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## Gibberellin and abscisic acid transporters facilitate endodermal suberin formation in *Arabidopsis*

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### Author contributions

J.B. performed the research and wrote the manuscript. N.W. performed the oocyte transporter assays. L.C., L.S.-A., J.-M.D. and P.A. carried out long-distance transport assays. K.K. performed the mathematical modelling. I.T. assisted in cloning overexpression and reporter lines. H.V. and A.A. quantified root GA and ABA content. M.A. helped with genotyping T-DNA mutant lines and profiling suberin patterning. Y.Z. helped with *npr1* mutant identification. D.R. and L.R. assisted in cross sectioning and staining. E.C. quantified hormone content in the phloem sap. E.M. and H.C. performed suberin monomer quantifications. S.L. and R.W. synthesized fluorescently tagged hormones. S.B. carried the qPCR and hormones treated reporter lines. V.N. and C.C. helped with nitrate and hormones quantification in oocyte assays, respectively. C. H. carried out hormone competition transport assays. L.B., P.A., H.H.N.-E. and ES designed and supervised the work and edited the manuscript. All authors discussed the results and commented on the manuscript.

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## Abstract

The plant hormone gibberellin (GA) regulates multiple developmental processes. It accumulates in the root elongating endodermis, but how it moves into this cell file and the significance of this accumulation are unclear. Here, we identified three NPF transporters required for GA and abscisic acid (ABA) translocation. We demonstrate that NPF2.14 is a subcellular GA/ABA transporter, the first to be identified in plants, facilitating GA and ABA accumulation in the root endodermis to regulate suberization. Further, NPF2.12 and NPF2.13, closely related proteins, are plasma membrane-localized GA and ABA importers that facilitate shoot-to-root GA<sub>12</sub> translocation, regulating endodermal hormone accumulation. This work reveals that GA is required for root suberization and that GA and ABA can act non-antagonistically. We demonstrated how the clade of transporters mediates hormone flow with cell-file-specific vacuolar storage at the phloem unloading zone, and slow release of hormone to induce suberin formation in the maturation zone.

## Introduction

The phytohormone gibberellin (GA) is essential for many developmental processes in plants. Among them are seed germination, organ elongation and expansion through cell growth and division, trichome development, the transition from vegetative to reproductive growth, and flower, seed, and fruit development<sup>1</sup>. GAs are produced mainly in the vasculature and move long distances in both acro- and basipetal directions<sup>2-5</sup>. The first reports of GA mobility through the phloem sap appeared over 50 years ago<sup>6-8</sup>. Grafting methods showed that biosynthesis of GA in the shoot can rescue GA biosynthesis mutations in the hypocotyl and root and *vice versa*<sup>6-8</sup>. In tobacco plants, defoliation results in an internode elongation, cambial activity, and fiber differentiation phenotypes that are similar to treatment with paclobutrazol, a GA biosynthesis inhibitor<sup>5</sup>. Another example of an organ dependent on an external source of GA are the petals, which require the anthers as their GA source<sup>9,10</sup>. The biosynthesis of active GA is a complex, multi-step process with diverse intermediates<sup>11</sup>. Regnault et al. conducted a series of grafting experiments using *Arabidopsis thaliana* mutant plants compromised at different stages of GA biosynthesis and identified GA<sub>12</sub> as the major GA form transported over a long distance through the vasculature<sup>12</sup>. GA<sub>12</sub> moves through the xylem in a root-to-shoot manner and in the phloem in a shoot-to-root direction to regulate plant growth<sup>12</sup>. The translocation of GA<sub>12</sub> from the root to the shoot is enhanced under ambient temperatures to induce shoot growth<sup>13</sup>. The transporters that regulate GA long-distance transport in the plant remain unknown.

In recent years, a number of *Arabidopsis* GA plasma membrane importers have been identified, two from the SWEET family and several others from the NITRATE TRANSPORTER1/PEPTIDE TRANSPORTER (NPF) family<sup>14-17</sup>. There are 53 NPFs in *Arabidopsis*, divided into eight subfamilies. These proteins are capable of transporting a large variety of substrates such as nitrate, glucosinolates, abscisic acid (ABA), auxin, and GAs<sup>14,18,19</sup>. Like auxin, GA is subjected to the ion-trap mechanism, limiting its ability to move out of cells. The existence of GA efflux transporters is therefore predicted to allow GA cell-to-cell movement<sup>20</sup>; however, no GA efflux transporters have been discovered<sup>21</sup>.

ABA, which regulates growth and stress responses, has been long thought to act antagonistically to GA in processes such as seed germination, seed maturation, and dormancy and in responses to external cues<sup>22–24</sup>. Both GA and ABA induce developmental responses specifically from the endodermis<sup>25–27</sup>. GA accumulates at high levels in the endodermal cells of the root elongation zone<sup>28</sup>. The process is dependent on the activity of NPF3.1 (NPF3, *AT1G68570*), a member of the NPF family, which has been shown to act as a dual-specificity GA and ABA importer<sup>16,29</sup>. The endodermis, the innermost cortical layer that surrounds the central vasculature, goes through two phases of differentiation: The first step involves polar localized lignin deposition, which results in the formation of the casparian strips, and the second is the deposition of suberin as a lamella below the primary cell wall<sup>30</sup>. The casparian strips restrict apoplastic diffusion of water and nutrients into the vascular tissues, whereas suberin limits backflow of nutrients from the stele<sup>31,32</sup>. The physiological relevance of GA accumulation in the endodermis remains unknown.

With this study, we identified a sub-clade of NPF transporters that orchestrate GA<sub>12</sub> long-distance shoot-to-root translocation and promote bioactive ABA and GA movement from the vasculature to the endodermis, which is required for endodermal suberization, a phenotype rescued by applying either ABA or GA. The importance of transporter distributions was evaluated via a multicellular mathematical model that suggest an intriguing slow-release mechanism whereby GA and ABA delivered to the root in the phloem unloading zone are loaded into pericycle vacuoles and then slowly released to induce suberin formation in the maturation zone. Together, our findings reveal the mechanism that facilitates long-distance shoot-to-root movement of GA<sub>12</sub> and explain how bioactive GA<sub>4</sub> and ABA are transported from the vasculature to the endodermis to mediate endodermal suberization.

## Results

### NPF2.14 is a tonoplast-localized GA and ABA transporter

To identify the missing GA exporters in plants, we tested whether NPF proteins were capable of GA<sub>4</sub> export in *Xenopus laevis* oocyte-based transport assays. All other GA transporters identified to date import GA<sup>14,33</sup>. The transport assays investigate whether NPFs lead to efflux of GA that is loaded into oocytes by three different approaches, namely diffusion, injection and import by a GA importer. Oocytes expressing various NPFs or control oocytes injected with water were exposed to an array of membrane-permeable GAs (GA<sub>4</sub>, GA<sub>7</sub>, GA<sub>9</sub> and GA<sub>12</sub>)<sup>34</sup>. The screen has identified NPF2.14 (*AT1G69860*) as a potential GA exporter. At the end of assay, NPF2.14-expressing oocytes contained significantly reduced GA levels compared to control oocytes (Fig. 1a). This suggests that NPF2.14 facilitates the export activity of GAs out of oocytes. This was confirmed in an injection-based assay in which GA<sub>3</sub>, a non-membrane-permeable form of GA, was injected directly into the oocyte. At the end of the assay, there was a lower GA<sub>3</sub> content in NPF2.14-expressing oocytes compared to control oocytes (Fig. 1b). Co-expression of both *NPF2.14* RNA and *NPF4.1* (*AT3G25260*), which encodes a known GA importer, led to a reduced accumulation of non-membrane permeable GA<sub>3</sub> compared to oocytes expressing only *NPF4.1* (Fig. 1c), further supporting that NPF2.14 exports GA out of *Xenopus laevis* oocytes.

