was only seen in the reduced structure of PcrAB, is the gating mechanism truly a redox-mediated process? Furthermore, the confirmation of the aforementioned discrete intermediates and the potential roles of other residues or components of the active site during catalysis are all currently being investigated.

Author Contributions—J. D. C. conceived of and coordinated the study. J. D. C. and M. D. Y. co-wrote the paper. M. D. Y. performed all protein purifications and kinetic analysis. C.-L. T. collected and solved all crystallographic data and contributed to writing the paper. I. C. C. performed and analyzed the phylogenetic data shown in Fig. 4. H. K. C., S. A. R., J. A. T., and J. D. C. made significant contributions to experimental design and interpretation. A. P. M. collected and analyzed data shown in Table 1. P. S. G.-P. and A. W. designed, constructed, and tested mutant enzymes. All authors reviewed the results and approved the final version of the manuscript.

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