Structural basis and evolution of redox regulation in plant adenosine-5'-phosphosulfate kinase

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Adenosine-5'-phosphosulfate (APS) kinase (APSK) catalyzes the phosphorylation of APS to 3'-phospho-APS (PAPS). In Arabidopsis thaliana, APSK is essential for reproductive viability and competes with APS reductase to partition sulfate between the primary and secondary branches of the sulfur assimilatory pathway; however, the biochemical regulation of APSK is poorly understood. The 1.8-Å resolution crystal structure of APSR from A. thaliana (AtAPSK) in complex with β , γ -imidoadenosine-5'-triphosphate, Mg²⁺, and APS provides a view of the Michaelis complex for this enzyme and reveals the presence of an intersubunit disulfide bond between Cys86 and Cys119. Functional analysis of AtAPSK demonstrates that reduction of Cys86-Cys119 resulted in a 17-fold higher k_{cat} / $K_{\rm m}$ and a 15-fold increase in $K_{\rm i}$ for substrate inhibition by APS compared with the oxidized enzyme. The C86A/C119A mutant was kinetically similar to the reduced WT enzyme. Gel- and activity-based titrations indicate that the midpoint potential of the disulfide in AtAPSK is comparable to that observed in APS reductase. Both cysteines are invariant among the APSK from plants, but not other organisms, which suggests redox-control as a unique regulatory feature of the plant APSK. Based on structural, functional, and sequence analyses, we propose that the redox-sensitive APSK evolved after bifurcation of the sulfur assimilatory pathway in the green plant lineage and that changes in redox environment resulting from oxidative stresses may affect partitioning of APS into the primary and secondary thiol metabolic routes by having opposing effects on APSK and APS reductase in plants.

enzyme kinetics | sulfur metabolism | X-ray crystallography | metabolic evolution

S ulfur is an essential element for all living organisms and is required for the biosynthesis of a diverse array of metabolites and macromolecules (1-4). Plants and prokaryotes are the primary assimilatory organisms that convert inorganic sulfate (SO_4^{2-}) , the predominant form of environmental sulfur, into physiologically useful forms of sulfur (5, 6). The metabolic organization of sulfur assimilation varies among plants and microbes (Fig. 1) (1-8). In yeast, fungi, and enterobacteria, including Escherichia coli, sulfate is incorporated into adenosine-5'-phosphate (APS), then converted to 3'-phospho-APS (PAPS) as a biologically "activated" compound that is reduced to sulfide (6, 7). In other sulfate-assimilating bacteria, such as Pseudomonas aeruginosa, APS can be used for reduction of sulfide (8). Plants possess bifurcated thiol metabolism pathways, which may reflect the metabolic needs of sessile organisms adapted to a range of environmental stresses and nutrient fluctuations (Fig. 1) (5). These pathways branch after formation of APS. The primary sulfur metabolic route in plants uses APS as the activated highenergy compound for sulfur reduction and the production of cysteine, which is crucial for the synthesis of proteins, methionine, iron-sulfur clusters, vitamin cofactors, and compounds that protect against oxidative stresses, including glutathione and phytochelatin peptides (9-12). Alternatively, APS can be converted into PAPS to provide a sulfate donor for the modification of multiple natural products, brassinosteroid and jasmonate hormones, phytosulfokines, and other sulfonated molecules (1315). Partitioning of sulfate at the branch point between primary (i.e., reductive) and secondary metabolic pathways in plants is controlled by APS reductase (APSR) and APS kinase (APSK), respectively (16, 17).

APSR catalyzes the glutathione-dependent reduction of APS to sulfite $(SO_3^{2^-})$ and AMP. Extensive studies of APSR in plants demonstrate a critical role for this enzyme in regulating flux through the primary sulfur assimilatory pathway (18–21). Modulation of activity by redox changes in the APSR from *Arabidopsis thaliana* (AtAPSR) occurs through a regulatory disulfide bond, which upon reduction attenuates activity (18). Additional evidence suggests that APSR is important for supplying reduced sulfur under stress conditions, such as nutrient deprivation and chilling (22). Conversely, application of cysteine and glutathione to *Arabidopsis* results in decreased APSR transcript levels and activity (23). In contrast to APSR, the role of APSK in plant thiol metabolism has only begun to be examined.

