

Metabolomic analysis of *Arabidopsis* reveals hemiterpenoid glycosides as products of a nitrate ion-regulated, carbon flux overflow

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An understanding of the balance between carbon and nitrogen assimilation in plants is key to future bioengineering for a range of applications. Metabolomic analysis of the model plant, *Arabidopsis thaliana*, using combined NMR-MS revealed the presence of two hemiterpenoid glycosides that accumulated in leaf tissue, to ~1% dry weight under repeated nitrate-deficient conditions. The formation of these isoprenoids was correlated with leaf nitrate concentrations that could also be assayed in the metabolomic data using a unique flavonoid–nitrate mass spectral adduct. Analysis of leaf and root tissue from plants grown in hydroponics with a variety of root stressors identified the conditions under which the isoprenoid pathway in leaves was diverted to the hemiterpenoids. These compounds were strongly induced by root wounding or oxidative stress and weakly induced by potassium deficiency. Other stresses such as cold, saline, and osmotic stress did not induce the compounds. Replacement of nitrate with ammonia failed to suppress the formation of the hemiterpenoids, indicating that nitrate sensing was a key factor. Feeding of intermediates was used to study aspects of 2-C-methyl-D-erythritol-4-phosphate pathway regulation leading to hemiterpenoid formation. The formation of the hemiterpenoids in leaves was strongly correlated with the induction of the phenylpropanoids scopolin and coniferin in roots of the same plants. These shunts of photosynthetic carbon flow are discussed in terms of overflow mechanisms that have some parallels with isoprene production in tree species.

For centuries mankind has relied on carbon, prehistorically fixed from the atmosphere by plants and algae, as a source of fuel and other materials. This is a dwindling resource and, to develop renewable sources of chemicals, knowledge of the biochemical pathways that link photosynthesis to the accumulation of useful materials will be paramount. Terpenoids (isoprenoids) are examples of carbon-rich products that occur prolifically across the plant kingdom. They are not only end products of chloroplast assimilation of photosynthetic carbon but also, by virtue of the phytol side chains of chlorophyll, key elements of the machinery of photosynthesis. Some terpenoids are ubiquitous and are classed as primary metabolites (sterols, carotenoids, and phytol side chains of chlorophyll); others are hormones (gibberellins, abscisic acid, brassinosteroids, and strigolactones). However, the majority of terpenoids, especially those that can accumulate to high levels in plants, are classed as secondary metabolites, although many have roles in chemical ecology. Manipulation of flux through the primary terpenoid pathways and the accumulation of terpenoids via the secondary pathways both have potential for exploitation.

Arabidopsis, the model plant, was, until recently, considered to be poorly endowed with terpenoids. For example, this species does not appear to contain oxidized sesqui- and diterpenes, compounds that are widespread in plants (1). There are some 30 terpene synthase genes in *Arabidopsis* (2) and recent analysis has revealed the presence of sesquiterpenes in flowers and monoterpenes in roots (1). Although the levels of terpenes are very low, the presence of the genes indicates that *Arabidopsis* remains a suitable model, especially for study of the central pathways of terpene biosynthesis.

Two pathways, producing the C₅ units that form terpenes, coexist in plants; most primary and hormonal terpenoids are

produced by the chloroplast-located 2-C-methyl-D-erythritol-4-phosphate (MEP) pathway (3) (Fig. S1). Notwithstanding the production of sterols and sesquiterpenes through the alternate mevalonate pathway, the MEP pathway is responsible for the majority of carbon flux into isoprenoids. The regulation of flux through the MEP pathway not only is crucial to plant growth and development but also has a role in the ability of the plant to react to changes in photosynthetic rate. Flux through the early part of the pathway varies with photosynthesis, by mechanisms involving gene and enzyme regulation in particular of 1-deoxy-D-xylulose-5-phosphate synthase (DXS) (4) and hydroxymethylbutenyl diphosphate reductase (HDR) (5). It is also known that plants need to balance N assimilation with photosynthesis-driven C metabolism (6). There is increasing evidence that flux through the MEP and phenylpropanoid (PP) pathways can be regulated by components of the nitrate (NO₃⁻) assimilation pathway, particularly via the partitioning of phosphoenolpyruvate between cytosolic processes and chloroplastidic supply of precursors for these pathways (Fig. S1) (7). In this paper we report unique work on the metabolomics of *Arabidopsis* under root nutrient deprivation and other stresses and the induction of the biosynthesis of two hemiterpenoid glycosides at surprisingly high levels.

