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Transceptor NRT1.1 and receptor-kinase QSK1 complex controls PM H⁺-ATPase activity under low nitrate

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Running title: NRT1.1-RK complex controls proton pump activity in low nitrate

1 Summary

NRT1.1, a nitrate transceptor, plays an important role in nitrate binding, sensing and 2 nitrate dependent lateral root (LR) morphology. However, little is known about NRT1.1-3 mediated nitrate signaling transduction through plasma membrane (PM)-localized 4 5 proteins. Through in-depth phosphoproteome profiling using membranes of Arabidopsis roots, we identified receptor kinase QSK1 and plasma membrane H⁺-6 7 ATPase AHA2 as potential downstream components of NRT1.1 signaling in a mild low nitrate (LN)-dependent manner. QSK1, as a functional kinase and molecular link, 8 9 physically interacts with NRT1.1 and AHA2 at LN, and specifically phosphorylates AHA2 at S899. Importantly, we found that LN, not HN, induces formation of NRT1.1-10 QSK1-AHA2 complex in order to repress the proton efflux into the apoplast by 11 increased phosphorylation of AHA2 at S899. Loss of either NRT1.1 or QSK1 thus 12 13 results in a higher T947/S899 phosphorylation ratio on AHA2, leading to enhanced pump activity and longer LRs under LN. Our results uncover a regulatory mechanism 14 15 in which NRT1.1, under LN conditions, recruits coreceptor QSK1 into a complex to transduce LN sensing to the PM H⁺-ATPase AHA2, controlling the phosphorylation 16 ratio of activating and inhibitory phosphorylation sites on AHA2. This then results in 17 altered proton pump activity, apoplast acidification, and regulation of NRT1.1-mediated 18 LR growth. 19

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21 Introduction

Nitrogen is an essential macronutrient and nitrate serves as an important signaling 22 molecule required for growth, gene expression, and metabolism in plants.¹ The nitrate 23 concentration in soil can influence plant root growth, shoot elongation, as well as crop 24 25 yield. Improving nitrate use efficiency in plants is critical for increasing agricultural production. The molecular mechanism of how nitrate is sensed and transported into 26 plants is well known since the characterization of the nitrate transporter and sensor 27 (transceptor) NRT1.1² (CHL1/NPF6.3) and the high-affinity nitrate transporter NRT2.1³ 28 in Arabidopsis thaliana. 29

NRT1.1 in Arabidopsis is essential for nitrate transport, sensing, binding and 30 signaling.⁴ Phosphorylation of NRT1.1 at T101 by the CIPK23-CBL9 calcium sensor-31 kinase complex switches NRT1.1 from a low-affinity to a high-affinity transporter and 32 increases NRT1.1-dependent nitrate uptake.⁵ Dephosphorylation of NRT1.1 is fine-33 tuned by protein phosphatase ABI2 (abscisic acid insensitive 2).⁶ NRT1.1 activation 34 also induces a rapid increase in cytoplasmic Ca²⁺ levels.⁷ Recently, the NRT1.1-35 CNGC15 module was found to be a molecular switch that controls calcium influx in a 36 nitrate-dependent manner.⁸ Calcium subsequently activates CPK10/30/32 to 37 phosphorylate transcription factor NLP7 at S205, promoting NLP7 retention in the 38 nucleus.⁹ Very recently, NLP7 was shown to be an intracellular nitrate sensor, as 39 nitrate can bind NLP7 directly.¹⁰ In rice, OsNRT1.1B-SPX4-NBIP-OsNLP3 was the first 40 module that demonstrated nitrate signaling from the plasma membrane (PM) to the 41 42 nucleus.¹¹ However, little is known about whether more regulatory components in NRT1.1-mediated nitrate signal transduction exist in the PM. 43

NRT1.1 also plays an important role in root morphology by directly influencing lateral 44 root (LR) growth. Under low nitrate (LN) availability, NRT1.1 as well as phosphorylation 45 of NRT1.1 at T101 plays a key role in repression of LR growth, and NRT1.1 is 46 considered to repress LR growth through its auxin transport capacity.^{12,13} Other studies 47 also revealed that there is a crosstalk between nitrate and auxin in modulating of 48 NRT1.1-dependent LR growth.¹⁴ Plasma membrane-localized H⁺-ATPases pump H⁺ 49 into the apoplast between the plasma membrane and cell wall, resulting in acidification 50 of the apoplast and increased cell elongation.^{15,16} However, it remains unknown 51

whether NRT1.1-mediated LR growth inhibition under LN relies on altered PM H⁺ ATPase activity and apoplastic pH.

54 Many studies have suggested that genes involved in nitrate signaling are regulated at 55 the transcriptional level.¹⁷⁻¹⁹ However, there are indications of posttranslational 56 regulation of nitrogen uptake, sensing, and metabolism through phosphorylation.²⁰ 57 Numerous nitrogen-dependent phosphoproteomic studies²¹⁻²⁴ identified PM-localized 58 transporters and receptor kinases, which showed nitrate-dependent phosphorylation 59 changes. However, whether, and by which molecular mechanisms these PM proteins 60 are involved in nitrate signaling and transport regulation still needs to be explored.

Here we carried out a systematic functional study of roots of WT and *nrt1.1-1* to generate mild low and high nitrate-induced membrane protein phosphorylation profiles. Using these profiles, we aimed to discover new regulatory components for NRT1.1mediated nitrate signal transduction. Ultimately, we uncovered a novel function of NRT1.1, which interacts with the receptor kinase QSK1 to constitute a molecular switch that controls PM H⁺-ATPase activity through phosphorylation balance on different phosphorylation sites (T947/S899) of AHA2 in a mild low nitrate dependent manner.

68 Results

Functional membrane phosphorylation profiling identifies downstream components of NRT1.1

71 We performed a comparative phosphoproteomic analysis of Arabidopsis root membrane protein comparing WT and *nrt1.1-1* under mild low nitrate (LN) and high 72 73 nitrate (HN)-induced conditions after nitrogen starvation (NS) (Figure 1A) to identify 74 downstream components that may be involved in NRT1.1-dependent LR growth and signal transduction. We identified 5,950 phosphosites corresponding to 1,945 proteins 75 76 (Figure S1A). Among these, there were 2,390 previously uncharacterized phosphosites and 3,246 new nitrate-induced phosphosites compared to the 77 PhosPhAt4.0 database²⁵ and published nitrate-induced phosphosites in Arabidopsis, 78 respectively²¹⁻²³ (Figure S1B and S1C). Among these were 2,295 phosphosites 79 80 corresponding to 1,280 proteins with localization probability >0.75 (class I sites), which were used for quantitative statistical analysis (Supplementary Table 1). The 81 82 phosphorylation sites distributed to peptides with 93.4% single, 6% double, and 0.6% triple or more phosphorylation (Figure S1D). Principal component analysis (PCA) 83

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revealed a strong separation of WT under LN and HN along the first principal component, and data points for WT at LN and *nrt1.1-1* at LN separated along the second component. Data points for WT and *nrt1.1-1* at HN clustered tightly and showed no separation (Figure 1B).

88 We performed stringent pairwise comparison analysis for genotypes and treatments to separate LN and HN specific responses. In the WT, 642 phosphosites exhibited 89 significantly different abundance between LN vs NS or HN vs NS. Of these, 579 and 90 36 phosphosites were specifically changed under LN and HN, respectively. 515 of 579 91 phosphosites exhibited a higher abundance under LN (Figure 1C). Overall, LN induced 92 larger changes in the membrane phosphoproteome compared to HN. Subsequently, 93 94 we compared the phosphoproteome of the WT and nrt1.1-1. Under LN, 661 phosphosites were found with altered abundance in *nrt1.1-1*, out of which 608 95 96 phosphosites showed significantly decreased phosphorylation in *nrt1.1-1* (Figure 1D). The asymmetry of the plots suggested that the absence of NRT1.1 had a strong impact 97 98 on the membrane phosphoproteome under LN. Overlap of phosphosites with significantly changed phosphorylation in WT under LN vs. HN, and under LN in 99 *nrt1.1-1* compared to WT identified 307 phosphosites corresponding to 231 proteins 100 as candidates for being downstream regulators of NRT1.1 (Figure 1E), indicating LN-101 induced phosphorylation of these proteins is dependent on the presence of NRT1.1. 102 103 These 231 proteins were enriched in functions of Signaling (bin30), Transport (bin34), 104 RNA (bin27), and Cell (bin31), and were overrepresented with subcellular locations at 105 the PM and in the N (nucleus) (Figure 1F).