Recent studies in *Arabidopsis* revealed that APSK is essential for plant reproduction and viability (17, 24–27). Use of T-DNA knockout lines targeting each of the four AtAPSK isoforms established a connection between this branch of sulfur metabolism and glucosinolate biosynthesis (24, 27). Subsequent analysis of triple AtAPSK knockout lines demonstrated that AtAPSK1 alone could maintain wild-type levels of growth and development in *Arabidopsis* (26). Moreover, measurements of flux through the primary sulfur assimilation pathway was increased when APSK activity was reduced, which implies that alterations in either APSR or APSK activity aids in partitioning of sulfur between the two branches of thiol metabolism (17).

In plants, the biochemical regulation of APSK is not well understood, and we are aware of no study to date that has linked stress conditions to the control of its activity. APSK catalyzes the transfer of the γ -phosphate from ATP to the 3'-hydroxyl group of the APS adenine ring to yield PAPS and ADP. The reaction sequence suggested by kinetic studies of the APSK from Penicillium chrysogenum, E. coli, and A. thaliana follows an ordered mechanism with substrate inhibition by APS (28-30). Although no structural information for a plant APSK is available, X-ray crystal structures of the P. chrysogenum APSK and bifunctional ATP sulfurylase-APSK from human, Aquifex aeolicus, and Thiobacillus denitrificans have been determined and reveal a canonical α/β purine nucleotide binding fold (31–36). For the bifunctional human enzyme, the substrate inhibition by APS on APSK activity is linked to an approximately 20-aa N-terminal loop of low sequence homology (36) (Fig. S1); however, the general role of this region across different species remains

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Data deposition: The atomic coordinates and structure factors reported in this paper have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 3UIE).

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Fig. 1. Sulfur assimilatory pathways in plants (Left) and bacteria/cyanobacteria (Right). Various bacteria and cyanobacteria use APS kinase and PAPS reductase (black arrows) or APS reductase (gray arrow and text marked with an asterisk) for formation of sulfite.

unclear. In addition, earlier studies of AtAPSK suggest that oxidation reduces specific activity (29).

To better understand the molecular function of the plant APSK, we determined the crystal structure of AtAPSK isoform 1, lacking its chloroplast localization leader, in complex with β , γ -imidoadenosine-5'-triphosphate (AMP-PNP), Mg²⁺, and APS. Although the overall structure is similar to that of the APSK from fungi and humans, the AtAPSK·AMP-PNP·Mg2+·APS complex new insight on the reaction sequence of this enzyme and reveals an intersubunit disulfide bond formed between Cys86 in the Nterminal loop of one subunit and Cys119 in the adjacent monomer. Kinetic analysis and redox titrations of WT and the C86A/ C119A mutant of AtAPSK indicate that the disulfide bond forms at a physiologically relevant redox potential and affects catalytic efficiency and substrate inhibition by APS. Because sequence comparisons suggest a unique regulatory element of the plant APSK, we also functionally compare AtAPSK and the APSK from the cyanobacterium Synechocystis sp. PCC 6803, which is evolutionarily related to plant chloroplasts (37), to demonstrate that a homologue lacking the cysteines is not redox-sensitive. Based on structural, functional, and sequence analyses, we propose that evolution of a thiol-based redox switch in the plant APSK evolved with bifurcation of the sulfur assimilatory pathway later in the plant lineage, and that changes in redox environment resulting from oxidative stresses may coordinate flux between the primary and secondary thiol metabolic routes by modulating APSR and APSK activity in plants.

Results

Overall Structure. AtAPSK isoform 1 (residues 78–276) (38) lacking the 77-aa N-terminal chloroplast localization sequence was used for protein crystallography. The structure of AtAPSK in complex with AMP-PNP, Mg^{2+} , and APS was solved by molecular replacement using the *P. chrysogenum* APSK (32) as a search model with three monomers in the asymmetric unit (Fig. 2 and Table S1). Chain A forms a crystallographic dimer, and chains B and C yield a second noncrystallographic dimer.