Results

Novel Metabolite Biomarkers in Leaves of Nutrient-Deprived *Arabidopsis*. Combined [¹H]-NMR and direct infusion electrospray ionization MS (DI-ESI-MS) is a technique that we have developed for rapid metabolite fingerprinting of plant extracts (8, 9). When applied to *Arabidopsis* tissue, the data obtained enable rapid assessment of concentrations and fluctuations of both primary and secondary metabolites. In this work, as a simple first experiment, we grew *Arabidopsis* in a hydroponic system with a full nutrient supply to the roots for 23 d to growth stage 1.12 (10) and then transferred the plants to water. Analysis of extracts of leaf tissue, harvested at 3 and 7 d after nutrient withdrawal, by parallel NMR and DI-ESI-MS revealed striking and repeatable differences in metabolite fingerprints between nutrient-deficient and nutrient-supplied plants (Fig. 1). Metabolite changes in the NMR spectra at 7 d are shown in the heatmap, Fig. 1A. Increases in carbohydrates, flavonoids (kaempferol glycosides), and in particular the amino acid phenylalanine (Phe) and decreases in sinapoyl malate and the amino acids Ala, Thr, Asp, and Glu were evident. However, the most striking feature of the NMR dataset was the presence of new signals for olefinic hydrogens at δ 5.701 (major isomer) and 5.608 and aliphatic methyl groups at δ 1.724 (major isomer), 1.710, and 1.680 in the nutrient-deprived plants. These signals are highlighted in Fig. 1C. These olefinic and methyl NMR signals were structurally linked to each other and also to signals at 4.18, 4.16, and 4.27

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terpenoid, (2*E*)-2-methylbut-2-en-1,4-diol glycoside, the major isomer being the 1-glycoside [1] and the minor isomer being the 4-glycoside [2] as shown in Fig. 2. 2-Methylbut-2-en-1,4-diol [3] was a by-product of the synthesis and was also available by deacetylation of the starting acetate. Small amounts of this compound were also detected in the NMR spectra (methyl group at δ 1.68) of nutrient-deprived *Arabidopsis* extracts (Fig. S3*B*). ^1H and ^{13}C NMR data for all three synthetic products agreed well (Table S1) with previously reported occurrences of these and related compounds from other plant species (14–17).

Nitrate Deprivation Is a Key Inducer of HTGs. To define further the relationship between nutrient deprivation and the HTGs, the hydroponics system was used in experiments where individual nutrients were omitted from the full medium. Using metabolomic screening it was demonstrated that the prime inducer of the HTGs in leaves was root NO_3^- deprivation. When NH_4^+ was substituted for NO_3^- as a nitrogen source, the HTGs were still produced, indicating that NO_3^- sensing was the underlying mechanism behind the synthesis of HTGs (Fig. 3). Alternating periods of NO_3^- deprivation and resupply led to a stepwise accumulation of HTGs under periods of starvation, rising to $\sim 1\%$ dry weight. The HTGs did not appear to be substantially reassimilated on nitrate resupply. Other primary metabolites also changed during NO_3^- resupply. Ala, Thr, Asp, Asn, Gln, and Glu which, were all depleted under NO_3^- deprivation, recovered during the resupply period. Conversely, levels of metabolites that increased during NO_3^- deprivation (malate, Phe, sucrose, glucose, stachyose, and maltose) were seen to drop during resupply, to a level similar to that observed in NO_3^- supplied plants (Fig. S4).

