Plants use nitrate, ammonium, and organic nitrogen in the soil as nitrogen sources. Since the elevated CO₂ environment predicted for the near future will reduce nitrate utilization by C₃ species, ammonium is attracting great interest. However, abundant ammonium nutrition impairs growth, i.e., ammonium toxicity, the primary cause of which remains to be determined. Here, we show that ammonium assimilation by GLUTAMINE SYNTHETASE 2 (GLN2) localized in the plastid rather than ammonium accumulation is a primary cause for toxicity, which challenges the textbook knowledge. With exposure to toxic levels of ammonium, the shoot GLN2 reaction produced an abundance of protons within cells, thereby elevating shoot acidity and stimulating expression of acidic stress-responsive genes. Application of an alkaline ammonia solution to the ammonium medium efficiently alleviated the ammonium toxicity with a concomitant reduction in shoot acidity. Consequently, we conclude that a primary cause of ammonium toxicity is acidic stress.
Nitrate and ammonium are the main sources of nitrogen (N) for most plants. Recent studies suggest that elevated CO₂ reduces nitrate reductase in C₃ species, such as wheat and Arabidopsis, whereas ammonium utilization does not decrease⁵. One estimate predicts that only a 1% drop in nitrogen use efficiency could increase worldwide cultivation costs for crops by about $1 billion annually⁶. Therefore, increasing ammonium use by crops is an important goal for agriculture as CO₂ levels rise in the world; however, millimolar concentrations of ammonium as the sole N source causes growth suppression and chlorosis in plants, compared with nitrate or lower concentrations of ammonium⁴⁻⁵. This phenomenon is widely known as ammonium toxicity, but the primary cause of impaired growth remains to be identified; in the present study, ammonium toxicity is defined as shoot growth suppression with toxic levels of ammonium as the sole N source compared with nitrate.

Plants grown in high ammonium conditions show several distinct characteristics from those grown in nitrate⁴⁻⁵. These toxic symptoms have evoked several hypotheses about the toxic causes, including futile transmembrane ammonium cycling, deficiencies in inorganic cations and organic acids, impaired hormonal homeostasis, disordered pH regulation, and the uncoupling of photophosphorylation; however, some of the symptoms are not directly associated with growth suppression by ammonium toxicity⁶, making it difficult to determine the toxic cause. Several efforts have isolated ammonium-sensitive mutants in Arabidopsis thaliana and determined their causative genes⁴⁻⁵. GMP1 is a causal gene whose deficiency causes stunted growth of primary roots under high ammonium conditions⁷. Given that GMP1 is crucial for synthesizing GDP-mannose as a substrate for N-glycosylation, a lack of N-glycoproteins could be involved in ammonium hypersensitivity. In accordance with this hypothesis, the ammonium-dependent inhibition of primary root growth was shown to be partly attenuated by the lack of a GDP-mannose pyrophosphohydrolase that hydrolyzes GDP-mannose to mannose-1-phosphate and GMP⁸. In another study, a genetic screen focusing on severely chlorotic Arabidopsis leaves identified AMOS1, a gene encoding a plastid metalloprotease, as a factor for improving ammonium tolerance⁹. Transcriptome analysis revealed that an AMOS1-dependent mechanism regulates more than half of the transcriptional changes triggered by toxic levels of ammonium. On the other hand, recent studies found that ammonium toxicity was partly alleviated by deficiencies in EIN2 and EIN3, regulators of ethylene responses, or by the application of ethylene biosynthesis and action inhibitors¹⁰,¹¹. This suggests that ammonium toxicity would be mediated via the ethylene signaling pathway.

The above-described genetic studies have succeeded in determining molecular components closely associated with ammonium toxicity. Nevertheless, the initial event that triggers ammonium toxicity remains to be identified and characterized. To address this question, we screened ammonium-insensitive Arabidopsis lines that were expected to attenuate the toxicity and isolated ammonium-insensitive 2 (ami2). Interestingly, the defect in ami2 was downregulation of the GLUTAMINE SYNTHETASE 2 (GLN2) gene encoding an ammonium assimilatory enzyme localized in the plastid. We identified that in the presence of toxic levels of ammonium, large levels of proton production, due to excessive assimilation of ammonium by GLN2, aggravate the acidic burden and lead to plant toxicity.

**Results**

A genetic screen isolated an ammonium-insensitive mutant. To find ammonium-insensitive lines, a gain-of-function population of the Arabidopsis FOX (full-length cDNA overexpressing) lines¹² was used. An apparent ammonium-insensitive mutant was identified that shows enhanced growth of cotyledons that are greener than wild-type (Col) when grown on 10 mM ammonium in the form of 5 mM (NH₄)₂SO₄ as the sole N source; the mutant was named ami2 (Fig. 1a). The fresh weights of ami2 11-d-old shoots were approximately double those of Col when grown on ammonium (Fig. 1b). A similar growth enhancement in ami2 was observed in the media containing 10 mM NH₄Cl as the sole N source (Supplementary Fig. 1a, b), confirming that ami2 is tolerant to ammonium. In the following experiments, we used 5 mM (NH₄)₂SO₄ as the ammonium source. In contrast, in media containing 10 mM nitrate or 5 mM ammonium plus 5 mM nitrate, the shoot fresh weights of ami2 were less than those of Col. In media containing 10 mM ammonium, the percentage increase in fresh weight of ami2 relative to Col was much larger for shoots (by ca. 110%) than for roots (by ca. 50%) (Fig. 1c). The greater shoot growth in ami2 was reduced in media with lower concentrations of ammonium (0.4, 2 mM), in which the shoot growth of Col was greater than that when grown on media containing 10 mM ammonium (Fig. 1d). Moreover, nitrate addition in the presence of 10 mM ammonium attenuated the deficiency in shoot growth more effectively in Col than in ami2, decreasing the growth difference in a concentration-dependent manner.

