The biosynthesis of many sulfur-containing molecules depends on cysteine as a sulfur source. Both the cysteine desulfurase (CD) and rhodanese (Rhd) domain–containing protein families participate in the trafficking of sulfur for various metabolic pathways in bacteria and human, but their connection is not yet described in plants. The existence of natural chimeric proteins containing both CD and Rhd domains in specific bacterial genera, however, suggests a general interaction between these proteins. We report here the biochemical relationships between two cystolic proteins from Arabidopsis thaliana, a Rhd domain–containing protein, the sulfurtransferase 18 (STR18), and a CD isoform referred to as ABA3, and compare these biochemical features to those of a natural CD–Rhd fusion protein from the bacterium Pseudorhodoferax sp. We observed that the bacterial enzyme is bifunctional exhibiting both CD and STR activities using l-cysteine and thiosulfate as sulfur donors but preferentially using l-cysteine to catalyze transpersulfidation reactions. In vitro activity assays and mass spectrometry analyses revealed that STR18 stimulates the CD activity of ABA3 by reducing the intermediate persulphide on its catalytic cysteine, thereby accelerating the overall transfer reaction. We also show that both proteins interact in planta and form an efficient sulfur relay system, whereby STR18 catalyzes transpersulfidation reactions from ABA3 to the model acceptor protein roGFP2. In conclusion, the ABA3–STR18 couple likely represents an uncharacterized pathway of sulfur trafficking in the cytosol of plant cells, independent of ABA3 function in molybdenum cofactor maturation.

Sulfur is an essential macronutrient playing pivotal roles in the physiology and development of all organisms as it is present not only in two amino acids, cysteine and methionine, but also in many other molecules, such as sulfolipids, thionucleosides, vitamins (thiamin, biotin, and lipoic acid), and iron–sulfur (Fe–S) clusters or molybdenum cofactors (Moco)

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ABA3 provides sulfur to STR18

In plants, mitochondrial NFS1 and plastidial NFS2 are the CDs providing the sulfur required for Fe–S cluster assembly in both mitochondria and cytosol or in chloroplasts, respectively (14, 15). The CD activity of NFS2 is relatively low in the absence of the specific SUFE1–3 activators (16–18). A third CD isoform, ABA3, is localized in the cytosol of plants and involved in Moco sulfuration, thus participating in the activation of aldehyde oxidase and xanthine dehydrogenase. These Moco-containing enzymes are involved in abscisic acid biosynthesis and purine degradation, respectively (19, 20). ABA3 is formed by two domains, an N-terminal aminotransferase class V domain (InterPro: IPR000192) responsible for the CD activity, as in NFS1 and NFS2. In addition, the protein possesses a C-terminal MOSC domain (InterPro: IPR005302 and IPR005303) responsible for the final incorporation of a sulfur atom into the Moco precursor (20, 21).

Similar transpersulfidation reactions between CDs and other sulfur carrier proteins occur during the biosynthesis of sulfur-containing molecules. Among these sulfur carrier proteins are sulfurtransferases (STRs), widespread enzymes present in bacteria, archaea, and eukarya. They possess a characteristic rhodanese (Rhd) domain usually containing a conserved catalytic cysteine present in a specific Cys-X-X-Gly-X-Arg signature (22, 23). This cysteine is mandatory for the catalytic activity of STRs since a cysteine persulfuration intermediate is formed during transpersulfidation reactions. Three different STR classes have been defined with respect to their modular organizations and substrate specificities (22–24). STRs with a single Rhd domain use preferentially thiosulfate as a sulfur donor in vitro and are referred to as thiosulfate-STRs (TSTs; InterPro: IPR001307). Those possessing two Rhd domains use preferentially 3-mercaptopyrurate (3-MP) as a sulfur donor in vitro and were named 3-MP-STRs (InterPro: IPR036873). Mammals possess an additional STR isoform with two Rhd domains, named Rhobov, which uses sulfite and glutathione persulfide to synthesize thiosulfate. Additional STR proteins contain one Rhd domain fused to one or several protein domains with another function conferring them specific roles (22, 23).

Examples of interaction between CDs and STRs in non-plant organisms have suggested a hub function for a cysteine persulfuration intermediate during the biosynthesis of Moco from IscS to the STRs, ThiI or YnjE, is required for thiamine and IPR036873). Mammals possess an additional STR isoform with two Rhd domains, named Rhobov, which uses sulfite and glutathione persulfide to synthesize thiosulfate. Additional STR proteins contain one Rhd domain fused to one or several protein domains with another function conferring them specific roles (22, 23).

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Results

The natural CD–Rhd fusion protein of Pseudorhodoferax sp. is a bifunctional enzyme

Genomic analyses like gene clustering, gene co-occurrence, or gene fusion are powerful tools to predict functional associations. For instance, the existence of natural fusions in some organisms often reflects a functional interaction in other organisms in which the constituting protein domains are expressed as separate proteins. By interrogating the STRING database (https://string-db.org/) using the COG1104 specific to CDs, we have noticed the existence of both adjacent CD and Rhd genes and natural CD–Rhd fusion genes/proteins in several bacteria. We focused our attention on a CD–Rhd isoform from Pseudorhodoferax sp. Leaf274. The presence of conserved catalytic cysteine residues in each protein domain suggests that this protein should possess both CD and TST-type STR activities (Fig. 1A) (8, 22). The corresponding His-tagged recombinant protein exhibited a yellow color after purification. In addition, the UV–visible absorption spectrum exhibited an absorption band at 418 nm characteristic for a bound-PLP cofactor as in characterized CDs (Fig. 1B) (9). Analytical gel filtration analysis demonstrated that CD–Rhd eluted predominantly in a peak corresponding to an apparent volume/molecular mass of 108 kDa (Fig. 1C). From the theoretical molecular mass of CD–Rhd (54 kDa), we concluded that this protein formed homodimers as observed for other CDs (7, 8).