It was previously shown that NPF3.1 transports both GA and ABA in *Xenopus laevis* oocytes<sup>16</sup>. To test whether NPF2.14 also has dual-substrate specificity, control and NPF2.14-expressing oocytes were exposed to ABA. NPF2.14-expressing oocytes accumulated less ABA than did control oocytes (Fig. 1d), indicating that NPF2.14 has dual-substrate transport activity. Several NPF transporters, including NPF6.3 (*AT1G12110*), also transport nitrate<sup>18</sup>, but NPF2.14 displayed no nitrate transport activity in oocyte assays (Fig. 1e).

Wulff et al. recently showed that NPF7.3 lowers the cytoplasmic pH in *Xenopus laevis* oocytes, which can indirectly influence the accumulation equilibrium of weak acids such as GA and ABA<sup>34</sup>. In order to assess whether NPF2.14 has a similar activity, we measured the intracellular oocyte pH using a proton-selective three-electrode voltage clamp setup. We showed that, unlike NPF7.3, NPF2.14 expression in the oocyte does not alter the internal oocyte pH (Fig. 1f). Another factor that theoretically can lead to a false-positive GA export result is alteration of membrane potential. Therefore, we measured the membrane potential of control and NPF2.14-expressing oocytes using a two-electrode voltage clamp setup. The membrane potential of control and NPF2.14-expressing oocytes were both approximately -15 mV (Fig. 1g). When oocytes were subjected to GA<sub>4</sub> for 60 min prior to membrane potential measurement, less GA was observed in NPF2.14-expressing oocytes than control oocytes (Fig. 1g). Thus, NPF2.14 does not shift oocyte membrane potential.

Many NPF proteins, including NPF2.14, contain the ExxE[K/R] motif<sup>35</sup>, which is involved in coupling substrate transport to the proton gradient across membranes<sup>36</sup>. Involvement of the ExxE[K/R] motif in NPF2.14-mediated effluxes would suggest antiporter function. To assess the involvement of the ExxE[K/R] motif, we generated C-terminal YFP-tagged NPF2.14 mutants substituted at each of the three charged residues with a polar but uncharged residue. The YFP-tag alone did not influence the apparent GA<sub>4</sub> transport by the wild-type NPF2.14 (Fig. 1h). When any of the charged residues of the ExxE[K/R] motif was replaced with the polar uncharged Gln residue, no significant difference in GA<sub>4</sub> transport was observed compared to wild-type NPF2.14 (Fig. 1h). Thus, the GA<sub>4</sub> transport mediated by NPF2.14 seems to be ExxE[K/R] motif independent.

In order to test whether the *Xenopus laevis* oocyte GA transport data is physiologically relevant *in planta* we isolated a homozygous T-DNA knockout line for *NPF2.14*. The single *npf2.14* mutant did not show significant shoot or root growth phenotypes (Sup. Fig. 1a-b). Several NPF family members transport nitrate, including NPF6.3<sup>37</sup>. Thus, despite having shown that NPF2.14 does not transport nitrate in oocytes (Fig. 1d), we checked whether *npf2.14* mutants display an impaired growth on low nitrate media. *npf2.14* T-DNA insertion mutants did not have a visible growth phenotype and did not differ from Col-0 plants under low nitrate conditions (Sup. Fig. 1c). To determine whether NPF2.14 is involved in GA distribution and accumulation in the root, we tested whether the distribution of a fluorescently tagged GA<sub>3</sub>-compound (GA<sub>3</sub>-Fl) was affected in the loss-of-function line. GA<sub>3</sub>-Fl has been developed in our lab to serve as a stable, bioactive reporter to study GA movement/accumulation *in planta*<sup>28</sup>. Accumulation of GA<sub>3</sub>-Fl in the endodermis was visible in the Col-0 plants, as previously shown<sup>28</sup>. The *npf2.14* mutants displayed a significantly stronger signal compared to the Col-0 control (Fig. 1i). This enhanced

accumulation was restored to normal levels when expressing *NPF2.14* driven by its native promoter (*pNPF2.14:NPF2.14-GFP*) on the background of the *npf2.14* T-DNA line (Fig. 1i), indicating that loss of NPF2.14 affects GA<sub>3</sub>-FI distribution in the plant. In agreement with this result, *npf2.14* mutants accumulated significantly higher levels of GA<sub>4</sub> in their roots (Fig. 1j).

To test whether NPF2.14 has a dual specificity function and can also import ABA, we tested the distribution of the fluorescently tagged ABA (ABA-FI) in the roots. ABA-FI is non-bioactive but can be utilized to estimate ABA movement in the plant<sup>33</sup>. In addition, we quantified endogenous levels of ABA in the roots. We found that similarly to GA-FI, *npf2.14* mutants accumulated significantly high levels of ABA-FI in their root endodermis cells (Sup. Fig. 2), but showed low levels of native ABA (extracted from the entire root) (Fig. 1j).

In order to study the subcellular localization of NPF2.14, we generated and imaged *35S:NPF2.14-YFP* lines. Interestingly, NPF2.14 localized to the tonoplast vacuole membrane (Fig. 1k). Therefore, although we had hypothesized that NPF2.14 was a GA exporter that transported GA from inside the cytosol to the apoplast, the protein is instead a tonoplast-localized transporter. To the best of our knowledge, this is the first report of a sub-cellular GA/ABA transporter.

### **NPF2.14 regulates suberin formation in the root endodermis**

To characterize NPF2.14 expression patterns in the plant, we generated NLS-YFP and GUS reporter lines driven by the *NPF2.14* promoter. Confocal imaging of the NLS-YFP lines indicated that NPF2.14 is expressed only in the pericycle of the root mature zone, mainly at the phloem poles, and not in the meristematic zone (Fig. 2a, Sup. Fig. 3a). In addition, GUS staining showed expression in the shoot vasculature in seedlings (Sup. Fig. 3b). To test if GA or ABA affects NPF2.14 expression patterns, we examined *pNPF2.14:GUS* lines after an exogenous treatment of GA<sub>3</sub> (5 μM) or ABA (1 μM). While GA did not affect *pNPF2.14:GUS* staining, we observed a stronger staining in the ABA treatment, yet expression remained in the vasculature (Sup. Fig. 4). In the mature stages, the *NPF2.14*-driven reporter was expressed in the periderm (Sup. Fig. 3c). The pericycle is a deep layer of post-embryonic meristematic cells encircling the vascular tissue<sup>38</sup>. In the root, it is required for lateral root emergence<sup>39</sup>, xylem loading<sup>40</sup> and phloem unloading<sup>41</sup>. At later stages it gives rise to the periderm, which serves as the outer protective layer when the surrounding tissue is sloughed off<sup>38</sup>. Both the endodermis and the cork, which is the outermost cell layer of the periderm are suberized tissues<sup>42</sup>.