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**Fig. 1 Enhanced shoot growth of ami2 in the presence of 10 mM ammonium.** a A representative photograph of shoots from the 11-d-old wild-type (Col) and ami2 grown on media containing 10 mM ammonium. The scale bar represents 5 mm. b Fresh weights (FW) of shoots from 11-d-old Col and ami2 grown on media containing 10 mM ammonium (mean ± SD; n = 10), 5 mM ammonium nitrate (mean ± SD; n = 5), or 10 mM nitrate (mean ± SD; n = 5). c FW of shoots and roots from 11-d-old Col and ami2 grown on media containing 10 mM ammonium (mean ± SD; n = 26). d FW of 11-d-old Col and ami2 shoots grown on media containing 0.4, 2, or 10 mM ammonium (mean ± SD, n = 5). e Violin plots of the differences in the expression of the ammonium stress-responsive genes between the 10-d-old Col and ami2 shoots 3 d after transfer to media containing 10 mM ammonium. The gene list was obtained from Li et al.⁹ (for further details, see Supplementary Data 1). Two independent experiments (Exp1 and Exp2) were performed. Nine shoots from three plates constituted a single biological replicate. An individual violin plot shows the median (black line) and the 25th to 75th percentiles (magenta line). b, d Six shoots from one plate constituted a single biological replicate. b, e *P < 0.05; **P < 0.01; ***P < 0.001 (Welch’s t-test). c, d Different letters denote significant differences at P < 0.05 (Tukey-Kramer’s multiple comparison test).

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2 NATURE COMMUNICATIONS | (2021) 12:4944 | https://doi.org/10.1038/s41467-021-25238-7 | www.nature.com/naturecommunications
manner (Supplementary Fig. 1c, d). A time-course analysis of shoot growth revealed that increased ammonium tolerance of the ami2 plants compared to Col was significant as soon as 5 d after culture initiation (Supplementary Fig. 1e). These results indicate that ammonium tolerance in ami2 shoots is manifested specifically under harsh ammonium conditions.

To corroborate this enhanced ammonium tolerance in ami2, we performed microarray experiments and compared the expression of genes responsive to toxic levels of ammonium9 between the Col and ami2 shoots growing in media containing 10 mM ammonium (Fig. 1e and Supplementary Data 1). The transcript levels of ammonium-inducible genes were significantly reduced in ami2 shoots compared with Col shoots, whereas those of ammonium-repressive genes showed the opposite trend. A reverse transcription-quantitative PCR (RT-qPCR) analysis confirmed that expression of CMCMU, MIOX2, and PDH2, ammonium-inducible genes9,13, was upregulated in Col shoots under 10 mM ammonium compared with 10 mM nitrate, and this ammonium induction was significantly suppressed in ami2 shoots (Supplementary Fig. 2a). We confirmed that the expression of a housekeeping gene TIP41 was little changed, and that of nitrate-inducible marker gene NIR14 was increased under 10 mM nitrate condition (Supplementary Fig. 2b, c). Collectively, these results indicate that ammonium toxicity is attenuated in ami2 shoots.

**GLUTAMINE SYNTHETASE 2** is a causative gene for ammonium toxicity. Next, to identify the causative gene in ami2, we recovered the transgene in a vector using specific primers and sequenced the construct. The gene was identified as GLUTAMINE SYNTHETASE 2 (GLN2), the sole plastidic isofrom in A. thaliana (Fig. 2a and Supplementary Fig. 3a). Because the transgene was driven by the cauliflower mosaic virus 3SS promoter, we expected that overexpression of GLN2 would enhance ammonium tolerance; however, in media containing 10 mM ammonium, the transcript levels of GLN2 in ami2 shoots were downregulated to about 5% of those in Col (Fig. 2b). In contrast, among the major cytosolic GLUTAMINE SYNTHETASE genes (GLN1s), GLN1;1 was upregulated in ami2 shoots, but GLN1;2 and GLN1;3 were slightly downregulated (Supplementary Fig. 3b). Also, an immunoblot analysis using anti-GLN antibodies15 confirmed that the protein levels of GLN2 were remarkably lower in ami2 shoots compared with Col, whereas the signal intensities corresponding to GLN1s were comparable between the mutant and wild type (Fig. 2c). These findings suggested that overexpression of GLN2 cDNA would result in a cosecretion event16, which makes it difficult to test for phenotypic complementation by introducing the GLN2 transgene. To ensure that reduced expression of GLN2 enhances ammonium tolerance, we obtained another GLN2-deficient line having a T-DNA insertion at the 3′-UTR region of GLN2 (SALK_051953, designated as gln2, Supplementary Fig. 3a). As expected, gln2 phenocopied ami2 in terms of the reduced GLN2 and GLN2 expression (Fig. 2b, c), the enhanced ammonium tolerance (Fig. 2d, e and Supplementary Fig. 3c, d), and the lowered induction of ammonium-inducible genes when grown on ammonium (Supplementary Fig. 2a). Furthermore, a similar ammonium tolerance was observed in the GLN2-knockout line (SALK_071292, designated as gln2-2, Supplementary Fig. 3a, e, f) which was recently isolated in another research group17. Thus, we concluded that GLN2 is a causative gene for ammonium toxicity.

**Shoot GLN2 causes while root GLN1;2 attenuates ammonium toxicity.** Previous studies had reported that mutants deficient in AtGLN1;2 were hypersensitive to millimolar concentrations of ammonium18–20. We also confirmed the ammonium hypersensitivity of gln1;2-1 and gln1;2-2 (Supplementary Fig. 4a–d). GLN2 and GLN1;2, therefore, have opposite effects on ammonium toxicity. To discover how GLN2 and GLN1;2 are involved in the toxicity, we evaluated the distribution of GLN2 and GLN1;2 expressions between shoots and roots in Col plants. In the presence of ammonium or nitrate, the steady-state levels of GLN2 expression were consistently higher in the shoots than in the roots, whereas expression of GLN1;2 was much higher in the roots (Fig. 3a, b and Supplementary Fig. 5a, b), implying that both shoot GLN2 and root GLN1;2 could affect ammonium toxicity. To support this hypothesis, we performed a growth analysis using reciprocally grafted plants between Col and ami2 (Fig. 3c and Supplementary Fig. 5c) and between Col and gln1;2-1 (Fig. 3d and Supplementary Fig. 5d). Prior to the analysis, we confirmed that shoot expression of GLN2 was lower in the ami2-derived shoots irrespective of root-genotype (Supplementary Fig. 6) because GLN2 mRNA is suggested to be root-to-shoot mobile21. Only
when the scion was derived from ami2 was shoot growth significantly enhanced in the presence of 10 mM ammonium (Fig. 3c). On the other hand, deficiency in root GLN1;2 content was sufficient to decrease shoot growth in ammonium (Fig. 3d). Further, we observed that in ammonium-grown plants, the total enzymatic activities of GLNs were significantly reduced by ca. 30–40% in 5-d-old shoots of ami2 and gln2 and by ca. 40–60% in 5-d-old roots of gln1;2-1 and gln1;2-2 compared with Col (Supplementary Fig. 7a, b). Additionally, partially compensatory inductions of other GLNs were found in the mutants (Supplementary Fig. 7c, d). Our findings demonstrate that although shoot GLN2 causes ammonium toxicity in the shoot, root GLN1;2 attenuates ammonium toxicity.