We have then evaluated the capability of the fusion protein to use L-cysteine or thiosulfate as substrates and determined the kinetic parameters of the reactions. The CD activity (i.e., cysteine desulfuration with the concomitant formation of a persulfide on catalytic cysteine) was monitored under steady-state conditions by measuring the release of H2S from the

NFS1 (27). Moreover, TUM1 proteins ensure sulfur transfer to participates in the biosynthesis of Moco, receiving sulfur from cytosol of yeast and human. The human STR isoform, TUM1, respectively (25, 26). A similar sulfur relay system exists in the different STR classes have been de
Figure 1. *Pseudorhodoferax* CD–Rhd fusion has a dual activity profile. A, modular organization of the *Pseudorhodoferax* CD–Rhd fusion (WP_056898193.1) presenting the position of the presumed catalytic cysteines of both CD and Rhd domains. B, UV–visible absorption spectrum of the purified N-terminal His-tagged recombinant CD–Rhd recorded in a 30 mM Tris–HCl (pH 8.0) buffer. C, analytical gel filtration (Superdex S200 10/300 column; GE Healthcare) of His-tagged recombinant CD–Rhd (100 μg). The presence of the polypeptide and PLP cofactor has been detected by measuring the absorbance at 280 nm (dark line) and 418 nm (green line), respectively. The apparent molecular weight of CD–Rhd was estimated from the separation of the indicated standards. D, steady-state kinetic parameters of the CD activity. Reactions were performed in the presence of 10 nM CD–Rhd, increasing concentrations of L-cysteine (0–2 mM), and in the presence of various reductants, either 5 mM of DTT (blue squares), or 5 mM GSH (green circles), or 5 mM β-mercaptoethanol (red triangles). The data are represented as mean ± SD of three independent experiments. E, steady-state kinetic parameters of the ABA3 provides sulfur to STR18.
ABA3 provides sulfur to STR18

persulfated protein in the presence of chemical or physiological reducing acceptors (Fig. 1D). Catalytic efficiencies ($k_{cat}/K_M$) of 2.2 × 10^8 M^-1 s^-1, 1.8 × 10^8 M^-1 s^-1, and 2.2 × 10^8 M^-1 s^-1 have been measured in the presence of DTT, GSH, and β-mercaptoethanol (β-ME), respectively (Table 1), thus validating the CD activity of the fusion. The activity of the Rhd domain was also evaluated by monitoring the release of H_2S in the presence of β-ME but providing thiosulfate as the canonical substrate of TST-type STRs. The catalytic efficiency of the reaction was 2.7 × 10^8 M^-1 s^-1, and the apparent $K_M$ value for thiosulfate was 756 ± 51 μM (Fig. 1E and Table 1), thus validating the STR activity of the fusion. Our results demonstrate that the CD–Rhd chimeric protein from *Pseudorhodoferax* is bifunctional having dual in vitro activities using l-cysteine or thiosulfate as a sulfur donor (Fig. 1F).

Rhd domain promotes CD activity and transpersulfidation reactions

Although the CD–Rhd protein is bifunctional, its CD activity was approximately eightfold higher than its TST activity. The Rhd domain only possesses the characteristic catalytic cysteine (Cys466) and thus could represent a sulfur acceptor for the CD domain. To determine the importance of the cysteine in the Rhd domain for the recorded CD activity, we analyzed the biochemical properties of a CD–Rhd C466S variant (Fig. S1 and Table 1). Similar to CD–Rhd, the His-tagged CD–Rhd C466S recombinant protein exhibited a UV–visible absorption spectrum with two absorption bands at 280 and 418 nm (Fig. S1A) and existed as a dimer in solution (Fig. S1B). The absence of TST activity confirmed that the cysteine of the Rhd domain is mandatory for this activity (Fig. S1C). Concerning CD activity, the CD–Rhd C466S variant is still active despite its catalytic efficiency was ~10-fold lower in the presence of GSH or β-ME (1.6 × 10^8 M^-1 s^-1) compared with the activity of CD–Rhd. This is notably explained by a decrease of the apparent $k_{cat}$ value by a factor of 10 (Fig. S1D and Table 1). On the contrary, CD activity did not significantly vary in the presence of DTT because the decrease of the apparent $k_{cat}$ by a factor of 6 is compensated by a change in the apparent $K_M$ value for l-cysteine (Fig. S1D and Table 1). The decrease in the turnover number of CD–Rhd C466S suggests that the Rhd domain stimulates the CD activity of the fusion with the catalytic cysteine probably serving as a persulfide-relay system.

To study the potential sulfur relay role of the Rhd domain, we investigated the capability of CD–Rhd and its C466S variant to transfer a sulfur atom to a protein substrate. In the absence of known CD–Rhd partners, we used roGFP2 that has been recently shown to act as an efficient sulfur acceptor for Rhd domain–containing proteins (30). We have first tested the oxidation of a prerduced roGFP2 in the presence of CD–Rhd or its C466S variant and l-cysteine (Fig. 2A). l-cysteine alone had no effect on roGFP2 oxidation. On the contrary, the combination of l-cysteine with CD–Rhd led to an efficient roGFP2 oxidation (Fig. 2A). The reaction is much slower in the presence of the CD–Rhd C466S variant. These results demonstrated that a functional Rhd domain is necessary for an optimal reaction. The catalytic cysteine of the Rhd domain likely promotes the sulfur transfer from the catalytic cysteine of the CD domain to roGFP2 catalyzing a transpersulfidation reaction between both proteins. Considering the TST activity of CD–Rhd, we performed similar experiments using thiosulfate instead of l-cysteine as a sulfur donor. Thiosulfate alone had no effect on roGFP2, and the CD–Rhd fusion was very poorly able to catalyze roGFP2 oxidation (Fig. 2B). No oxidation was observed with the CD–Rhd C466S variant (Fig. 2B). Altogether, these findings indicate that CD–Rhd preferentially uses l-cysteine as sulfur donor, and the Rhd domain promotes transpersulfidation reaction between CD domain and protein partner.

A. thaliana STR18 stimulates the CD activity of ABA3

The results obtained with this chimeric protein prompted us to investigate the interaction between *A. thaliana* orthodoxers that exist as separate proteins, focusing on the cytosolic ABA3 and STR18, a single Rhd domain–containing protein (21, 23, 31–33). STR18 possesses two cysteine residues, Cys47 and Cys89, the latter corresponding to the catalytic cysteine present in the Cys-X-Gly-X-Arg signature typical of the Rhd domain (23, 32). TST activity assays were first performed with STR18 and both C47S and C89S variants. The STR18 C89S variant was inactive, whereas the catalytic efficiency of the STR18 C47S variant was only marginally affected (Fig. S2 and Table 2). This indicated that STR18 exhibits TST activity, Cys89 is mandatory, and Cys47 is dispensable. The influence of STR18 and its variants on CD activity of ABA was then evaluated. The turnover for cysteine desulfuration by ABA3 was approximately twofold higher in the presence of STR18 (0.66 versus 1.39 mol sulfur mole enz^-1 min^-1), indicating a stimulating effect of STR18 on the CD activity of ABA3 (Fig. 3A). As expected, the STR18 C47S variant also stimulated the CD activity of ABA3, whereas the STR18 C89S variant did not (Fig. 3A).