Flavonoid Glycoside–Nitrate Adducts Formed in DI-ESI-MS Are Indicators of Tissue NO_3^- Levels. The m/z 640 ion routinely seen in untreated *Arabidopsis* leaf DI-ESI-MS negative ion spectra was absent in the NO_3^- -deprived plants (Fig. S5). From MS-MS experiments it was determined that this ion was a conjugate of the flavonoid glycoside, KRR, the first product ion being m/z 577 (KRR, M-H). Thereafter, fragmentation was identical to that of KRR (product ions at m/z 431 and 285). The accurate mass of the m/z 640 ion was 640.1541 kDa. This result is consistent (-9.8 ppm accuracy) with an empirical formula of $\text{C}_{27}\text{H}_{30}\text{O}_{17}\text{N}$. The data indicated that m/z 640 arises from [KRR + NO_3^-] and that it is an in-source formed [+62] adduct. The absence of m/z 640 in the extracts from the NO_3^- -deprived tissue was consistent with lack of NO_3^- in the plant. The titer of the m/z 640 ion also serves as a second biomarker for NO_3^- starvation in *Arabidopsis*. Independent measurement of [NO_3^-] in samples of nutrient-deprived [3.6 ± 0.8 mg/g dry weight (d.w.) after 7 d] and nutrient-supplied plants (30–40 mg/g d.w.) (Table 1) provided support for

this hypothesis. To confirm this, we isolated pure KRR from *Arabidopsis* by HPLC and collected DI-ESI-MS data from solutions of this compound spiked with 0.02 mM NaNO_3 (Fig. 4*A*). We also added increasing concentrations of NaNO_3 to solutions of the extracts of leaf tissue from NO_3^- -depleted plants. The effect of [NO_3^-] on the formation of m/z 640 is shown in Fig. 4*B*, demonstrating that the adduct intensity in the spectra can be correlated to [NO_3^-]. A weak negative correlation of intensity of +303 with -640 ($r = -0.465$; $P < 0.0001$) and +287 with -640 ($r = -0.390$; $P = 0.0025$) provided a semiempirical means of relating high HTG to low NO_3^- . There are a number of flavonoid glycosides in *Arabidopsis*. We also detected the unusual [M + NO_3^-] for several of these in negative ion DI-ESI-MS. Thus, we observed [M+62] $^-$ at m/z 656.1489 associated with KRG, [M+62] $^-$ at m/z 672.1413 associated with quercetin-3-glucoside-7-rhamnoside, and [M+62] $^-$ at m/z 802.2092 associated with kaempferol-3-rhamnosylglucoside-7-rhamnoside.

Analysis of the Biomarkers in Nitrate Reductase (NR) Mutants. Mutants with deletions in NR genes can contain higher levels of free NO_3^- and thus served as a means to further validate the relationship between HTGs and NO_3^- starvation. We used the null mutant, *nia2* (10% of wild-type NR) (18) and the double mutant *nia1nia2* (0.5% of wild-type NR) (19) and measured HTG [1] by NMR along with the KRR- NO_3^- adduct (m/z 640) in ESI-MS of leaves of plants grown on full media and then starved of NO_3^- for 4 d. The data (Table 2) showed that the single mutant *nia2* produced less (69%) HTG than wild type when NO_3^- was withdrawn and retained more NO_3^- (higher m/z 640 intensity), whereas the double mutant *nia1nia2* did not produce HTGs at all and maintained high NO_3^- levels, despite the period of starvation.

Analysis of the HTG Biomarker Production in Other Stress Treatments. The Arapronics system and the ready detection of the HTG biomarkers allowed us to survey the effect of a range of stresses (*SI Materials and Methods*). The HTGs were quantified from the leaf NMR spectra. The data are shown in Table 1, along with measurements of [NO_3^-] in the same extracts. Wounding of roots, but not of leaves, induced the HTGs in leaves. Other root stresses such as NaCl treatment, osmotic stress (delivered through treatment with polyethylene glycol), and cold did not induce HTGs. K^+ deprivation induced HTGs, but the magnitude was less than that of NO_3^- starvation or NH_4^+ substitution. However, oxidative stress induced by peroxide treatment of roots induced the HTGs in leaves, to levels similar to those from NO_3^- starvation. In general high HTG was always associated with low foliar NO_3^- (Table 1).