106 We then ranked the phosphosites with significantly changed phosphorylation in nrt1.1-107 1 under LN (based on p values) and highlighted phosphosites on 19 transport-related 108 proteins (Figure S1E). Among the highly-ranked phosphorylation sites, T947 and S899 109 from the plasma membrane H⁺-adenosine triphosphatase 2 (PM H⁺-ATPase isoform, 110 AHA2) were also identified (Figure S2A and S2B). AHA2 is the major proton pump in 111 roots. These two residues were reported as resulting in activation (T947) and inactivation (S899) upon phosphorylation^{26,27}, and are located in the C-terminal 112 113 autoinhibitory domain.²⁸ AHA2 is known to promote both lateral root and primary root growth under LN.²⁹ T947 and S899 in AHA2 exhibited significantly higher 114 115 phosphorylation in WT in LN than in HN, and phosphorylation at both sites, but especially S899 was drastically decreased in *nrt1.1-1* under LN, but not HN (Figure 2A) 116

and 2B). AHA2 protein abundances showed no change (Figure 2C). These results 117 suggested that the presence of NRT1.1 is required for phosphorylation of AHA2 under 118 119 LN. Since phosphorylation at T947 and S899 have contrasting effects on AHA2 activity, 120 we used the ratio of T947 and S899 phosphorylation as an indication of the balance of 121 AHA2 activity. Interestingly, the phosphorylation ratio of T947/S899 was significantly 122 increased in WT at LN compared to HN, and an even more increased ratio was found 123 in nrt1.1-1 compared to WT under LN, while no difference was observed under HN (Figure 2D). This suggests that loss of NRT1.1 disrupted the balance of 124 125 phosphorylation between T947 and S899 at LN due to a stronger reduction of 126 phosphorylation at S899.

127 NRT1.1 represses proton efflux at LN

128 PM-localized ATPases fine-tunes proton efflux into the apoplast, resulting in apoplastic 129 acidification. To assess whether NRT1.1 is required for H⁺-ATPase activation under LN, we applied membrane-impermeable 8-hydroxypyrene-1,3,6-trisulfonic acid 130 trisodium salt (HPTS) as a ratiometric fluorescent pH indicator for assessing changes 131 in the apoplastic pH at cellular resolution in the LR elongation zone.³⁰ Two different 132 forms of HPTS (protonated and deprotonated) were observed in two channels using a 133 confocal microscope with excitation wavelengths of 405 nm and 488 nm, respectively. 134 135 The ratiometric value (458/405) represents the apoplastic pH, and a high ratio represents a relatively high pH. The WT exhibited a lower pH in LN compared to HN. 136 137 Moreover, an even lower pH was observed in *nrt1.1-1* under LN (Figure 2E and 2F), 138 uncovering that the higher phosphorylation ratio of T947/S899 (Figure 2D) resulted in 139 activation of H⁺-ATPase in WT under LN compared to HN, and an even more increased 140 activation of the H⁺-ATPase in *nrt1.1-1* than in WT under LN. These results suggested that PM H⁺-ATPases function downstream of NRT1.1, mediating the LN-induced 141 142 proton efflux into the apoplast space. In turn, the phosphorylation ratio of T947/S899 143 controls the AHA2 activity.

As a result of enhanced proton pump activity, the average cortical cell length of LRs in the WT was longer under LN than HN, and an even longer cortical cell length was observed in *nrt1.1-1* under LN, but not under HN (Figure 2G and 2H). Consequently, WT displayed longer LR length under LN, and *nrt1.1-1* exhibited more enhanced LR growth (Figure S2C and S2D). Taken together, our results suggested that the lower

apoplastic pH in the LR elongation zone led to longer cortical cell length and LRs in 149 WT in LN, and that the even lower apoplastic pH in *nrt1.1-1* produced even longer 150 cortical cell length and LRs. The longer LRs in *nrt1.1-1* under HN was not correlated 151 with the apoplastic pH. Under LN, the reduced inhibition of AHA2 at S899 in *nrt1.1-1* 152 153 seems to override the co-occurring slightly decreased phosphorylation at T947, the 154 activating site. Direct interaction of NRT1.1 and AHA2 was not observed by ratiometric bimolecular fluorescence complementation (rBiFC) assays (Figure 2I and S3A), the 155 known interaction of CBL9-CIPK23 and the known absent interaction of CBL9-CIPK14 156 157 were used as positive control and negative control, respectively. As NRT1.1 itself cannot phosphorylate AHA2 at different sites, we postulate a kinase exists as a 158 159 molecular link between NRT1.1 and AHA2 under LN.

160 **Coreceptor QSK1 functions downstream of NRT1.1 to repress LR growth in LN**

To assess the potential kinases involved in this NRT1.1-mediated regulation, we 161 ranked phosphorylation sites (based on p value) on receptor kinases (Figure 3A and 162 163 Figure S3B), and found that phosphorylation of the coreceptor QSK1 at S621/S626 was detected in the WT under LN, however, no phosphorylation of QSK1S621/S626 164 was detectable either in the WT under HN or in *nrt1.1-1* (Figure 3B), while the protein 165 abundance of QSK1 was unchanged (Figure 3C). Therefore, we conclude that NRT1.1 166 167 is required for the phosphorylation of QSK1 at S621/S626 under LN. QSK1 phosphorylation at S621/S626 was also found in previous experiments under LN 168 169 condition.^{22,23}

170 QSK1 is strongly expressed in the root elongation zone and LR primordia³¹, which overlaps with the expression pattern of NRT1.1.¹² To investigate whether QSK1 as well 171 172 as its phosphorylation sites regulates LN-induced LR growth in similar way as NRT1.1, 173 we generated transgenic plants overexpressing QSK1, phospho-dead (AA) and phosphorylation-mimic QSK1 (DD) at S621S626 into *qsk1*. Compared to WT, the LR 174 length was significantly longer in *qsk1* under LN, and QSK1-OE/*qsk1* can complement 175 176 *qsk1* phenotype. Consistent with the observation of *qsk1* under LN, LR of QSK1-AA-OE/qsk1 were significantly longer when compared with the WT, while QSK1-DD-177 OE/gsk1 displayed the similar phenotype as WT under LN (Figure 3D and 3E), implying 178 that phosphorylation-mimic QSK1 can rescue the phenotype of *qsk1*, repressing LR 179 180 growth under LN.

Published protein-protein interaction networks from Arabidopsis roots suggested that 181 QSK1 may form a complex with NRT1.1 and AHA2 in a LN-dependent manner.³² rBiFC 182 assays confirmed that NRT1.1 directly interacts with QSK1 in the plasma membrane 183 (Figure 4A and Figure S4A). H356 of NRT1.1 is critical for nitrate binding and 184 uptake.^{33,34} To test whether nitrate binding is required for interaction of NRT1.1 with 185 QSK1, we mutated H356 to H356A. Indeed, interaction between NRT1.1H356A and 186 QSK1 was weaker compared to NRT1.1-QSK1 (Figure 4A and Figure S4A). To further 187 test whether LN-induced phosphorylation of NRT1.1 at T101² and QSK1 at S621S626 188 189 (Figure 3B) are essential for NRT1.1-QSK1 interaction, we analyzed the interaction of different combinations of NRT1.1 and QSK1 mutants. We found that the NRT1.1-QSK1 190 191 interaction was not affected by either the phosphorylation-dead (NRT1.1-A) or phosphorylation-mimic version (NRT1.1-D) of NRT1.1T101. The phospho-dead 192 193 QSK1S621/S626 (QSK1-AA) still interacted with NRT1.1-A, but the interaction of NRT1.1 and QSK1 was significantly enhanced by phosphorylation mimic QSK1 194 195 (QSK1-DD) and NRT1.1 (NRT1.1-D) (Figure 4A and Figure S4A). Importantly, coimmunoprecipitation (Co-IP) assays using membrane proteins isolated from tobacco 196 197 leaves expressing NRT1.1-eGFP and QSK1-FLAG showed that association between 198 NRT1.1-eGFP and QSK1-FLAG was enhanced when LN is present, not HN, 199 suggesting that LN promotes the NRT1.1-QSK1 interaction (Figure 4B). Altogether these results demonstrated that NRT1.1 directly interacts with QSK1, and this 200 interaction is facilitated by the perception of LN by NRT1.1 and subsequent 201 202 phosphorylation events.

