Kinase SnRK1.1 regulates nitrate channel SLAH3 engaged in nitrate-dependent alleviation of ammonium toxicity

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Abstract
Nitrate ($NO_3^-$) and ammonium ($NH_4^+$) are major inorganic nitrogen (N) supplies for plants, but $NH_4^+$ as the sole or dominant N source causes growth inhibition in many plants, known as ammonium toxicity. Small amounts of $NO_3^-$ can significantly mitigate ammonium toxicity, and the anion channel SLAC1 homolog 3 (SLAH3) is involved in this process, but the mechanistic detail of how SLAH3 regulates nitrate-dependent alleviation of ammonium toxicity is still largely unknown. In this study, we identified SnRK1.1, a central regulator involved in energy homeostasis, and various stress responses, as a SLAH3 interactor in Arabidopsis (Arabidopsis thaliana). Our results suggest that SNF1-related protein kinase 1 (SnRK1.1) functions as a negative regulator of SLAH3. Kinase assays indicate SnRK1.1 strongly phosphorylates the C-terminal of SLAH3 at the site S601. Under high-$NH_4^+/low-pH$ condition, phospho-mimetic and phospho-dead mutations in SLAH3$S601$ result in barely rescued phenotypes and fully complemented phenotypes in slah3. Furthermore, SnRK1.1 migrates from cytoplasm to nucleus under high-ammonium stress. The translocation of SnRK1.1 from cytosol to nucleus under high-ammonium stress releases the inhibition on SLAH3, which allows SLAH3-mediated $NO_3^-$ efflux leading to alleviation of high-$NH_4^+/low-pH$ stress. Our study reveals that the C-terminal phosphorylation also plays an important role in SLAH3 regulation and provides additional insights into nitrate-dependent alleviation of ammonium toxicity in plants.

Introduction
Nitrogen (N) is an essential nutrient element for all living organisms on earth. For plants, N is an indispensable component of proteins, nucleic acids, and chlorophylls. Except for legumes, most plants are not capable of directly utilizing the gaseous N abundant in the atmosphere and can only take up organic forms of N such as amino acids (AAs) and inorganic forms of N mainly including ammonium ($NH_4^+$)
and nitrate (\(\text{NO}_3^-\)) through specific transportation systems in root (Ludewig et al., 2007; Wang et al., 2012).

In Arabidopsis (Arabidopsis thaliana), the uptake of \(\text{NH}_4^+\) is mediated by ammonium transporters (AMTs) localized at the plasma membrane (PM; Ludewig et al., 2007). When the root absorbs \(\text{NH}_4^+\) from the environment, the AMT-coupled \(\text{H}^+\)-ATPase pumps protons out of the root cell in order to maintain balanced intracellular charge and stabilize membrane potential (Meier et al., 2020; Motte and Beeckman, 2020). \(\text{NO}_3^-\) is transported by PM-localized nitrate transporters (NRTs; Wang et al., 2012). Unlike AMT-mediated \(\text{NH}_4^+\) absorption that often causes \(\text{H}^+\) efflux, most NRTs are \(\text{NO}_3^-/\text{H}^+\) symporters, mediating \(\text{NO}_3^-\) uptake accompanied by \(\text{H}^+\) influx (Mengel et al., 1983; Pearson and Stewart, 1993; Wang et al., 2012; Xu et al., 2012).

After absorption, \(\text{NO}_3^-\), the oxidized form of inorganic N, needs to be reduced to \(\text{NH}_4^+\) before further assimilation, catalyzed by nitrate reductase and nitrite reductase (Crawford, 1995). Therefore, \(\text{NH}_4^+\) is theoretically the preferred inorganic N source for plants (Salsac et al., 1987). However, when \(\text{NH}_4^+\) is the dominant N source, it often significantly inhibits plant growth, featured by the phenotypes of severely inhibited root growth and chlorosis, known as ammonium toxicity (Kronzucker et al., 1997; Britto and Kronzucker, 2002). To date, little is known about the detailed molecular mechanisms of ammonium toxicity. Nevertheless, nitrate effectively alleviates ammonium toxicity (Domene and Ayres, 2001; Britto and Kronzucker, 2002; Houdusse et al., 2005). Likewise, how nitrate lessens ammonium toxicity is largely unclear.

The proteins in slow anion channel/SLAC1 homolog (SLAC/SLAH) family in vascular plants function as efflux channels for anions such as \(\text{Cl}^-\) and \(\text{NO}_3^-\) (Geiger et al., 2009; Wang et al., 2012). In Arabidopsis, the SLAC/SLAH gene family consists of five members, SLAC1 and SLAH1 to 4, encoding PM proteins with predicted 10 transmembrane helices with both amino termini (NT) and carboxyl termini (CT) in cytoplasm (Negi et al., 2008; Vahisalu et al., 2008; Hedrich and Geiger, 2017). SLAC1 is specifically expressed in guard cells and its encoding protein product is an important regulator of stomatal movement (Negi et al., 2008; Vahisalu et al., 2008). SLAH3 plays the same role as SLAC1 in regulating stomatal movement (Geiger et al., 2011). In guard cells, the channel activity of SLAH3 is regulated by reversible phosphorylation mediated by protein kinase CPK21 and protein phosphatase ABI1 (Geiger et al., 2011; Demir et al., 2013). Compared to SLAC1, SLAH3 has a stronger selectivity for \(\text{NO}_3^-\), with an about 20 times higher affinity over \(\text{Cl}^-\), and is therefore considered as a nitrate efflux channel (Geiger et al., 2011; Wang et al., 2012).

SLAH3 is also expressed in roots, suggesting it may serve distinct physiological functions in addition to regulating stomatal movement (Zheng et al., 2015). Our previous studies show that slah3 mutants are extremely sensitive to the culture conditions with higher concentrations of \(\text{NH}_4^+\) and lower concentrations of \(\text{NO}_3^-\) compared with wild-type (WT) Columbia-0 (Col-0) plants (Zheng et al., 2015). We also found that under high-ammonium conditions, elevating the medium pH significantly lessens ammonium toxic phenotypes in plants, and more importantly, wipes out the phenotypic differences between slah3 mutants and Col-0. These results indicate medium acidification is likely one of the important causes of ammonium toxicity and that SLAH3 may function through regulating the pH of the rhizosphere. Our study thus revealed SLAH3 is involved in the process of nitrate-dependent alleviation of ammonium toxicity likely via mediating nitrate efflux under high-\(\text{NH}_4^+\)/low-pH conditions (Zheng et al., 2015). We speculate that under high-\(\text{NH}_4^+\) conditions, the absorption of \(\text{NH}_4^+\) leads to accumulation of apoplastic \(\text{H}^+\), causing rapid acidification in the rhizosphere. In response to this condition, SLAH3 mediates the outflow of \(\text{NO}_3^-\) to the rhizosphere. \(\text{NO}_3^-/\text{H}^+\) symporter NRTs then mediate influx of both \(\text{NO}_3^-\) and \(\text{H}^+\), reducing the medium acidification caused by absorption of \(\text{NH}_4^+\) and effectively alleviating ammonium toxicity.

Sucrose non-fermenting 1 (SNF1)-related protein kinase 1 (SnRK1) proteins, the paralogs of SNF1 in yeasts and AMP-activated protein kinases (AMPK) in animals, are a group of Ser/Thr protein kinases widely found in plants (Polge and Thomas, 2007; Crozet et al., 2014). SnRK1/SNF1/AMPK kinases play central roles in sensing sugar signals and regulating energy homeostasis (Baena-González and Sheen, 2008; Crozet et al., 2014). In plants, SnRK1s also play key roles in regulating biotic and abiotic stress responses as well as plant growth and development (Baena-González et al., 2007; Polge and Thomas, 2007; Baena-González and Sheen, 2008; Hulsmans et al., 2016). SnRK1 is a heterotrimeric complex composed of a catalytic subunit and two regulatory subunits (Crozet et al., 2014; Hulsmans et al., 2016). The catalytic subunit is encoded by SnRK1.1 (Kin10) or SnRK1.2 (Kin11) in Arabidopsis. SnRK1.1 and SnRK1.2 are highly functionally redundant. Single mutation in SnRK1.1 or SnRK1.2 causes no obvious phenotypic defects but the snrk1.1 snrk1.2 double mutant is lethal (Baena-González et al., 2007).