The overall fold of each monomer in the dimer consists of a variable N-terminal region (residues 80–98), which includes $\alpha 1$ and entwines the adjacent monomer, a canonical α/β -purine nucleotidebinding domain ($\beta 1$ - $\alpha 2$ - $\beta 2$ - $\alpha 3$ - $\beta 3$ - $\alpha 4$ - $\beta 4$ - $\alpha 7$), and a small domain, which includes $\alpha 5$ and $\alpha 6$, that caps the nucleotide binding sites (Fig. 24). Although AtAPSK shares a common fold with the bifunctional PAPS synthetase from human (1.2 Å rmsd for 187 C_{α} atoms; 51% sequence identity) (35) and the *P. chrysogenum* APSK (1.3 Å rmsd. for 176 C_{α} atoms; 55% sequence identity) (32), the positioning of the N-terminal region differs in these structures (Fig. 2*B*). This region is characterized by an unstructured loop followed by a approximately 7-aa α -helix (α 1) in



Fig. 2. Structure of AtAPSK. (A) Ribbon diagram of the dimer. Each monomer is colored in pink and blue, respectively. The positions of AMP-PNP and APS are shown as stick molecules. The position of the C86-C119 disulfide bond is indicated in the left monomer. Secondary structure features are labeled in the right monomer. (*B*) Structural comparison of APSK structures. The overall AtAPSK structure is shown in gray. The position of the N-terminal region of the AtAPSK domain from the human PAPS synthetase-ADP-PAPS complex is nearly identical to that of AtAPSK. The N-terminal of *P. chrysogenum* APSK apoenzyme is in yellow and the position in the *P. chrysogenum* APSK in the AtAPSK active site. (*D*) The 2F₀-F_c omit map (1.5 σ) for Cys86-Cys119 disulfide bond. Note that the cysteine residues are found in alternate conformations.

various APSK structures (32–36) and displays sequence divergence between the APSK homologues from bacteria, cyanobacteria, green algae, fungi, plants, and human (Fig. S14).

In the initial $2F_o$ - F_c maps of the AtAPSK structure, clear electron density for AMP-PNP, APS, and Mg²⁺ was observed (Fig. 2*C*). Additional contiguous density was observed between Cys86 in monomer B and Cys119 of α 2 in monomer C (Fig. 2*D*), Cys86 in monomer C and Cys119 in monomer B, and Cys86 of monomer A and Cys119 in the crystallographic symmetry-mate of monomer A. The distance between the sulfur atoms of each cysteine falls within the range of expected bond length (~2.05 Å) for a disulfide linkage. The Cys86(A)-Cys119(A) and Cys86(B)-Cys119(C) disulfides were fully oxidized and the Cys86(C)-Cys119(B) disulfide was partially oxidized. The cysteines forming the disulfide are invariant across the APSK from plants and mosses, but are generally missing in the homologues from other organisms (Fig. S1).

Structure of Active Site and N-Terminal Loop. The structure of AtAPSK with a nonhydrolyzable ATP analogue (i.e., AMP-PNP), Mg^{2+} , and APS bound in the active site provides a view of the Michaelis complex in the APSK reaction sequence (Fig. 3). APS is locked in the active site through multiple binding contacts (Fig. 3). The adenine ring is stacked between Phe150 and Phe232. Arg141 and Asn158 provide bridging interactions between the sulfate and phosphate groups of the substrate. Arg155 and the backbone nitrogen of Ile181 also interact with the sulfate interaction with the ribose hydroxyl groups to position the 3'-OH proximal to the γ -phosphate group of AMP-PNP for the ensuing phosphoryl transfer reaction.

Binding of AMP-PNP is mediated almost exclusively through the phosphate moieties of the ligand to residues in the active site and an extensive water network centered on the bound Mg^{2+} ion (Fig. 3). The adenine ring is positioned between Pro257 and Arg215. Extending into the active site, phosphate groups of AMP-PNP form multiple contacts with main-chain atoms of residues 110 to 116 in the P-loop (39). The α -phosphate group is bound by interactions with the side-chain hydroxyl group and backbone nitrogen of Thr116 and the backbone nitrogen of Gly113. Lys114 contacts both the β - and γ -phosphates with Ser110 and Lys228, forming additional interactions to the γ phosphate. The β - and γ -phosphates of AMP-PNP, three water molecules, the hydroxyl-groups of Ser115 and Ser179, and the carboxylate of Asp136 octahedrally coordinate the Mg^{2+} ion. Asp136 and Asp138 form part of the DGDN-loop that bridges the APS and $ATP \cdot Mg^{2+}$ binding sites and are critical for catalytic activity in APSK (31, 35, 40).