Feeding of MEP Pathway Intermediates. Previously Page et al. (20) using gene silencing in *Nicotiana benthamiana* showed that deletion of HDR activity caused a diversion of HMBPP to the diol [3]. This was confirmed (20) by feeding labeled DXP to excised leaves. Using this experimental setup, we fed pathway intermediates (DXP, HMBPP, and diol [3]) to leaves of *Arabidopsis*, excised from plants that had been subjected to 4 d NO_3^- starvation (via both water only and NO_3^- deplete treatments). The metabolism of exogenous substrates was quantified by NMR after 8.5 and 24 h of incubation. DXP was metabolized efficiently (42% conversion after 24 h) to the diol and the HTGs, when plants had been starved of NO_3^- . (Fig. 5 and Table S2). No DXP or other products [e.g., deoxyxylulose (DX) or HMBPP] were detected. In NO_3^- sufficient leaves, DXP turnover to HTG and diol was observed at a much lower level (8% after 24 h). These results confirmed the biosynthesis of the diol and HTGs from HMBPP and a significant effect of NO_3^- starvation on flux into these compounds. Conversely, exogenous HMBPP was converted to diol [3] and HTGs in all plants regardless of nutrient status. Similarly, exogenous diol [3] was efficiently converted to the HTGs in all conditions (Fig. 5 and Table S2). These data indicate that the metabolism of exogenous HMBPP and diol is not subject to regulation, unlike HMBPP generated within the chloroplast from the DXP feed.

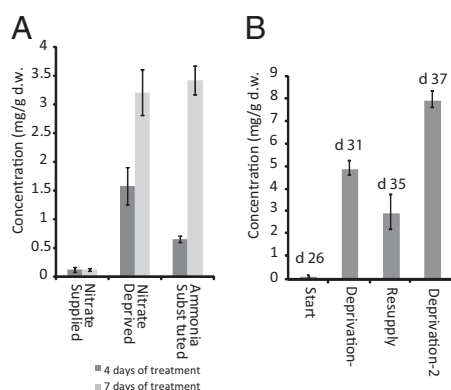


Fig. 3. Effect of nitrogen supply on HTG major isomer production. (A) Effect of NO_3^- deprivation and NH_4^+ ion replacement; (B) NO_3^- resupply experiment. Twenty-six-day-old plants were deprived of NO_3^- for 5 d, resupplied for 4 d, and deprived again for a further 2 d. Data are represented as mean \pm SD.

Table 1. Concentrations of NO₃⁻ and HTGs in leaf tissue under different stress conditions

No days of treatment	Concentration of nitrate, mg/g d.w.		Concentration of HTG [1]*, μg/g d.w.		Concentration of HTG [2]*, μg/g d.w.	
	4	7	4	7	4	7
Full nutrient supplied	49.6 ± 4.8	36.1 ± 1.5	n.d.	n.d.	n.d.	n.d.
Total nutrient deprived	15.8 ± 1.0	3.6 ± 0.8	1155 ± 328	4729 ± 298	120 ± 61	1145 ± 298
Nitrate deprived	7.4 ± 2.4	0.2 ± 0.2	731 ± 240	2395 ± 366	383 ± 211	431 ± 152
Ammonium substituted	15.5 ± 4.4	2.7 ± 0.6	80 ± 41	2480 ± 218	n.d.	486 ± 56
Potassium deprived	32.6 ± 14.7	29.0 ± 1.9	44 ± 40	695 ± 170	n.d.	90 ± 57
Salt stress	36.3 ± 3.8	27.9 ± 2.4	n.d.	n.d.	n.d.	n.d.
Oxidative stress	41.2 ± 6.2	12.9 ± 1.7	170 ± 75	2583 ± 518	21 ± 20	571 ± 137
Osmotic stress	61.6 ± 4.0	72.3 ± 10.4	n.d.	n.d.	n.d.	n.d.
Root damage	23.0 ± 4.9	14.5 ± 2.7	119 ± 76	1920 ± 21	23 ± 21	529 ± 220
Leaf damage	39.1 ± 1.9	47.5 ± 11.5	n.d.	n.d.	n.d.	n.d.
Cold treatment	49.9 ± 11.6	21.8 ± 5.8	0.25 ± 0.25	n.d.	n.d.	n.d.