The expression of the reporter driven by the *NPF2.14* promoter in a tissue that undergoes suberization, taken together with the ability of NPF2.14 to transport ABA, which has been previously shown to regulate suberin deposition<sup>43</sup>, led us to hypothesize that this transporter might facilitate root suberization.

To test this, we analyzed suberization in *npf2.14* T-DNA mutants using Nile red and Fluorol yellow, which are suberin dyes<sup>44,45</sup>. Suberization commences in the endodermis of the upper part of the maturation zone of the root, and, as the plant matures more cells undergo

suberization<sup>46</sup>. Quantification of Nile red and Fluorol yellow fluorescence intensity in the uppermost part of 5-day-old roots revealed that the mutant *npf2.14* plants had significantly lower levels of endodermal suberin than Col-0 plants (Fig. 2b, Sup. Fig. 5). In addition, Col-0 plants roots showed a typical pattern of suberin formation with a non-suberized zone, followed by a suberizing zone where only patches of endodermal cells are suberized (patchy suberization) and a continuous suberized zone. We found significant reduction in the continuously suberized zone for *npf2.14* T-DNA, and an additional CRISPR *npf2.14* allele we generated (Fig. 2c). Suberin levels remained lower compared to Col-0 at later stages of development, including 10-day-old roots and 3-week-old hypocotyls (Sup. Fig. 6), showing that the reduction of suberin deposition in the endodermis is stable over time and that also cork suberin is affected.

Suberin is a complex polyester based on glycerol and long-chain  $\alpha,\omega$ -diacids and  $\omega$ -hydroxyacids<sup>47</sup>, which is primarily found in structures such as the periderm, endodermis, and seed coat<sup>30</sup>. To examine changes in root suberin composition between *npf2.14* and Col-0, we analyzed their suberin monomer profiles via gas chromatography-mass spectrometry (GC-MS). We found significant reductions in ferulic acid, the predominant aromatic component of suberin, as well as in C22 fatty acid and C18:1(9)  $\omega$ -hydroxyacid, two of the most abundant suberin building blocks in the Arabidopsis root endodermis (Fig. 2d). In addition, C18 ester was lower in *npf2.14* roots. These reductions accompanied by lower levels of other monomers resulted in 35% less total suberin contents in the mutant roots (Fig. 2d). Overall, the findings provide several lines of evidence that alteration of NPF2.14 has a substantial effect on suberin deposition in the root endodermis.

Similar to the *npf2.14* mutant, *npf3.1* mutant plants, which have been shown to have impaired GA and ABA delivery to the endodermis<sup>16</sup>, displayed reduced suberization levels compared to Col-0 plants (Sup. Fig. 7). GA and ABA treatments completely rescued the *npf2.14* mutant suberization levels (Fig. 2b).

Endodermal suberization is regulated by ABA perception, both under normal and stress conditions<sup>48,49</sup>. ABA treatment was previously reported to induce endodermal suberization<sup>50</sup>. In agreement with this report, ABA significantly upregulated the Nile red fluorescence intensity in the root endodermis and rescued the reduced suberization observed in the *aba2-1* mutant (*AT1G52340*), which is deficient in ABA biosynthesis<sup>50</sup> (Fig. 2e). To the best of our knowledge, GA has not been previously associated with endodermal suberization. To establish whether or not GA regulates root suberization in *Arabidopsis* seedlings, we quantified suberization levels in GA-treated wild-type and *gal-13* mutant plants using the Nile red dye. GA1 (*AT4G02780*) catalyzes the first committed step in the GA biosynthetic pathway<sup>51</sup>. The GA biosynthesis mutant *gal-13* displayed a significant reduction in suberization levels, which was restored by the exogenous application of GA<sub>3</sub> (Fig. 2f). Furthermore, while ABA treatment, but not GA treatment, can rescue and induce the reduced suberin levels of *aba2-1*, both ABA and GA can rescue the reduced suberin levels of *gal-13* (Fig. 2e-f). Notably, GA treatment could only rescue the suberin back to WT levels, and did not induce it to a higher level as ABA treatment did. GA and ABA have long been thought to have completely antagonistic functions<sup>52</sup>. Our results indicate that this antagonistic activity is more complex and that GA and ABA induce root suberization. Taken

together, our data suggest that NPF2.14 is a pericyclespecific GA and ABA transporter, localized to the tonoplast, and involved in regulating GA and ABA accumulation in the endodermis to promote suberization.

### NPF2.12 and NPF2.13 are plasma membrane-localized GA and ABA importers

*NPF2.12* (*AT1G27080*) and its close paralog *NPF2.13* (*AT1G69870*) form a phylogenetic sub-clade with *NPF2.14* (Fig. 3a, Sup. Fig. 8). We hypothesized that due to this proximity on the phylogenetic tree, NPF2.12 and NPF2.13 might also contribute to GA and ABA accumulation in the endodermis. Both transporters were previously characterized as low-affinity nitrate transporters<sup>53,54</sup> and more recently, were shown to promote GA import activity in heterologous systems<sup>14,34</sup>. Neither have been characterized in plants as GA or ABA transporters. To test for direct GA transport activity of NPF2.12 and NPF2.13, we performed *Xenopus laevis* oocyte-based transport assays. Oocytes expressing NPF2.12 or NPF2.13 accumulated significantly higher levels of GA<sub>1</sub>, GA<sub>3</sub>, GA<sub>4</sub>, GA<sub>7</sub>, GA<sub>9</sub>, GA<sub>12</sub>, GA<sub>19</sub>, and GA<sub>24</sub> compared to control oocytes over the course of 60 min (Fig. 3b, Sup. Fig. 9a). This suggests that NPF2.12 and NPF2.13 are promiscuous GA importers. Both NPF2.12- and NPF2.13-expressing oocytes also had higher levels of ABA accumulation than controls (Fig. 3c).

NPF2.12 and NPF2.13 both contain an ExxE[K/R] motif, which couples the substrate transport to the proton gradient<sup>36</sup>. To test whether NPF2.12 and NPF2.13 substrate transport activity is coupled with external proton concentration, NPF2.12- and NPF2.13-expressing oocytes were exposed to membrane-impermeable GA1 in solutions ranging from pH 5 to 7 in 0.5 pH unit increments. In both NPF2.12- and NPF2.13-expressing oocytes, GA1 accumulation increased as pH was lowered. (Fig. 3d). These data indicate that GA transport by NPF2.12 and NPF2.13 is likely proton-coupled.