Lowered GLN2 activity reduces the conversion of ammonium to amino acids. It is generally held that ammonium per se is a toxic compound22. On the other hand, a deficiency in GLN2 content should lead to ammonium accumulation in the shoot. Our determination of shoot ammonium content revealed that ami2 and gln2 shoots grown on 10 mM ammonium both accumulated more than 100 µmol g⁻¹ fresh weight of ammonium (Fig. 4a), albeit the two mutants accumulated more fresh weight than Col (Supplementary Fig. 8). In Col shoots, the ammonium concentrations were 12.9 ± 0.9 µmol g⁻¹ and 2.2 ± 0.4 µmol g⁻¹ (mean ± SD) under 10 mM ammonium and 10 mM nitrate, respectively (Fig. 4a). This result indicates that ammonium assimilation by GLN2, rather than ammonium accumulation, triggers ammonium toxicity in the shoot.

An ample supply of ammonium increases the concentrations of amino acids compared with nitrate supply alone.6,23. In particular, the molar ratios of Gln to Glu are elevated at higher ammonium levels, suggesting that Gln synthesis by glutamine synthetase (GLN) overflows glutamate synthase (GOGAT) capacity. Our hierarchical cluster analysis of amino acid content in shoots clearly demonstrated that the type of N source, i.e., 10 mM ammonium or nitrate, was the strongest determinant for plant amino acid composition (Fig. 4b and Supplementary Fig. 9a). In this analysis, Col and the GLN2-deficient lines are categorized into separate clusters depending on the N source. The molar ratio of Gln to Glu (Fig. 4c), total amino acid-N content per amino acid (Supplementary Fig. 9b), total amino acid-N content per fresh weight (Supplementary Fig. 9c), and the molar ratios of N to C in total amino acids (Supplementary Fig. 9d) were consistently larger in ammonium-grown shoots than nitrate-grown shoots, and this large ammonium-N input was partly but significantly attenuated by GLN2 deficiency. These findings suggest that the GLN2 reaction leads to excessive incorporation of ammonium-N into amino acids in shoots when toxic levels of ammonium are present. In addition, we observed that total N and protein concentrations were generally larger in ammonium-grown shoots than nitrate-grown shoots (Supplementary Fig. 10a, b). In ammonium-grown shoots, the total N concentrations were significantly reduced by GLN2 deficiency (Supplementary Fig. 10a), and the protein concentrations were marginally decreased (Supplementary Fig. 10b). Collectively, the shoot GLN2 would facilitate the accumulation of assimilated N in the shoot under toxic ammonium condition.

Ammonium assimilation by GLN2 causes acidic stress. The amino acid profiles suggested that metabolic imbalances due to excessive ammonium assimilation by GLN2 could be a cause of ammonium toxicity. This hypothesis is supported by the observation that shoot growth of Col was much lower under 10 mM ammonium than under 2 mM ammonium and the shoot growth enhancement by GLN2 deficiency was determined only under 10 mM ammonium as already explained above. We have previously demonstrated that nitrate addition at adequate concentrations mitigates ammonium toxicity without reducing amino acid accumulation5. Therefore, a phenomenon triggered by some GLN2-mediated process other than amino acid accumulation should be a cause of ammonium toxicity. Notably, the GLN reaction is a proton-producing process24. The stoichiometry of this reaction is two protons per each glutamine incorporated, one proton of which is derived from ATP hydrolysis and the other is from deprotonation of NH₄⁺. Conversely, the subsequent ferredoxin-dependent glutamate synthase (Fd-GOGAT) reaction consumes two protons per one glutamine incorporated. Given that the molar ratio of Gln to Glu was about 10 in the ammonium condition but close to 1 in the nitrate condition (Fig. 4c), proton production in the ammonium condition could proceed beyond its consumption. Strikingly, a previous study found that 43% of ammonium-inducible genes correspond to acidic stress-inducible genes in Arabidopsis roots13,25. We found that ammonium-inducible genes whose expression was reduced by GLN2 deficiency under 10 mM ammonium condition (Supplementary Fig. 2a) are known as acid-inducible genes25. Thus, we hypothesized that excessive ammonium assimilation by GLN2 causes acidic stress to the plants growing in ammonium.

We re-surveyed our microarray data by focusing on previously identified acidic stress-responsive genes25 (Fig. 5a and
larger in ammonium-grown shoots than nitrate-grown shoots, and these larger ratios were significantly reduced by GLN2 deficiency (Supplementary Fig. 11b), implying facilitated proton consumption by GAD1 in a GLN2-dependent manner under ammonium condition. These results support our hypothesis associating ammonium assimilation with acidic stress. Moreover, in Col and ami2 reciprocally grafted plants growing in the ammonium condition, ALMT1 expression was significantly lower in the ami2-derived shoots than the Col-derived shoots (Supplementary Fig. 11c), indicating that shoot GLN2 locally causes acidic stress to the shoot. Furthermore, ALMT1 expression was analyzed using grafted plants between Col and a mutant lacking the STOP1 transcription factor (stop1-KO) that induces ALMT1 to respond to acidic stress26 (Supplementary Fig. 11d). In the stop1-KO-derived shoots, the ammonium-dependent induction of ALMT1 disappeared, confirming the notion that acidic stress occurs in plants growing in ammonium.

It is widely accepted that the reduction from nitrate to ammonium consumes a proton, suggesting that nitrate reduction could attenuate acidic stress caused by excess ammonium and might explain why nitrate addition alleviates ammonium toxicity. To verify this hypothesis, we analyzed shoot expression of ALMT1 using grafted plants between Col and the NITRATE REDUCTASE-null mutant (designated as NR-null)44 (Supplementary Fig. 11e). The addition of 2.5 mM nitrate diminished the ammonium-dependent ALMT1 induction in the Col-derived shoots but not in the NR-null-derived shoots, thereby supporting the above hypothesis.