*Values represent average (± SD) of three biological and two technical replicates ($n = 6$); integration was achieved by summation of the NMR regions: δ 5.725–5.685 [1] and δ 5.635–5.585 [2]. Where HTG concentration was <100 μg/g, values stated are considered to be less accurate and are given as indications of the presence of a small peak. However, in all cases the presence of the HTG was confirmed by the presence of the relevant methyl group singlet in NMR and the 287/303 ions in positive ion ESI. n.d., not detected.

Reprogramming of Root and Leaf Metabolism During a Variety of Root Stresses.

Data for key primary and secondary metabolites, including the HTGs were extracted from the metabolomic screen of roots and leaves under the various stress treatments and were compared by hierarchical cluster analysis as shown in Fig. 6. HTGs were not observed in roots. However, major changes in root metabolism as a consequence of NO₃⁻ deprivation were apparent, including an increase in coniferin and the induction of scopolin, a known coumarin in *Arabidopsis* roots, and also glucoraphanin. These compounds were identified by comparison with authentic samples isolated from *Arabidopsis* and characterized by NMR (Fig. S6). Levels of these metabolites in root tissue across all of the treatments, correlated with levels of HTG in leaf tissue ($R^2 > 0.6$, $P < 0.001$). Associated metabolite changes in roots included small increases in sucrose and glucose and significant decreases in adenosine, malate, Ala, and Thr. Also evident from the metabolite map in Fig. 6 is the specificity of the changes induced by the different stresses. The accumulation of amino acids in leaves during root salt and osmotic stress is particularly striking as is the specificity of the accumulation of coniferin and scopolin in roots under NO₃⁻ deprivation.

Discussion

We have demonstrated how NMR-MS fingerprinting of hydroponically grown *Arabidopsis* plants can be used to discover and monitor metabolite biomarkers, as plants react to changing root environments. ¹H NMR signals of the HTGs and their associated DI-ESI-MS ions were analyzed alongside a negative mode ion (m/z 640), reflecting NO₃⁻ status in the sample, to provide a “three-point check” allowing the tracking of HTG content in relation to NO₃⁻ status across roots and leaves. The production of HTGs in leaves was highly correlated with the production of

scopolin and coniferin in roots of stressed plants, suggesting that a coordinated modulation of MEP and PP pathways in the two tissues was occurring. This phenomenon was observed for plants subjected to root oxidative stress, root wounding, or NO₃⁻ deprivation and to a lesser extent under K⁺ deficiency, but was not evident at all for other stresses, such as root saline or osmotic shock, which gave rise to a different palette of metabolite changes known to arise under these conditions (21). This result indicates that there are specific root–leaf signaling mechanisms that result in HTG and scopolin/coniferin coproduction. As all samples containing the HTG had lower foliar NO₃⁻, the data suggests a role for NO₃⁻ sensing. Root wounding and oxidative stress both resulted in low leaf NO₃⁻ (Table 1), and it is assumed that root damage prevents the uptake of sufficient NO₃⁻ to leaves. The oxidative stress effect may be more complex than simple damage to the root NO₃⁻ assimilation pathway, but the observed outcome is low leaf NO₃⁻ (Table 1) and coproduction of the HTGs and PPs.

Studies using NH₄⁺ as a replacement nitrogen source suggested that the induction of HTGs was related to NO₃⁻ rather than nitrogen per se and repeated deprivation and resupply indicated that *Arabidopsis* is able to switch back and forth from NO₃⁻ poor and NO₃⁻ rich states. Analysis of leaf tissue from the *nia1nia2* double mutant in which NR activity was reduced to 0.5% (19) also pointed to a direct NO₃⁻ relationship with HTG formation. When grown on a NO₃⁻ source, foliar NO₃⁻ accumulated to higher than wild-type levels and thus no HTG was produced under the 4-d starvation period. This result was in contrast to that for *nia2* plants that retain 10–15% NR activity (18) and did not appear to retain enough NO₃⁻ before starvation to prevent HTG buildup on NO₃⁻ removal. These data also suggest that the observed NO₃⁻ effect is not attributable to an NR product such as nitrite.