To investigate whether the transport of GA by NPF2.12 and NPF2.13 is electrogenic, we used two-electrode voltage clamp electrophysiology (TEVC) to evaluate oocytes that express the transporters. Subtracting the currents elicited by oocytes at different membrane potentials in the absence of GAs from the currents elicited by oocytes in the presence of 500  $\mu$ M GA<sub>3</sub> revealed that GA<sub>3</sub> transport by both NPF2.12 and NPF2.13 is associated with negative currents relative to control oocytes (Fig. 3e). As the negative currents reflect a net positive influx of charges, this indicates at least a 2:1 proton:GA<sub>3</sub> stoichiometry if GA<sub>3</sub> is transported in its anionic form. However, it cannot be excluded that GA<sub>3</sub> may be transported in its neutral form. In non-clamped conditions, NPF2.13 displayed significantly higher uptake levels of GA1 and GA<sub>4</sub> compared to NPF2.12. Similarly, currents elicited by NPF2.13-expressing oocytes were significantly higher compared to NPF2.12-expressing oocytes when oocytes were clamped at membrane potentials that mimic the membrane potential in the non-clamped uptake assays (Fig 3e, f). However, at high negative membrane potential (-120 mV), GA<sub>3</sub> induced-currents in NPF2.12-expressing oocytes were of the same magnitude as those of NPF2.13-expressing oocytes (Fig. 3e, f). This suggests that the transport activity of NPF2.12 is sensitive to alternations in the membrane potential (Fig. 2b). In agreement with previous publications<sup>53,54</sup>, we detected nitrate import into oocytes that expressed NPF2.12 or NPF2.13, but this transport did not interfere with the GA



transport capabilities as GA accumulation was not affected in GA/nitrate competition assays (Sup. Fig. 9b). In continuation, we performed equimolar GA/ABA competition transport assays, which showed ABA does not affect GA transport, whereas a slight but significant enhancement in ABA uptake was seen in NPF2.13-expressing oocytes when exposed simultaneously to GA<sub>3</sub>. The data shows that NPF2.12 and NPF2.13 are multi-specific towards nitrate, ABA and GA and suggests that GA might enhance NPF2.13 ABA transport activity (Sup. Fig. 10). Multi-specificity has emerged as an inherent property of the NPF family that is suggested to enable the tantalizing integration of environmental information to the availability of the different nutrients<sup>18</sup>. Exhaustive structure–function studies are needed to decipher the molecular basis of the selectivity of these transporters.

To elucidate whether these transporters are part of the GA transport mechanisms in the plant, we treated T-DNA knockout lines, which did not display any visible phenotypes (Sup. Fig. 11), with GA<sub>4</sub>-Fl. *npf2.12* and *npf2.13* single mutant plants treated with GA<sub>4</sub>-Fl displayed a significant reduction in accumulation in the endodermis compared to Col-0 plants (Fig. 3g), similar to reduced levels detected in *npf3.1* mutants. To verify that the reduction in GA-Fl accumulation was not due to an off-target mutation, we repeated the test with an additional *npf2.12* T-DNA insertion line (*npf2.12-2*) and obtained similar results (Sup. Fig. 12a). In addition, homozygous *pNPF2.13:NPF2.13-YFP* plants completely rescued the root GA<sub>3</sub>-Fl phenotype (Sup. Fig. 12b).

We next generated plants ectopically expressing *NPF2.12* and *NPF2.13* fused to YFP (*p35S:NPF2.12-YFP* and *p35S:NPF2.13-YFP* respectively) and treated them with GA<sub>3</sub>-Fl. These lines showed a remarkably strong accumulation of GA<sub>3</sub>-Fl in all root cells (Fig. 3h), supporting the hypothesis that NPF2.12 and NPF2.13 are GA transporters *in planta*. In order to test whether NPF2.12 and NPF2.13 have a dual specificity function and can also import ABA (such as NPF2.14), we treated plants with fluorescently tagged ABA (ABA-Fl)<sup>33</sup>. Overexpression of NPF2.12 or of NPF2.13 resulted in extreme ABA-Fl accumulation compared to Col-0 plants (Fig. 3h), implying that both transporters import GA and ABA *in planta*.

Finally, in order to address the subcellular localization of NPF2.12 and NPF2.13, we imaged the root epidermis cells of *p35S:NPF2.12-YFP* and *p35S:NPF2.13-YFP* lines. Both NPF2.12 and NPF2.13 are localized to the plasma membrane (Fig. 3i). Together, the *in planta* and oocyte results indicate that NPF2.12 and NPF2.13 are dual-substrate, plasma membrane-localized GA and ABA importers.

### NPF2.12 and NPF2.13 regulate root suberization

To further elucidate the biological function of NPF2.12 and NPF2.13, we generated NLS-YFP and GUS reporter lines to map NPF2.12 and NPF2.13 expression patterns. Confocal microscopy of plants expressing *NLS-YFP* driven by *NPF2.12* and *NPF2.13* native promoters revealed that in the root, NPF2.12 was expressed in the pericycle of the whole root and subsequently in the periderm of mature plants; NPF2.13, on the other hand, was expressed only in the shoot (Fig. 4a, Sup. Fig. 13). Analysis of *pNPF2.12:GUS* and *pNPF2.13:GUS* lines showed that the two transporters are expressed in the shoot vasculature (Fig. 4b). GA or ABA treatment did not have a major effect on *NPF2.12* and *NPF2.13*

expression level, detected by qPCR and GUS reporter lines (Sup. Fig. 14). The fact that the *NPF2.13* translational fusion construct (*pNPF2.13:NPF2.13-Venus*) introduced into the *npf2.13* mutant background rescued the root GA<sub>3</sub>-FI accumulation phenotype indicated that the promoter region we cloned is sufficient and that NPF2.13 expression is restricted to the shoot (Sup. Fig. 12b). Similar to *npf2.14* mutants, *npf2.12* and *npf2.13* mutant plants displayed a reduction in suberization. Mutant roots stained with Nile red or Fluorol yellow showed a weaker fluorescence intensity than detected in Col-0 plants (Fig. 4c, Sup. Fig. 15). In agreement, additional mutant alleles for *npf2.12* and *npf2.13* showed reduction in suberin level and patterning (Sup. Fig. 16). Similar to *npf2.14* mutants, we detected a reduction in endodermal suberin levels of *npf2.12* and *npf2.13* 10-day-old roots and a lower levels of cork suberin in 3-week-old *npf2.12 npf2.13* double mutant hypocotyls stained with Fluorol yellow (Sup. Fig. 17).

In order to test if *NPF2.12* and *NPF2.13* have partially redundant activities, we generated the *npf2.12 npf2.13* double mutant. The phenotype of the *npf2.12 npf2.13* double mutant line was not enhanced compared to the single *npf2.12* and *npf2.13* mutants (Fig. 4c-d). We further generated additional double and triple mutant combinations with the *npf2.14* mutant via CRISPR genome editing (as *NPF2.14* is genetically linked to *NPF2.13*), and tested their activity. We analyzed suberin patterning for all genotypes and found that the phenotypes of the higher order mutant knockouts were not enhanced compared to the single mutants (Sup. Fig. 18). This result is in line with the fact that there is limited overlap between the three genes in terms of expression pattern or protein localization. Notably, GA<sub>3</sub> or ABA treatment completely rescued *npf2.12* low-suberin phenotype. The phenotypes of *npf2.13* and of the *npf2.12 npf2.13* double mutant were completely rescued by ABA and largely rescued by GA<sub>3</sub> (Fig. 4c). Quantification of suberin monomer content revealed that both *npf2.12* and *npf2.13* mutant roots accumulated ~40% less total suberin contents compared to the Col-0 attributed to lower levels of vanillic and ferulic acids, C22 fatty acid, C20 fatty alcohol, and C18:1(9), C22 and C26 ω-hydroxyacids (Fig. 4e).