To obtain direct evidence for ammonium-dependent proton production, we measured the proton concentrations of water extracts from the Col and ami2 shoots incubated in media containing 10 mM ammonium or nitrate with or without MSX (Fig. 5c). The ammonium-treated Col shoots contained the highest concentrations of protons; proton content was significantly decreased by GLN2 deficiency and by MSX treatment to levels comparable to those in nitrate-treated shoots. A similar trend was observed among the Col, ami2, and gln2 shoots grown on ammonium- or nitrate-containing media (Supplementary Fig. 11f).

The presence of ammonium in cultures generally acidifies the external media33. Thus, we quantified the proton influx from the Col and ami2 shoots incubated in media containing 10 mM ammonium or nitrate with or without MSX (Fig. 5d). Incubation of the Col shoots in the presence of ammonium strongly acidified the external media, which was alleviated by GLN2 deficiency and by MSX treatment. A similar tendency was observed by qualitative measurements with a pH indicator of proton effluxes from mesophyll cells where GLN2 is predominantly expressed (Supplementary Fig. 11g). Thus, we conclude that ammonium assimilation by GLN2 without nitrate increases shoot acidity.

Ammonium toxicity is closely associated with acidic stress. If acidic stress rather than ammonium accumulation has a dominant effect on ammonium toxicity, an application of alkaline ammonia should reduce the toxicity. Given that the GLN2 reaction is a primary cause of increased acidic stress, an elevation in medium pH may increase the shoot growth of Col more effectively than that of the GLN2-deficient mutants. As expected, the addition of an ammonia solution to media containing 10 mM ammonium elevated the pH from 5.7 to 6.7 and significantly improved shoot growth with a concomitant decrease in acidity (Fig. 6a, b). Fresh weights of Col shoots grown at pH 6.7 increased by ca. 180% compared with those grown at pH 5.7, whereas fresh weights of ami2 and gln2 shoots only increased by ca. 30% and 60%, respectively (Fig. 6c). In addition, the acid-
Fig. 5 Ammonium assimilation by GLN2 causes acidic stress. a Violin plots of the differences in expression of the acidic stress-responsive genes between the 10-d-old Col and ami2 shoots 3 d after transfer to media containing 10 mM ammonium. The gene list was obtained from Lager et al.25 (For further details, see Supplementary Data 2). Two independent experiments (Exp1 and Exp2) were performed. Nine shoots from three plates constituted a single biological replicate. An individual violin plot shows the median (black line) and the 25th to 75th percentiles (magenta line). b Effects of MSX treatment on the relative transcript level of ALMT1 in the 10-d-old Col and ami2 shoots 3 d after transfer to media containing 10 mM ammonium or 10 mM nitrate. The transcript levels were evaluated both by RT-qPCR (mean ± SD; n = 3) and semi-quantitative RT-PCR with agarose gel electrophoresis. ACTIN2 (ACT2) was the internal standard. Three shoots from one plate constituted a single biological replicate. c Effects of MSX treatment on proton concentrations in water extracts from the 10-d-old Col and ami2 shoots 3 d after transfer to media containing 10 mM ammonium or 10 mM nitrate (mean ± SD; n = 3). Three shoots from one plate constituted a single biological replicate. d Effects of MSX treatment on proton efflux rates from the 10-d-old Col and ami2 shoots 3 d after transfer to media containing 10 mM ammonium or 10 mM nitrate (mean ± SD; n = 3). Three shoots from one plate constituted a single biological replicate. a, b *P < 0.05; **P < 0.01; ***P < 0.001 (Welch’s t-test). c, d Different letters denote significant differences at P < 0.05 (Tukey-Kramer’s multiple comparison test).

Fig. 6 Ammonium toxicity is closely linked with acidic stress. a, b Effects of NH₃ application on shoot FW and proton concentrations in water extracts of 5-d-old Col grown on media containing 10 mM ammonium or 10 mM nitrate (mean ± SD; n = 3). Thirty-seven shoots from one plate constituted a single biological replicate. The pH was adjusted to pH 5.7 with 1 N KOH; subsequently, a commercial 25% (v/v) ammonia solution was added to adjust the pH from 5.7 to 6.7. A representative photograph of 11-d-old shoots grown on 10 mM ammonium (12 plants per plate) is shown. c Effects of higher pH on the FW of shoots from 11-d-old Col, ami2, and gln2 grown on media containing 10 mM ammonium for 11 d (mean ± SD; n = 6). Three shoots from one plant constituted a single biological replicate. The pH was adjusted to pH 5.7 with 1 N KOH; subsequently, 1 N NaOH was used to adjust the pH from 5.7 to 6.7 to maintain the potassium concentration constant among all samples. A representative photograph of 11-d-old shoots is shown. d FW of shoots from 11-d-old Col, stop1-KO mutant (stop1-k), and the stop1 mutant (stop1-m) grown on media containing 10 mM ammonium (mean ± SD; n = 20) or 10 mM nitrate (mean ± SD; n = 5) for 11 d. Six shoots from one plate constituted a single biological replicate. *P < 0.05; **P < 0.01; ***P < 0.001 (Welch’s t-test). e FW of shoots from 17-d-old plants grafted between Col (C) and the NR-null mutant (nr) 7 d after transfer to media containing 10 mM ammonium (NH₄⁺) or 2.5 mM nitrate and 10 mM ammonium (NH₄⁺-NO₃⁻) conditions (mean ± SD; n = 3). One shoot from one plate constituted a single biological replicate. A representative photograph of 17-d-old shoots 7 d after transfer to media is shown. a–e Different letters denote significant differences at P < 0.05 (Tukey-Kramer’s multiple comparison test). The scale bar represents 10 mm.

sensitive STOP1-deficient mutants had slightly but significantly lower shoot growth when grown in 10 mM ammonium (Fig. 6d), although their acid-hypersensitivity has been described only in roots to date26,27. Moreover, the NR-null-derived shoots that lack a proton-consuming nitrate reduction capacity failed to attenuate ammonium toxicity by nitrate addition (Fig. 6e). Collectively, our results lead to the conclusion that acidic stress is one of the primary causes of ammonium toxicity.