Although not immediately in contact with ligands in the AtAPSK active site, part of the N-terminal loop (residues 80–84) of the adjacent monomer abuts the DGDN-loop and the side-chain of

F150

N158

F150

R141

1183-

N158



Fig. 3. Stereoview of the AtAPSK active site. The N-terminal loop of the adjacent monomer is shown as the dark gray loop, which includes His84.

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His84 hydrogen bonds to a water molecule in the network that coordinates the Mg²⁺ ion (Fig. 3). The structure of the AtAPSK Nterminal region is nearly identical to that of the APSK domain from human PAPS synthetase in complex with ADP and PAPS, but differs from the P. chrysogenum APSK apoenzyme and ADP APS complex (Fig. 2B). Changes in the N-terminal region of PAPS synthetase were suggested to promote conformational changes that stabilize interactions between the DGDN-loop and substrates during the catalytic cycle (33). Interestingly, the average B-factors for the N-terminal loop residues (i.e., 80–90) in the monomers containing the fully oxidized Cys86(A)-Cys119(A) and Cys86(B)-Cys119(C) disulfides were 39 $Å^2$ and 47 $Å^2$, respectively, which were lower than the average B-factor of 56 Å² in the monomer with the partially oxidized Cys86(C)-Cys119(B) disulfide bond. Crystallographic analysis of the Cys86-Cys119 disulfide bond in AtAPSK suggests that redox changes may alter the positioning and/ or mobility of the N-terminal loop near the active site and potentially affect enzymatic activity.

Cys86-Cys119 Disulfide and Redox Regulation. Formation of the Cys86-Cys119 disulfide linkage in the N-terminal region of AtAPSK may have a functional role in modulating enzymatic activity. Nonreducing SDS/PAGE analysis of AtAPSK incubated with either reduced DTT (DTT_{red}) or trans-4,5-dihydroxy-1,2-dithiane (DTT_{ox}) showed the enzyme as reduced monomeric (22 kDa) and oxidized dimeric (44 kDa) forms, respectively (Fig. 44). The same analysis using the C86A/C119A mutant confirmed that mutation of the cysteines prevented cross-linking of the two monomers (Fig. 44). Both the WT and C86A/C119A mutant AtAPSK migrated as dimers upon size-exclusion chromatography in the presence or absence of reducing agents.

Kinetic analysis of AtAPSK incubated in the presence of DTT_{red} or DTT_{ox} demonstrated that reduction of the disulfide bond resulted in enhanced catalytic efficiency (i.e., k_{cat}/K_m) and a decreased effect of substrate inhibition by APS (Table 1 and Fig. S2). Reduced AtAPSK was 17-fold more efficient and had



Fig. 4. Functional analysis of redox activity. (A) SDS/PAGE of WT and C86A/ C119A AtAPSK incubated in the absence or presence of 5 mM DTT_{red} or DTT_{ox}. Protein (0.3–1 µg) was stained with Coomassie blue. Arrows on the left side of the gel indicate positions of the molecular weight markers. Arrows on the right indicate positions of the oxidized dimeric and reduced monomeric forms. (*B*) Gel-based redox titration of AtAPSK. Protein was in cubated in increasing ratios of oxidized:reduced glutathione (GSSG:GSH; 20 mM total; E_{hr} –240 to –340 mV). (C) Comparison of gel-based (squares) and activity-based (circles) redox titrations. Solid lines represent fits to the Nernst equation.