### **NPF2.12 and NPF2.13 regulate shoot-to-root GA translocation**

Our previous work showed that GA<sub>12</sub>, though not bioactive, is the primary GA form transported over long distances through the vasculature in *Arabidopsis thaliana*<sup>55</sup>. GA<sub>12</sub> can move through the xylem in a root-to-shoot manner and in the phloem in a shoot-to-root direction to regulate adaptive plant growth<sup>12,13</sup>. However, the mechanism regulating this process remains unknown<sup>21</sup>. NPF2.13 was expressed strictly in the shoot (Fig. 4a, b), yet the knockout led to a phenotype in the root endodermis (Fig. 4c). This led us to hypothesize that the transporters are involved in the long-distance shoot-to-root translocation of GA. To test whether NPF2.12 and NPF2.13 facilitate shoot-born GA loading into the phloem, we quantified GA content in phloem exudates collected from leaf petioles. The double *npf2.12 npf2.13* loss-of-function mutant showed a striking reduction in GA<sub>12</sub> content in the collected phloem exudates (Fig. 5a). Other GA metabolites (GA15, GA24, GA9, GA4, GA34, and GA51) were not significantly reduced (Fig. 5a). ABA levels showed a mild decrease, significant when compared to Col-0 using two-tailed t-tests but not significant when using Dunnett's multiple comparisons test ( $p > 0.05$ ). The results imply that NPF2.12 and NPF2.13 regulate GA, and possibly ABA, loading into the shoot phloem. In agreement,

quantification of active GA<sub>4</sub> content in the root which is downstream to GA<sub>12</sub> in the biosynthesis pathway, showed a significant reduction in the *npf2.12* and *npf2.13* single and double mutant lines (Fig. 5b). ABA content was also significantly lower in mutant roots compared to Col-0 (Fig. 5b).

To further test this hypothesis, we examined the expression pattern of the two transporters using reporter lines. In mature rosette, *NPF2.12* and *NPF2.13* were expressed in the shoot apex and the main vascular vein (Fig. 5c). Cross-sections of *pNPF2.12:GUS* and *pNPF2.13:GUS* leaf petioles showed that both genes were expressed in the phloem companion cells (Fig. 5c). Next, we investigated whether these transporters are involved in long-distance GA movement from the shoot to the root. For this purpose, 16-day-old plants were grown on paclobutrazol, a GA biosynthesis inhibitor, for 4 days to deplete the plants of native GA, and GA<sub>12</sub> was applied to a single leaf. We then quantified the abundance of the DELLA growth repressing protein REPRESSOR OF GA1-3 (RGA, *AT2G01570*) in the root. DELLA proteins are central inhibitors of GA-regulated processes and GA relieves their inhibiting activity by activating their degradation<sup>56</sup>. Time-course experiments in Col-0 plants showed a significant reduction in RGA accumulation in the root after GA<sub>12</sub> treatment, indicating GA<sub>12</sub> movement from the shoot to the root (Fig. 5d). On the other hand, in the *npf2.12 npf2.13* double mutant, RGA abundance remained stable (Fig. 5d), signifying a reduced GA accumulation in the root.

To understand how the NPF2.12/2.13/2.14-mediated long-distance GA transport influences root growth and suberin formation, we performed a series of micrograftings between Col-0, different combinations of *npf* double mutants and GA-deficient *gal-3* mutants and quantified both root elongation rate and Nile red fluorescence intensity. As expected, *gal-3* self-grafts had shorter roots and displayed a significantly lower Nile red fluorescence intensity than the Col-0 self-grafts (Fig. 5e, Sup. Fig. 19). Remarkably, *gal-3* roots grafted to Col-0 shoots were indistinguishable from that of Col-0 self-grafts (Fig. 5e), supporting our hypothesis that GA can be transported from the shoot to the root to induce suberin formation. In comparison, *gal-3* roots grafted to *npf2.12 npf2.13* shoots (*gal-3/npf2.12 npf2.13*) were smaller and showed a reduction in suberization levels compared to *gal-3/Col-0*, indicating a partial requirement of NPF2.12 and NPF2.13 in long-distance GA transport (Fig. 5e, Sup. Fig. 19). Similarly, *npf2.12 npf2.14* and *npf2.13 npf2.14* double mutant's shoots grafted to *gal-3* roots, showed reduced suberin levels compared to *gal-3/Col-0*. In addition, we found that GA<sub>3</sub>-Fl application, specifically to the shoots of *npf2.12 npf2.14* double mutant or *npf2.14*, resulted in lower levels of GA<sub>3</sub>-Fl in the roots of seedlings growing on paclobutrazol (Sup. Fig. 20). The results further suggest that NPF2.12/13/14 contribute to long-distance, shoot-to-root, GA translocation.

We hypothesize that once GA<sub>12</sub> is translocated to the roots, it can be converted to the bioactive GA<sub>4</sub> by the GA20ox and GA3ox enzymes, which are expressed in the root<sup>57</sup>. Profiling the expression pattern of the *GA3ox* promoters (*AT1G15550*, *AT1G80340*), catalyzing the last step in bioactive GA<sub>4</sub> hormone synthesis, showed that expression of these enzymes is restricted to the stele as previously reported<sup>57</sup> (Fig. 5f). Together, these results imply that NPF2.12 and NPF2.13 function in GA<sub>12</sub> loading into the phloem for

long-distance transport from the shoot to the root, with conversion to GA<sub>4</sub> taking place in the root stele.

### Hormone storage in the phloem unloading zone facilitates suberization

To broaden our knowledge of GA and ABA distribution in the root and how it affects suberization, we created a mathematical model to simulate hormone distributions within the root cross-section, extending a modeling framework previously developed to study auxin transport<sup>58</sup>. Using a multicellular template segmented from a root-cross-sectional image (Sup. Fig. 20), we incorporated into the model experimentally observed transporter distributions: NPF3.1 on endodermal cell membranes<sup>16,29</sup>, NPF2.12 on pericycle cell membranes (Fig. 3i, 4a), and NPF2.14 on the pericycle tonoplasts (Fig. 1k, 2a). The model simulated active hormone transport via NPF3.1, NPF2.12, and NPF2.14, passive hormone transport across both plasma membrane and tonoplast, hormone synthesis and degradation, and hormone diffusion within the apoplast with significantly reduced diffusion in the endodermal apoplast due to the presence of the Casparian strip. To parameterize the model, permeabilities associated with each passive and active transport component were estimated using the oocyte data, and transport rates were then specified based on established pH and membrane potential values for plant cells<sup>58–60</sup> (Sup. Table 5). An important factor is the source of the hormone. *ABA2* and *AAO3*, which encode enzymes necessary for ABA biosynthesis, were previously shown to be expressed in the vasculature<sup>61</sup>. Bioactive GA<sub>4</sub> is also synthesized at high levels in the *Arabidopsis* stele<sup>57</sup> (Fig. 5f). Considering these data, together with the phloem unloading zone<sup>41</sup>, the docking belt for the long-distance shoot-to-root transported hormones, led us to specify the stele as the source of active GA and ABA in the model.