GLN2 causes ammonium toxicity independently of NRT1.1. We have already reported that Arabidopsis NITRATE TRANSPORTER 1.1 (NRT1.1), acting as a nitrate transceptor, also aggravates ammonium toxicity28. This result was recently confirmed by another group11. Thus, we investigated whether NRT1.1 and GLN2 increase the sensitivity to ammonium through a common mechanism. A RT-qPCR analysis revealed that deficiency of either NRT1.1 or GLN2 did not downregulate the expression of the other gene (Supplementary Fig. 12a), and that GLN2 expression was almost 3-times higher in nrt1.1 than in Col. Therefore, the enhanced ammonium tolerance of nrt1.1 cannot be explained by reduced GLN2 expression as in gln2. Moreover, a homozygous double mutant of NRT1.1 and GLN2 showed slightly but significantly larger shoot fresh weight, leaf number, shoot diameter, and chlorophyll content compared with any of the single mutants (Supplementary Fig. 12b–d). These findings suggest that NRT1.1 and GLN2 would be implicated in ammonium-sensitivity independently.

Discussion
Although ammonium is believed to be a toxic compound for plant growth, our results demonstrate that ammonium assimilation by shoot GLN2 rather than ammonium accumulation is a major cause of ammonium toxicity. In plants growing in toxic levels of ammonium as a sole N source, assimilation of
ammonium by GLN2 would occur largely due to bypassing nitrate reduction as the rate-limiting step for N assimilation. The resultant increase in the ratio of Gln to Glu content (Fig. 4c) corresponds to the preferential enhancement of the proton-producing GLN reaction over the proton-consuming GOGAT reaction. The metabolic imbalances exerted by the GLN2 reaction and GLN2-related pathways could lead to the production of large amounts of protons in shoot cells that stimulate proton effluxes to the apoplasm; however, the volume of the shoot apoplasm is probably too small to accommodate such a large proton efflux. Thus, in the presence of toxic levels of ammonium, the GLN2 reaction causes acidic stress inside and outside the cells and triggers acidic stress responses that modulate gene expression (Fig. 5 and Supplementary Fig. 11). Given that the wild-type grown at a higher pH phenocopies the ammonium-insensitive lines at lower pH, the acidic stress-sensitive mutants show ammonium-hypersensitivity, and proton-consuming nitrate reduction alleviates ammonium toxicity (Fig. 6), we conclude that acidic stress is one of the primary causes for ammonium toxicity in A. thaliana. In this framework, upregulation of Arabidopsis GLN1;1 and NIA2 genes encoding nitrate reductase (NR)25,26 and activation of spinach NR29 responding to acidic stress are understandable regulatory responses in the context of maintaining cellular pH homeostasis.

The present study does not address how ammonium-dependent acidification triggers growth deficiency at the cellular and subcellular scales. The chloroplastic localization of GLN2 indicates that proton production must occur within chloroplasts in the elevated ammonium condition. A previous study reported abnormal chloroplast membrane structure including swollen compartments at late stages of ammonium toxicity30; however, we did not find any similar structural changes in the shoots of ammonium-grown plants (Supplementary Fig. 13a), where the intermediates of the Calvin-Benson cycle were not depleted compared with nitrate-grown shoots (Supplementary Fig. 13b). These observations do not support a deficiency in chloroplast function as a primary cause of ammonium toxicity. Apoplastic pH in sunflower leaves and cytosolic pH in carrot cell suspensions decrease after application of millimolar levels of ammonium31,32. The ammonium-inducible genes whose expression is down-regulated by GLN2 deficiency, PGIP1 and PGIP2 (Supplementary Fig. 11a), contribute to cell wall stabilization under acidic stress33, implying apoplastic acidification as a target of ammonium toxicity. On the other hand, in the presence of toxic levels of ammonium, the expression of GABA shunt-related genes (Supplementary Fig. 11a), the molar ratio of GABA to Glu (Supplementary Fig. 11b), and oxygen uptake rates33 are increased as biochemical pH-stats26,34 that may represent an intracellular acidic burden. Given that changes in pH environments influence a wide spectrum of physiological processes, elucidating the relationship between ammonium-dependent acidification and growth deficiency awaits future study.

At the whole-plant scale, our grafting work demonstrated that root GLN1;2 activity attenuates ammonium toxicity in the shoots, whilst shoot GLN2 activity causes the condition (Fig. 3). Considering that GLN1;2 is the ammonium-inducible low-affinity enzyme expressed in the epidermis and cortex of roots, and its deficiency elevates ammonium levels in xylem sap when ammonium is supplied20, root GLN1;2 could act as a barrier to prevent the shoot-to-root transport of ammonium, thus avoiding ammonium assimilation by shoot GLN2. In oilseed rape plants, replacing 3 mM nitrate in a nutrient solution with 10 mM ammonium increased the ammonium levels in xylem sap linearly with time, attaining concentrations >5 mM35, which could indicate breaking through the barrier. On the other hand, we observed that shoot expression of GLN1;2 and GLN1;3 encoding low-affinity GLN1 isoforms was significantly larger under 10 mM ammonium than 10 mM nitrate (Supplementary Fig. 3b). Moreover, a larger protein signal corresponding to shoot GLN1s was detected when plants received ammonium rather than nitrate nutrition (Fig. 2c and Supplementary Fig. 14a, b), implying the above-mentioned barrier function of GLN1;2 and GLN1;3 in the shoot. Whilst a large activity of shoot GLN1s could reduce ammonium assimilation by shoot GLN2, shoot GLN1s reaction per se may produce an excessive proton, thereby causing ammonium toxicity. Thus, we compared the proton concentrations of water extracts from the Col, ami2, and gln1;2 gln1;3 shoots incubated in media containing 10 mM ammonium (Supplementary Fig. 14c). Interestingly, we found that double knockout of GLN1;2 and GLN1;3 did not significantly reduce the proton concentrations, whereas it decreased shoot GS activity by ca. 35%, being comparable to GLN2 deficiency (Supplementary Fig. 14c, d). This suggests that shoot GLN1;2 and GLN1;3 could assimilate excessive ammonium without causing proton accumulation in the shoot, supporting their barrier function. It is a challenging issue to address how GLN1s and GLN2 have different effects on pH regulation in the future.