These results prompted us to investigate the interaction of ABA3 with STR18 and its monocysteinic variants and to determine the steady-state kinetic parameters associated with CD activity (Fig. 3B and Table 3). In the absence of STR18, the apparent $K_M$ values of ABA3 for l-cysteine was 23 ± 6 μM, and the deduced catalytic efficiency was 680 M^-1 s^-1 (Fig. 3B and Table 3). In the presence of STR18, ABA3 was fivefold more efficient, and this is notably explained by a decrease of the thiosulfate sulfurtransferase activity. Reactions were performed in the presence of 100 nM CD–Rhd, increasing concentrations of thiosulfate (0–5 mM), and 5 mM β-mercaptoethanol. The data are represented as mean ± SD of three independent experiments. F, proposed mechanism for the CD and sulfurtransferase activities of the CD–Rhd fusion. For the CD activity, the catalytic cysteine (Cys334) of the CD domain (in blue) catalyzes the cysteine desulfuration, leading to its persulfidation and the concomitant release of alanine (1). Then, the sulfur atom is transferred to the catalytic cysteine (Cys466) of the Rhd domain (in green) through a transpersulfidation reaction (2). Cys466 is also responsible for thiosulfate sulfurtransferase activity through the conversion of thiosulfate into sulfite (3). Under persulfidated form, Rhd domain promotes sulfur transfer to acceptors leading to the reduction of CD–Rhd (4). CD, cysteine desulfurase; PLP, pyridoxal 5’-phosphate; Rhd, rhodanese.
apparent \( K_M \) value for L-cysteine by a factor of 3 (Fig. 3B and Table 3). Similar kinetic parameters were obtained using the STR18 C47S variant, whereas, as expected, the CD activity of ABA3 was not stimulated in the presence of the STR18 C89S variant as illustrated by the apparent \( K_M \) value for L-cysteine and the \( k_{cat}/K_M \) value close to those determined for ABA3 alone (Fig. 3B and Table 3).

Altogether, these data indicate that STR18 stimulates the CD activity of ABA3 by increasing ABA3 affinity for L-cysteine and of increasing STR18 concentrations. This allowed us to determine an apparent \( K_M \) value of ABA3 for STR18 of 1.2 ± 0.2 μM (Fig. 3C and Table 3). This \( K_M \) value in the low micromolar range indicates that the ABA3–STR18 interaction may be physiologically relevant.

**STR18 is persulfidated upon reaction with ABA3**

As L-cysteine is not a sulfur donor for STR18 (Fig. S3), we assumed that STR18 stimulated ABA3 activity by reducing the persulfide formed on ABA3 more efficiently than the

![Figure 2](image)

**Figure 2.** CD–Rhd catalyzes the oxidation of roGFP2 via transpersulfidation. Persulfide-dependent oxidation kinetics of 1 μM roGFP2 in the presence of 5 μM CD–Rhd or the C466S variant and 1 mM L-cysteine (A) or 1 mM thiosulfate (B). Arrows indicate the addition of the respective substrate after 3 min. The fully reduced or oxidized roGFP2 used as references were obtained after incubation with 10 mM DTT or \( \text{H}_2\text{O}_2 \), respectively. The ratio of 400/480 nm excitation was normalized to the respective value of maximal oxidation by \( \text{H}_2\text{O}_2 \). The data are represented as mean ± SD (dotted lines) of three independent experiments. CD, cysteine desulfurase; Rhd, rhodanese.

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**Table 1**

| Donor       | Reductant | \( K_M \) (μM) | \( k_{cat} \) (s\(^{-1}\)) | \( k_{cat}/K_M \) (M\(^{-1}\) s\(^{-1}\)) |
|-------------|-----------|----------------|----------------|---------------------------------|
| CD–Rhd      | L-cysteine| 310 ± 38       | 6.8 ± 0.5       | 2.2 ± 0.2 \( \times 10^4 \)       |
|             | GSH       | 476 ± 146      | 8.2 ± 1.2       | 1.8 ± 0.2 \( \times 10^4 \)       |
|             | β-ME      | 273 ± 26       | 6.1 ± 0.2       | 2.2 ± 0.1 \( \times 10^4 \)       |
|             | Thiosulfate| 756 ± 51      | 2.0 ± 0.1       | 2.7 ± 0.3 \( \times 10^3 \)       |
| CD–Rhd C466S| DTT       | 31 ± 5        | 1.01 ± 0.01     | 3.3 ± 0.6 \( \times 10^4 \)       |
|             | GSH       | 597 ± 168      | 0.93 ± 0.12     | 1.6 ± 0.2 \( \times 10^3 \)       |
|             | β-ME      | 346 ± 79       | 0.53 ± 0.03     | 1.6 ± 0.3 \( \times 10^3 \)       |
| Thiosulfate | β-ME      | ND             | ND              | ND                              |
| Thiosulfate | ND        | ND             | ND              | ND                              |

The apparent \( K_M \) and turnover values (\( k_{cat} \)) were calculated by nonlinear regression using the Michaelis–Menten equation. The data are represented as mean ± SD of three independent experiments.

Abbreviation: ND, not detected.
ABA3 provides sulfur to STR18

reductants used in the activity assay. In other words, this implied the transfer of sulfur atoms from ABA3 to STR18. To test this assumption, we analyzed by mass spectrometry the molecular mass of STR18 before and after incubation with a catalytic amount of ABA3 and an excess of L-cysteine. An increase of the molecular mass of STR18 by 31.3 Da, corresponding to the mass of a sulfur atom, was observed after the reaction as compared with a prereduced STR18. As this mass difference disappeared after DTT treatment, we concluded that STR18 was monopersulfated upon reaction with ABA3 in the presence of L-cysteine (Table 4 and Fig. S4).

To firmly establish which cysteine of STR18 is persulfated by ABA3, similar incubation of STR18 variants with ABA3 and L-cysteine has been performed and analyzed by mass spectrometry. A DTT-reversible increase of 33 Da was detected for the C47S variant but not the C89S variant (Table 4 and Figs. S5 and S6). This indicated that STR18 was persulfated on Cys89. Altogether, these data demonstrated the persulfidation of the Cys89 of STR18 by ABA3 in the presence of L-cysteine and the dispensable role of Cys47 for both the TST activity and the ABA3-mediated persulfidation of STR18.