Parameter	$k_{\rm cat}$, s ⁻¹	<i>K</i> _m , μΜ	<i>K</i> i, μΜ	$k_{\rm cat}/K_{\rm m}$, M ⁻¹ s ⁻¹ × 10 ⁸
AtAPSK _{red}	272 ± 39	0.48 ± 0.41	37.5 ± 6.9	5.67
AtAPSK _{ox}	14.1 ± 2.3	0.43 ± 0.26	2.51 ± 0.84	0.328
C86A/C119A _{red}	239 ± 44	1.90 ± 0.53	34.8 ± 2.7	1.25
C86A/C119A _{ox}	203 ± 31	0.87 ± 0.49	41.6 ± 7.1	2.33
SynAPSK _{red}	18.6 ± 4.5	0.33 ± 0.17	27.6 ± 3.5	0.56
SynAPSKov	15.9 ± 2.2	0.57 ± 0.26	21.6 ± 3.7	0.28

Average values \pm SE (n = 3) are shown.

a K_i value for APS 15-fold higher compared with the oxidized protein (Table 1). Moreover, the K_i values of the reduced protein increased dramatically with decreasing ATP concentrations (Fig. S2D); however, substrate inhibition by APS was comparable at all ATP concentrations for the oxidized enzyme (Fig. S2E). Steady-state kinetic analysis of the C86A/C119A mutant showed similar kinetic parameters for both forms of the enzyme that were comparable to those of the reduced AtAPSK (Table 1). The C86A/C119A mutant treated with DTT_{red} showed little difference in kinetic parameters with respect to AtAPSK_{red}. Similarly, treatment of the mutant with DTT_{ox} caused a modest 1.5-fold decrease in activity compared with DTT_{red}, but had little effect on either K_m or K_i . These results suggest that redox environment may have a role in regulating the activity of AtAPSK.

To determine the redox midpoint potential (E_m) for the Cys86-Cys119 disulfide, titrations using a gel-based system and activity assays were performed (Fig. 4 B and C). AtAPSK was incubated in solutions containing varied ratios of oxidized:reduced glutathione (GSSG:GSH; total concentration 20 mM). Aliquots from incubations (E_h of -340 to -240 mV) were analyzed by nonreducing SDS/PAGE and activity assays. Triplicate titrations of AtAPSK at pH 7.5 yielded average E_m values of -295 ± 12 mV and -286 ± 18 mV in the gel- and activity-based assays, respectively. At pH 7, the titrations yielded average $E_{\rm m}$ values of -249 ± 9 mV and -260 ± 17 mV in the gel- and activity-based assays, respectively. Compared with the Em values of other redox-regulated plant enzymes, which range from -390 to -237 mV (18, 41-45), these values suggest that changes in redox environment may modulate APSK activity in vivo, which could provide a strategy for modulating flux through the primary and secondary sulfur metabolism pathways in plants.

Characterization of APSK from *Synechocystis* **sp. PC 6803.** Comparison of APSK sequences indicates that the residues corresponding to Cys86 and Cys119 are invariant in the enzymes from monocot and dicot plants and the mosses *Selaginella moellendorffii* and *Physcomitrella patens* (Fig. S1*B*). Cys119 is conserved in the APSK from green alga *Chlorella variabilis* and *Chlamydomonas reinhardtii*, but is not found in the enzyme from cyanobacteria *Synechocystis* and *Cyanothece* (Fig. S1*A*). These sequence alignments suggest that redox-regulation of APSK appeared later in the lineage of photosynthetic organisms. To confirm that an evolutionarily earlier homologue is not redoxsensitive, the APSK from *Synechocystis* sp. PC 6803 was cloned, expressed, purified, and assayed. Steady-state kinetic analysis of SynAPSK in the presence of DTT_{red} or DTT_{ox} indicates that its activity does not change with redox environment (Table 1).

Discussion

The sulfur assimilation pathway in plants supports sulfur reduction and the synthesis of cysteine in the primary metabolic branch and the generation of sulfonated molecules for specialized metabolism in the secondary branch. Partitioning of sulfate at the branchpoint between primary and secondary metabolic pathways in plants is balanced by the activities of APSR and APSK (Fig. 1) (16, 17). Recent work reveals an essential role of APSK for the reproductive viability of *Arabidopsis* (17, 24–27), but the biochemical regulation of this enzyme in plants is not well understood. Structural and functional studies of AtAPSK provide insights on the molecular basis for PAPS formation and redox-control of this important metabolic branchpoint enzyme.