203 We next created a *nrt1.1-1gsk1* double mutant to analyze the effect on LR growth. 204 Compared to WT, the LRs length was significantly longer in *qsk1*, *nrt1.1-1*, and the 205 *nrt1.1-1qsk1* double mutant in LN, while the LR length in the double mutant resembled 206 that of the two single mutants. At HN, only the *nrt1.1-1* and the *nrt1.1-1qsk1* double mutant showed longer LRs length, with no effect observed in *qsk1* (Figure 4C and 4D). 207 208 These results suggested that QSK1 and NRT1.1 function in the same signaling 209 pathway to repress LR growth in LN. To elucidate whether QSK1 contributes to nitrate 210 uptake under LN, we performed nitrate influx assays on N starved plant or after 211 induction with 1 mM nitrate for 1 or 4 h (Figure 4E). Nitrate influx was measured after 212 supplying a nutrient solution containing 200µM K¹⁵NO₃ during 5 minutes. In N starved 213 plants, higher uptake rates were observed in *qsk1* compared to WT. This difference was not observed when plants were treated with 1 mM NO₃⁻. These results indicated

that QSK1 inhibits LR growth and nitrate uptake in LN-dependent manner.

QSK1 interacts with and phosphorylates AHA2S899 and represses proton pump activity in LN

To investigate association of QSK1 and AHA2, we then performed rBiFC and Co-IP assays. rBiFC assays confirmed the interaction of QSK1-DD and AHA2, while QSK1-AA and AHA2 interaction was not obvious (Figure 5A and Figure S4B). Using Co-IP assays, a much stronger association was observed between QSK1-DD-FLAG and AHA2-MYC compared to QSK1 or QSK1-AA with AHA2 (Figure 5B).

223 We next examined the phosphorylation status of AHA2 at T947 and S899 in WT and *qsk1* roots. LN-induced phosphorylation of AHA2 at T947 and S899 was significantly 224 225 decreased in *qsk1*, while under HN no changes were observed for either phospho-site (Figure 5C and 5D), suggesting that under LN, QSK1 is required for phosphorylation 226 227 of both sites on AHA2. The protein abundances of AHA2 displayed no change (Figure 5E). The phosphorylation ratio of T947/S899 was also disrupted in *gsk1*, being 228 229 significantly higher in *qsk1* compared to WT under LN, but not under HN (Figure 5F). The AHA2 phosphorylation patterns found in *qsk1* resemble T947/S899 ratio found in 230 231 nrt1.1-1 under LN, implying an increased activity of H⁺-ATPase in also gsk1. Furthermore, an in vitro peptide phosphorylation assay using the recombinant 232 233 intracellular domain of QSK1 (QSK1C) and synthesized peptides covering S899 or T947 site as substrates, confirmed that QSK1C could phosphorylate S899, but not 234 235 T947 on the synthesized peptide (Figure 5G).

To verify the role of QSK1 in impeding PM H⁺-ATPase activity under LN, we again used HPTS as a pH indicator on the LR elongation zone of WT and *qsk1. qsk1* showed lower apoplastic pH in the LR elongation zone under LN compared to WT (Figure 5H and 5I), which correlates with longer cortical cell (Figure 5J and 5K) and LR length (Figure 3E). These results suggested that QSK1 could serve as a downstream component of NRT1.1 and that QSK1 has a key role in repressing apoplast acidification through direct phosphorylation of AHA2 at S899 under LN, not HN condition.

243 NRT1.1-QSK1 complex coordinately transduce LN sensing to the proton pump

Our results show that LN enhanced NRT1.1-QSK1 interaction (Figure 4B), induced 244 phosphorylation of QSK1 (Figure 3B), and phosphorylated QSK1 displayed stronger 245 association with AHA2 (Figure 5B). Thus, we propose that NRT1.1-mediated LN 246 sensing may modulate the QSK1-AHA2 interaction. To test this hypothesis, we 247 performed Co-IP assays using membrane protein mixture extracted from Nicotiana 248 benthamiana leaves harboring NRT1.1-eGFP, QSK1-FLAG and AHA2-MYC in 249 presence of LN, HN, or KCI. A strong interaction between QSK1-FLAG and AHA2-250 MYC was observed in the presence of LN-NRT1.1, with the interaction becoming 251 252 weaker in the presence of HN-NRT1.1 or KCI- NRT1.1 (Figure 6A). We also analyzed 253 the effect of LN on apoplast pH and cortical cell length of LRs in double mutant nrt1.1-254 1qsk1. As expected, nrt1.1-1qsk1 double mutant showed lower apoplastic pH in LR 255 elongation zone and longer cells when compared to WT, but no differences when 256 compared to respective single mutants under LN (Figure 6B and 6C). Together, these results suggested that LN promotes NRT1.1-QSK1-AHA2 complex in order to 257 258 transduce the LN signal from NRT1.1 to AHA2 through QSK1. Formation of this complex then ultimately results in repression of the proton pump activity through 259 260 phosphorylation at inhibitory S899, which ultimately then affects cell elongation and 261 nitrate uptake.

262 Discussion

NRT1.1 is a plasma membrane-localized nitrate transceptor and critical for nitrate 263 binding and sensing. However, it is still poorly understood how NRT1.1 mediates 264 265 transduction of nitrate signals within plasma membrane localized proteins. Recent findings that NRT1.1 can form a complex with Ca²⁺channel CNGC15 suggest that 266 267 protein-protein interactions are the molecular mechanism coupling nitrate sensing and signaling to downstream processes such as Ca²⁺ signaling.⁸ Here, based on in-depth 268 269 membrane phosphoproteomics analysis, our data provide a first unbiased view of 270 NRT1.1-dependent phosphorylation events under LN (Figure 1). A noteworthy finding 271 is that phosphorylation of several receptor kinases and transporters under LN depends on the presence of NRT1.1 (Figure 3A and Figure S1E), for example, three of the 272 273 known ammonium and nitrate-regulated phosphosites, T460 and S480/S489, in the cytosolic C-terminal region of AtAMT1;1 were identified.³⁵ This opens the possibility 274 that some of these proteins could be involved in further complexes with NRT1.1 275 276 integrating, for example, nitrate and ammonium signaling.

277 Our work describes a new role of NRT1.1 in modulation of AHA2 activity by formation of a protein-protein interaction complex. This is unlike the ability of NRT1.1 to directly 278 transport auxin.^{12,13} Since NRT1.1 is not directly associated with AHA2 (Figure 2I) we 279 postulated that kinases exit between NRT1.1 and AHA2, and at the plasma membrane, 280 receptor kinases are likely candidates linking NRT1.1 and proton pump AHA2 (Figure 281 282 3A). Receptor kinases play critical roles in perception and transduction of extracellular signals, as well as in plant growth and defense.³⁶ Several studies have uncovered that 283 RKs could directly interact with PM-localized transport proteins and regulate their 284 activity. For example, TMK1, PSY1R, and BAK1 are well characterized in activation of 285 AHA2 through phosphorylation modification at different sites on AHA2^{27,37-40}, while 286 FERONIA can induce phosphorylation of S899 on AHA2, inhibiting proton efflux.²⁶ 287 FLS2 interacts with Ca²⁺ -ATPase ACA8 to regulate Ca²⁺ bursts⁴¹, and SIRK1 can 288 phosphorylate and activate aquaporins.^{42,43} These short, posttranscriptional and direct 289 regulatory circuits between RKs and transport proteins indicates a rapid and economic 290 291 mechanism, allowing plants to efficiently respond to local signals and rapidly adjust to changes in the environment. Our findings provide a first evidence for a transceptor 292 293 being linked to another transmembrane transport proteins (here: AHA2) by a receptor 294 kinase, QSK1.

QSK1 ("thousand-hand" giān shou kinase 1) is a highly expressed co-receptor kinase 295 in Arabidopsis thaliana. Depending of the physiological state of the plant, it is able to 296 directly phosphorylate specific substrate proteins, such as an ABC camalexin 297 transporter upon fungal infection⁴⁴ and potassium channel TPK1 during stomatal 298 closure.⁴⁵ It can also recruit substrates such as the aquaporin PIP2;4 to other receptor 299 300 kinases, such as SIRK1, forming a complex ultimately resulting in substrate phosphorylation.^{42,43} Therefore, QSK1 is proposed to be a coreceptor and responsible 301 302 for recruiting different substrates to the respective signaling complexes. QSK1 is known to be able to change localization, being depleted from ordered low-density 303 304 membranes (DRM) upon treatment with cytochalasin D or oryzalin.⁴⁶ Here, we show 305 that phosphorylated QSK1 recruits and phosphorylates AHA2 under LN (Figure 5).

306 Posttranslational regulation of PM H⁺-ATPases by phosphorylation and 307 dephosphorylation plays a crucial role in the activation or inactivation in response to 308 stress.⁴⁷⁻⁴⁹ Several phosphorylation sites within the autoinhibitory C-terminal domain 309 of PM H⁺-ATPases, which differentially affect pump activity, have been identified.