STR18 promotes transpersulfidation reaction between two proteins

In the absence of a known sulfur acceptor for STR18, we have investigated the capability of STR18 to transfer a sulfur atom to a protein by using roGFP2. We have thus tested the oxidation of a prereduced roGFP2 in the presence of STR18 and thiosulfate (Fig. 4A). Whereas thiosulfate alone had no effect, the presence of STR18 promoted roGFP2 oxidation (Fig. 4A). This result validated a transpersulfidation reaction between thiosulfate, STR18, and roGFP2. Then, we investigated roGFP2 oxidation by STR18 in the presence of ABA3 and L-cysteine (Fig. 4B). We first analyzed whether STR18 or ABA3 alone was able to oxidize roGFP2 with L-cysteine. The obtained results confirmed that L-cysteine is not a sulfur donor for STR18 and indicated that ABA3 is unable to promote roGFP2 oxidation. On the contrary, roGFP2 was oxidized by the whole sulfur relay system (L-cysteine, ABA3, and STR18). Similar results were obtained using the STR18 C47S variant, whereas the STR18 C89S did not promote the ABA3-dependent roGFP2 oxidation (Fig. 4B). These results demonstrated that STR18 mediates sulfur transfer from ABA3 to

Table 2

| Protein   | Km (μM) | kcat (s⁻¹) | kcat/Km (M⁻¹ s⁻¹) |
|-----------|---------|------------|--------------------|
| STR18     | 527 ± 8  | 5.1 ± 0.1  | 9.7 ± 0.2 × 10³    |
| STR18 C47S| 333 ± 15 | 2.5 ± 0.1  | 7.5 ± 0.2 × 10³    |
| STR18 C89S| ND      | ND         | ND                 |

The TST activity of STR18, C47S, and C89S variants was monitored in the presence of varying concentrations of thiosulfate as described in the "Experimental procedures" section. The apparent Km and turnover values (kcat) were calculated by nonlinear regression using the Michaelis–Menten equation. The data are represented as mean ± SD of three independent experiments. Abbreviation: ND, not detected.

Figure 3. STR18 stimulates ABA3 activity. A, effect of STR18 on ABA3 cysteine desulfurase (CD) activity. CD activity was measured using 1 μM ABA3, 500 μM L-cysteine, and 1 mM DTT in the presence or the absence of 5 μM STR18. The data are represented as mean ± SD of five independent experiments. B, steady-state kinetic parameters of the CD activity of ABA3 alone or in the presence of STR18 or its cysteinic variants. CD activity was measured in the presence of 1 μM ABA3, increasing concentrations of L-cysteine (0–1 mM), 5 mM DTT, and 5 μM STR18 when present. C, steady-state kinetic parameters of the CD activity of ABA3 in the presence of STR18. Activity was measured in the presence of 0.5 μM ABA3, 500 μM L-cysteine, 5 mM DTT, and increasing concentrations of STR18 (0–5 μM). The data are represented as mean ± SD of three independent experiments. STR18, sulfurtransferase 18.
ABA3 provides sulfur to STR18

The multiple properties of CD proteins are also evident from the existence of fusion proteins containing a CD domain associated with diverse protein domains. The plant ABA3 possesses a MOSC domain, which links the protein function with Moco maturation. According to the known interaction between E. coli IscS and the Thil or YnjE STRs (25, 26), CD–Rhd chimera exist in several bacteria. Here, we described that a Pseudorhodoferax CD–Rhd is a PLP-containing homodimer exhibiting a dual activity profile, as it catalyzes cysteine and thiosulfate desulphurization using its respective functional domains. Noteworthy, this Pseudorhodoferax CD–Rhd is the most efficient CD characterized so far, with a rate of sulfide formation of 7600 nmol min⁻¹ mg⁻¹ in the presence of l-cysteine and DTT. This activity is between 20-fold and 600-fold higher as compared with bacterial (A. vinelandii NiFs and IscS, B. subtilis SufS, E. coli IscS and SufS, and Erwinia chrysanthemi SufS), and eukaryotic CDs (A. thaliana NFS2 and human and yeast NFS1) (Table 5). This remains true if we consider the activity of group II CD members in the presence of their respective activators. Indeed, the rate of sulfide formation ranged from 550 nmol min⁻¹ mg⁻¹ for A. thaliana NFS2–SUFE1 to 750 and 900 nmol min⁻¹ mg⁻¹ for E. chrysanthemi and E. coli SufS–SUFE, respectively (Table 5) (11, 16). The CD activity of the CD–Rhd C466S variant, in which the Rhd domain is inactive, remains high despite it decreased by ~10-fold in the presence of GSH and β-ME compared with CD–Rhd. In this case, with a rate of sulfide formation of 553 nmol min⁻¹ mg⁻¹, this CD activity is in the same range as those determined for E. coli IscS, SufS–SufE, and A. thaliana NFS2–SUFE1 (Table 5). All these results indicate that (i) the CD domain of Pseudorhodoferax CD–Rhd is highly active and (ii) the fusion to an Rhd domain increases its activity with the catalytic cysteine of the Rhd domain acting as a sulfur acceptor, as observed for group II CD isoforms and their specific activators.

The natural CD–Rhd fusion protein of Pseudorhodoferax sp. represents a highly efficient CD isoform

| Protein | Theoretical mass (Da) | Theoretical mass without Met (Da) | Prereduced with DTT | Treatment with ABA3 and l-cysteine | Δ mass (Da) |
|---------|-----------------------|----------------------------------|---------------------|----------------------------------|-----------|
| STR18   | 17117.0               | 16985.8                          | 16985.4             | 17016.7                          | 31.3      |
| STR18 C47S | 17100.9             | 16969.8                          | 16968.2             | 17001.2                          | 33.0      |
| STR18 C89S | 17100.9             | 16969.8                          | 16970.0             | 16969.0                          | –1.0      |

Reduced proteins and proteins incubated with l-cysteine and ABA3 were analyzed by mass spectrometry. The mass accuracy is generally ±0.5 to 1 Da. Note that the mass decrease of ca 131 Da compared with the theoretical molecular masses indicated that the methionine was cleaved off in Escherichia coli.