We first used the model to test the hypothesis that the discovered clade of transporters is necessary and sufficient to explain the observed endodermal hormone accumulation. With all transporters present, the model predicts high levels of both cytoplasmic and vacuolar hormone in the endodermis (Fig. 6a for GA and Sup. Fig. 21), consistent with the wildtype GA-FI and ABA-FI observations (Fig. 1i, Sup. Fig. 2)<sup>16</sup> and where suberization later occurs. Mutations in *npf3.1* and *npf2.12* were predicted to have reduced endodermal hormone cytoplasmic concentrations (Fig. 6a, Sup. Fig. 21), in agreement with the loss of endodermal GA-FI in these mutants (Fig. 3g)<sup>16</sup> and explaining their loss of suberization (Fig. 4c, Sup. Fig. 7, Sup. Fig. 15–18). We also considered NPF2.13 which is not expressed in the root (Fig. 4a) but contributes to the long-distance translocation of GA<sub>12</sub> (Fig. 6a, Sup. Fig. 21). Reducing long-distance translocation via the *npf2.13* mutation can be simulated by reducing the stele-specific synthesis rate, which leads to reduced predicted hormone concentrations throughout the root cross-section, again providing an explanation of the reduction in suberization (Fig. 4c, Sup. Fig. 15–18). We concluded that NPF2.12, NPF2.13 and NPF3.1 all play distinct and necessary roles in creating the hormone accumulation within the endodermis that mediates suberization.

We then used the model to investigate the role of the tonoplast-localized NPF2.14. In contrast to the *npf2.12*, *npf2.13* and *npf3.1* mutations, simulations of the *npf2.14* mutation predicted endodermal cytoplasmic hormone concentrations that are higher than those in

the wild-type (Fig. 6a, Sup. Fig. 22), in agreement with the GA-FI accumulation at the elongation zone (Fig. 1i). Why *npf2.14* exhibits reduced suberization when endodermal accumulation is higher remained unclear. However, the model predictions revealed that the vacuolar concentrations in the *npf2.14* pericycle are much lower than in the wild-type (Fig. 6a, Sup. Fig. 23). These predictions led us to hypothesize that NPF2.14 regulates hormone levels both inside and outside of vacuoles which could provide a hormone source in the maturation zone, where the hormone is no longer supplied by the phloem yet is required for suberization. To test this hypothesis, we simulated the hormone dynamics after cells leave the phloem unloading zone and found that the presence of NPF2.14 leads to higher predicted endodermal cytoplasmic concentrations as cells mature (Fig. 6b, Sup. Fig. 24). This suggests that the tonoplast-pericycle-localized NPF2.14 ensures that endodermal hormone concentrations are at levels necessary to mediate suberization. Thus, based on the model, we propose a pericycle-specific slow-release GA and ABA mechanism that explains how the two hormones are loaded into the pericycle vacuoles at the phloem unloading zone<sup>41</sup> and released from these vacuoles later on when the cells are mature (Fig. 6b). In this mechanism, GA and ABA unloaded from the phloem are transported into the pericycle and loaded into the vacuole to form a storage pool. When these cells reach the maturation zone, the GA and ABA that were stored in the pericycle vacuoles are transported by NPF3 into the endodermis to induce suberization.

In conclusion, the mathematical model revealed that the discovered clade of transporters is sufficient to explain the observed hormone distributions, with NPF3.1, NPF2.12 and NPF2.13 playing distinct and necessary roles for endodermal accumulation. Furthermore, the model revealed that the tonoplast-localized NPF2.14 facilitates vacuolar hormone accumulation within the pericycle which provides a source of hormone, enabling the cross-section to maintain high endodermal hormone levels after cells leave the phloem unloading zone<sup>41</sup>. Thus, the model predictions provide mechanistic explanations for the suberization phenotypes observed in the NPF mutants.

## Discussion

In this work, we identified *NPF2.14*, a previously uncharacterized transporter, as a dual-specificity GA and ABA vacuolar transporter. To the best of our knowledge, NPF2.14 is the first known sub-cellular GA/ABA transporter. We showed that *NPF2.14* is expressed in the pericycle to facilitate endodermal root suberization. Oocyte experiments showing that NPF2.14 exports GA from the cytosol, combined with NPF2.14's localization to the tonoplast, indicates that NPF2.14 transports GA and ABA from the cytosol into the vacuole.

The results presented here suggest that the pericycle serves as a buffer zone, regulating the transitions of hormones from the stele to the endodermis (Fig. 7). The stele acts as the source of bioactive GA and ABA in the root<sup>57,61</sup>. Both hormones have been shown to accumulate and affect their respective response specifically in the endodermis<sup>25,28,62,63</sup>. In addition, it appears that the GA and ABA do not simply flow through the pericycle but rather are loaded into the vacuole by NPF2.14 to form a reservoir for later developmental stages. We propose that the high levels of GA and ABA present in the phloem unloading zone are taken into the pericycle by NPF2.12. Once in the pericycle

cells, NPF2.14 facilitates their import into the vacuole for storage. We speculate that a slow-release mechanism feeds the differentiating cells with GA and ABA, thus allowing suberin formation in the mature root (Fig. 7). Vacuoles have been proposed to act as storage, modification, or degradation compartments for plant hormones<sup>64</sup>. It is possible that tonoplast-localized NPF2.14 mediates the hormonal homeostasis balance that is needed for the proper execution of the developmental plan in the neighboring cell file. At this point, it is not clear if GA and ABA are stored at their bioactive form in the vacuole.

Confocal imaging of the NLS-YFP lines indicated that both *NPF2.12* and *NPF2.14* are expressed in the pericycle, mainly at the phloem poles (Fig. 2a, 4a). It is therefore possible that the hormone uptake is not carried out uniformly throughout the pericycle ring, but rather amplified at the pericycle phloem poles cells. If so, do the two hormones retain polar distribution in the pericycle and in the subsequent endodermis layer? Or do the hormones have the ability to move within the two cell file rings? Future mathematical models and genetic work is required to address these questions.

The significantly reduced GA<sub>12</sub> content in *npf2.12 npf2.13* double knockout phloem extracts implies that these transporters are required for GA<sub>12</sub> loading into the phloem and translocation of GA<sub>12</sub> from the shoot to the root. Thus, we hypothesize that NPF2.12 and NPF2.13, which are plasma membrane-localized importers expressed in the shoot phloem companion cells, are part of the long-sought mediators of long-distance GA shoot-to root translocation. Our results are in agreement with the previous finding that GA<sub>12</sub> is the main form of GA that is transported long distances through the plant<sup>12</sup>. In addition, we showed that GA<sub>3ox1</sub> and GA<sub>3ox2</sub>, which catalyze the final step of active GA production, are expressed in the root stele<sup>57</sup> (Fig. 5f). Thus, once GA<sub>12</sub> docks at the root stele, it is converted by GA<sub>3ox1</sub> and GA<sub>3ox2</sub> into the bioactive GA<sub>4</sub> form, which is then delivered from the stele to the endodermis by NPF2.12 and NPF3.1 (Fig. 6c).