The discovery of the high levels of HTGs in *Arabidopsis* was unexpected, yet adds another dimension to our knowledge of isoprenoid metabolism in the model plant. The detection of the diol [3] and both isomers of the HTG suggested that the bio-

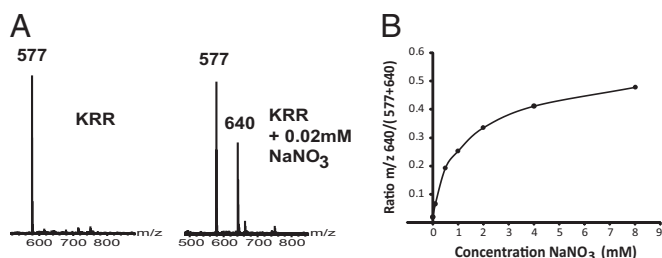


Fig. 4. DI-ESI-MS analysis of KRR and NO₃⁻ ion adducts. (A) DI-ESI-MS (–) ion data showing effect of NO₃⁻ addition to pure KRR; (B) effect of increasing NO₃⁻ addition to an extract from an NO₃⁻-depleted plant.

Table 2. Concentration of HTG [1] and intensity of nitrate status ion (m/z 640) in nitrate reductase mutants subjected to nitrate and starvation

	NO ₃ ⁻ status	HTG [1], mg/g d.w.	m/z 640, intensity × 10 ³
Col-0	+	n.d.	6.81 ± 0.7
Col-0	–	1.54 ± 0.13	0.11 ± 0.16
<i>nia1nia2</i>	–	n.d.	8.13 ± 0.67
<i>nia2</i>	–	1.074 ± 0.055	1.57 ± 0.0103

n.d., not detected. Values represent mean ($n = 3$) ± SD.

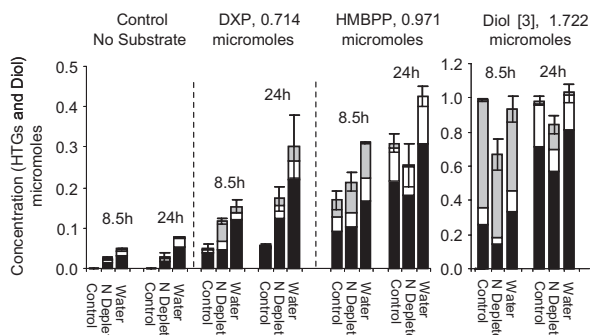


Fig. 5. Metabolism of DXP, HMBPP, and diol [3] by leaves of nutrient-starved plants. Shown is turnover of substrates, quantified by NMR, after 8.5 and 24 h of feeding, to excised leaves, of control (N replete) and nutrient-deprived (4 d) (N deprived or water only) *Arabidopsis*. Stacked plots represent mean concentrations ($n = 2$). Concentrations of individual metabolites are depicted within the bars (solid, HTG major [1]; open, HTG minor [2]; shaded, diol [3]). Numerical data relating to the plots are given in Table S2. Error bars represent the SD of the combined HTGs.

synthesis involves dephosphorylation of HMBPP followed by glycosylation at either end of the diol so formed. The likely biosynthetic route was investigated by feeding intermediates to detached leaves. Conversion of exogenous DXP to the HTGs was efficient in NO_3^- deplete plants (Fig. 5), inferring that the substrate entered the chloroplast where it was converted to HMBPP. Our DXP feeding results align with those of Page et al. (20), who observed conversion of labeled DXP to the diol [3] in intact leaves of *N. benthamiana* plants containing gene-silenced HDR. However, it is generally regarded that phosphorylated substrates such as DXP are not taken up by plant cells and that DX is a more suitable substrate for MEP pathway feeding studies (22). The incorporation of exogenous DXP into HMBPP in intact leaves (ref. 20 and this work) thus requires dephosphorylation of DXP to DX before uptake into the cell and then rephosphorylation to DXP as shown in our working model (Fig. 7). Such a phosphorylation of DX in the cytosol has been demonstrated in *Arabidopsis* (23), as has the existence of chloroplast import of DXP by a translocator (24).