**Table 3**

Kinetic parameters of CD activity of ABA3

| Protein | Km (μM) | k₉₅ (s⁻¹) | kcat/Km (M⁻¹ s⁻¹) |
|---------|---------|-----------|-------------------|
| l-cysteine |          |           |                   |
| ABA3    | 23 ± 6  | 0.015 ± 0.001 | 6.8 ± 1.6 × 10² |
| ABA3 + STR18 | 8 ± 1   | 0.024 ± 0.001 | 3.2 ± 0.5 × 10³ |
| ABA3 + STR18 C47S | 6 ± 1   | 0.019 ± 0.002 | 3.1 ± 0.4 × 10³ |
| ABA3 + STR18 C89S | 19 ± 3  | 0.017 ± 0.002 | 9.3 ± 2.2 × 10² |
| STR18   | 1.2 ± 0.2 | 0.006 ± 0.001 | 5.1 ± 0.8 × 10³ |
| ABA3    |          |           |                   |

The CD activity of ABA3 was monitored in the presence of varying concentrations of l-cysteine and with or without STR18 as described in the "Experimental procedures" section. The apparent Km and turnover values (k₉₅) were calculated by nonlinear regression using the Michaelis–Menten equation. The data are represented as mean ± SD of three independent experiments.

**Table 4**

Electrospray ionization mass spectrometry analysis of the redox state of STR18 and its monocysteinic variants

**Discussion**

In plants, CDs are key enzymes involved in the maturation of both Fe–S cluster and Moco-containing proteins (14, 15, 19). As CDs act in the early steps of the maturation process and these metalloproteins fulfill important roles, the deletion of CD-encoding genes in plants, more particularly NFS1 and NFS2, is generally lethal or strongly affects development (14, 15). Considering that CDs serve as a central hub for sulfur mobilization and subsequent transfer to various metabolic pathways in nonphotosynthetic organisms, we postulate that the strong phenotypes of mutant plants have thus so far prevented the identification of other sulfur-dependent pathways in which CDs are involved.

A. thaliana STR18 and ABA3 interact in planta

To test whether an ABA3–STR18 interaction could be detected in planta, we performed split-luciferase complementation assays in transiently transformed tobacco leaves (Fig. 5). The bioluminescence emission corresponding to the activity of reconstituted luciferase was tested for different combinations with either candidates fused to the N terminus of the N-terminal (nLuc) domain of luciferase or to the C terminus of C-terminal (cLuc) domain of luciferase. An intense luciferase signal was detected when both ABA3 and STR18 were fused either to nLuc or cLuc domains and coexpressed in tobacco leaves (Fig. 5). On the contrary, no signal was detected when ABA3 or STR18 was coexpressed with a subunit of the ATP citrate lyase 1 (ACL1) known to be localized in the cytosol like ABA3 and STR18 (Fig. 5). In combination with free-nLuc and free-cLuc controls, the latter finding confirmed the specificity of the bioluminescence signal detected after coexpression of ABA3 and STR18. Taken together, these results strongly suggest that ABA3 and STR18 interact in the cytosol of plant cells.

**Table 5**

| Protein | Theoretical mass (Da) | Theoretical mass without Met (Da) | Prereduced with DTT | Treatment with ABA3 and l-cysteine | Δ mass (Da) |
|---------|-----------------------|----------------------------------|---------------------|----------------------------------|-----------|
| STR18   | 17117.0               | 16985.8                          | 16985.4             | 17016.7                          | 31.3      |
| STR18 C47S | 17100.9             | 16969.8                          | 16968.2             | 17001.2                          | 33.0      |
| STR18 C89S | 17100.9             | 16969.8                          | 16970.0             | 16969.0                          | –1.0      |

Reduced proteins and proteins incubated with l-cysteine and ABA3 were analyzed by mass spectrometry. The mass accuracy is generally ±0.5 to 1 Da. Note that the mass decrease of ca 131 Da compared with the theoretical molecular masses indicated that the methionine was cleaved off in Escherichia coli.

**Table 4**

Electrospray ionization mass spectrometry analysis of the redox state of STR18 and its monocysteinic variants

**Table 3**

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|---------|---------|-----------|-------------------|
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| ABA3    | 23 ± 6  | 0.015 ± 0.001 | 6.8 ± 1.6 × 10² |
| ABA3 + STR18 | 8 ± 1   | 0.024 ± 0.001 | 3.2 ± 0.5 × 10³ |
| ABA3 + STR18 C47S | 6 ± 1   | 0.019 ± 0.002 | 3.1 ± 0.4 × 10³ |
| ABA3 + STR18 C89S | 19 ± 3  | 0.017 ± 0.002 | 9.3 ± 2.2 × 10² |
| STR18   | 1.2 ± 0.2 | 0.006 ± 0.001 | 5.1 ± 0.8 × 10³ |
| ABA3    |          |           |                   |

The CD activity of ABA3 was monitored in the presence of varying concentrations of l-cysteine and with or without STR18 as described in the "Experimental procedures" section. The apparent Km and turnover values (k₉₅) were calculated by nonlinear regression using the Michaelis–Menten equation. The data are represented as mean ± SD of three independent experiments.
isozymes, GlpE and PspE, ($K_{M,app}$ of 756 μM versus 78 and 2.7 mM) (34, 35) but a fourfold lower catalytic efficiency than STR18 (Table 2). Hence, CD–Rhd is a bifunctional enzyme using both L-cysteine and thiosulfate as sulfur donors. Nevertheless, considering catalytic efficiencies of both CD and TST activities (2.2 × 10^4 M^−1 s^−1 versus 2.7 × 10^3 M^−1 s^−1 in the presence of β-ME) and the fast and specific CD domain–dependent oxidation of roGFP2, L-cysteine and the associated CD activity represent the preferential substrate and activity of Pseudorhodoferax CD–Rhd.

The efficient cysteine-dependent oxidation of roGFP2 through transpersulfidation reaction catalyzed by CD–Rhd (Fig. 2) also suggests that a role in persulfidation of target proteins may be physiologically relevant. Moreover, considering H₂S release measured in the presence of various reductants and notably GSH, Pseudorhodoferax CD–Rhd might be also involved in the synthesis of H₂S and/or of low–molecular-weight persulfides.

ABA3–STR18 represents a new cytosolic pathway of sulfur trafficking in plant cells

The existence of such natural fusion proteins prompted us to analyze whether the CD activity of ABA3 is enhanced by a STR or in other words if a persulfide transfer reaction is possible between these proteins. In the presence of L-cysteine and DTT, ABA3 displayed an activity and $k_{cat}$ value in the range of the values reported for other CD isoforms (Table 5) (29). Concerning the impact of STR18, the catalytic efficiency of ABA3, measured under steady-state conditions, increased fivefold in the presence of STR18, an effect mostly due to a threefold lower apparent $K_M$ value for L-cysteine. Similar effects were reported for the plastidial SUFE1 protein, which decreased by a factor 2 the $K_M$ value of NFS2 for L-cysteine and increased 42-fold the rate of sulfide formation by NFS2 (16). Furthermore, the low $K_M$ value of 1.2 μM of ABA3 for STR18 determined under steady-state conditions is consistent with the values obtained for the B. subtilis SufS–SufU and E. coli SufS–SufE couples (Table 5) (13, 36, 37). Of interest, in all these examples, the apparent $K_M$ values of the CDs for their protein partners are lower than their apparent $K_M$ values for L-cysteine (eightfold lower for ABA3–STR18 and 20-fold lower for B. subtilis SufS–SufU and E. coli SufS–SufE couples) (13, 36, 37). The physical interaction between both proteins observed with split-luciferase complementation suggests a specific and physiologically relevant ABA3–STR18 interaction.