It is widely believed that the plastidic isofrom of GS (GS2) plays a central role in re-assimilating photorespiratory ammonium. The mutants lacking GS2 in barley cannot survive under photorespiratory air conditions36. In Lotus japonicus, the leaves of GS2-deficient mutants show severe necrotic phenotype when transferred from high to low CO2 condition37. On the other hand, a recent study reported that Arabidopsis GS2-deficient mutants can complete their life cycle under air conditions, albeit smaller and slightly chlorotic than the wild-type17. Even under conditions that lower photorespiration, the GLN2-deficient mutants show growth impairment, suggesting an important role of GLN2 in primary nitrogen assimilation. Also, we did not find a strong suppression of vegetative growth of ami2 and gln2 plants grown in the soil under photorespiratory conditions compared with the wild-type (Supplementary Fig. 15a), although their seedling growth during 1 week after imbibition in the half-strength Murashige and Skoog media was significantly retarded (Supplementary Fig. 15b, c). Moreover, ami2 and gln2 plants propagated the seeds whose total N and C concentrations and germination rates were comparable to those of the wild-type (Supplementary Fig. 15d–f). Thus, in A. thaliana, GLN2, GLN1s, and GDHs proteins could redundantly contribute to the re-assimilation of photorespiratory ammonia. This may be why no mutant lacking GLN2 has been isolated as a photorespiratory mutant from Arabidopsis plants. On the other hand, the present study did not clarify how improved ammonium insensitivity in Arabidopsis GLN2-deficient mutants is related to photorespiration. To elucidate this, it awaits many efforts in comparison between photorespiratory and non-photorespiratory conditions.

The present study demonstrated that GLN2 and NRT1.1 reduce ammonium tolerance via separate mechanisms when plants experience high ammonium conditions (Supplementary Fig. 12). By contrast, these genes are nitrate-inducible genes that are crucial for plant adaptation to nitrate-dominant environments38,39. This observation suggests that the adaptive traits to nitrate and ammonium could be exclusive, and therefore, breeding ammonium-tolerant crops might sacrifice their adaptability to nitrate. Arabidopsis growth is enhanced by elevating CO2 more remarkably with ammonium as the sole N source than with nitrate, although, even under elevated CO2, the absolute value of plant biomass is still greater when receiving nitrate4. Overcoming ammonium toxicity could lead to elevated CO2-adapted crops in terms of their mode of N utilization.
Methods

Plant materials and growth conditions. A. thaliana, accession Columbia (Col) was the wild-type line used in this study. The T-DNA insertion mutants gln2 (SALK_051953), gln2-2 (SALK_071292)17, gln1-1-2 (SALK_145235)19, gln1-2-2 (SALK_102291)18, stop1-KO (SALK_114108)16, and nrt1.1 (SALK_097431)20 were purchased from the European Arabidopsis Stock Centre. The Arabidopsis FOX lines and the ethyl methylsulfonate-mutagenized stop1 mutant (pss0087)16 were obtained from the Riken BioResource Research Center. Seeds of the NR-null line stop1-KO were obtained from Dr. Nigel M. Crawford (University of California, San Diego) and Dr. Soichi Kojima (Tohoku University, Sendai), respectively.

Seeds were surface-sterilized and sown in plastic Petri dishes (diameter 90 mm, depth 20 mm, Iwaki, Tokyo, Japan) containing about 30-ml N-modified Murashige and Skoog medium, supplemented with 10 mM MES, 2% (w/v) sucrose, and 0.25% (w/v) gellan gum (pH 5.7). The N sources were 10 mM KNO3 (10 mM nitrate condition), 5 mM NH4NO3 with 10 mM KCl (5 mM ammonium nitrate condition), or 5 mM (NH4)2SO4 with 10 mM KCl (5 mM ammonium condition). The seeds were placed in the dark at 4 °C for 3 d. Plants were grown in a horizontal position under a photosynthetic photon flux density of 100−130 μmol m−2 s−1 (16 h light/8 h dark cycle) at 23 °C. For the transfer experiments, surface-sterilized seeds were sown in plastic Petri dishes (length 140 mm, width; 100 mm; depth; 20 mm; Iwaki Chemical Co. Ltd., Taito-ku, Tokyo, Japan) containing 50-ml half-strength N-modified Murashige and Skoog medium containing 2.5 mM ammonium as the sole N source, supplemented with 4.7 mM MES, 1% (w/v) sucrose, and 0.4% (w/v) gellan gum (pH 6.7)40. This culture condition allowed almost uniform growth of different lines, alleviating ammonium toxicity. The seeds were placed in the dark at 4 °C for 3 d. The plants were grown in a vertical position for 7 d under a photosynthetic photon flux density of 100−130 μmol m−2 s−1 (16 h light/8 h dark cycle) at 23 °C. The plants were transferred with sterilized tweezers to varying N conditions and grown in a horizontal position for further experiments. Other details of plantlet cultivation are presented in the Results section and the figure legends.

Isolation of ammonium-insensitive lines. Forty-five seedlings per plate of Arabidopsis FOX lines17 were grown on the above-mentioned N-modified Murashige and Skoog media containing 5 mM (NH4)2SO4 with 10 mM KCl as the sole N source for 11 d in a horizontal position. The ammonium-insensitive candidate lines were visually selected based on shoot size and color of the cotyledons or true leaves. Then, the next generation of each candidate line was obtained to refresh the seeds. In the first screen for an ammonium-insensitive line, the 10 mM ammonium condition was placed in the dark at 4 °C for 3 d. The plants were grown in a horizontal position for 7 d. Plants grown in a 1:1,000 dilution of the polyphosphates antibiotics raised maize cytosolic glutamine synthetase15 that was obtained from Dr. Hitoshi Sakakibara (Nagoya University, Nagoya, Aichi, Japan). After rinsing with a buffer containing 0.1% Tween-20, 50 mM Tris–HCl, and 150 mM NaCl (pH 7.6), the blocked membrane was then incubated with a 1:1,000,000 dilution of the polyphosphates antibodies raised maize cytosolic glutamine synthetase15 that was obtained from Dr. Hitoshi Sakakibara (Nagoya University, Nagoya, Aichi, Japan). After rinsing with a buffer containing 0.1% Tween-20, 50 mM Tris–HCl, and 150 mM NaCl (pH 7.6), the antigen–antibody complex was detected using a 1:100,000 dilution of horseradish peroxidase-conjugated goat anti-rabbit IgG (NA935, GE Healthcare) and visualized by chemiluminescent detection (ECL Prime, GE Healthcare) using ImageQuant LAS 3000 mini (Fujifilm, Tokyo, Japan). The signal intensities for each band corresponding to GLN1s and GLN2 isoproteins were quantified using Image J software, version 10.2. After detection, the membrane was stained with the GelCode Blue Stain Reagent (ThermoFisher Scientific).