Among them, phosphorylation at T947 and T881 of AHA2 increases pump 310 activity^{27,37,38}, while phosphorylation at S899 and S931 inhibits pump activity.^{26,50} flg22, 311 a peptide derived from pathogenic bacterial flagella, elicits an increase in 312 phosphorylation at S899 and a decrease at T881 and T947, and leads to a rapid 313 apoplastic alkalinization.⁵¹ PM-localized ATPases regulates H⁺ efflux into the apoplast, 314 resulting in apoplastic acidification, which in turn affects LR growth. The proton 315 gradient is also used for ion uptake as well as cell elongation. Also, nitrate uptake has 316 been found to be coupled to protons.³³ We found that the elevated phosphorylation 317 ratio T947/S899 on AHA2 in nrt1.1-1 and gsk1 at LN results in higher AHA2 activity 318 and subsequently longer LRs (Figure 2 and Figure 5). Our results show that the PM 319 320 H⁺-ATPase activity is controlled coordinately by the balance of phosphorylation at 321 different sites. Since QSK1 specifically phosphorylates the S899 on AHA2 (Figure 5G), thus, we propose there exist other proteins (e.g., RKs or other Kinases) downstream 322 323 of NRT1.1 which phosphorylate T947 on AHA2.

324 It is interesting to note that QSK1 is a coreceptor, not a ligand-binding receptor kinase. Further research will be necessary to identify possible receptor kinase, as well as their 325 ligands, may function in a NRT1.1 dependent or independent manner. As NRT1.1 326 plays a critical role in the assembly of the here described signaling module, one may 327 speculate that NRT1.1, as a transceptor, functions as a main receptor to transduce the 328 329 LN signal to co-receptor QSK1. Mild nitrate deficiency activates BR signaling and induces LR elongation⁵², while the role of auxin in LN-mediated LR elongation has 330 been well studied.^{12,13,53} Additional research will be directed in exploring whether BR 331 and auxin are involved in regulating the NRT1.1-QSK1-AHA2 module. 332

333 In summary, we demonstrate that at LN, nitrate binding to NRT1.1 is able to enhance 334 the NRT1.1-QSK1 interaction and phosphorylation of S621S626 on coreceptor QSK1. 335 This is the first example for a nutrient sensing transporter to recruit a co-receptor for signal transduction. Phosphorylated QSK1 then recruits AHA2 into the complex in 336 order to phosphorylate S899 on AHA2. Since also phosphorylation of AHA2 at T947 337 can be affected by NRT1.1 or QSK1, we suspect further yet unknow kinases to be 338 339 involved in regulation of AHA2 activity directly or indirectly. (Figure 6D). Our results 340 have not only identified the key role of RKs and proton pump AHA2 under LN signaling 341 but has also coupled LN sensing from NRT1.1 to the activity of the proton pump.

342 Methods

343 Plant Material and growth conditions

For liquid growth cultures, Arabidopsis seeds of WT (Col-0), gsk1 mutant 344 (SALK_019840), and nrt1.1-1 mutant (SALK_097431) were sterilized and grown in 345 basal medium composed of micro- and macronutrients with a total of 1 mM KNO₃. After 346 19 days, seedlings were starved for 2 days by changing the growth medium to a 347 nitrogen-free medium. Nitrate was then resupplied to a final concentration of 0.45 mM 348 (mild low nitrate, LN) and 9.4 mM (high nitrate, HN) for 15 minutes before harvesting 349 350 roots for microsomal protein preparation. All experiments consisted of at least three 351 biological replicates.

For LR growth analysis, WT (Col-0), qsk1, 35S::QSK1-GFP/qsk1 (QSK1-OE/ qsk1), 353 35S::QSK1S621AS626A-GFP/qsk1 (QSK1-AA-OE/qsk1), 35S::QSK1S621DS626D-354 GFP/qsk1 (QSK1-DD-OE/qsk1), and nrt1.1-1 seeds were grown on medium 355 mentioned above with 1% agar and 0.45 mM or 9.4 mM KNO₃ for 10 days in a growth 356 chamber and positioned vertically (16/8 hours light/night). Seedlings were scanned as 357 pictures on the 10th day and LR length was measured using Fiji software.

358 Constructs

For ratiometric bimolecular fluorescence complementation (rBiFC) of Arabidopsis 359 proteins, cDNAs of the following genes were cloned into rBiFC plasmids ⁵⁴: NRT1.1, 360 NRT1.1H356A, NRT1.1T101A, NRT1.1T101D and AHA2 were cloned as fusions with 361 the C-terminal half of YFP, whereas QSK1, QSK1 QSK1S621A/S626A, 362 363 QSK1S621D/S626D were cloned as fusions with the N-terminal half of YFP. All 364 constructs were transformed into Agrobacterium tumefaciens strain GV3101 by electroporation. Positive colonies were confirmed by spectinomycin and rifampicin 365 366 resistance and colony PCR.

For purification of the cytoplasmic domain of QSK1 (QSK1C, amino acid 276–627) were cloned into *Escherichia coli* BL21(DE3) expressing plasmid pETGST1a and fused with His and GST tags, resulting in the plasmid His-GST-SIRK1C and His-GST-QSK1C.

371 **Protein expression and purification**

Plasmids His-GST-SIRK1C and His-GST-QSK1C were transformed into *Escherichia coli* BL21 (DE3). After 5 hours induction by IPTG (isopropyl-_D-thiogalactopyranoside),
cells were harvested and lysed using lysis buffer (20mM Tris-HCl pH 7.4, 1mM EDTA,
200mM NaCl, 1mM PMSF, 1mM DTT), soluble fractions were used over gravity flow
Ni²-NTA Sepharose columns for His-GST-SIRK1C and His-GST-QSK1C protein
purification.

378 **Co-immunoprecipitation assays**

Membrane protein was extracted from infiltrated Tobacco (*Nicotiana Benthamian*) 379 leaves with IP buffer (50 mM Tris-HCI (pH 7.5), 150 mM NaCI, 1 mM EDTA, 1% Triton 380 X-100, 10% glycerol, 1 mM PMSF, 1× protease inhibitor cocktail), 0.45 mM potassium 381 382 nitrate or potassium chloride was then added and incubated with anti-GFP magnetic 383 agarose beads (Chromotek, gta-20) for 3 h at 4 °C. The beads were washed five times with wash buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 1 mM EDTA, 1× protease 384 inhibitor cocktail). The precipitated proteins were eluted with 2×SDS buffer loading 385 386 buffer at 95 °C for 10 min for immunoblot analysis using the indicated antibodies 387 (Abcam).

388 Microsomal membrane preparation, tryptic digestion, and phosphopeptide 389 enrichment

Microsomal membrane (MF) preparation and phosphopeptide enrichment were 390 performed as described in the "ShortPhos" workflow ⁵⁵. A total of 1 g of roots (fresh 391 392 weight) was homogenized in 10 ml extraction buffer (330mM sucrose, 100 mM KCl, 1 mM EDTA, 50 mM Tris-MES, fresh 5 mM DTT, and 1 mM phenylmethylsulfonylfluoride, 393 394 pH 7.5) in the presence of 0.5% v/v proteinase inhibitor mixture (MedChemExpress) and phosphatase inhibitors (25 mM NaF, 1 mM Na₃VO₄, 1 mM benzamidin, 3 µM 395 396 leupeptin) in Dounce homogenizers. The homogenate was centrifuged for 15 minutes at 7500 × g at 4 °C. The pellet was discarded, and the supernatant was centrifuged for 397 398 75 minutes at 48,000 × g at 4 °C. The microsomal pellet was re-suspended in 100 µl 399 UTU (6 M urea, 2 M thiourea, pH 8). Protein concentrations were determined using a 400 Bradford (Sigma–Aldrich) assay with BSA (Bovine serum albumin) as protein standard. 401 Samples were stored at -80°C.

402 150 μg MF were aliquoted for tryptic digestion and phosphopeptides enrichment. MF 403 were predigested for 3 h with endoproteinase Lys-C (0.5 μg/μl; Wako Chemicals) at

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404 room temperature. After a 4-fold dilution with 10 mM Tris-HCI (pH 8), samples were 405 digested with 3 μ l sequencing-grade modified trypsin (0.5 μ g/ μ l; Promega) overnight 406 at 37 °C. After overnight digestion, 10% v/v trifluoroacetic acid (TFA) was added (until 407 the pH was 3 or less) to stop digestion. Digested peptides were dried in a vacuum 408 concentrator.