In plants, the SnRK1 family has expanded to a larger family containing SnRK2 and SnRK3 subfamilies (Halford and Hardie, 1998; Hrabak et al., 2003). SnRK2s mainly function in ABA signaling pathway and SnRK3s, also known as calcium-in B-like (CBL)-interacting protein kinas (CIPKs), interact with CBL proteins upon \(\text{Ca}^{2+}\) signal to regulate various stress responses (Fujii and Zhu, 2009; Luan, 2009; Coello et al., 2011). In plants, calcium-dependent protein kinases (CPKs) share similar structures with the SnRK family and are included in the CPK–SnRK superfamily (Hrabak et al., 2003). Intriguingly, a number of CPK–SnRK members have been identified as key regulators in controlling the activities of ion transporters and channels (Kudla et al., 2018).

To understand the molecular mechanisms of how SLAH3 is regulated in response to high-\(\text{NH}_4^+\) stress, we identified SnRK1.1 as a SLAH3 interactor through a yeast-two-hybrid screen. Distinct from the previously identified kinases that
associate with and phosphorylate the NT of SLAH3 and subsequently activate the SLAH3 channel, SnRK1.1 interacts with and phosphorylates the CT of SLAH3. By using a combination of molecular, biochemical, and genetic approaches, we demonstrated that SnRK1.1 likely negatively regulates SLAH3 in modulating the process of nitrate-dependent alleviation of ammonium toxicity. Moreover, we found that SnRK1.1 translocates from cytoplasm to nucleus upon high-ammonium conditions, suggesting the release of the inhibition of SnRK1.1 on SLAH3 in response to high-ammonium conditions. Thus, our study revealed an additional mechanism of how nitrate efflux channel SLAH3 is regulated in response to high-ammonium stress.

Results

Protein kinase SnRK1.1 interacts with the C-terminus of SLAH3

Our previous study showed that an anion channel SLAH3 contributes to the process of nitrate-dependent alleviation of ammonium toxicity in Arabidopsis (Zheng et al., 2015). We analyzed the phenotypes of WT Col-0 and the loss-of-function SLAH3 mutants under the condition of high-ammonium, low-nitrate, and low-pH [5 mM NH\textsubscript{4}\textsuperscript{+}, 1mM NO\textsubscript{3}\textsuperscript{-} (ASN1), and pH 4.5]. Compared to Col-0, slah3-3 and slah3-4 mutants showed strong ammonium toxic phenotypes under higher concentrations of NH\textsubscript{4}\textsuperscript{+} and lower concentrations of NO\textsubscript{3}\textsuperscript{-}. More importantly, elevated pH not only greatly promoted growth in all plants but also eliminated the phenotypic differences between Col-0 and slah3 mutants. In addition, merely increasing the concentration of NO\textsubscript{3}\textsuperscript{-} also abolished the ammonium toxic phenotypes of slah3 mutants (Figure 1A; Zheng et al., 2015). Quantitative analyses performed by measuring the fresh weight were consistent with the phenotypes (Figure 1B). Our results thus indicated nitrate efflux is important for the process of nitrate-dependent alleviation of ammonium toxicity, in which the medium pH plays an essential role. Topologic analyses suggested that SLAH3 has 10 transmembrane helices with both its NT and CT in the cytoplasm (Figure 1C). The NT of SLAH3, which is around three times larger than the CT, was reported to be important for the regulation of SLAH3 channel activity. For instance, CPK21 phosphorylates the NT of SLAH3 to activate the channel in guard cells (Geiger et al., 2011). However, the reported SLAH3-targeting CPKs show little expression in roots, suggesting additional components are responsible for SLAH3 regulation in roots.

To identify more protein kinases involved in SLAH3 regulation, we used a yeast-two-hybrid system to screen the potential interactors of SLAH3. By using the NT and CT of SLAH3 as the baits, we screened the protein collection including all members in the CPK–SnRK superfamily previously constructed by our lab. We identified SnRK1.1 as a potential SLAH3 interactor. Interestingly, unlike the CPKs that interact with the NT of SLAH3, SnRK1.1 showed strong interaction with the CT, suggesting SnRK1.1 functions differently from CPKs (Figure 1D). SnRK1.1 has two isoforms, SnRK1.1-Long (L) and SnRK1.1 (Figure 1E). SnRK1.1-L has additional 23 AA residues at the NT compared to SnRK1.1. SnRK1.1 contains three major domains including kinase, ubiquitin-associated (UBA) and kinase-associated (KA1) domains. The lysine at site 48 (K48) is essential for SnRK1.1 activity and was shown to abolish kinase activity of SnRK1.1 when substituted with methionine (M) (Figure 1E; Cho et al., 2016; Ramon et al., 2019). We next analyzed which domains of SnRK1.1 were responsible for the interaction with SLAH3. Using a yeast-two-hybrid system, we tested the interactions between SLAH3-CT and different truncated SnRK1.1 protein fragments (Figure 1F). Our results indicated the CT of SnRK1.1 including KA1 domain is sufficient to interact with SLAH3-CT. The NT of SnRK1.1 including the kinase domain is not involved in the association with SLAH3-CT. Furthermore, the kinase-dead variant (K48M) of SnRK1.1 (mSnRK1.1) still associated with SLAH3-CT, indicating the kinase activity of SnRK1.1 is not required for its interaction with SLAH3-CT (Figure 1F). Thus, our results showed that SnRK1.1, a central regulator in energy signaling and stress responses, interacts with the CT of SLAH3.

SnRK1.1 interacts with SLAH3 in vivo

To verify the interaction between SnRK1.1 and SLAH3, we utilized bimolecular fluorescence complementation (BiFC) to test the association of SnRK1.1 with full length SLAH3 in vivo. After being co-expressed in the leaf epidermal cells of Nicotiana benthamiana, SnRK1.1 interacted with SLAH3 (Figure 2A). Furthermore, the kinase-dead variant (K48M) of SnRK1.1 (mSnRK1.1) still associated with SLAH3, confirming that kinase activity of SnRK1.1 has no effect on the SnRK1.1–SLAH3 interaction. After isolating mesophyll protoplasts of N. benthamiana co-expressed with SnRK1.1 and SLAH3, we found SnRK1.1–SLAH3 interaction occurred at the PM (Figure 2B). Next, an assay using the mating-based split ubiquitin system (mbSUS) also confirmed that SnRK1.1 interacted with full length SLAH3 in yeast cells (Figure 2C). Moreover, we used fluorescence resonance energy transfer (FRET) to further verify the interaction between SLAH3 with SnRK1.1 in planta. In the acceptor photobleaching experiment, the images of the same regions before and after photobleaching were acquired to determine the FRET signal when SnRK1.1/mSnRK1.1-CFP (cyan fluorescent protein) was co-expressed with SLAH3-YFP (yellow fluorescent protein). The fluorescence intensity of the acceptor decreased while that of the donor increased with acceptor bleaching (Figure 2, D and E). In the sensitized emission experiment, the FRET signal was detected when SLAH3-YFP was coexpressed with SnRK1.1/mSnRK1.1-CFP (Supplemental Figure S1). Both of the FRET experiments demonstrated that SLAH3 interacts with SnRK1.1 and mSnRK1.1 in planta (Figure 2, D and E and Supplemental Figure S1).

SnRK1.1 and SLAH3 are expressed in roots

We next analyzed whether SnRK1.1 can regulate SLAH3 spatially. The expression patterns of SnRK1.1 and SLAH3 were determined by promoter-GUS analyses (Figure 3). SLAH3

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SnRK1.1 was mainly expressed in root, guard cells, and floral organs. SnRK1.1 was ubiquitously expressed in the plants and showed high expression in roots, suggesting SnRK1.1 can potentially function as a SLAH3 regulator in roots.