ABA3–STR18 couple catalyzes transpersulfidation reactions

Both the TST activity and the positive effect on ABA3 activity of STR18 underlined the ability of STR18 to form an intermediate persulfide as demonstrated previously for

Figure 4. STR18 catalyzes the oxidation of roGFP2 via transpersulfidation. A, persulfide-dependent oxidation kinetics of 1 μM roGFP2 in the presence of 5 μM STR18 and 5 mM thiosulfate. B, persulfide-dependent oxidation kinetics of 1 μM roGFP2 in the presence of 1 mM L-cysteine, 1 mM PLP, 1 μM ABA3, and 5 μM STR18 or its cysteine variants. The arrows indicate the addition of STR18, if present, after 2 min. The fully reduced or oxidized roGFP2 used as references were obtained after incubation with 10 mM DTT or H₂O₂, respectively. The ratio of 400/480 nm excitation was normalized to the respective value of maximal roGFP2 oxidation by H₂O₂. The data are represented as mean ± SD (dotted lines) of three independent experiments. PLP, pyridoxal 5′-phosphate; STR18, sulfurtransferase 18.
A. vinelandii Rhd isoform RhdA in the presence of E. coli IscS (38). This was also expected from the sulfur transfer observed from E. coli SufS and CsdA to SufE and CsdE, respectively (10, 39). The ABA3-dependent persulfdation of the catalytic Cys89 of STR18 was indeed demonstrated by mass spectrometry after incubation of prereduced STR18 with both L-cysteine and ABA3 (Table 4). By accepting the sulfur atom, STR18 stimulates the CD activity of ABA3 and regenerates its active form being able to bind the next cysteine molecule (40). In the absence of known STR18 partner(s), we further demonstrated the capacity of STR18 to perform transpersulfdation reactions from either thiosulfate or ABA3 and L-cysteine to roGFP2.

Table 5

| Protein name | Organism                     | Activity (nmol per min per mg) | $K_M$ L-cysteine (µM) | $K_M$ acceptor (µM) | Reference |
|--------------|------------------------------|--------------------------------|-----------------------|---------------------|-----------|
| NifS         | Azotobacter vinelandii       | 168                            | —                     | —                   | (4)       |
| IscS         | Azotobacter vinelandii       | 124                            | —                     | —                   | (4)       |
| SuS          | Bacillus subtilis            | 7                              | —                     | —                   | (13)      |
| SuS–SufU     | Bacillus subtilis            | 240a                           | 86                    | 3                   | (36)      |
| SuS          | Bacillus subtilis            | 153.5                          | 49.7                  | —                   | (36)      |
| SuS–SufU     | Bacillus subtilis            | 93.4                           | 2.6                   | —                   | (36)      |
| SufS         | Erwinia chrysanthemi         | 21                             | —                     | —                   | (11)      |
| SuS–SufE     | Erwinia chrysanthemi         | 750                            | —                     | —                   | (11)      |
| CsdA         | Escherichia coli             | 1.2                            | 140                   | —                   | (12)      |
| CsdA–CsdE    | Escherichia coli             | 2.5                            | 540                   | —                   | (12)      |
| IscS         | Escherichia coli             | 51.7f                          | —                     | —                   | (37)      |
| IscS         | Escherichia coli             | 312.8f                         | —                     | —                   | (37)      |
| SuS          | Escherichia coli             | 380                            | —                     | —                   | (52)      |
| SuS          | Escherichia coli             | 19                             | —                     | —                   | (52)      |
| SuS–SufU     | Escherichia coli             | 900                            | —                     | —                   | (11)      |
| SuS          | Escherichia coli             | 25                             | —                     | —                   | (39)      |
| SuS–SufE     | Escherichia coli             | 700                            | —                     | —                   | (39)      |
| SuS          | Escherichia coli             | 7.9f                           | —                     | —                   | (37)      |
| SuS–SufE     | Escherichia coli             | 54.3f                          | 43.5                  | —                   | (37)      |
| SuS–SufE     | Escherichia coli             | 85.4f                          | —                     | 1.9                 | (37)      |
| CD–Rhd       | Pseudomonas rhodoferax       | 7600                           | —                     | 310                 | (This study) |
| NFS1         | Homo sapiens                 | 6.4                            | —                     | —                   | (53)      |
| NFS1–ISD11   | Homo sapiens                 | 435                            | —                     | —                   | (42)      |
| NFS1         | Saccharomyces cerevisiae     | 12.7                           | —                     | —                   | (54)      |
| NFS2         | Arabidopsis thaliana         | 13                             | —                     | 100                 | (16)      |
| NFS2–SUFE1   | Arabidopsis thaliana         | 550                            | —                     | 43                  | (16)      |
| ABA3         | Arabidopsis thaliana         | 16                             | —                     | —                   | (19)      |

*SuS–SufU activity with 10-fold excess of SufU.*

*CD activity with 2 mM L-cysteine and 2 mM DTT.*

*CD activity with 12 mM L-cysteine and 50 mM DTT.*
ABA3 provides sulfur to STR18

(Fig. 4). From an experimental point of view, the roGFP2 assay enables us to study the ability of a candidate protein to catalyze transpersulfdation reaction in the absence of known partners. It was recently demonstrated that STR1 and STR2, which possess two Rhd domains, efficiently transfer a persulfide to roGFP2 (30). Arabidopsis STR16, another single Rh domain–containing protein, is also able to catalyze roGFP2 oxidation in the presence of thiosulfate (Fig. S7). All these results suggest that the catalysis of transpersulfdation reaction might be a conserved function of STRs. From the apparent \( K_M \) values of STR18 for thiosulfate (527 ± 8 \( \mu \)M) and of ABA3 for STR18 (1.2 ± 0.2 \( \mu \)M), and the ability of ABA3 to promote STR18 persulfidation more efficiently than thiosulfate, ABA3 may be seen as the preferential sulfur donor for STR18.