Dry peptides were dissolved in 200 µl of 1 M glycolic acid in 80% v/v acetonitrile (ACN) 409 410 and 5% v/v trifluoroacetic acid (TFA). Phosphopeptides were enriched over titanium 411 dioxide (TiO₂) (GL Sciences). 1.5 mg TiO₂ beads per sample were washed once with 412 100 µl of 1% v/v ammonia solution and equilibrated three times with 50 µl of 1 M 413 glycolic acid in 80% v/v ACN and 6% v/v TFA. 200 µl digested peptides were mixed 414 with equilibrated TiO₂ for 30 min incubation. Peptides and TiO₂ beads mixture were washed once with 100 µl of 1 M glycolic acid in 80% v/v ACN and 6% v/v TFA, and 415 416 three times with 100 µl of 80% v/v ACN and 1% v/v TFA. Phosphopeptides were eluted 417 from TiO₂ beads three times with 1% v/v ammonia solution. Eluates were immediately acidified with 70 µl of 10% v/v formic acid. Acidified phosphopeptides were desalted 418 419 over a C18 stage tip prior to mass spectrometric analysis.

420 LC-MS/MS of Peptides and Phosphopeptides

421 Enriched phosphopeptides were resuspended in 5 µl resuspension buffer (0.2% v/v TFA, 5% v/v ACN) and analyzed via LC-MS/MS using standard setting as described³³ 422 423 with nanoflow Easy-nLC 1200 (Thermo Scientific) as an HPLC system and an Orbitrap hybrid mass spectrometer (Orbitrap exploris 480, Thermo Scientific) as a mass 424 425 analyzer. Peptides were eluted from a 75 µm homemade 25 cm analytical column on 426 a linear gradient running from 4 to 64% acetonitrile over 130 min and sprayed directly 427 into the mass spectrometer. Peptides were identified via MS/MS based on the 428 information-dependent acquisition of fragmentation spectra of multiple charged 429 peptides. A data-dependent acquisition MS method was used, and for full-scan spectrum and MS/MS scan acquired at resolution of 60,000 and 15,000 at m/z 200. 430

431 **Protein Identification and Ion Intensity Quantitation**

Raw data acquired by the mass spectrometer were processed using MaxQuant ⁵⁶,
version 1.6.5.0 using settings as described for protein identification and label-free
intensity quantitation (LFQ). Spectra were matched against the *Arabidopsis* proteome
UP000006548 (UniProt database, 39,324 entries). Common contaminants (trypsin,

keratin, etc.) were included during database searches. Carbamidomethylation of 436 437 cysteine was set as a fixed modification, and the oxidized methionine (M), acetylation (protein N-term) and phosphorylation (STY) were set as variable modifications. Trypsin 438 439 was specified as the digesting protease, and up to two missed cleavages were allowed. 440 The mass tolerance for the database search was set to 20 ppm for full scans and 0.5 441 Da for ions fragment. The multiplicity was set to 1. For label-free quantitation, retention time matching between runs was chosen within a time window of 0.7 min. False 442 discovery rate cutoffs were set to 0.01 for peptide and protein identification, and to 0.05 443 444 for phosphorylation site assignment. Hits to contaminants (e.g., keratins) and reverse hits identified by MaxQuant were excluded from further analysis. 445

446 **Bioinformatics and statistical Analysis**

Bioinformatics analysis was performed with Perseus software (version 1.6.4.0).⁵⁷ For the analysis of phosphoproteomic results, reported label free intensity (LFQ) values (Phospho (STY)Sites. txt) were used for data analysis. Missing values were imputed from a normal distribution around the detection limit of the mass spectrometer. For each phosphosite, imputed values from five or six biological replicates in different genotypes or treatment were used for student *t* tests (FDR<0.05, S0=0.1).

Annotations were extracted from MapMan⁵⁸, subcellular locations were obtained from SUBA3.⁵⁹ Functional enrichment analysis was done via Fisher's exact test, *p* values were adjusted using Bonferroni correction. *p* value indicates a degree of significance and enrichment factor represent the level of enrichment with respect to the background.

457 Ratiometric bimolecular fluorescence assay

458 Agrobacterium tumefaciens harboring the relevant constructs described above were injected into 5 to 6 weeks old Nicotiana benthamiana leaves for 2 days before 459 460 observation. Fluorescence was observed using a Zeiss LSM800 confocal microscope (20× water immersion objective). In all cases, excitation intensities, filter settings, 461 462 photomultiplier gains and other parameters were standardized. The YFP and RFP 463 fluorochromes were excited with 488 nm and 561 nm, respectively. Emitted light was 464 collected at a range of 500-560 nm for YFP and 575-625 nm for RFP. All images 465 throughout all experiments were collected using the same settings. The collected 466 images were processed and both YFP and RFP intensity was measured using the FIJI software and the YFP/RFP ratio was calculated. RFP is internal control. At least 20 467

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different cells from leaves of the 2 plants were analyzed. Statistical significance was determined using one-way ANOVA test with a Tukey test. To calibrate YFP/RFP ratios, known interaction of CBL9 with CIPK23 was used as positive control, whereas the interaction of CBL9 with CIPK14 was used as negative control.⁶⁰ YFP/RFP ratio greater than 1 indicate interaction of two proteins, while YFP/RFP ratio smaller than 1 suggest weak or no interaction.

474 HPTS assay

HPTS staining was performed as described³⁰ with minor modifications. 10 days old 475 seedlings grown on LN and HN medium were transferred into liquid LN and HN 476 medium supplemented with 1mM HPTS for 30 min. The seedlings were subsequently 477 478 mounted in the same growth medium supplemented with HPTS on a microcopy slide. 479 HPTS stained images of lateral roots elongation zone was performed using an inverted Zeiss LSM800 confocal microscope equipped with a highly sensitive GaAsP detector. 480 Fluorescent signals for the protonated HPTS form (Excitation 405 nm, emission peak 481 482 514 nm), as well as the deprotonated HPTS form (excitation 488 nm, emission peak 514 nm) were detected with a 20× (water immersion) objective. The image analysis 483 was performed using the Fiji macro. The ratiometric value (488/405) correlates with 484 485 the apoplastic pH. The higher value represents higher pH.

486 ¹⁵NO₃ influx experiment

Root NO3-influx was assayed as described by Laugier et al.⁶¹ Five weeks old 487 hydroponically-grown plants were nitrate-starved for 5 days, then subjected to 1 mM 488 NO³ for 0h, 1h or 4h. After washing with 0.1 mM CaSO4, roots were subsequently 489 transferred into a nutrient solution containing 200µM ¹⁵NO₃ (99 atom% excess 15N) 490 for 5 minutes. Roots were harvested and dried at 70°C for 48 h, and samples were 491 492 analyzed for total N and atom% 15N using a continuous flow isotope ratio mass spectrometer coupled with a C/N elemental analyzer (model Euroflash; Eurovector, 493 494 Pavia, Italy). 6-12 replicates were performed in each genotype and condition.

495 An in vitro peptide phosphorylation assay

In vitro peptide phosphorylation assays were performed as described in published
 paper⁴³ with minor modification. In brief, kinase activity assays were performed with 10
 pmol of the peptides GLDIETPSHYTV (covering T947) or EAVNIFPEKGSYR

- 499 (covering S899) as a substrate and His-GST-QSK1-Cterminal Domain (QSK1C) as
 500 kinases in 30 μl kinase reaction buffer (20 mM Tris/HCl pH 7.5, 10 mM MgCl₂, 1 mM
- 501 MnCl₂, 0.1% BSA, 2 mM DTT, 100 µM ATP). After incubation for one hour, the reaction
- 502 mixture was acidified with TFA. Acidified mixture was then desalted over C18 column.
- 503 The phosphorylation of peptides was analyzed vis mass spectrometry and quantified
- 504 based on the intensity of signature fragment ion.

505 Author contributions

- 506 Z.Z and L.K performed most experiments. X.N.W, L.K and W.X.S designed research;
- 507 S.L ran samples in mass spectrometry; X.L and T.W assisted with confocal microscope

508 for rBiFC; Z.L, Z.H, X.Y and S.G for phosphoproteomic sample preparation; H.L, M.G.

- 509 M.H and J.Z for partial data analysis. B.N, A.J and L.L for ¹⁵N uptake assay. X.N.W,
- 510 W.X.S and L.K wrote paper.

511 Data availability

512 The raw MS data from this study was deposited at the ProteomeXchange Consortium 513 via the PRIDE partner repository with the identifier PXD045818 (User name: 514 reviewer_pxd045818@ebi.ac.uk Password: KiHE8I6e).