It is intriguing that NPF2.12 and NPF2.13 act as GA<sub>12</sub> shoot-to-root transporters and also promote the delivery of the bioactive forms of GA and ABA to the endodermis. In both cases, a plasma membrane import activity is involved, but the substrate specificity differs. Since we showed that NPF2.12 and NPF2.13 are able to transport both intermediates of- and the bioactive forms of GA (Fig. 3b, Sup. Fig. 9-10) and that GA<sub>12</sub> is present at high levels in the shoot phloem (Fig. 5a), we speculate that GA<sub>12</sub> is the primary substrate of the transporters. In the root stele, NPF2.12 recognizes bioactive ABA and GA<sub>4</sub>, which are present in high concentrations due to being synthesized there<sup>61</sup>.

The presented work showed for the first time that GA deficiency results in lower endodermal suberization and can be complemented by GA or ABA. This result is interesting at multiple points of view. First, it may explain for the first time, the physiological importance of GA accumulation in the endodermis. Second, it suggests that GA and ABA function non-antagonistically to promote endodermis suberization. At this stage, it is not clear whether GA promotes endodermis suberization directly by, for example, direct binding and activation of suberin biosynthesis factors by DELLA or DELLA co-partners, or whether suberization is a secondary effect of maturation or signaling through the ABA components. Our result in oocytes, suggesting that GA enhances ABA import, may point to another possibility

where the co-activity in regulating suberin formation at the transport level of the two hormones. While GA and ABA are considered antagonistic hormones<sup>65</sup>, it is possible that the two hormones can act non-antagonistically to promote root suberization by dual-transport activity of the hormones. How does GA promote ABA import biochemically from the transport structure/function point of view is not clear at the moment. The non-antagonistically results between the two hormones are in agreement with the growth defects displayed by biosynthesis mutants of these hormones, which result in small dark green plants<sup>66–68</sup>. Thus, it could be that, in a specific context where GA and ABA transport plays a role, the two hormones act synergistically.

## Methods

### Plant material and growth conditions

All *Arabidopsis thaliana* lines are in a Col-0 background. T-DNA insertion lines were supplied by the *Arabidopsis* Biological Resource Center. PCR genotyping for homozygous lines was performed using the primers listed in Sup. Table 1. To generate *npf2.12* *npf2.13* double mutant, *npf2.12* (SALK\_138987) was crossed to *npf2.13* (SALK\_022429) to obtain an F1 population. F3 homozygous plant was selected by PCR genotyping. Additional T-DNA lines described in this study: *npf2.12* (SALK\_104042c), and *npf2.13*: (SALK\_053264). To generate the triple *npf2.12* *npf2.13* *npf2.14* mutant, a CRISPR/Cas9 construct targeting *NPF2.14* was introduced into the *npf2.12* *npf2.13* double mutant background.

Plants were sown on vertical plates containing 0.5 × Murashige-Skoog (MS) medium, 1% sucrose, and 0.8% agar, pH 5.7, stratified for 2 days at 4 °C in the dark, then transferred to growth chambers (Percival CU41L5) at 21 °C, 100 μE m<sup>-2</sup> s<sup>-1</sup> light intensity under long day light (16 h light/8 h dark). All plants in suberin quantification experiments were grown on 0.5 × MS medium, 1% sucrose, and 0.8% agar, pH 5.7 for 3 days and subsequently moved to 0.5 × MS medium without sucrose supplementation due to phenotype masking by the sucrose treatment. For low-nitrate experiments, plants were sown on MS with vitamins, nitrate free (Caisson labs MSP07-50LT), which was supplemented with 0.01 mM (Low nitrate) or 10 mM (High nitrate) KNO<sub>3</sub>.

### CRISPR

To generate the CRISPR/Cas9 vector targeting *NPF2.14* the MoClo system was implemented. Cloning of the *NPF2.14* specific guide (Guide sequence: ATTGTTGTCTCGTTCGTTAAATCCG) into the system was done according to Engler et al.<sup>69</sup>.

### Hypocotyl cross sections

Sectioning and clearing were performed as describe in Ursache et al<sup>44</sup>. 3-Week-old hypocotyls were fixed in 4% PFA for an hour, rinsed twice in 1xPBS embedded in 5% agarose and sectioned to 150 μM slices using a Leica VT1000S vibratome. Slices were cleared using a ClearSee solution for 5 days. Following clearing sections were counterstained with 0.1% Calcofluor White in ClearSee solution for 30 min. Next, the

seedlings were washed in ClearSee for 30 min with gentle shaking. For imaging, sections were mounted directly in ClearSee and imaged using a Zeiss LSM 780 inverted microscope.

### Hormone application

Hormone was added to the agar medium at concentrations indicated in the figure legends. Seedlings were either germinated on media or moved after germination to treatment plates. GA-F1 (5  $\mu$ M) was applied in liquid MS media for 16 h prior to imaging. For *gal* experiments, both Col-0 and *gal* seeds were imbibed in sterile water containing 5  $\mu$ M GA<sub>3</sub> for 16 h to induce uniform germination. Following imbibition, seeds were washed three times in sterile water to wash away excess GA and were sown on MS plates.

### Cloning of NPFs overexpression and reporter lines

*NPF2.12* and *NPF2.14* coding sequences were synthesized by Bio Basic Inc., cloned into pENTR/D-TOPO (Invitrogen K2400), and subsequently cloned into the pH7YWG2 destination vectors using the LR Gateway reaction (Invitrogen 11791). *NPF2.12* and *NPF2.14* promoters were amplified with the primers listed in Sup. Table 2 using a Phusion high-fidelity polymerase (New England Biolabs), cloned into pENTR/D-TOPO, and then cloned into pMDC7 vector for *NLS-YFP* reporters and pGWB3 vector for GUS reporters.

To generate *pNPF2.13:GUS* reporter, the promoter of *NPF2.13* (1.7-kb fragment) was PCR amplified from Col-0 genomic DNA with appropriate primers listed in Sup. Table 2 and inserted into pDONR221 (Invitrogen) by Gateway cloning and recombined with pGWB633

### Imaging and analysis

Seedlings were stained in 10 mg/L<sup>-1</sup> propidium iodide (PI) for 5 min, rinsed, and mounted in water. Seedlings were imaged using a laser scanning confocal microscope (Zeiss LSM 780 inverted microscope), with argon laser set at 488 nm for fluorescein, 514 nm for YFP, and 561 nm for PI excitation. Emission filters used were 493-548 nm for fluorescein derivatives, 508-570 nm for YFP, and 583-718 nm for PI emission. Image analysis and signal quantification were done with the measurement function of ZEN lite software. The number of quantified biological repeats and sampling points is indicated for each graph in figure legends. All statistical analyses and graphs were made using GraphPad Prism V. 8.