SnRK1.1 is involved in the process of nitrate-dependent alleviation of ammonium

To date, two SnRK1s have been found in the Arabidopsis genome, SnRK1.1 and SnRK1.2, which are highly redundant in terms of both structures and functions. The snrk1.1 snrk1.2 double null mutants are lethal, supporting their essential functions in plants (Baena-González et al., 2007). To further investigate the potential regulation of SLAH3 by SnRK1.1, we generated overexpression lines of SnRK1.1 and mSnRK1.1, driven by a constitutively active 35S promoter. RT-qPCR results indicated SnRK1.1 and mSnRK1.1 were highly expressed in the overexpression plants (Supplemental Figure S2). We analyzed the phenotypes of two independent lines of SnRK1.1 or mSnRK1.1 under the conditions of high-ammonium, low-nitrate, and low-pH (A5N1, pH 4.5), supplemented with different concentrations of sucrose (Figure 4A). The slah3-3 and slah3-4 mutant plants, exhibiting no phenotypic difference compared with Col-0 in 1/2 Murashige and Skoog (MS) medium, showed obvious defective phenotypes in A5N1, pH 4.5 medium when supplied with 1% or 2% sucrose. Under same A5N1, pH 4.5 condition, the phenotypic differences between Col-0 and slah3 mutants were
greatly reduced when sucrose was decreased to lower concentrations (0.5%, 0.2%, or 0%). These results indicated the presence of sugars affects the ammonium toxic phenotypes of slah3 mutants. When A5N1, pH 4.5 conditions was applied with 1% or 2% sucrose, SnRK1.1-OE lines and slah3 mutants showed obvious growth defects compared with Col-0. Under same A5N1, pH 4.5 conditions, the phenotypic differences between SnRK1.1-OE lines, slah3 mutants, and Col-0 were nearly abolished when sucrose was decreased to 0.2%. SnRK1.1-OE plants showed inhibited growth in 1/2 MS medium when higher concentrations of sucrose (1% or 2%) were supplemented, but exhibited similar phenotypes to Col-0 when lower concentrations of sucrose (0.5%, 0.2%, or 0%) were applied. Whereas, under A5N1, pH 4.5 condition with no sucrose provided, SnRK1.1-OE plants showed even longer root length compared with Col-0. mSnRK1.1-OE plants showed inhibited growth in A5N1, pH 4.5 condition with no sucrose supplemented, but exhibited similar phenotypes to Col-0 when sucrose was applied. Quantitative analyses were performed by measuring the fresh weight and
root length of the plants (Figure 4, B and C). These results suggested SnRK1.1 plays a role under high-ammonium/low-pH conditions, and sugar is involved in this process. Similar phenotypic characteristics were also observed in the media with different concentrations of glucose, which further indicated that SnRK1 indeed participates in the responses to high-ammonium stress through the regulation of sugar signals (Supplemental Figure S3). In addition, we analyzed the phenotypes in mediums supplemented with different concentrations of palatinose (a nonmetabolizable analog of sucrose) for 20 d. In 1/2 MS mediums, all plants showed no difference when supplemented with palatinose. Under the conditions of A5N1, pH 4.5, compared to wildtype (WT) plants, SnRK1.1-OE plants showed enhanced sensitivity to higher concentrations of sucrose but less sensitivity to higher concentrations of palatinose. Meanwhile, supplemented with all concentrations of palatinose, slah3 mutants showed similar phenotypes to Col-0 (Supplemental Figure S4). Furthermore, the expression of SnRK1.1 affected the phenotype in a dosage-dependent manner when treated with metabolic sugar (sucrose or glucose). The transgenic lines expressing lower levels of SnRK1.1 showed weaker phenotype than the lines expressing higher level of SnRK1.1. In contrast, the treatment of palatinose had no dosage effect on the phenotype in the plant lines with different SnRK1.1 expression levels. These results suggested metabolizable sugar such as sucrose or glucose plays a role in SLAH3-mediated nitrate-dependent alleviation of ammonium toxicity. Next, the SnRK1.1-OE or mSnRK1.1-OE plants were crossed to slah3-4 background. Under A5N1, pH 4.5 condition, the phenotypes caused by the overexpression of SnRK1.1 or mSnRK1.1 were abolished in slah3-4 background (Supplemental Figure S5, A and C). Statistical analyses were performed by analyzing the fresh weight and root length
SLAH3 is required for SnRK1.1 to regulate hypocotyl growth in dark
SnRK1 kinases play important roles in regulating carbohydrate metabolism and are involved in sucrose-induced hypocotyl elongation of plants in dark (Simon et al., 2018; Hamasaki et al., 2019). Overexpression lines of SnRK1.1 (SnRK1.1-OE) or mSnRK1.1 (mSnRK1.1-OE) in Col-0 or slah3-4 backgrounds were next analyzed in dark condition. Without sucrose supplement, SnRK1.1 overexpression caused no phenotypic differences in Col-0 or slah3-4 under either 1/2 MS or AS5N1, pH 4.5 condition. Supplemented with 1% sucrose, however, SnRK1.1-OE in Col-0 showed shortened hypocotyl compared with Col-0 in both 1/2 MS and AS5N1, pH 4.5 conditions. SnRK1.1-OE in Col-0 was crossed to slah3-4 to generate SnRK1.1-OE in slah3-4. Interestingly, although slah3-4 mutant showed similar phenotypes with Col-0 under all conditions tested, the shortened hypocotyl phenotype caused by SnRK1.1-OE was suppressed in slah3-4 (Figure 5, A and C). Different from SnRK1.1-OE plants, mSnRK1.1-OE lines exhibited no difference with their Col-0 and slah3-4 backgrounds when 1% sucrose was supplemented, but showed shortened hypocotyl when no sucrose was supplied (Figure 5, B and D). Similarly, the short hypocotyl phenotype caused by mSnRK1.1 overexpression in Col-0 disappeared in slah3-4 background on the medium without sucrose. The results surprisingly showed that SnRK1.1 seems to require SLAH3 to regulate hypocotyl growth during skotomorphogenesis in response to sugar signals.

SnRK1.1 phosphorylates SLAH3-CT in vitro
Given that SnRK1.1 interacts with the CT of SLAH3 and may contribute to SLAH3-mediated physiological process, we next analyzed the detailed molecular mechanism of how SnRK1.1 regulates SLAH3. Kinase assay based on ³²P-labeled ATP showed SnRK1.1 strongly phosphorylated SLAH3-CT while SLAH3-NT only showed a basal phosphorylation upon SnRK1.1 presence, indicating SnRK1 mainly phosphorylates SLAH3-CT (Figure 6A). The phosphorylation of SLAH3-CT was further confirmed by another kinase assay using a Pro-Q diamond phosphoprotein gel stain by which the phosphorylated proteins on the protein gel can be directly revealed by fluorescence scanner detection. SnRK1.1 caused band shifting of SLAH3-CT in a Pro-Q gel but the shifting band disappeared when protein phosphatase (λ-pp) was added (Figure 6B). Thus, the kinase assays, together with protein interaction results, indicated SnRK1 interacts and strongly phosphorylates SLAH3-CT.

Liquid chromatograph-mass spectrometer (LC-MS/MS) analysis was next used to determine the specific sites of SLAH3-CT phosphorylated by SnRK1.1 (Figure 7A). The site S601 was identified as the only AA residue of SLAH3-CT phosphorylated by SnRK1.1 (Figure 7B). We then verified the result from MS by using Pro-Q kinase assay. It was previously reported that the T513 of SLAC1 in its CT functions as a crucial phosphorylation site for the activation of SLAC1 (Dreyer et al., 2012; Maierhofer et al., 2014). The conserved serine in SLAH3, S580, was therefore used as a control. When S601 was substituted with A (S601A), the phosphorylation of SLAH3-CT mediated by SnRK1.1 was completely abolished in a Pro-Q kinase assay (Figure 7C). In contrast, S580A substitution did not affect the phosphorylation of SLAH3 by SnRK1.1. Thus, the biochemical and MS assays revealed that SnRK1.1 phosphorylates S601 at the CT of SLAH3.