Data from NO_3^- deprivation and NH_4^+ replacement studies, from repeated NO_3^- depletion and resupply experiments, and from the NO_3^- -accumulating *nial1/nia2* double mutant all provide evidence for involvement of (low) NO_3^- sensing systems in the regulation of the MEP pathway in leaves and an associated diversion of lignin precursors to scopolin and coniferin in roots. It is known that plants need to balance C and N metabolism (6, 25) and transcriptomic studies of NO_3^- -limited *Arabidopsis* support this scenario (26). Involvement of light in the regulation of HDR gene expression is known (5) and the potential of a more complex regulation of this activity, which integrates photosynthesis with NO_3^- availability, can now be considered. This system would also be capable of regulating the production of coniferin and scopolin in *Arabidopsis* roots, which is NO_3^- sensitive (this work) and has previously been associated with light treatment (27). Indeed, scopolin production in tobacco roots was associated with low NO_3^- as early as 1970 (28).

The feeding study (Fig. 5 and Table S2) of DXP to intact leaves supports this model in that the pathway diverts HMBPP, generated in situ, to the HTGs via the diol, in NO_3^- -depleted leaves (42% conversion after 24 h), whereas in control leaves DXP turnover to HTGs was some five times lower (8% conversion after 24 h). This result is consistent with a repression of flux through HDR in NO_3^- -depleted plants. The observation of low (but not zero) conversion of DXP to HTGs in excised leaves of control plants can be attributed to an overflow due to an unnaturally large HMBPP pool, arising from the applied DXP.

In our working model (Fig. 7) HMBPP pool size determines the overflow and NO_3^- depletion restricts flow through HDR, causing HMBPP to accumulate and overflow to the shunt products. It is

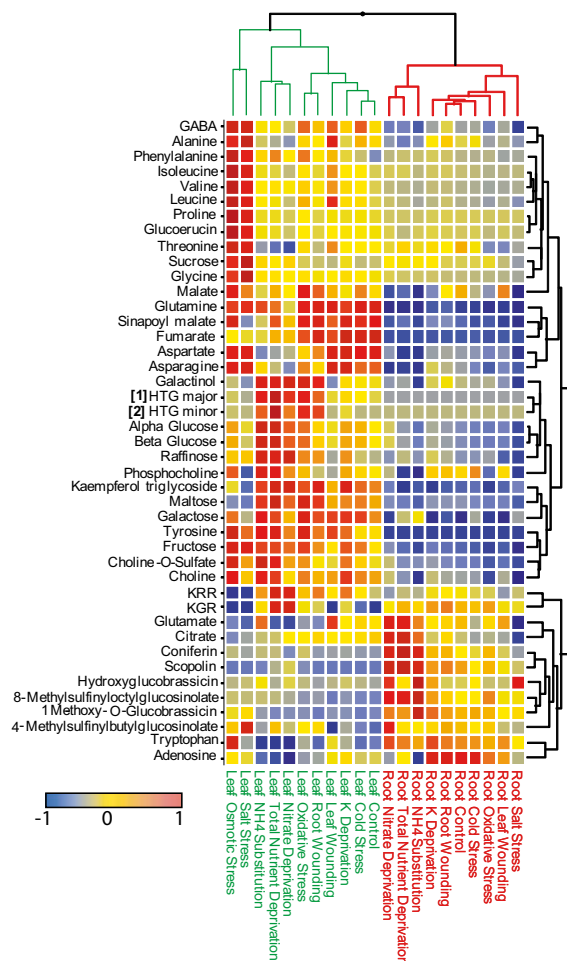


Fig. 6. Relative levels of key metabolites in root and leaf tissue of stressed *Arabidopsis*. Comparisons were generated via HCA using the complete linkage method with similarity based on Euclidian distance. Metabolite levels are based on characteristic chemical shift ranges from NMR or m/z intensities from DI-ESI-MS (–ve ion mode). For each metabolite, data have been normalized to unit variance and mean centered. ^1H NMR shift ranges used are given in SI Materials and Methods.

not clear whether dephosphorylation of excess HMBPP occurs in the chloroplasts and the resultant diol [3] moves to the cytosol where it is glycosylated or whether HMBPP can be exported from the chloroplasts and metabolized in the cytosol. Transporter function analysis (24) indicates that HMBPP, unlike IPP, is not