Microarray analysis. Total RNA was extracted as described above. RNA quality was assessed using an Agilent 2100 bioanalyzer (Agilent Technologies). RNA amplification, labeling, hybridization, and scanning with the 3′ IVT Express Kit (Affymetrix) and the GeneChip Arabidopsis Genome ATH1 Array (Affymetrix) were conducted according to the manufacturer’s instructions. The data set from the microarray chips was normalized by the Microarray Suite 5.0 (MASS) method (Acturus). When the signal of a transcript was “Present,” “Marginal,” or “Absent,” the sequence was omitted from the subsequent quantitative analysis. The raw data are shown in Supplementary Data 4. Changes in respective gene expression levels between Col and ami2 were represented as logarithms to base 2 of the ratios in signal intensities.

Extraction of total RNA. Shoots and roots were harvested, immediately frozen with liquid N2, and stored at −80 °C until use. Frozen samples were ground with a Multi-Beads Shocker (Yasui Kikai Corp., Osaka Prefecture, Osaka, Japan) using zirconia beads (diameter, 5 mm). Total RNA was extracted using the RNaseasy Plant Mini Kit (Qiagen) and on-column DNase digestion according to the manufacturer’s instructions.

Reverse transcription and quantitative (real-time) PCR. Reverse transcription was performed using a ReverTra Ace qPCR RT Master Mix with gDNA Remover (Toyobo Co., Ltd., Tokyo, Japan) according to the manufacturer’s instructions. The synthesized cDNA was diluted 10-fold with distilled water and used in quantitative PCR (qPCR). Transcript levels were measured using a StepOnePlus Real-Time PCR System (ThermoFisher Scientific). The obtained cDNA (2 μL) was amplified in the presence of 10-μL KAPA SYBR FAST qPCR Kit (Kippon Genetics Co., Ltd., Tokyo, Japan), 0.4-μL specific primers (0.2 μM final concentration), and 7.2-μL sterile water. Transcript levels were quantified using a relative standard curve with ACTIN2 as the internal standard. A dilution series of total cDNAs were used as templates to generate the standard curves. The primer sequences used in the experiments are shown in Supplementary Data 3.

Immunoblot analysis of GLN isoproteins. Immunoblot analysis of GLN proteins was performed based on a previously described method48. Shoots were harvested, frozen with liquid N2, and stored at −80 °C until use. Frozen samples were ground with a Multi-Beads Shocker (Yasui Kikai Corp.) using zirconia beads (diameter, 5 mm). Tissue extracts were extracted with 2% (w/v) trichloroacetic acid in 1 N HCl. Ferric chloride (ferric chloride (I 1 N HCl)). The extracts were incubated at 95 °C for 5 min followed by cooling on ice and centrifugation at 20,400 × g at room temperature (20−25 °C) for 10 min. The supernatant (equivalent to 1 mg fresh sample) was subjected to SDS-PAGE in a 12% (w/v) TEB (TG) FastCast Acrylamide Kit, Bio-Rad Laboratories, Hercules, CA, USA) and transferred to a PVDF membrane (Trans-Blot Turbo Mini PVDF Transfer Packs, Bio-Rad Laboratories, Hercules, CA, USA) using HIGH W2 with a semi-dry blotting system (Trans-Blot Turbo Transfer System, Bio-Rad Laboratories). The membrane was incubated overnight in blocking buffer containing 5% (w/v) ECL Prime Blocking Agent (GE Healthcare, Little Chalfont, UK), 0.1% (v/v) Tween-20, 50 mM Tris–HCl, and 150 mM NaCl (pH 7.6). The blocked membrane was then incubated with a 1:1,000,000 dilution of the polyclonal antibodies raised against maize cytosolic glutamine synthetase15 that was obtained from Dr. Hitoshi Sakakibara (Nagoya University, Nagoya, Aichi, Japan). After rinsing with a buffer containing 0.1% Tween-20, 50 mM Tris–HCl, and 150 mM NaCl (pH 7.6), the antigen–antibody complex was detected using a 1:100,000 dilution of horseradish peroxidase-conjugated goat anti-rabbit IgG (NA935, GE Healthcare) and visualized by chemiluminescent detection (ECL Prime, GE Healthcare) using ImageQuant LAS 3000 mini (Fujifilm, Tokyo, Japan). The signal intensities for each band corresponding to GLN1s and GLN2 isoproteins were quantified using Image J software, version 10.2. After detection, the membrane was stained with the GelCode Blue Stain Reagent (ThermoFisher Scientific).

Determination of glutamine synthetase activity. GLN activity was determined as the ADP-dependent conversion rate of L-glutamine to γ-glutamylhydroxamate48. Shoots and roots were harvested, immediately frozen with liquid N2, and stored at −80 °C until use. Frozen samples were ground with a Multi-Beads Shocker (Yasui Kikai Corp., Osaka Prefecture, Osaka, Japan) using zirconia beads (diameter, 5 mm). Metabolites were extracted with 50% (v/v) methanol containing 50 mM sodium acetate (pH 5.5) as the extraction standard. After the extraction, the solution was centrifuged (21,500 × g, 5 min, 4 °C), the supernatant was transferred to a 3.5-kDa cut-off filter (UFC300324, Millipore, Billerica, MA, USA) and centrifuged again (13,700 × g, 30 min, 4 °C). The filtered extract was used for the electrophoresis–mass spectrometry or cation spectrometry (CE-MS) analysis.