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528 **Declaration of interests**

529 The authors declare no competing interest.

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Figure legends

Figure 1. The global membrane phosphoproteome of Arabidopsis roots was changed in the *nrt1.1-1* mutant under LN condition

- (A) The work flow of High-throughput phosphoproteomics to identify NRT1.1-mediated
- 771 LN and HN-induced signaling components.
- (B) Principal components analysis (PCA) of WT and *nrt1.1-1* under LN and HNconditions. The first and second components are shown.
- (C) Student *t*-test difference between LN-induced and nitrogen starvation (NS) at membrane phosphoproteome (*x* axis) and the difference between HN-induced and nitrogen starvation (NS) in membrane phosphoproteome (*y* axis) in WT. Significant phosphosites were determined using a permutation-based false discovery rate calculation (FDR \leq 0.05, S0 = 0.1). The phosphosites were colored for significant changed abundance in x axis (orange), y axis (red), and both (purple), respectively.
- (D) Student *t*-test difference between *nrt1.1-1* and WT in LN-induced membrane phosphoproteome (*x* axis) and the difference between *nrt1.1-1* and WT in HN-induced membrane phosphoproteome (*y* axis). Significant phosphosites were determined using a permutation-based false discovery rate calculation (FDR \leq 0.05, S0 = 0.1). The phosphosites were colored for significant changed abundance in x axis (green), y axis (blue) and, both (black), respectively.
- (E) Venn diagram showing overlap for phosphosites with significantly changed
 phosphorylation under LN compared to HN (C) and *nrt1.1-1* compared to WT in LN
 (D).
- 789 (F) Fisher exact test (2% FDR) on the group of 307 phosphosites corresponding to 231
- 790 proteins that are indicated in (E). Enriched Mapman terms and SUBA are displayed.

Figure 2. NRT1.1 represses T947/S899 phosphorylation ratio on AHA2 and proton pump activity in LN

- (A and B) Difference of phosphorylation levels induced by LN and HN at T947 andS899 sites of PM H+-ATPase AHA2 in WT and nrt1.1-1.
- (C) Difference in protein levels of AHA2 induced by LN and HN in WT and nrt1.1-1.
- (D) Box plots show T947/S899 phosphorylation ratio for WT and nrt1.1-1 under LN and
- HN. Center lines of boxes represent medians, vertical line indicate theminimal/maximal value.
- Data are mean + SD, at least 4 biological replicates (A-D), statistical analysis (A-D)
 was performed using Student t-test (*****P*<0.0001; NS, not significant).
- 801 (E and F) Quantification (E) and confocal images (F) of apoplastic pH in the lateral root
- elongation zone (EZ) of WT and nrt1.1-1 grown in LN and HN medium. Ratiometric
 value (488 nm/405 nm) of fluorescent HPST was used to monitor pH in the EZ of lateral
- roots, $n \ge 38$ individual root for each. For F, scale bar, 20 µm.
- 805 (G and H) Quantification (G) and images (H) of cortical cell length of lateral roots
- 806 mature zone of WT and *nrt1.1-1* grown in LN and HN medium, $n \ge 63$ cells for each.
- For H, scale bar, 50 μ m. Statistical analysis (E and G) was performed using one-way ANOVA with a Tukey test (*****P*<0.0001; NS, not significant).
- 809 (I) Quantification of the in vivo interaction of NRT1.1, with AHA2. Around 20 randomly
- selected cells were quantified. The known interaction of CBL9-CIPK23 and the known
- absent interaction of CBL9-CIPK14 were used as positive control and negative control,
- 812 respectively. Scale bar, 20 μm. Different letters represent significant differences at
- p<0.05 according to one-way ANOVA with a Tukey test.

Figure 3. LN promotes NRT1.1-QSK1 interaction and QSK1 phosphorylation

- (A) Ranking of phosphosites significantly upregulated in WT under LN and
 downregulated in *nrt1.1-1* in LN. 10 named receptor-like kinases are highlighted in red.
 (B) Phosphorylation differences of S621/S626 on QSK1 in WT and *nrt1.1-1* induced
 by LN and HN.
- (C) Differences of protein levels in QSK1 induced by LN and HN in WT and *nrt1.1-1*.
- B20 Data are mean + SD, at least 4 biological replicates (B-C), statistical analysis (B-C)
- 821 was performed using Student *t*-test (different letter indicates P<0.05).
- 822 (D) Representative images of WT, *qsk1*, transgenic line overexpressing QSK1, QSK1
- phospho-dead and as well as phosphorylation-mimic version in *qsk1* mutant (QSK1-
- 824 OE/*qsk1*, QSK1-AA-OE/*qsk1* and QSK1-DD-OE/*qsk1*) in LN and HN medium, scale 825 bar 1cm.
- (E) Total lateral root length of WT, *qsk1*, QSK1-OE/*qsk1*, QSK1-AA-OE/*qsk1* and
 QSK1-DD-OE/*qsk1* in LN and HN medium. Bar plots indicate means ± SEM (at least
 17 independent seedlings). Lateral roots were measured after 8 days on LN and HN.
 Statistical analysiswas performed using one-way ANOVA (*****P*<0.0001; NS, not
- 830 significant).

831 Figure 4. QSK1 functions downstream of NRT1.1 to modulate LRs growth in LN

- (A) Quantification of the *in vivo* rBiFC interaction of QSK1, QSK1SASA, and
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 calibration is shown in Figure S3. 18-35 randomly selected cells were quantified.
- (B) Co-IP assays using membrane protein mixture extracted from tobacco leaves
 (*Nicotiana benthamiana*) expressing NRT1.1-eGFP and QSK1-FLAG in the presence
 of LN (0.45 mM KNO₃), HN (9.4 mM KNO₃) or potassium chloride (0.45 mM).
- 838 (C) Representative images of WT, *nrt1.1-1*, *qsk1* and *nrt1.1-1qsk1* in LN and HN 839 medium, scale bar, 1cm. Different letters (A and D) represent significant differences at 840 p<0.05 according to one-way ANOVA with a Tukey test.
- (D) Total lateral root length of WT, *nrt1.1-1*, *qsk1*, and *nrt1.1-1qsk1* in LN and HN
 medium. Bar plots indicate means ± SEM (16-20 independent seedlings). Lateral roots
 were measured after 10 days on LN and HN medium.
- Different letters (A and D) represent significant differences at p<0.05 according to oneway ANOVA with a Tukey test.
- 846 (E) ¹⁵Nitrate uptake activity under LN. Bar plots indicate means ± SEM (at least 10
- 847 biological replicates). Statistical analysis was performed using two-tailed Student *t*-test
- 848 (*P*<0.05; NS, not significant).
- 849

Figure 5. phosphorylated QSK1 forms a strong complex with AHA2 and represses proton pump activity in LN

- (A) Quantification of the *in vivo* rBiFC interaction of QSK1, QSK1SASA and QSK1SDSD with AHA2. rBiFC calibration is shown in Figure S3. 50 randomly selected cells were quantified. Different letters represent significant differences at p<0.05 according to one-way ANOVA with a Tukey test.
- (B) Co-IP assays using membrane protein mixture extracted from tobacco leaves
 (*Nicotiana benthamiana*) expressing QSK1-FLAG, QSK1-AA-FLAG and QSK1-DD FLAG with AHA2-MYC.
- (C and D) Difference of phosphorylation levels of T947 (C) and S899 (D) on AHA2
 induced by LN and HN in WT and *qsk1*.
- (E) Difference of protein levels of AHA2 under LN and HN in WT and *qsk1*.
- (F) Box plots show T947/S899 phosphorylation ratio for WT and *qsk1* under LN. Center
- 863 lines of boxes represent medians, vertical line indicates the minimal/maximal value.
- Bata (C, D and E) are mean+SD. Statistical analysis (C-F) was performed using two tailed Student *t*-test (*****P*<0.0001; NS, not significant).
- (G) *In vitro* peptide phosphorylation assay using GLDIETPSHY**T**V (covering T947) and
- 867 EAVNIFPEKG**S**YR (covering S899) from AHA2 as substrates for the QSK1C terminus.
- 868 nd indicates not detected.
- 869 (H and I) Quantification (H) and confocal images (I) of apoplastic pH in the lateral root
- 870 EZ of WT and *qsk1* grown on LN and HN medium. Ratiometric value (488 nm/405 nm)

- of fluorescent HPST was used to monitor pH in the EZ of lateral roots, n≥39 individual
- 872 root for each. For I, scale bar 20 μ m.
- 873 (J and K) Quantification (J) and images (K) of cortical cell length of the lateral roots
- mature zone of WT and *qsk1* grown on LN and HN medium. $n \ge 61$ cells for each. For

875 K, scale bar 50 μm.

- 876 Statistical analysis (H and J) was performed using one-way ANOVA with a Tukey test
- 877 (*****P*<0.0001; NS, not significant).