Phosphorylation of S601 of SLAH3 by SnRK1.1 plays an important role in high-ammonium response
Since S601 of SLAH3 was phosphorylated by SnRK1.1 in the in vitro assays, we next verified the importance of S601 for SLAH3 function. To understand the biological role of S601, we generated transgenic plants harboring SLAH3 phosphorylation-mimetic (S601D) or phosphorylation-dead (S601A) mutation driven by its native promoter in the slah3-4 background. Reverse transcription-polymerase chain reaction (RT-PCR) indicated the transcripts of SLAH3 S601D or S601A in complemented lines showed similar levels as those in Col-0 (Supplemental Figure S6). S601A mutation fully rescued the phenotypes of slah3-4 under AS5N1 condition. In contrast, S601D mutation can barely complement the phenotypes of slah3-4 under AS5N1 condition (Figure 8A). Quantitative analyses were performed by measuring the fresh weights and root lengths of the plants (Figure 8, B and C). Yeast-two-hybrid assay showed S601A or S601D mutation did not affect the interaction between SLAH3-CT and SnRK1.1 (Supplemental Figure S7). In contrast, SLAH3 completely complemented the phenotypes of slah3-4 under AS5N1 condition (Supplemental Figure S8). The genetic results, therefore, supported the notion that S601 phosphorylation mediated by SnRK1.1 may negatively regulate SLAH3 function in regulating the process of nitrate-dependent alleviation of ammonium toxicity.

To further understand the physiological role of the SLAH3 S601 site, we analyzed the nitrate fluxes by using non-invasive micro-test technology (NMT) method. To test whether Cl⁻ interferes with the measurements of NO₃⁻ in the NMT system, the NO₃⁻ concentrations and net fluxes in measuring solution 1 (1 mM KNO₃, 0.1 mM KCl, 0.1 mM CaCl₂, 0.3 mM MES, pH 6.0) and measuring solution 2 (1 mM KNO₃, 1.1 mM KCl, 0.1 mM CaCl₂, 0.3 mM MES, pH 6.0) were detected without seedlings. The presence of Cl⁻ did not interfere with the measurements of NO₃⁻ (Supplemental Figure S9). Next, we detected the net NO₃⁻ fluxes at the maturation zone next to the primary

(Supplemental Figure S5, B and D). These results indicated SLAH3 is required for SnRK1.1 in regulating high-ammonium stress adaption. Taken together, these results suggested sugar signaling, in which SnRK1.1 functions as a central regulator, is involved in the process of nitrate-dependent alleviation of ammonium toxicity. SnRK1.1 likely contributes to this process via negatively regulating SLAH3.
root tip (Figure 9A, Supplemental Figure S10A). Seedlings from Col-0, slah3-3, slah3-4, and all complementary lines that were grown on 1/2 MS medium for 8 d and transferred to high-NH$_4^+$/low-pH condition (1 mM NO$_3^-$, 10 mM NH$_4^+$, pH 4.5) or non-high-NH$_4^+$/low-pH condition (1 mM NO$_3^-$, 1 mM NH$_4^+$, pH 5.7) for 2 h were measured. NO$_3^-$ fluxes from the plant roots were detected over 6 min (Figure 9B; Supplemental Figure S10B). Under high-NH$_4^+$/low-pH stress, the mean value of NO$_3^-$ effluxes from Col-0 was around 93 pmol cm$^{-2}$ s$^{-1}$, and the mean value of NO$_3^-$ effluxes from slah3 mutants was around 0 pmol cm$^{-2}$ s$^{-1}$. Complementary lines of SLAH3 and SLAH3 S601A in slah3-4 background showed nitrate effluxes similar to Col-0 plants. In contrast, complementary lines of SLAH3 S601D in slah3-4 background showed nitrate effluxes similar to slah3 mutants (Figure 9B). In addition, under non-high-NH$_4^+$/low-pH stress, nitrate effluxes were similar among all plants, and the mean value of NO$_3^-$ effluxes were around 0 pmol cm$^{-2}$ s$^{-1}$ (Supplemental Figure S10).

These results suggested that S601 of SLAH3 plays an important role in regulating nitrate efflux under high-NH$_4^+$/low-pH stress.

High-ammonium stress causes accumulation of SnRK1.1 in nucleus
Under normal condition, SnRK1.1 locates in both cytoplasm and nucleus. A recent study demonstrated that cytosol-localized SnRK1.1 migrates to nucleus under metabolic, hypoxia, DCMU and dark stresses (Ramon et al., 2019). Therefore, we tested whether SnRK1.1 also accumulates in the nucleus under high-ammonium stress. We analyzed the distribution of SnRK1.1 protein upon high-ammonium treatments. In the Col-0 plants grown on 1/2 MS or ASN1, pH 4.5 medium with 0.9% agarose culture condition for 14 d, the cellular contents in cytoplasm and nucleus were separated. An anti-SnRK1.1 antibody was used to immunoblot SnRK1.1 protein in cytoplasmic or nuclear fraction. Anti-PEPC and anti-Histone H3 antibodies were used for positive

**Figure 5** SLAH3 is required for SnRK1.1 to regulate hypocotyl growth in dark. A, The phenotypes of Col-0, slah3-4, and mSnRK1.1-OE lines cultivated for 8 d under different culture conditions in dark. White scale = 2 cm. B, The phenotypes of Col-0, slah3-4, and mSnRK1.1-OE lines cultivated for seven days under different culture conditions in the dark. White scale = 2 cm. C and D, The statistical analyses of hypocotyl length on the plants presented in (A) and (B). Data are shown as means ± SE, n = 20. Statistical analyses were performed by two-way ANOVA analysis with Tukey’s multiple comparisons test. Significant differences are indicated with different letters. P < 0.05. S, Sucrose.
indicators for cytoplasmic and nuclear sections, respectively. SnRK1.1 was detected in both cytoplasm and nucleus when the plants were grown in 1/2 MS (Figure 10, A and B). In contrast, when cultivated in A5N1, pH 4.5 media, the abundance of SnRK1.1 protein significantly decreased in cytoplasm but increased in nucleus accordingly (Figure 10, A and B). Quantitative analyses were performed by measuring the relative intensity of SnRK1.1 bands (Figure 10, C and D). The images of original western blots are included in supplementary data (Supplemental Figure S11).

Furthermore, we analyzed the responses of SnRK1.1 protein to short-term high-ammonium stress. The Col-0 plants expressing YFP-SnRK1.1 were grown in the 1/2 MS medium for 5 d. The transgenic plants were transferred to the liquid 1/2 MS medium or liquid A5N1, pH 4.5 medium, and the YFP signal was checked at different time courses. In the control plants that were transferred to the liquid 1/2 MS medium, the distribution of SnRK1.1 was not altered and was detected in both cytoplasm and nucleus. In contrast, in the plants subjected to A5N1, pH 4.5 condition, the SnRK1.1 protein gradually accumulated in nucleus. Within 10 h, most SnRK1.1 protein was detected in nucleus under A5N1, pH 4.5 conditions (Figure 10E). Our results indicated SnRK1.1 accumulates in nucleus under both long-term and short-term high-ammonium stress conditions.