Relationships between ABA3–STR18 and other cytosolic STR isoforms

All these results represent the first evidence of a functional relationship between CD and STR in plant cells. Of interest, the sulfur transfer pathway from ABA3 to STR18 may be independent of a sulfur transfer to the MOC5 domain and thus independent of Moco sulfuration (Fig. 6). In which physiological context such a pathway is relevant remains to be demonstrated because other cytosolic STRs are present in A. thaliana. In addition to STR18, A. thaliana possesses at least two other cytosolic STR isoforms, the 3-MP-STR isoform STR2 and the two-domain-containing protein STR13 also referred to as CNX5/MOCS3 (23). Noteworthy, STR2 and STR13 are present in all eukaryotic photosynthetic organisms, whereas STR18 is present only in dicotyledonous plants (23). The physiological function(s) of STR2 and STR18 are yet unknown in planta unlike STR13, which possesses a dual function, delivering the sulfur needed for the thiomodification of cytosolic tRNAs and for Moco biosynthesis owing to its N-terminal domain (31, 41). In human cells, a cytosolic form of NFS1 was proposed to provide sulfur to MOCS3 eventually involving a relay by the cytosolic version of TUM1, the ortholog of plant STR2 (27, 42, 43).

While similar actors are present in plants, it may be that this cytosolic sulfur trafficking pathway is different between human and plant cells. Indeed, A. thaliana str2 null mutant lines have no phenotype, whereas str13 null mutants (cnx5-1 and cnx5-2) are sterile and exhibit a severe dwarf phenotype with slightly green and morphologically aberrant leaves (41, 44). On the contrary, aba3 mutants (aba3-1, aba3-2, los5-1, los5-2, aba3-7, and aba3-8) have distinct and less severe phenotypes (45–47) than str13 mutants. This suggests either that STR13 persulfidation would be independent of ABA3 or that STR13 possesses additional functions.

From these results, we propose that in addition to its role in the maturation of the Moco-containing proteins, xanthine dehydrogenase and aldehyde oxidase, ABA3 acts as a sulfur donor to STR proteins (either STR18 as demonstrated here or other cytosolic members such as STR2). The transpersulfidation pathway involving cysteine and an ABA3–STR couple might thus represent an uncharacterized sulfur trafficking pathway in the cytosol of plants.

![Figure 6. Relationships between ABA3 and STR18 in sulfur trafficking in cytosol.](image)

ABA3 catalyzes the desulfuration of cysteine, leading to the persulfidation of the catalytic cysteine of the CD domain and the concomitant release of alanine (1). Then, the persulfidation of STR18 might occur either via the CD domain (2A) or the MOC5 domain (2B) of ABA3. In the next step, STR18 catalyzes a transpersulfidation reaction leading to persulfidation of the target protein (3). This pathway might be independent of the involvement of ABA3 in the sulfuration of Moco-containing proteins (4). CD, cysteine desulfurase; Moco, molybdenum cofactor; STR18, sulfurtransferase 18.
**Experimental procedures**

**Materials**

3-MP (sodium salt) was purchased from Santa Cruz Biotechnology; lead (II) acetate, pyridoxine hydrochloride, L-cysteine, thiosulfate, GSH, and β-ME were from Sigma–Aldrich.

**Cloning and site-directed mutagenesis**

The sequences coding for A. thaliana STR16 (At5g66040), STR18 (At5g66170), and ABA3 (At1g16540) were cloned into the Ndel and BamHI restriction sites of pET15b. Catalytic cysteine (Cys80) of STR16 and both cysteine residues (Cys47 and Cys89) of STR18 were individually substituted into serines to generate pET15b-STR16 C80S, pET15b-STR18 C47S, and pET15b-STR18 C89S recombinant plasmids. A synthetic complementary DNA (GeneCust) coding for CD–Rhd fusion protein (WP_056898193.1) from Pseudomonas aeruginosa sp. Leaf274 was cloned into the Ndel and BamHI restriction sites of pET15b. The cysteine in position 466 was substituted to serine to generate a pET15b–CD–Rhd C466S recombinant plasmid. All primers used in this study are listed in Table S1.

**Heterologous expression in E. coli and purification of recombinant proteins**

For protein expression, the E. coli BL21 (DE3), C41 (DE3), and Rosetta2 (DE3) pLysS strains were transformed, respectively, with pET15b AtSTR16, AtSTR18, AtABA3, Pseudorhodoferax CD–Rhd, and CD–Rhd C466S. The BL21 (DE3) and C41 (DE3) strains also contained the pSBET plasmid, which allows expression of the tRNA needed to recognize the AGG and AGA rare codons. Cell cultures were progressively amplified up to 2.4 l, for STR16, STR16 C80S, STR18, STR18 C47S, STR18 C89S, CD–Rhd and CD–Rhd C466S, and 4.8 l for ABA3, in LB medium supplemented with 50 μg/ml of ampicillin and kanamycin for BL21 and C41 strains or with 50 μg/ml of ampicillin and 34 μg/ml of chloramphenicol for Rosetta2 strain and grown at 37 °C. STR18 expression was induced at exponential phase by adding 100 μM IPTG for 4 h at 37 °C. For ABA3, CD–Rhd, and CD–Rhd C466S, the culture protocol was modified. At exponential phase, the cultures were supplemented with ethanol 0.5% (v/v) and 100 μM pyridoxine hydrochloride and placed at 4 °C for 2 h. Protein expression was then induced by adding 100 μM IPTG for 18 h at 20 °C. After centrifugation (20 min at 6380g), the cell pellets were resuspended in about 20 ml of 50 mM Tris–HCl (pH 8.0), 300 mM NaCl, 10 mM imidazole buffer, and stored at −20 °C.

Cell lysis was completed by sonication (3 × 1 min with intervals of 1 min), and the soluble and insoluble fractions were separated by centrifugation for 30 min at 27,210g. For all proteins, the soluble fraction was loaded on Ni<sup>2+</sup> affinity column (Sigma–Aldrich). After extensive washing, proteins were eluted by a 50 mM Tris–HCl (pH 8.0), 300 mM NaCl, and 250 mM imidazole buffer. The recombinant proteins were concentrated by ultrafiltration under nitrogen pressure and dialyzed (Amicon; YM10 membrane) and finally stored in a 30 mM Tris–HCl (pH 8.0) and 200 mM NaCl buffer supplemented with 5 mM DTT and 50% glycerol at −20 °C. Protein concentrations were determined spectrophotometrically using a molecular extinction coefficient at 280 nm of 10,095 M<sup>−1</sup> cm<sup>−1</sup> for STR16 and 9970 M<sup>−1</sup> cm<sup>−1</sup> for its monocysteinic variant, 11,585 M<sup>−1</sup> cm<sup>−1</sup> for STR18 and 11,460 M<sup>−1</sup> cm<sup>−1</sup> for its monocysteinic variants, 97,845 M<sup>−1</sup> cm<sup>−1</sup> for ABA3, and 47,690 M<sup>−1</sup> cm<sup>−1</sup> for CD–Rhd and CD–Rhd C466S, respectively. The roGFP2 recombinant protein used in this study has been purified as described previously (48).