Measurements of metabolites were performed using an Agilent 6400 series triple quadrupole CE-MS system (CE; 7100 Capillary Electrophoresis, MS; 6420 Triple Quad LC/MS, Agilent Technologies, Santa Clara, CA, USA) in a multi reaction monitoring (MRM) mode14,15. A DB-Wax capillary (polystyrene glycol–coated) column (30 μm i.d., Agilent Technologies Inc., Santa Clara, CA, USA) was used for the separation of samples. The carrier buffer (pH 8.5) as the running buffer was used for measuring anionic compounds (e.g., organic acids and phosphorylated compounds), and an uncoated fused silica
Determination of ammonium. Ammonium was extracted and its concentration was determined with slight modifications to the method reported by Hachiya et al.42. Shoots were harvested, immediately frozen with liquid N<sub>2</sub> and stored at −80 °C until use. Frozen samples were ground with a Multi-Beads Shocker (Yasui Kikai Corp.) using zirconia beads (diameter, 5 mm). One mL of 0.1 N HCl and 500 µL of chloroform were added to the frozen powder, followed by vortexing for 15 min. The mixture was centrifuged at 12,000 × g at 8 °C for 10 min. The aqueous phase was transferred to a microtube containing 50 mg of acid-washed activated charcoal (No. 035-18081; Wako, Osaka, Japan). The mixture was vortexed and centrifuged at 20,400 × g at 8 °C for 10 min. The ammonium content of the supernatant was spectrophotometrically determined using an Ammonia Test Kit (No. 277-14401, Wako) according to the manufacturer’s instructions.

Determination of H<sup>+</sup> concentration in a water extract of shoots. For Fig. 5c and Supplementary Fig. 14c, plants grown on media containing 10 mM ammonium or 10 mM nitrate were submerged in 5 mL of the corresponding liquid medium without MES buffer in the presence or absence of 1 mM methionine sulfoximine (MSX). The plants were returned to their original position on each medium and incubated for 5 h in the light. Shoots were harvested, immediately frozen with liquid N<sub>2</sub> and stored at −80 °C until use. For Fig. 6a and Supplementary Fig. 11f, the shoots grown on media containing 10 mM ammonium or 10 mM nitrate were harvested without prior incubation in liquid media, immediately frozen with liquid N<sub>2</sub> and stored at −80 °C until use. Ten volumes of H<sub>2</sub>O for Fig. 5c and Supplementary Figs. 11f, 14c and 40 volumes of H<sub>2</sub>O for Fig. 6a were added to the frozen powder followed by centrifugation at 10,000 × g at room temperature for 10 min. The pH of the supernatant was measured with a portable pH meter (B212, Horiba, Ltd.). The values were converted to proton concentrations.

Determination of H<sup>+</sup> efflux from shoots. Excised shoots from the plants grown on media containing 10 mM ammonium or 10 mM nitrate were submerged in 100 volumes of the corresponding liquid medium without MES buffer in the presence or absence of 1 mM methionine sulfoximine (MSX). The submerged shoots were incubated in liquid media for 5 h in the light. The pH changes in the incubation media were measured with a portable pH meter (B212, Horiba, Ltd.). The changes in pH were converted to changes in proton concentrations.

Determination of H<sup>+</sup> efflux from mesophyll protoplasts. Mesophyll protoplasts were prepared with slight modifications to the method reported by Endo et al.35. In one dish, 20 plants were grown for 7 d on media containing 2.5 mM ammonium as the sole N source at pH 6.720 in a vertical position. Ten plants were transferred to fresh medium with the same composition and grown for another 7 d. Two plants were further transferred to a medium containing 10 mM ammonium and grown for 3 d in a horizontal position. The adaxial epidermal surface was affixed to plastic tape, whereas the abaxial epidermal surface was attached to a strip of 15-mm-wide Scotch transparent tape (3 M, St. Paul, MN, USA). The Scotch tape was then carefully pulled away from the plastic tape, peeling away the abaxial epidermal surface cell layer. The plastic tape samples with adhering peeled leaves were transferred to 1.5 mL microtubes containing 1 mL of an enzyme solution [0.75% (w/v) cellulase “Onozuka” R10 (Yakult, Tokyo, Japan), 0.25% (w/v) cymomezyme “Onozuka” R10 (Yakult), 0.4 M mannitol, 8 mM CaCl<sub>2</sub>, and 5 mM MES, pH 5.6]. The tubes were slowly rotated for 20 min at room temperature. The liberated protoplasts were filtered through a 30-µm nylon filter (NY30-HD, SEMITEC, Osaka, Japan), and collected by centrifugation at 100 × g at 4 °C for 5 min. The protoplast suspensions of leaf samples from 16 plants were pooled. The supernatant was removed, and the protoplasts were resuspended in 1 mL of a wash solution [0.4 M mannitol, 8 mM CaCl<sub>2</sub>, pH 6.5] followed by centrifugation at 100 × g at 4 °C for 5 min. This washing step was repeated once more, after which the protoplasts were suspended in 100 µL of the wash solution. Ten µL of protoplast suspension was mixed with 90 µL of an assay solution [0.4 M mannitol, 8 mM CaCl<sub>2</sub>, 0.02% brom cresol purple (BCP) pH 6.5] supplemented with 5 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and 10 mM KC<sub>2</sub> in the presence or absence of 1 mM MSX, 10 mM KCl or 10 mM KNO<sub>3</sub>. The mixture was incubated at 23 °C in the light until color changes were observed.

Observation of chloroplast ultrastructure in mesophyll cells with TEM. For ultrastructural analyses, true leaves from Col, ami2, and glr5 d after the transfer to media containing 10 mM ammonium or 10 mM nitrate were analyzed by transmission electron microscopy according to a previously described method46 with modification. Leaves were fixed with 4% (w/v) paraformaldehyde and 2% (v/v) glutaraldehyde in 50 mM sodium cacodylate buffer (pH 7.4) overnight at 4 °C. After the samples were post-fixed with 1% (w/v) osmium tetroxide for 3 h and dehydrated in a graded methanol series, the samples were embedded in Epon812 resin. Micrographs were recorded using a JEM-1400 (JEOL) transmission electron microscope.

Statistical and cluster analyses. All statistical analyses were conducted using R software, version 3.6.2. Two-sided tests were used for Welch’s t-test. Adjustments were made for Tukey–Kramer’s multiple comparison test. Hierarchical clustering was performed using Gene Cluster software, version 3.0, with “Correlation (uncentered)” as the similarity metric and “Average linkage” as the clustering method. The results were visualized using Java TreeView software, version 1.1.6r4. Other details of analyses are provided in the “Results” section and in the table and figure legends.

Data availability. The raw microarray data used in this study are available in the ArrayExpress database at EMBL-EBI under accession number E-MTAB-10796. Source data are provided with this paper.
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