878 Figure 6. NRT1.1-QSK1 complex regulates AHA2 activity in LN

- (A) Co-IP assays using membrane protein mixture extracted from tobacco leaves
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 presence of LN (0.45 mM KNO₃), HN (9.4 mM KNO₃) or potassium chloride (0.45 mM).
- (B) Quantification of apoplastic pH in the lateral root EZ of WT, nrt1.1-1, qsk1 and nrt1.1-1qsk1 grown in LN and HN medium. Ratiometric value (488 nm/405 nm) of fluorescent HPST was used to monitor pH in the EZ of lateral roots, n≥16 individual
- root for each.
- (C) Quantification of the cortical cell length of lateral roots mature zone of WT, *nrt1.1- n, gsk1* and *nrt1.1-1gsk1* grown on LN and HN medium, n≥152 cells for each.
- 888 Statistical analysis (B and C) was performed using one-way ANOVA with a Tukey test 889 (*****P*<0.0001; NS, not significant).
- (D) Proposed model for LN signal transduction through NRT1.1-QSK1 to AHA2.
 Perception of LN leads to NRT1.1 interacting with and phosphorylation of QSK1.
 Subsequently QSK1 recruits AHA2 and directly phosphorylates AHA2S899. NRT1.1
 and QSK1 also induces AHA2T947 phosphorylation indirectly through unknown
 proteins (e.g. Kinases). As a result, phosphorylation ratio of T947/S899 and thus H+ATPase activity is affected. In both the *nrt1.1-1* and *gsk1* mutants, phosphorylation
- ratio of T947/S899 is dramatically increased due to a stronger reduction of phosphorylation at S899, resulting in an even higher proton activity, longer cells and longer LRs.
- 899

900 Supplementary Figures and Tables

Figure S1 The global phosphoproteome of *Arabidopsis* roots of WT and *nrt1.1-1* under LN and HN-induced conditions.

- 903 (A) Distribution of the assigned phosphorylated amino acid residues for class I904 phosphosites.
- (B) Venn diagram showing the overlap of the phosphosites identified in this study andpublished in previous nitrate-induced experiments.
- 907 (C) Venn diagram showing the overlap of the phosphosites identified in this study and 908 published in previous nitrate-induced experiments.
- 909 (D) Summary of identified and quantified class I phosphosites (localization probability
- 910 of >0.75) corresponding to number of proteins in this study.

- 911 (E) Ranking of phosphosites significantly upregulated in WT under LN and 912 downregulated in *nrt1.1-1* in LN. 19 named transport proteins are highlighted in blue.
- Figure S2 (A and B) Representative spectra of AHA2 C-terminal peptides
 GLDIETPSHYTV (containing phosphorylated T947 (A) and EAVNIFPEKGSYR
 (containing phosphorylated S899) (B) directly exported from MaxQuant version 1.6.4.0.
 The phosphorylated amino acid is indicated by (ph).
- 917 (C) Representative images for total lateral root length of WT and *nrt1.1-1* in LN and 918 HN, scale bar 1cm.
- 919 (D) Total lateral root length of WT and *nrt1.1-1* in LN and HN. Bar plots indicate means 920 \pm SEM (at least 16 independent seedlings). Lateral roots were measured after 10 days 921 on LN and HN medium. Different letters represent aignificant differences at *B*<0.05
- on LN and HN medium. Different letters represent significant differences at *P*<0.05
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- **Figure S3** (A) Representative images of the in vivo interaction of NRT1.1, with AHA2.
- 924 The known interaction of CBL9-CIPK23 and the known absent interaction of CBL9-
- 925 CIPK14 were used as positive control and negative control, respectively. Cartoons 926 show the respective T-DNA of the rBiFC-2in1-CC vector, harboring different versions
- of CIPK23-nYFP and CIPK14-nYFP with cYFP-CBL9, NRT1.1-cYFP and AHA2-nYFP.
 Scale bar, 20µm.
- (B) Representative spectra of QSK1 C-terminal peptides LIEEVSHSSGSPNPVSD
 (containing phosphorylated S621S626) directly exported from MaxQuant version
 1.6.4.0. The phosphorylated amino acid is indicated by (ph).
- **Figure S4** (A) Representative images of the in vivo interaction of QSK1, QSK1SASA, and QSK1SDSD with NRT1.1, NRT1.1H356A, NRT1.1T01A, and NRT1.1T01D.
- 934 (B) Representative images of the in vivo interaction of QSK1, QSK1SASA and
 935 QSK1SDSD with AHA2 (C).
- 936 Cartoons show the respective T-DNA of the rBiFC-2in1-CC vector, harboring different
- 937 versions of QSK1s-nYFP, NRT1.1s-cYFP and AHA2-cYFP. Scale bar, 20µm.
- 938
- 939 **Supplementary Table 1** Quantitative phospho-profiling in WT and *nrt1.1-1* mutant.
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- 941

Figures:

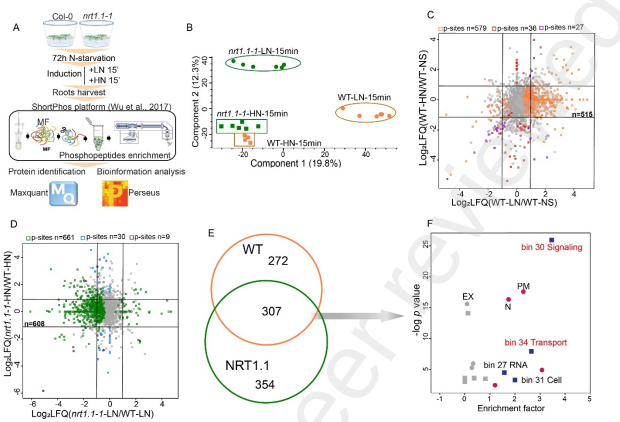


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(D) Student *t*-test difference between *nrt1.1-1* and WT in LN-induced membrane phosphoproteome (*x* axis) and the difference between *nrt1.1-1* and WT in HN-induced membrane phosphoproteome (*y* axis). Significant phosphosites were determined using a permutation-based false discovery rate calculation (FDR ≤ 0.05 , S0 = 0.1). The phosphosites were colored for significant changed abundance in x axis (green), y axis (blue) and, both (black), respectively.

(E) Venn diagram showing overlap for phosphosites with significantly changed phosphorylation under LN compared to HN (C) and *nrt1.1-1* compared to WT in LN (D).

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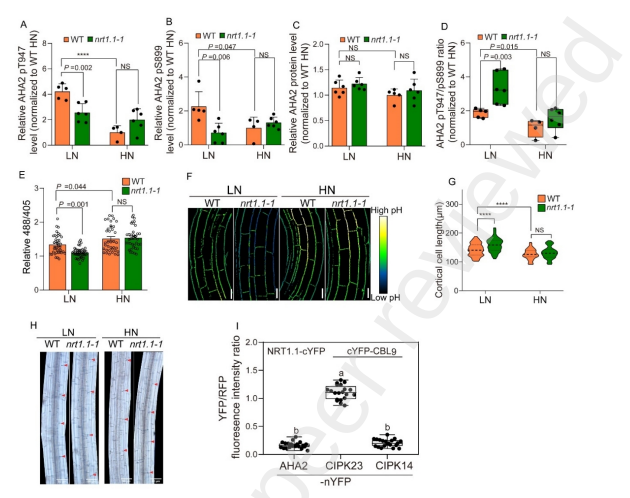


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(C) Difference in protein levels of AHA2 induced by LN and HN in WT and *nrt1.1-1*.

(D) Box plots show T947/S899 phosphorylation ratio for WT and *nrt1.1-1* under LN and HN. Center lines of boxes represent medians, vertical line indicate the minimal/maximal value.

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(G and H) Quantification (G) and images (H) of cortical cell length of lateral roots mature zone of WT and *nrt1.1-1* grown in LN and HN medium, $n \ge 63$ cells for each. For H, scale bar, 50 µm. Statistical analysis (E and G) was performed using one-way ANOVA with a Tukey test (*****P*<0.0001; NS, not significant).