Discussion

SnRK1.1 acts as a central regulator responsible for multiple stress-related signaling. In this study, SnRK1.1 is identified as an important component contributing to nitrate-dependent alleviation of ammonium toxicity via negatively regulating SLAH3. In our hypothetical model, relatively lower levels of sugar lead to activation of SnRK1.1, likely in the sink organs such as root. SnRK1.1 is localized to both cytoplasm and nucleus under normal condition. Upon
Figure 8 S601 of SLAH3 plays a role in regulating nitrate-dependent alleviation of ammonium toxicity. A, 20-d-old plants of Col-0, slah3-4 and slah3-4 complemented with different variants of SLAH3 sequences are presented under different culture conditions. The phospho-dead variant (S601A) and phospho-mimetic variant (S601D) were driven by its native promoter. White scale = 2 cm. B, The statistical analyses of fresh weight on the plants presented in (A). Fresh weights of five seedlings per plate for each line were analyzed. Bars indicate SE, \( n = 3 \). C, The statistical analyses of main root length on the plants presented in (A). Root lengths of five seedlings per plate for each line were analyzed. Bars indicate SE, \( n = 3 \). Statistical analyses were performed by two-way ANOVA analysis with Dunnett’s multiple comparisons test. *\( P < 0.05; **P < 0.01 \). S, Sucrose.
activation, the cytosol-localized SnRK1.1 phosphorylates SLAH3-CT at S601, leading to SLAH3 inhibition. Under normal condition, the activation of SnRK1.1 is regulated by cellular energy status. The negative regulation of SLAH3 by SnRK1.1 may serve as a strategy to reduce nitrate loss in root under normal condition. It is likely an important mechanism of how plants coordinate energy homeostasis and nutrient usage. When the plants are subjected to high-ammonium stress, cytosol-localized SnRK1.1 migrates to nucleus. The translocation of SnRK1.1 releases its inhibitory effect on SLAH3, which in turn allows SLAH3 activation mediated by an unknown kinase(s). At the same time, high-ammonium/low-pH stress condition may cause severe energy deficiency, which leads to strong activation of nucleus-localized SnRK1.1 that regulates downstream components such as transcription factors responsible for the expression of energy/stress-related genes. Nitrate efflux mediated by SLAH3 ultimately causes nitrate accumulation in rhizosphere, which is sufficient to detoxify high-ammonium condition (Figure 11). Unlike the previously identified SLAH3 regulators that phosphorylate the NT of SLAH3 and function as SLAH3 activators, our study revealed that stress-related protein kinase SnRK1.1 phosphorylates the CT of SLAH3 and potentially acts as a SLAH3 inhibitor.

It is puzzling that for many plants NH$_4^+$ is toxic N source when it is dominant over NO$_3^/$C0$_3$ because NH$_4^+$ can be directly assimilated after absorption into the cell (Britto et al.,

**Figure 9** SLAH3 S601 influences the net NO$_3^-$ fluxes at the maturation zone in primary roots under high-ammonium/low-pH condition. A, The net NO$_3^-$ fluxes in Col-0, slah3-3, slah3-4, and complementary lines seedlings that were grown on 1/2 MS medium for eight days and transferred to high-NH$_4^+$ conditions (1 mM NO$_3^-$, 10 mM NH$_4^+$, pH 4.5) for 2 h were measured by NMT. The NO$_3^-$ fluxes at the maturation zone next to root tip in primary roots of Arabidopsis were measured. Each point represents the mean ± se of three individual plants. B, Mean values of NO$_3^-$ fluxes from (A) are presented. Data are shown as means ± se, n = 3. Statistical analyses were performed by one-way ANOVA with Tukey’s multiple comparisons test. Significant differences among all plants are indicated with different letters. P < 0.05.

**Figure 10** SnRK1.1 transfers from cytoplasm to nucleus under high-ammonium/low-pH conditions. A and B, The abundance of SnRK1.1 protein is decreased in cytoplasm (A) and increased in nucleus (B) under high-ammonium/low-pH condition. In Col-0 plants grown for 14 d in 1/2 MS and A5N1, pH 4.5 conditions, the protein contents in cytoplasm and nucleus were separated. SnRK1.1 protein abundance was analyzed by immunoblotting using an anti-SnRK1.1 antibody. Anti-PEPC and anti-Histone H3 antibodies were used to detect PEPC and Histone H3, marker proteins for cytoplasmic and nuclear sections, respectively. These assays were performed three times and showed similar results. C and D, The quantitative values of the band intensities of SnRK1.1. The ratios of SnRK1.1 to PEPC (C) and Histone H3 (D) were calculated. Data are shown as means ± se, n = 3. Statistical analyses were performed by one-way ANOVA with Tukey’s multiple comparisons test. Significant differences are indicated with different letters. P < 0.05. E, Transient treatment of high-ammonium/low-pH conditions causes accumulation of SnRK1.1 in nucleus. The Col-0 plants expressing YFP-SnRK1.1 grown for 5 d in 1/2 MS were transferred to liquid 1/2 MS or liquid A5N1, pH 4.5 condition. Starting from 2 h, in the plants that were subjected to A5N1 condition, YFP-SnRK1.1 signal was clearly detected to be accumulated in nucleus. White scale = 50 μm.
NO3− needs to be reduced into NH4+ via a two-step process, which is energy-consuming, before it can be utilized by plants (Crawford, 1995; Xu et al., 2012). In contrast to NH4+, NO3− usually does not cause obvious toxicity in plants when it is the sole N nutrient. The detailed mechanisms how a theoretically preferred nitrogen source inhibits plant growth are still largely unclear. The proposed causes of ammonium toxicity include futile ammonium cycling, inorganic cation depletion, organic acid depletion, accumulation of ROS, and medium acidification (Britto et al., 2001; Koyama et al., 2001; Britto and Kronzucker, 2002). However, no sufficient evidence is provided to clearly describe these mechanisms so far. Small amounts of nitrate can efficiently detoxify ammonium with unknown mechanisms (Domene and Ayres, 2001; Britto and Kronzucker, 2002; Houdusse et al., 2005). Our previous study revealed that a nitrate efflux channel, SLAH3, contributes to nitrate-dependent ammonium detoxification (Zheng et al., 2015). SLAH3-mediated nitrate efflux contributes to nitrate-dependent alleviation of ammonium toxicity, in which medium pH is an essential factor.

SLAC1 was the first characterized gene in the SLAC/SLAH family. SLAC1 regulates stomatal movement upon ABA signal or PAMP stimulus through Ca2+-dependent and Ca2+-independent pathways (Brandt et al., 2012; Maierhofer et al., 2014; Zheng et al., 2018). SLAH3 functions redundantly with SLAC1 in guard cells (Negi et al., 2008; Geiger et al., 2011; Demir et al., 2013). Nevertheless, distinct from SLAC1 that is restricted to guard cells, SLAH3 is expressed ubiquitously in plant tissues such as root and pollen tube, suggesting additional functions of SLAH3 besides controlling stomatal movement (Gutermuth et al., 2013; Zheng et al., 2015). Protein kinases were identified to phosphorylate SLAH3 to activate the SLAH3 channel. CPK21 activates SLAH3, a

![Figure 11](image-url)
component of nanodomain in mesophyll cells, which is regulated by an ABA-signaling mediator, phosphatase AB1 (Geiger et al., 2011; Demir et al., 2013). SLAH3 also functions in pollen tube tip likely regulating anion homeostasis during pollen tube growth (Gutermuth et al., 2013). Earlier studies have shown that the NT phosphorylation of SLAH3 or SLAC1 is involved in channel activation mediated by CPKs, SnRK2s, and CIPKs (Geiger et al., 2009; Maierhofer et al., 2014; Kudla et al., 2018). CPK21 interacts with the NT of SLAH3 and gates the opening of SLAH3. Therefore, the phosphorylation of NT of SLAH3 serves as a positive control for SLAH3 channel activity. A recent report showed an RLCK, PBL27, also phosphorylates the NT and opens SLAH3. However, whether the CT phosphorylation of SLAH3 also regulates SLAH3 activity is still unclear (Liu et al., 2019). Despite the discoveries that the CT of SLAH3 can be phosphorylated by different protein kinases, the physiological roles of the phosphorylation of SLAH3-CT are still beyond our understandings. In this study, we demonstrate that SnRK1.1 kinase can strongly phosphorylate the CT of SLAH3. More importantly, by using a combination of biochemical, molecular, genetic and NMT analyses, we show that this SnRK1.1-mediated CT phosphorylation likely serves as a negative regulation on SLAH3 (Figures 8 and 9; Supplemental Figure S8). Our study thus provides additional insights into the regulatory mechanisms of SLAH3.