The reaction sequence of APSK is proposed to follow an ordered mechanism with ATP·Mg²⁺ binding first, followed by APS, catalysis, release of PAPS, and release of ADP·Mg²⁺ (46). Structures of the *P. chrysogenum* APSK apoenzyme (32), the *P. chrysogenum* APSK-ADP·Mg²⁺. APS dead-end complex (33), the ADP·Mg²⁺. PAPS product complex of the APSK domain from human PAPS synthetase (35), and AtAPSK·AMP-PNP·Mg²⁺. APS Michaelis complex (Figs. 2 and 3) clearly define the active site changes accompanying ATP-dependent phosphorylation of APS to PAPS. Shifts in the side chains of residues in the P-loop (Ser110-Thr116 in AtAPSK) position the phosphate backbone of ADP or AMP-PNP into the active site with additional conformational changes in residues of the APS site allowing for binding of this ligand. These alterations are accompanied by ordering of the "lid"-domain, which in AtAPSK contains α 5 and α 6, over the active site to constrain the flexibility of the structure.

These crystal structures suggest critical roles for the Mg^{2+} ion in catalysis and organization of the active site architecture. The interactions centered on the Mg^{2+} ion intricately link the ATP binding site, the active site water network, and residues in the catalytically essential DGDN-loop, which includes Asp136 and Asp138 of AtAPSK. Sekulic et al. (35) noted that Mg^{2+} binding induces the DGDN-loop to switch from an inactive to active conformation. Contacts from Asp136 to the Mg^{2+} and active site water molecules position Asp138 in proximity to the ribose hydroxyl groups of APS. In the AtAPSK structure, the carboxylate oxygens of Asp138 are 2.67 Å and 3.15 Å from the 3'-OH and 2'-OH groups of the APS ribose, respectively (Fig. 3). This suggests that the negatively charged oxygen of Asp138 serves as a general base in the catalytic mechanism to abstract a proton from the 3'hydroxyl group of APS and promotes its nucleophilic attack on the γ -phosphate group of ATP.

Phosphorylation of APS to yield PAPS requires an orchestrated series of structural changes to organize the APSK active site for catalysis. Comparisons of various APSK in complex with different ligands (Fig. 2B) suggest that the conformational flexibility of the N-terminal loop region likely affects these structural changes. Within the active site, this loop sterically contacts the DGDN-loop (Fig. 3) and is proposed to position catalytically essential residues to promote ligand binding and phosphoryl transfer (35, 36). Deletion of the N-terminal region from the APSK domain of human PAPS synthetase indicates that this region, which is not in direct contact with ligands in the active site, is responsible for substrate inhibition by APS (36). The AtAPSK structure reveals a regulatory control feature in the Nterminal loop, which is unique to plants.

Formation of a disulfide bond between Cys86 and Cys119 in the N-terminal loop region cross-links subunits of the AtAPSK dimer (Figs. 2D and 4), decreases catalytic efficiency, and enhances substrate inhibition by APS (Table 1). Comparisons of the kinetic properties of the reduced and oxidized forms of AtAPSK indicate that reduction of the disulfide yields a more active enzyme. Mutagenesis of the two cysteines in AtAPSK also mimics reduction of the disulfide, as demonstrated by activity assays (Table 1) and gel-based analysis (Fig. 4A). Moreover, redox titrations of the disulfide bond using gel-based and activitybased assays indicate that the E_m is within a physiological range (Fig. 4 *B* and *C*) (18, 41–45, 47). The redox E_m of the Cys86-Cys119 disulfide in AtAPSK is comparable to that determined for AtAPSR (E_m of -255 at pH 7). Interestingly, both AtAPSK and AtAPSR are closer to the redox midpoint for GSH/GSSG than to the reported values associated with plant thioredoxins (48–50). Structurally, changes in the oxidation state of Cys86 and Cys119 in AtAPSK likely alter the dynamic movements of the N-terminal loop to affect catalytic efficiency and substrate inhibition. Physiologically, the Cys86-Cys119 disulfide bond may act as a regulatory switch to coordinate flux between the primary and secondary branches of sulfur assimilation in *Arabidopsis* and other plants.