**Determination of the oligomerization state of CD–Rhd**

The oligomerization state of CD–Rhd and CD–Rhd C466S variant was analyzed by analytical size-exclusion chromatography as described previously (49). The detection was recorded by measuring absorbances at 280 and 418 nm. The column was calibrated using the following molecular weight standards: thyroglobulin (669 kDa, 8.8 ml), β-amylose (200 kDa, 12 ml), bovine serum albumin (66 kDa, 13.6 ml), and cytochrome c (12.4 kDa, 16.8 ml).

**CD activity assays**

The CD activity was assayed at 25 °C in a final volume of 400 μl of 30 mM Tris–HCl (pH 8.0) buffer, 10 μM PLP, 5 mM reductant (DTT, GSH, or β-ME), and 10 nM CD–Rhd, 100 mM CD–Rhd C466S, or 1 μM ABA3. To assess the impact of STR18 on ABA3 activity, 5 μM STR18 was added in the reaction mixture. The reaction was initiated by adding 1-cysteine and stopped after 30 min by adding 50 μl of 20 mM N,N,N-dimethyl-p-phenylene diamine dihydrochloride (prepared in 7.2 M HCl). The addition of 50 μl of 30 mM FeCl<sub>3</sub> (prepared in 1.2 M HCl) followed by a 20 min incubation led to formation of methylene blue, which was then measured at 670 nm. Sulfide in the range of 1 to 100 μM was used for standard curve calibration.

**Thiosulfate STR activity assays**

The thiosulfate STR activity of CD–Rhd, STR18, and their variants was assayed at 25 °C in a final volume of 500 μl of 30 mM Tris–HCl (pH 8.0) buffer, 5 mM β-ME, 0.4 mM lead (II) acetate, various concentrations of thiosulfate ranging from 0 to 5 mM, and 100 nM enzyme. The reaction was initiated by adding CD–Rhd or STR18, and the rate of lead sulfide formation was monitored at 390 nm using a molar extinction coefficient of 5500 M<sup>−1</sup> cm<sup>−1</sup>.

**Detection of persulfidated STR18 by mass spectrometry**

In a final volume of 150 μl of 30 mM Tris–HCl (pH 8.0), 200 mM NaCl buffer, 150 μM of prereduced STR18, STR18 C47S, and STR18 C89S were incubated 30 min in the presence of 300 μM 1-cysteine, 2 μM ABA3, and 5 μM PLP at 25 °C. After extensive dialysis, samples were split in two parts and treated or not with 1 mM DTT. Mass spectrometry analysis of these samples was performed using a Bruker microTOF-Q spectrometer (Bruker Daltonik), equipped with Apollo II electrospray ionization source with ion funnel, operated in the
ABA3 provides sulfur to STR18

negative ion mode. The concentrated samples in formic acid were injected at a flow rate of 10 to 20 μl min⁻¹. The potential between the spray needle and the orifice was set to 4.5 kV. Before each run, the instrument was calibrated externally with the Tunemix mixture (Agilent Technologies) in quadratic regression mode. Data were analyzed with the DataAnalysis software (Bruker).

roGFP2 oxidation experiments

The capacity of CD–Rhd, ABA3, STR16, and STR18 to oxidize roGFP2 was analyzed in vitro by ratiometric time-course measurements on a fluorescence plate reader (EnSight multimode plate reader; PerkinElmer) with excitation at 400 ± 10 and 480 ± 10 nm and detection of emitted light at 520 nm with a bandwidth of 10 nm. The maximum oxidation and reduction of roGFP2 were defined using 10 mM H₂O₂ and DTT. Prereduced roGFP2 was obtained by incubation with 10 mM DTT for 1 h and subsequent desalting on a G25 column to remove excess DTT. In a final volume of 400 μl of 30 mM Tris–HCl (pH 8.0) and 200 mM NaCl, the reaction mixtures contained 1 μM prereduced roGFP2 and either 5 mM thiosulfate and 5 μM STR16/STR18/CD–Rhd or 1 mM l-cysteine, 5 μM CD–Rhd or 1 mM l-cysteine, 10 μM PLP, 1 μM ABA3, and 5 μM STR18.

Split-luciferase complementation assays

Full-length coding sequences of ABA3, STR18, and the negative control ACL1 (At1g10670) were selectively amplified with primers defined in Table S1 and cloned via KpnI and BamHI restriction endonucleases into the pCAMBIA1300-cLuc or via KpnI and MluI restriction endonucleases into the pCAMBIA1300-nLuc vectors described (50). The resulting fusion constructs were named cLuc-ABA3, cLuc-STR18, ABA3-nLuc, STR18-nLuc, and cLuc-ACL1. Different combinations of cLuc-fusion and nLuc-fusion constructs were coexpressed in tobacco (Nicotiana benthamiana) leaves after Agrobacterium inoculation, plants were kept for 24 h in the dark and subsequently grown for 2 days under long day conditions (16 h light 250 μE, 8 h dark, temperature of 25°C, and humidity of 50%) to allow expression of the protein of interest in fusion with the N-terminal (nLuc) or C-terminal (cLuc) fragment of luciferase. The abaxial sides of the transformed leaves were sprayed with luciferase substrate (1 mM luciferin), and the substrate was allowed to enter the leaf for 5 min. The resulting luciferase signal was detected with the digital camera system "ImageQuant LAS 4000" (GE Healthcare) and visualized with the open access software suite “Image J” (National Institutes of Health).

Data availability

All data are presented in the article.

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Abbreviations—The abbreviations used are: β-ME, β-mercaptoethanol; 3-MP, 3-mercaptopyruvate; ACL1, ATP citrate lyase 1; CD, cysteine desulfurase; Fe–S, iron–sulfur; Moco, molybdenum cofactor; PLP, pyridoxal 5’-phosphate; Rhd, rhodanese; STR, sulforhodamine; STR18, sulforhodamine transferase; STR18, sulforhodamine transferase 18; TST, thiosulfate-STR.

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ABA3 provides sulfur to STR18