SnRK1/SNF1/AMPK protein kinases are proposed to function as essential energy sensors in regulating energy homeostasis in eukaryotes (Polge and Thomas, 2007; Crepin and Rolland, 2019). SnRK1/SNF1/AMPK function as heterotrimERIC complexes consisting of three subunits consisting of a catalytic α-subunit, a scaffold β-subunit and an adenine nucleotide AMP or ADP-associating γ subunit (Polge and Thomas, 2007; Crozet et al., 2014; Crepin and Rolland, 2019). In plants, SnRK1 complex contains α kinase subunit SnRK1.1 or SnRK1.2, a plant-specific β subunit and a unique hybrid βγ subunit which lacks AMP binding domain (Broeckx et al., 2016; Crepin and Rolland, 2019). Different from SNF1 and AMPK that require adenine nucleotide-binding to be activated, SnRK1 α subunit possesses auto-phosphorylation in a complex-independent manner (Broeckx et al., 2016; Crepin and Rolland, 2019). The activation of SnRK1, distinct from SNF1 and AMPK, involves the removal of repressors but not association with adenine nucleotides (Crozet et al., 2014; Maya-Bernal et al., 2017; Emanuelle et al., 2018; Ramon et al., 2019). Intriguingly, this negative regulation of SnRK1.1 seems to be in line with its negative role in regulating SLAH3. Negative regulations seem to be plausible in plants, which often cause rapid downstream responses (Crepin and Rolland, 2019).

In Xenopus oocytes, SLAH3 is silent without activators and extracellular NO$_3^-$ (Geiger et al., 2011). However, in the natural conditions, the activators of SLAH3 can always be dynamically activated and there is always nitrate present. Therefore, SLAH3 can be activated and its activity needs to be repressed to prevent nitrate loss under normal conditions. In our model, the repression of SLAH3 activity can be released upon stressed conditions. Under higher concentrations of extracellular NO$_3^-$, the activity of SLAH3 should be increased (Geiger et al., 2011). However, we believe that extracellular NO$_3^-$-mediated SLAH3 activation barely contributes to nitrate-dependent alleviation of ammonium toxicity. Under high-ammonium/low-nitrate condition (ASN1, pH 4.5), slah3 mutants showed severe ammonium toxic phenotype compared to WT (Figure 1A). Whereas, when the concentration of nitrate was increased (ASNs, pH 4.5), the phenotypic difference between slah3 mutants and WT disappeared (Figure 1A). These results indicated SLAH3-mediated ammonium detoxification occurs under low concentrations of nitrate. Given that higher extracellular NO$_3^-$ will cause stronger SLAH3 activity, the result that SLAH3 can only detoxify ammonium under low extracellular NO$_3^-$ demonstrated that nitrate-activated SLAH3 activation unlikely contributes to SLAH3-mediated nitrate-dependent alleviation of ammonium toxicity. Thus, we speculated that the activation and inactivation of SLAH3 by intracellular regulators including unknown kinases and SnRK1.1, instead of extracellular NO$_3^-$, play a more important role in SLAH3-mediated nitrate-dependent alleviation of ammonium toxicity. Even though nitrate alleviation of ammonium toxicity could be in part resulting from the activation of SLAH3 by nitrate itself, it plays much less of a role in this process.

The activities of channels and transporters are fine-tuned by regulators such as kinases and phosphatases in order to fulfill physiological activities and to respond to various stress conditions (Köhler and Blatt, 2002; Geiger et al., 2009; Hedrich, 2012). Interestingly enough, a number of the members in the CPK-SnRK superfamily are engaged in these regulations (Hedrich, 2012). CPK–SnRK superfamily mainly contains CPKs family and SnRK family that comprises SnRK1, SnRK2, and SnRK3 subfamilies (Hrabak et al., 2003). Numerous reports have shown that CPKs, SnRK2s, and SnRK3s are extensively involved in phosphorylating and regulating channels and transporters (Kudla et al., 2018). Despite that SnRK1s have overlapping substrate specificity with CPKs and SnRK2/3, they have not been shown to act as channel or transporter regulators. Our study indicated SnRK1.1, like other members in the CPK–SnRK superfamily, also functions in regulating ion channel via phosphorylation. Besides its role in sugar signaling, SnRK1.1 is also suggested to mediate multiple stress responses (Baena-González and Sheen, 2008; Hulsmans et al., 2016). The functional analyses of SnRK1.1 demonstrate a complicated network in which a variety of stress signaling is connected by SnRK1.1. Our results suggested that high-ammonium/low-pH stress links with other stress response pathways, in which SnRK1 may function as a key regulator in the crosstalk.

A recent study revealed that PBL27 phosphorylates both NT (S127 and S189) and CT (S601) of SLAH3 during PAMP-induced stomatal closure (Liu et al., 2019). PBL27 can phosphorylate S127 and S189 to activate SLAH3. S601A or S601D
did not affect SLAH3 activation by PBL27. However, when SLAH3 S127D (or S189D)/S601D double mutation and PBL27 were coexpressed in oocyte, the SLAH3 S127D (or S189D)/S601D activation seemed to be reduced compared to coexpression of WT SLAH3, S127D, S601D, or S189D with PBL27 (Liu et al., 2019). These results indicated that although S601D single mutation does not affect SLAH3 activity, S601D may function together with other phosphorylation sites to regulate SLAH3 activity. Interestingly, based on their results, S601D may negatively regulate SLAH3 activation which is consistent with our results. Taken together, we conjecture that SLAH3 may be activated via N-terminal phosphorylation but be inhibited by S601 phosphorylation at C-terminal mediated by SnRK1.1. However, the activators that regulate SLAH3 in the process of nitrate-dependent alleviation of ammonium toxicity have not been identified yet. Whether S601 phosphorylation directly regulates the channel activity of SLAH3 remains to be clarified further. In addition, it is also possible that S601 phosphorylation leads to recruiting additional regulatory components, which are directly involved in regulating SLAH3 channel activity.

**Materials and methods**

Primer sequences used for all experiments are available in the Supplemental Table 1.

**Plant materials and growth conditions**

The slah3-3 (SALK_106054) and slah3-4 (SALK_111623) T-DNA insertion alleles were obtained from the ABRC. Overexpression plants were in the Columbia (Col-0) background. The overexpression plants were crossed to slah3-4 to create the overexpression lines in slah3-4 background. All complementary lines were in slah3-4 background.

Arabidopsis (*Arabidopsis thaliana*) grown in soil were cultivated in a green house at 22°C under a 16-h-light/8-h-dark photoperiod. Sterilized plump seeds were vernalized in the dark at 4°C for 3 d and sowed on plates. 1/2 MS medium (Phytotech) and Nitrogen-free MS medium (Caisson) were supplemented with different concentrations of ammonium and nitrate were used. Plates were sealed and incubated in rinse chambers at 22°C under a 16-h-light/8-h-dark photoperiod. For phenotypic analyses, and extraction and quantification of gene expression in seedlings, total RNA extraction was performed with RNAprep pure Plant Kit (TIANGEN) according to the manufacturer’s instructions. Two micrograms of total RNA were used for reverse transcription with the RevertAid Reverse Transcription System (Takara). *ACTIN2* was used as a reference gene. RT-qPCR was performed according to the manufacturer’s instructions. The PCR program was applied by a StepOne Plus Real Time PCR system (Applied Biosystems). The expression of *ACTIN2* was used as an internal control to normalize all data. Each experiment was performed using at least three independent biological replicates.

**GUS staining**

Different developmental stages and different tissues of transgenic plants were collected and stained. Tissues were first incubated in rinse solution (34.2 mM Na₂HPO₄, 15.8 mM NaH₂PO₄, 0.5 mM K₃Fe(CN)₆, and 0.5 mM K₄Fe(CN)₆·3H₂O) for 5 min, then incubated in stain solution (34.2 mM Na₂HPO₄, 15.8 mM NaH₂PO₄, 0.5 mM K₃Fe(CN)₆, 0.5 mM K₄Fe(CN)₆·3H₂O, and 2 mM X-Gluc) at 37°C for appropriate time. After GUS staining, the plant tissues were immersed in 30%, 50%, 75%, and 95% ethanol for 1 h in turn, and then immersed in 75% ethanol. After being decolorized, the tissues were observed with the Stereomicroscope (Leica).