(I) Quantification of the *in vivo* interaction of NRT1.1, with AHA2. Around 20 randomly selected cells were quantified. The known interaction of CBL9-CIPK23 and the known absent interaction of CBL9-CIPK14 were used as positive control and negative control, respectively. Different

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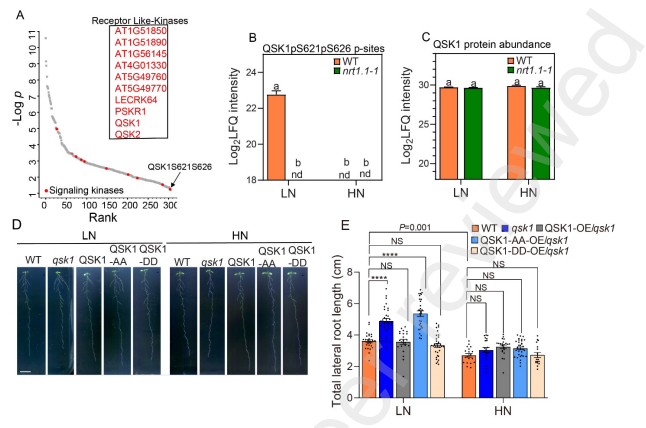


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(A) Ranking of phosphosites significantly upregulated in WT under LN and downregulated in *nrt1.1-1* in LN. 10 named receptor-like kinases are highlighted in red.

(B) Phosphorylation differences of S621/S626 on QSK1 in WT and *nrt1.1-1* induced by LN and HN.

(C) Differences of protein levels in QSK1 induced by LN and HN in WT and nrt1.1-1.

Data are mean + SD, at least 4 biological replicates (B-C), statistical analysis (B-C) was performed using Student *t*-test (different letter indicates P<0.05).

(D) Representative images of WT, *qsk1*, transgenic line overexpressing QSK1, QSK1 phospho-dead and as well as phosphorylation-mimic version in *qsk1* mutant (QSK1-OE/*qsk1*, QSK1-AA-OE/*qsk1* and QSK1-DD-OE/*qsk1*) in LN and HN medium, scale bar 1cm.

(E) Total lateral root length of WT, *qsk1*, QSK1-OE/*qsk1*, QSK1-AA-OE/*qsk1* and QSK1-DD-OE/*qsk1* in LN and HN medium. Bar plots indicate means \pm SEM (at least 17 independent seedlings). Lateral roots were measured after 8 days on LN and HN. Statistical analysiswas performed using one-way ANOVA (*****P*<0.0001; NS, not significant).

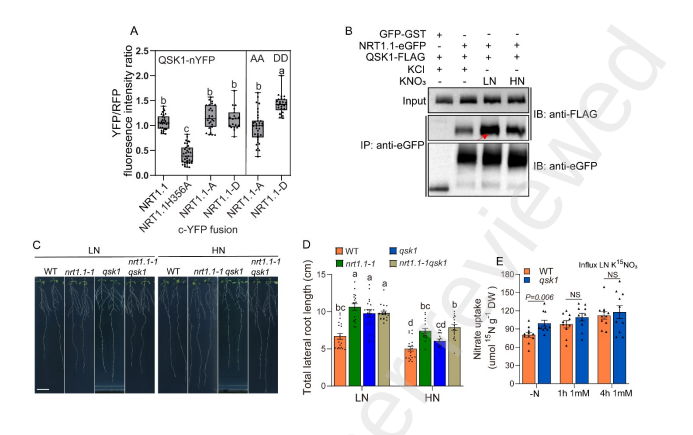


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(C) Representative images of WT, *nrt1.1-1*, *qsk1* and *nrt1.1-1qsk1* in LN and HN medium, scale bar, 1cm. Different letters (A and D) represent significant differences at p<0.05 according to one-way ANOVA with a Tukey test.

(D) Total lateral root length of WT, *nrt1.1-1*, *qsk1*, and *nrt1.1-1qsk1* in LN and HN medium. Bar plots indicate means ± SEM (16-20 independent seedlings). Lateral roots were measured after 10 days on LN and HN medium.

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(E) ¹⁵Nitrate uptake activity under LN. Bar plots indicate means \pm SEM (at least 10 biological replicates). Statistical analysis was performed using two-tailed Student *t*-test (*P*<0.05; NS, not significant).

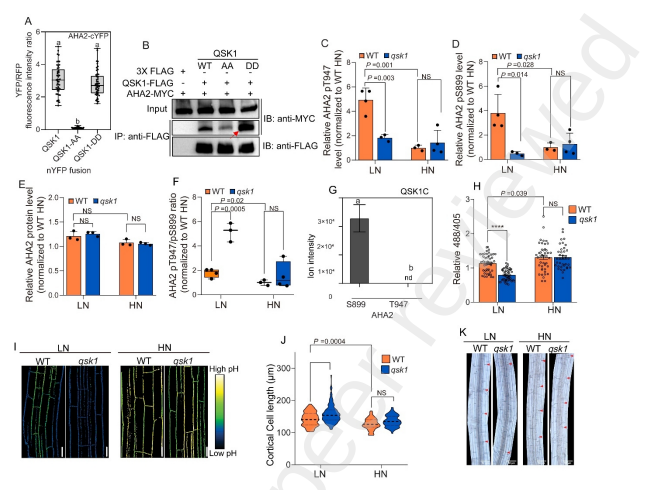


Figure 5. phosphorylated QSK1 forms a strong complex with AHA2 and represses proton pump activity in LN

(A) Quantification of the *in vivo* rBiFC interaction of QSK1, QSK1SASA and QSK1SDSD with AHA2. rBiFC calibration is shown in Figure S3. 50 randomly selected cells were quantified. Different letters represent significant differences at p<0.05 according to one-way ANOVA with a Tukey test.

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(E) Difference of protein levels of AHA2 under LN and HN in WT and qsk1.

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Data (C, D and E) are mean+SD. Statistical analysis (C-F) was performed using two-tailed Student *t*-test (*****P*<0.0001; NS, not significant).

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(J and K) Quantification (J) and images (K) of cortical cell length of the lateral roots mature zone of WT and *qsk1* grown on LN and HN medium. $n \ge 61$ cells for each. For K, scale bar 50 µm.

Statistical analysis (H and J) was performed using one-way ANOVA with a Tukey test (*****P*<0.0001; NS, not significant).

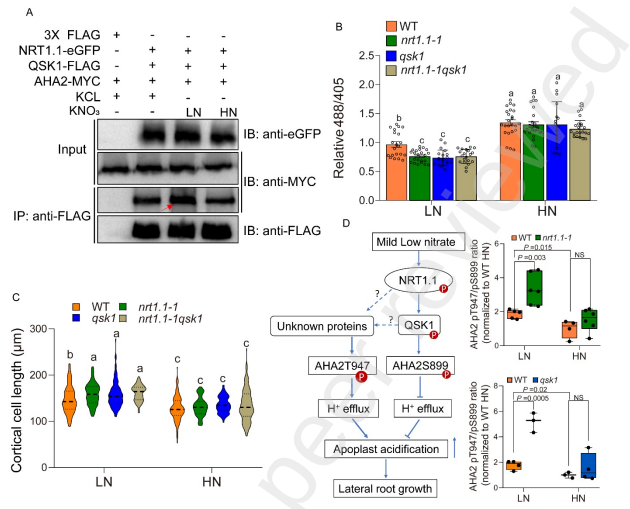


Figure 6. NRT1.1-QSK1 complex regulates AHA2 activity in LN

(A) Co-IP assays using membrane protein mixture extracted from tobacco leaves (*Nicotiana benthamiana*) expressing NRT1.1-eGFP, QSK1-FLAG and AHA2 in the presence of LN (0.45 mM KNO₃), HN (9.4 mM KNO₃) or potassium chloride (0.45 mM).

(B) Quantification of apoplastic pH in the lateral root EZ of WT, *nrt1.1-1*, *qsk1* and *nrt1.1-1qsk1* grown in LN and HN medium. Ratiometric value (488 nm/405 nm) of fluorescent HPST was used to monitor pH in the EZ of lateral roots, $n \ge 16$ individual root for each.

(C) Quantification of the cortical cell length of lateral roots mature zone of WT, *nrt1.1-1*, *qsk1* and *nrt1.1-1qsk1* grown on LN and HN medium, $n \ge 152$ cells for each.

Statistical analysis (B and C) was performed using one-way ANOVA with a Tukey test (*****P*<0.0001; NS, not significant).

(D) Proposed model for LN signal transduction through NRT1.1-QSK1 to AHA2. Perception of LN leads to NRT1.1 interacting with and phosphorylation of QSK1. Subsequently QSK1 recruits AHA2 and directly phosphorylates AHA2 at S899. NRT1.1 and QSK1 also induces AHA2T947 phosphorylation indirectly through unknown proteins (e.g. Kinases). As a result, phosphorylation ratio of T947/S899 and thus H⁺-ATPase activity is affected. In both the *nrt1.1-1* and *qsk1* mutants, phosphorylation ratio of T947/S899, resulting in an even higher proton activity, longer cells and longer LRs.