Several plant thiol metabolism enzymes [i.e., APSK, APSR, and glutamate-cysteine ligase (GCL); Fig. 1] possess cysteines that form disulfide bonds to regulate enzymatic activity in response to changes in redox environment (18, 43-45). This suggests that oxidative stresses may simultaneously affect multiple proteins across sulfur assimilation and metabolism in plants. Because APSR and APSK share similar redox E_m values and partition the flow of APS into different branches of sulfur metabolism (16, 17), coordinated biochemical regulation of their activities may be important to meet metabolic demands for sulfur reduction and/or production of PAPS for specialized sulfonation reactions. Redox regulation of these branchpoint proteins would provide a mechanism for controlling sulfur allocation between the two metabolic routes. For example, under oxidative stress conditions that increase demand for sulfur reduction to support glutathione synthesis for maintaining redox state, formation of the disulfides in APSR and APSK would increase and decrease their activities, respectively, and result in direction of APS into cysteine and glutathione synthesis and away from synthesis of PAPS. Moreover, previous studies demonstrate that formation of a key disulfide bond in the plant GCL under multiple oxidative stress conditions activates this enzyme to increase glutathione synthesis (42–45). Thus, both APSR and GCL in the primary thiol metabolic pathway would be active in response to stress and APSK activity attenuated in the secondary pathway. Further experimental studies in plants that analyze metabolite distribution and sulfur pathway flux in response to biotic and abiotic stresses that alter oxidation state is required to explore this potential biochemical regulatory mechanism.

Sequence comparisons of the APSK from plants, mosses, fungi, human, green algae, and cyanobacteria show that active site residues are highly conserved (24-29, 31-36) and that the Nterminal regions are divergent (Fig. S1), which may reflect the need for specialization of APSK regulation in organisms with differing metabolic demands for sulfur assimilation. Residues corresponding to Cys86 and Cys119 of AtAPSK are nearly invariant across the homologues from plants and mosses with some unicellular alga and yeast retaining Cys119; however, in the APSK from organisms lacking plastids, neither cysteine is found (Fig. S1B). In addition, of the four APSK isoforms in Arabidopsis (22), the disulfide cysteines are found in the three plastid-localized isoforms, but only Cys119 is conserved in the cytosolic isoform. Given that the chloroplast is a highly redox-active organelle (9, 10, 12), this difference between AtAPSK isoforms suggests the evolution of a specialized regulatory control in the enzymes of this organelle. Through the lineage from cyanobacteria to green algae to mosses and "higher" plants, there is also a shift in the organization of the sulfur assimilation pathway from a linear one in cyanobacteria to a branched pathway in the later evolving mosses and plants (Fig. 1) (5, 37). Thus, the redoxsensitive disulfide observed in the plant and moss APSK may be a later adaptation to the development of chloroplasts and branched sulfur metabolism pathways in these multicellular photosynthetic organisms, and reflects a greater need for regulatory control in response to more specialized metabolic demands for coordination of flux between the primary and secondary sulfur pathways by reciprocally modulating APSR and APSK activity in plants.

Materials and Methods

Protein Expression and Purification. Generation of the pET-28a-AtAPSKA77 bacterial expression vector, which encodes AtAPSK isoform 1 lacking the plastid localization sequence and with an N-terminal hexahistidine tag, was previously described (38). Transformed E. coli BL21(DE3) cells were grown at 37 °C in Terrific broth containing 50 µg mL⁻¹ kanamycin until an absorbance at 600 nm of approximately 0.8. After induction with 1 mM isopropyl 1-thio- β -D-galactopyranoside, the cultures were grown at 20 °C overnight. Cells were pelleted by centrifugation and resuspended in 50 mM Tris (pH 8.0), 500 mM NaCl, 20 mM imidazole, 1 mM β -mercaptoethanol (β ME), 10% (vol/vol) glycerol, and 1% Tween-20. After sonication and centrifugation, the supernatant was passed over a Ni²⁺-nitriloacetic acid (Qiagen) column. The column was washed with buffer minus Tween-20 and the bound protein eluted by using 250 mM imidazole in wash buffer. The eluent was dialyzed overnight at 4 °C against 25 mM Hepes (pH 7.5), 200 mM KCl, 5% glycerol, and 5 mM $\beta \text{ME},$ and then loaded onto a Superdex-200 26/60 HiLoad FPLC size-exclusion column equilibrated in the same buffer. Fractions corresponding to the major protein peak were pooled, and judged to be more than 95% pure by SDS/PAGE. Protein was concentrated (Amicon) to 10 mg mL⁻¹ with protein concentration determined by using a molar extinction coefficient (ϵ_{280nm} of 22,430 M⁻¹·cm⁻¹) calculated in ProtParam (http://web. expasy.org/protparam). Protein was dialyzed against 25 mM Hepes (pH 7.5), 200 mM KCl, 10% glycerol, and 5 mM BME, flash-frozen in liquid nitrogen, and stored at -80 °C. For crystallization and redox titrations, β ME was removed by buffer exchange.