**Yeast two-hybrid assays**

The Yeastmaker Transformation System 2 (Clontech) was used according to the manufacturer’s manual. The interaction tests were performed in yeast (*Saccharomyces cerevisiae*) strain AH109 (Clontech). To identify SLAH3-interacting proteins, the N-terminal and C-terminal region of SLAH3 coding sequences were cloned into pGBT9BS as the bait. cDNAs of the genes in CPK-SnRK superfamily were cloned from into pGADGH as prey. For analyzing the interaction promoter sequences about 1.6 kb were cloned into binary vector pBIB-BASTA-GWR-GUS. For constructing complementary plants, SLAH3 CDSs with the stop codon were cloned into binary vector pBIB-BASTA-pSLAH3-GWR. For BifC assay, the coding sequences of target genes were cloned into vector pEG202-GWR-nYFP or pEG202-GWR-cYFP. For constructs used in the yeast two-hybrid assays, the target gene fragments were cloned in vector pGBT9BS-GWR or pGADGH-GWR. For mbSUS assay, the CDSs were cloned in pMetYCgate and pX-NubWTgate plasmids. For prokaryotic expression, the target gene fragments were cloned into the vector pDEST15-GST-GWR. Site-directed mutagenesis was performed with PCR using complementary mutation primer pairs extending 15 nucleotides on either side of the modification. DpnI was used to digest the methylated template DNA. For YFP signal detection, the coding sequences of target genes were cloned into vector pEarleyGate-BASTA-35S-YFP-GWR. For CFP signal detection, the coding sequence of target gene was cloned into vector pBIB-BASTA-35S- GWR-CFP.
between SnRK1.1 and C-terminal of SLAH3, the full-length coding sequence and various truncated coding regions of SnRK1.1 were cloned into pGADGH. Empty vector combinations were used as negative controls. mbsUS was performed as previously described (Obrdlik et al., 2004). The coding sequence of SnRK1.1 was cloned into pX-NubWTgate vector to transform yeast strain THY.AP4, and the full-length coding sequence of SLAH3 was cloned into pMetYCgate vector to transform yeast strain THY.AP4. The empty vectors were used as negative control. Interactions were examined by growing diploid yeast cells on a synthetic dextrose (SD) minimal medium containing 0, 150, 300, or 500 μM Met.

**BiFC and subcellular localization assay**

BiFC assay was performed as previously described (Cui et al., 2018). For isolating leaf mesophyll protoplasts, 1.5 cm diameter infiltrated leaves were treated with d-Mannitol (500 mM) for 30–60 min, then incubated with 500 μL protoplast solution containing 500 mM d-Mannitol, 10 mM CaCl₂, 5 mM MES (pH 5.6), 3% (w/v) Cellulase-Onozuka, and 0.75% (w/v) Macerozyme in vacuum for 5 min. The samples were slightly vibrated in the darkness for 4 h, then transferred to the tubes and added half volume of CaCl₂ (200 mM). The materials were centrifuged at 40g for 5 min. The precipitates resuspended with 1 mL Mannitol (500 mM) and 1 mL CaCl₂ (200 mM) were centrifuged at 40g for 5 min. The protoplasts were resuspended with 200–500 μL WS solution (125 mM CaCl₂, 5 mM KCl, 154 mM NaCl, 5 mM glucose, 1.5 mM MES, adjust pH 5.6 with KOH). Leaf mesophyll protoplasts were observed using a two-photon laser confocal microscope (Olympus FV1000 MPE). For YFP visualization, excitation at 514-nm and detection between 525-nm and 620-nm were used. Laser power and PMT gain were adjusted appropriately. Excitation and emission settings of all samples were kept constant. Images were analyzed with image FV10-ASW software.

Subcellular localization of SnRK1.1 was determined by observing the YFP fluorescence signal. The YFP-SnRK1.1 overexpression plants were grown on 1/2 MS medium for 5 d, and then transferred to 1/2 MS or ASNI, pH 4.5 media with 0.9% agarose for 2 weeks, respectively. The seedlings were immediately frozen and ground well in liquid nitrogen. Cells were lysed with 3-mL lysis buffer containing 10 mM Tris–HCl (pH 7.5), 10 mM NaCl, 10 mM MgCl₂, 10% (v/v) Glycerol, 10 mM β-mercaptoethanol, and 0.1% protease inhibitor cocktail (Roche). Lysates were then filtered twice through a 100-mesh filter membrane. The filtrates were centrifuged at 2,500g for 10 min at 4°C. The supernatants were cytoplasmic crude extracts and the precipitates were nuclear crude extracts. The supernatants were transferred to new tubes and centrifuged at 13,000g for 15 min at 4°C. The supernatants were cytoplasmic fractions. The nuclear crude extracts were resuspended in 20 μL nuclear resuspension buffer containing 10 mM Tris–HCl (pH 7.5), 10 mM NaCl, 10 mM MgCl₂, 10 mM β-mercaptoethanol, 1 M hexylene glycol, 0.5% (v/v) Triton X-100, and 1 mM protease inhibitor cocktail, centrifuged at 1,500g for 10 min at 4°C. The supernatant was discarded. The resuspension process was repeated five times using nuclear resuspension buffer without protease inhibitor cocktail. The precipitate resuspended in 100-μL resuspension buffer was the nuclear fraction. All steps were performed at 4°C or on ice. The antibodies used in the experiments include anti-AKIN10 (1:500, Agrisera), anti-Histone H3 (1:5,000, Abcam), anti-PEPC (1:5,000, Rockland), and anti-rabbit (1:5,000, SeraCare).

**FRET assay**

To perform sensitized emission method, a set of three filter configurations were preset and three samples were prepared: one containing donor only (SnRK1.1/mSnRK1.1-CFP), one containing acceptor only (SLAH3-YFP), and one containing both donor and acceptor. Each of the samples was imaged with a donor filter set, a FRET filter set, and an acceptor filter set. The FRET filter set was configured to have an exciter that matched with the donor filter set and an emitter filter that matched with the acceptor filter set. To visualize emission, the three tracks were analogous to a donor (CFP) filter set (track 1: laser, 458-nm; emission 463–510 nm), a FRET filter set (track 2: laser, 458-nm; emission 520–620 nm), and an acceptor (YFP) filter set (track 3: laser, 514-nm; emission 520–620 nm). Laser power and PMT gain were adjusted appropriately. Excitation and emission settings of the three samples were kept constant. Images from each track were displayed in separate image channels.

To perform acceptor photobleaching method, regions of interest were chosen and bleaching parameters were defined, and then the predetermined regions were photobleached. The prebleach and postbleach images were acquired. The fluorescence intensity was recorded in each region before, during, and after selectively photobleaching. All FRET experiments were performed using the ZEN 3.1 system combined with a Zeiss LSM 880 confocal microscope.

**Expression and purification of recombinant SnRK1.1 and SLAH3-CT in Escherichia coli**

Recombinant GST-SnRK1.1, GST-SLAH3-CT, GST-SLAH3-CT (S601A), and GST-SLAH3-CT (S580A) were expressed in E. coli (Rosetta). Protein induction and purification was performed as previously described (Cui et al., 2018). The glutathione agarose beads (Sangon Biotech) was used according to the manufacturer’s manual.
**In vitro kinase assays**

For autoradiography, the experiment was performed as previously described (Lee et al., 2013).

Pro-Q diamond phosphoprotein gel stain provides a more efficient and safe method than autoradiography, and the phosphorylated proteins on the protein gel can be directly revealed by fluorescence scanner detection. The stain is compatible with a visible light-scanning instruments, imaging equipment with appropriate filters, blue-LED transilluminators, or 300-nm UV transilluminators and also can be used for mass spectrometry analysis. For this assay, the protein combinations were incubated in 20-μl kinase reaction solution containing 20 mM Tris–HCl (pH 8.0), 5 mM MgCl₂·6H₂O, 1 mM CaCl₂, 1 mM DTT, and 0.5 mM ATP for 25 min at 30°C. The reaction solutions added with 5 × SDS loading buffer were boiled at 95°C for 5 min. Then they were loaded into the sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) gel for separating proteins. The gel was first fixed by fixation solution (50% methanol, 10% acetic acid) for 30 min. The gel was then put in fixation solution and fixed overnight. After washing twice with ddH₂O for 15 min each time, the gel was stained with Pro-Q solution containing 1/3 volume of Pro-Q diamond phosphoprotein gel stain (Invitrogen) and 2/3 volume of ddH₂O for 2 h in the dark. All of the following experiments were performed in dark. The gel was washed four times by destaining solution (20% acetonitrile, 50 mM sodium acetate, pH 4.0) for 30 min and washed twice by ddH₂O for 20 min each time. Then the gel was observed with the Molecular Imager PharosFX (Bio-rad). Coomassie Brilliant Blue (CBB) staining of proteins was used for a loading control.