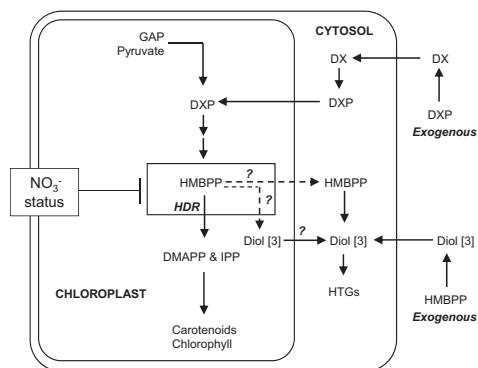


Fig. 7. Proposed routes to HTGs from endogenous and exogenous substrates.

transportable across the chloroplast membrane. Thus, dephosphorylation of HMBPP within the chloroplasts may be more likely. Further work, beyond the scope of this paper, is needed to identify a specific phosphatase activity in chloroplasts. Exogenous HMBPP, applied to leaves via cut petioles, produced similar levels of shunt products in both control and NO_3^- -starved plants. Because exogenous HMBPP is very unlikely to reach chloroplasts, an explanation (Fig. 7) of the feeding results of this compound (Fig. 5) involves nonspecific phosphatase action outside the cell and then glycosylation of the diol [3] in the cytosol. This “unnatural” metabolism of HMBPP was not influenced by NO_3^- status of the leaves (Fig. 5) and thus equally high levels of HTGs were produced in both control and NO_3^- -starved plants.

Although further evidence of involvement of NO_3^- in the regulation of HDR, at the gene or the enzyme level, will be needed, the data presented provide a compelling case for such sensing mechanisms playing a fundamental role in the reprogramming of metabolism that is necessary for plants to balance photosynthetic carbon flux with NO_3^- availability. Given the known role of light in regulation of the MEP pathway at HDR (5), an integrated NO_3^- and light signal transduction chain is a possibility that requires further investigation.

The biosynthetic shunt of HMBPP observed in *Arabidopsis* under stress has parallels with the formation of isoprene from dimethylallyl alcohol diphosphate (DMAPP) under stress in trees (29). Although contrasting hypotheses concerning the role of isoprene as a carbon overflow or as a thermoprotectant have been the subject of much discussion (29, 30), some corollaries with HTG formation in *Arabidopsis* can be made. In *Populus* sp. a link between low NO_3^- and isoprene has been established (7) and other work indicates that DMAPP levels may govern the rate of emission (31, 32).

Conclusion

Our results suggest that an overflow mechanism is present in *Arabidopsis* that deals with undesired buildup of HMBPP via dephosphorylation and glycosylation. The overflow hypothesis

put forward for isoprene emission from trees under stress (7, 30) can thus possibly be extended to other plants and other hemiterpenoids arising from HMBPP. This mechanism, in *Arabidopsis*, does not occur as a generalized stress response but instead appears to be induced by NO_3^- deprivation itself or scenarios that cause a lower NO_3^- status within the leaf, whether directly or indirectly. The high levels of HTGs and the coproduction of coniferin and scopolin under NO_3^- starvation provide ideal biomarkers to unravel the signaling pathways using the genetic resources available for *Arabidopsis*. Such efforts will benefit enormously from the methodology to rapidly quantify NO_3^- , HTGs, phenylpropanoids, and other key metabolites that we have described.

Materials and Methods

Plant Growth and Treatments. *Arabidopsis* (Col-0) and mutant (*nia2* and *nia1/ nia2*) plants were grown using the Arapronics system. For nutrient deprivation, resupply, and stress treatments, plants were moved to appropriate new media as detailed in *SI Materials and Methods*. Shoot and root tissues were harvested separately into liquid N_2 and then lyophilized. Feeding studies were as described in Page et al. (20). Outer leaves excised from treated *Arabidopsis* were incubated with solutions of DXP, HMBPP and diol [3] introduced via the cut petioles.

Metabolomics. Samples for NMR-MS screening were prepared and analyzed according to the protocols described in ref. 9. Full details are in *SI Materials and Methods*.

Synthesis of HTGs [1 and 2] and Diol [3]. Full details of the synthesis and associated structural characterization by NMR are provided in *SI Materials and Methods*.

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