Crystallography. Crystals of the AtAPSK·AMP-PNP·Mg²⁺·APS complex grew at 4 °C in hanging drops from a 1:1 ratio of protein preincubated with 5 mM APS, 5 mM AMP-PNP, and 10 mM MgCl₂ and crystallization buffer (100 mM Hepes, pH 7.5, 200 mM MgCl₂, and 15–17.5% PEG 2000). For data collection, crystals were transferred to cryoprotectant (mother liquor containing ligands and 20% glycerol) and then frozen in liquid nitrogen. Diffraction data (100 K) were collected at beam 19-ID of the Advanced Photon Source/ Argonne National Laboratory. Diffraction intensities were integrated, merged, and scaled by using the HKL3000 software suite (51). The structure of AtAPSK was determined by molecular replacement with PHASER (52) by using the structure of APSK from *P. chrysogenum* (PDB ID 1M7G) (32) as the search model. After iterative rounds of model building in COOT (53) and refinement in PHENIX (54), the R-factors converged to those reported in Table S1. Coordinates and structure factors for the AtAPSK-AMP-PNP-Mg²⁺.APS complex have been deposited in the Protein Data Bank.

Mutagenesis. The AtAPSK C86A/C119A mutant was generated using the QuikChange PCR method (Stratagene) with pET-28a-AtAPSK Δ 77 as the template. Two pairs of oligonucleotides (Table S2) were used to sequentially generate the double-mutant. Mutations were confirmed by sequencing (Washington University DNA Sequencing Facility). Protein expression and purification were performed as described for WT protein.

Enzyme Assays and Redox Titrations. Enzymatic activity of purified WT and mutant AtAPSK was determined spectrophotometrically by using a coupled assay (30). Steady-state kinetic parameters were determined by initial velocity experiments with data fitted to a general substrate inhibition model:

$$V = (V_{max}[S])/K_M + [S]*((1 + [S])/K_i)$$
 [1]

The effect of redox potential on AtAPSK activity was examined with assays containing defined ratios of reduced (i.e., GSH) and oxidized (i.e., GSSG) glutathione (Sigma-Aldrich). Protein (5 μ M) was equilibrated in degassed 100 mM Hepes (pH 7.5 or 7.0), 200 mM KCl, 5% glycerol, and 20 mM GSH/GSSG for 60 min at 25 °C. Aliquots were removed and transferred to the coupled assay (final protein concentration of 10–30 nM), and initial velocities measured. Values for E_m were determined by fitting titration data to the Nernst equation by using Kaleidagraph (Synergy Software):

$$\mathbf{E}_{h} = \mathbf{E}_{m} + (\mathbf{R}\mathbf{T}/\mathbf{n}\mathbf{F}) \left(\mathbf{In} \left([\mathbf{G}\mathbf{S}\mathbf{S}\mathbf{G}] / [\mathbf{G}\mathbf{S}\mathbf{H}]^{2} \right) \right)$$
[2]

with an RT/F of 25.7 mV (44, 47) and n = 2. The effect of redox potental on AtAPSK was also evaluated by nonreducing SDS/PAGE. Protein was equilibrated with varying ratios of GSSG/GSH, DTT_{red}, or DTT_{ox}, and then separated into reduced and oxidized forms by SDS/PAGE. The fraction of reduced vs. oxidized AtAPSK was calculated by quantifying the fractions of oxidized cross-linked (44 kDa) and reduced monomeric (22 kDa) species by densitometry (Quantity One software; BioRad).

Cloning, Expression, Purification, and Analysis of APSK from *Synechocystis* **sp. PCC 6803.** The cDNA encoding APSK from *Synechocystis* **sp** PCC. 6803 was PCRamplified by using PfuUltraHF polymerase and gene-specific oligonucleotides (Table S2). The PCR product was digested and cloned into the Nhel and EcoRI sites of pET28a. Protein expression, purification, and activity assays were performed as described for AtAPSK.

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