**Measurement of the net NO₃⁻ fluxes using the NMT system**

The net fluxes of NO₃⁻ were measured non-invasively using the NMT with Non-invasive Micro-test system (NMT150S, YoungerUSA LLC, Amherst, MA 01002, USA) and imFluxes V2.0 (YoungerUSA LLC, Amherst, MA 01002, USA) Software. The principle of this technology and instruments has been described previously (Sun et al., 2009). Measurements were performed at room temperature (25°C). Pre-pulled and silanized 8–10 μm aperture glass NO₃⁻-microsensor (XY-CGQ-01) were first filled with a backfilling solution (10 mM KNO₃) to a length of ~1.0 cm from the tip. LIX Holder (XY-LIX-01) was used to add NO₃⁻-LIX (NMT selective liquid ion-exchange cocktails, XY-SJ-NO₃-25) to the front of the microelectrode. An Ag/AgCl wire microsensor holder (YG003-Y11) was inserted in the back of the microsensor to make electrical contact with the electrolyte solution used as the reference microsensor. Prior to the flux measurements, the ion selective electrode was calibrated using the calibration solution 1 (0.2 mM KNO₃, 0.1 mM KCl, 0.1 mM CaCl₂, 0.3 mM MES, pH 6.0) and the calibration solution 2 (2 mM KNO₃, 0.1 mM KCl, 0.1 mM CaCl₂, 0.3 mM MES, pH 6.0) successively. To test whether Cl⁻ interferes with the measurements of NO₃⁻ in the NMT system, the NO₃⁻ concentrations and net fluxes in measuring solution 1 (1 mM KNO₃, 0.1 mM KCl, 0.1 mM CaCl₂, 0.3 mM MES, pH 6.0) and measuring solution 2 (1 mM KNO₃, 1.1 mM KCl, 0.1 mM CaCl₂, 0.3 mM MES, pH 6.0) were detected without seedlings. To analyze the net NO₃⁻ fluxes, the background was recorded by the vibrating electrode in measuring solution 1 without seedlings. After culturing on 1/2 solid medium for 8 d, the seedlings were treated with treatment solution (1 mM KNO₃, 10 mM NH₄Cl, pH 4.5) or control solution (1 mM KNO₃, 1 mM NH₄Cl, pH 5.7) for 2 h. Then the seedlings were transferred and fixed into small plastic dishes (6 cm diameter), and the measuring solution 1 was added into plastic dishes for measurement. The microelectrode vibrated between 5 and 35 μm from the root surface along an axis perpendicular to the root in the solution. The net fluxes of NO₃⁻ were measured individually at the mature zone next to the root tip. Each plant was measured for over 6 min. The final flux values for each treatment represent the average of 3 plants. The microelectrode, LIX Holder, NO₃⁻-LIX, and Ag/AgCl wire microsensor holder used in the experiment were purchased from Xuyue (Beijing) Science and Technology Co., Ltd., Beijing, China.

**Mass spectrometry analysis**

Protein samples separated by SDS–PAGE gels and stained with the CBB were cut to small pieces and washed twice by ddH₂O for 10 min each time; 200-μL 100 mM NH₄HCO₃/ACN (1:1) was added to decolorize the gel at 37°C for 2 min. Then the samples were dehydrated by 200-μL ACN and evaporated to dryness in rotary vacuum evaporator for 5 min. The gel particles were incubated in 50-μL 10 mM TCEP (NH₄HCO₃ attenuating) at 56°C for 1 h, and then incubated in 50-μL 40 mM CAA (NH₄HCO₃ attenuating) in the dark for 45 min. Dried micelles were washed with 200-μL 25 mM NH₄HCO₃, 50% ACN and 100% ACN for completely dehydration. The gel particles were incubated with 50 μL ACN for 15 min, and evaporated to dryness in rotary vacuum evaporator for 5–10 min after removing ACN. Then the gel particles were incubated with 50-μL 0.01 mg/mL trypsin solution (50 mM NH₄HCO₃ attenuating) at 37°C overnight. The peptides were extracted with 50-μL 50% ACN/0.1% TFA twice for 10 min each time by ultrasonic instrument, evaporated to dryness in rotary vacuum evaporator and stored at −80°C. The samples were added 15-μL 0.1% FA, and centrifuged at 18000g for 30 min twice. The samples were analyzed by an Orbitrap Fusion Lumos mass spectrometer (Thermo Fisher Scientific) connected to an EASY-nLC 1200 system. The software Proteome DiscovererDaemon 2.2 (Thermo Fisher Scientific) was used for data analysis.

**Plant fresh weight and root length assays**

Plant fresh weight for quantitative analysis was measured by Ten-thousandth electronic analytical balance. The sum of fresh weights of 5 seedlings per plate for each line was analyzed. The final values for each treatment represent the average of 3 plates. As another indicator of plant growth,
primary root length was measured by the National Institutes of Health Image software ImageJ. The vertical distance between hypocotyl junction and primary root tip was defined as root length. The average of root lengths of five seedlings per plate for each line was analyzed. The final values for each treatment represent the average of three plates.

Reproducibility and statistics
All experiments were repeated at least three times. Statistical analyses were indicated in each figure legend, and performed using the program GraphPad Prism.v8. Statistical significance is indicated by the P-value (*P < 0.05; **P < 0.01). Statistical analyses were performed by one-way analysis of variance (ANOVA with Tukey’s multiple comparisons test, two-way ANOVA analysis with Tukey’s multiple comparisons test or two-way ANOVA analysis with Dunnett’s multiple comparisons test.

Protein structure analysis website
The topology of SLAH3 protein was predicted by http://wlab.ethz.ch/protter/start/.

Accession numbers
Sequence data from this article can be found in the Arabidopsis Information Resource database (TAIR, http://www.arabidopsis.org/) under the following accession numbers: SnRK1.1, AT3G01090; SnRK1.2, AT3G29160; SLAH3, AT5G24030.

Supplemental data
The following materials are available in the online version of this article.

Supplemental Figure S1. The sensitized emission method of FRET indicates that SnRK1.1/mSnRK1.1 interacts with SLAH3 in planta.

Supplemental Figure S2. Expression analyses of SnRK1.1/mSnRK1.1 in SnRK1.1/mSnRK1.1 overexpression lines in Col-0.

Supplemental Figure S3. Glucose contributes to SnRK1.1-mediated high-ammonium responses.

Supplemental Figure S4. Palatinose does not contribute to SnRK1.1-mediated high-ammonium responses.

Supplemental Figure S5. SnRK1.1 regulates ammonium toxicity depending on SLAH3.

Supplemental Figure S6. Expression analyses of SLAH3 gene in slah3-4 mutant complemented with different SLAH3 variants and SLAH3.

Supplemental Figure S7. The phosphorylation status of S601 does not affect the interaction of SLAH3 with SnRK1.1.

Supplemental Figure S8. SLAH3 can complement the sensitivity phenotype of slah3-4 to high-ammonium/low-nitrate and low-pH condition.

Supplemental Figure S9. The presence of Cl− does not interfere with the measurements of NO3− in the NMT analyses.

Supplemental Figure S10. SLAH3 S601 does not influence the net NO3− fluxes under non-high-ammonium/low-pH condition.

Supplemental Figure S11. Original western blot data of Figure 10.

Supplemental Table S1. Primer sequences used in this study.

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Conflict of interest statement
The authors declare that they have no conflict of interest.

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