### Root length characterization

For root length measurements, seedlings were imaged using Zeiss Stemi 2000-C stereo microscope and measured using ImageJ software (<http://rsbweb.nih.gov/ij/index.html>).

### Histochemical GUS staining

Plants were immersed in 100 mM sodium phosphate buffer (pH 7.0) containing 0.1% Triton X-100, 1 mM 5-bromo-4-chloro-3-indolyl- $\beta$ -D-glucuronic acid cyclohexylammonium salt (Sigma-Aldrich), 2mM potassium ferricyanide, and 2 mM potassium ferrocyanide. Plants subject to vacuum treatment for 10 min and then incubated at 37 °C for 16 h. Tissues were cleared with 30%, 50%, and 70% ethanol for 30 min in each concentration and imaged using an AxioZoom 16, Zeiss binocular microscope.



For cross-sectioning of GUS-stained leaf petioles, after clearing in 70% ethanol, the samples were fixed in FAA solution (3.2% formaldehyde, 5% acetic acid, 50% ethanol) for 30 min and kept overnight at 4°C. The samples were then dehydrated in an ethanol gradient ranging from 50% to 96%, and incubated in 2% eosin overnight at 4°C. After several washes in 96% ethanol, the samples were progressively rehydrated in ethanol/HISTO-CLEAR II (Electron Microscopy Sciences) solution, incubated in 50% HISTO-CLEAR II 50% PARAPLAST PLUS (McCormick Scientific) at 60°C for 2 h, and embedded in 100% PARAPLAST PLUS. Paraffin-embedded samples were cross-sectioned with LEICA RM2155 microtome and imaged using a Leica Leitz Dmrb microscope.

### Nile red suberin staining, imaging and quantification

Nile red suberin staining was performed as described by Ursache et al.<sup>44</sup> In short, 5-day-old seedlings were fixed in paraformaldehyde for 1 h under gentle agitation and washed twice in phosphate-buffered saline, pH 7.4. Plants were covered in filtered 0.05% Nile red (Acros Organics, 7385-67-3) solution dissolved in ClearSee for 16 h. Following staining, plants were washed three times in ClearSee for 30 min each wash. Next, plants were counterstained with 0.1% calcofluor white (Glentham Life Sciences, 4404-43-7) dissolved in ClearSee for cell wall imaging. After 30 min, plants were washed in ClearSee for 30 min. Plants were mounted directly in ClearSee on slides and imaged with a Zeiss LSM780 confocal microscope. Images were taken from the upper part of the root and under the root-hypocotyl junction with an argon laser set at 514 nm for Nile red excitation and 405 nm for calcofluor excitation. Emission filters used were 561-753 nm filter for Nile red and 410-511 nm filter for calcofluor emission. Fluorescence intensity was assessed using the Zen software from 5 endodermal cells per root.

For root patterning, following Nile red staining, roots were imaged using an AxioZoom 16, Zeiss binocular microscope. Root length and length of continuously- and patchy-suberized zones was measured using ImageJ. Percent of suberized area was normalized to total root length. All statistical analyses and graphs were made using GraphPad Prism V. 8.

### Quantitative RT-PCR

Total RNA was isolated from the indicated plant materials using RNeasy Plant Mini Kit (QIAGEN 74,904). DNA was removed by RQ1 RNase-free DNase (Promega M6101). Total RNA (2 µg) converted to complementary DNA (cDNA) using M-MLV Reverse Transcriptase (Promega M1701) with oligo(dT)<sub>15</sub> primer according to manufacturer protocols. Quantitative RT-PCR was performed with 40 ng cDNA in a final volume of 10 µl with Fast SYBR Green Master Mix (ABI 4385612) using Step One Plus System and software (ABI). The reaction conditions included 40 amplification cycles, (3 s at 95 °C; 30 s at 60 °C). Three technical repeats were performed for each cDNA sample, and at least three biological repeats were used for each treatment. The relative quantification was calculated with the  $\Delta\Delta C_t$  method, PP2A used as the reference gene. Primers are specified in Supplementary Table 3.

## Phylogenetic tree

Protein sequences for *Arabidopsis thaliana* NPF family members were retrieved from TAIR (<https://www.arabidopsis.org>). Phylogenetic relationships were defined using Phylogeny.fr (<http://www.phylogeny.fr/>) and visualized with FigTree software (<http://tree.bio.ed.ac.uk/software/figtree/>).

## Transport assays in *Xenopus oocytes*

Coding sequences were cloned into the pNB1u vector, and complementary RNA (cRNA) was produced as described in Wulff et al.<sup>34</sup> *Xenopus* oocyte assays were performed as described previously.<sup>34</sup> Defolliculated *Xenopus laevis* oocytes (stage V-VI) were purchased from Ecocyte Biosciences and were injected with 25 ng cRNA in 50.6 nl using a Drummond Nanoject II and incubated for 2-4 days at 16 °C in HEPES-based kulori (90 mM NaCl, 1 mM KCl, 1 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 5 mM HEPES, pH 7.4) before use. Oocytes were pre-incubated in MES-based kulori (90 mM NaCl, 1 mM KCl, 1 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 5 mM MES, pH 5) for 4 min and were then transferred to phytohormone-containing MES-based kulori for 60 min. After washing three times in 25 ml HEPES-based kulori followed by one wash in 25 ml deionized water, oocytes were homogenized in 50% methanol and stored for >30 min at -20 °C. Following centrifugation (25000 g for 10 min 4 °C), the supernatant was mixed with deionized water to a final methanol concentration of 20% and filtered through a 0.22-µm filter (MSGVN2250, Merck Millipore) before analytical LC-MS/MS as described below. For nitrate assays, sodium chloride in kulori was substituted for equimolar sodium nitrate in order not to affect the membrane potential.

## Quantification of phytohormone content by LC-MS/MS

Compounds in the diluted oocyte extracts were directly analyzed by LC-MS/MS. The analysis was performed with modifications from the method described in Tal et al.<sup>16</sup> In brief, chromatography was performed on an Advance UHPLC system (Bruker). Separation was achieved on a Phenomenex Kinetex 1.7u XB-C18 column (100 x 2.1 mm, 1.7 µm, 100 Å) with 0.05% v/v formic acid in water as mobile phase A and acetonitrile with 0.05% formic acid (v/v) as mobile phase B. The gradients used for elution of GAs were 0-0.5 min, 2% B; 0.5-1.3 min, 2-30% B; 1.3-2.2 min 30-100% B, 2.2-2.8 min 100% B; 2.8-2.9 min 100-2% B; and 2.9-4.0 min 2% B. The gradients used for elution of ABA were 0-0.5 min, 2% B; 0.5-1.2 min, 2-30% B; 1.2-2.0 min, 30-100% B; 2.0-2.5 min, 100%; 2.5-2.6 min, 100-2% B; and 2.6-4.0 min, 2% B. The mobile phase flow rate was 400 µl min<sup>-1</sup>, and column temperature was maintained at 40 °C. The liquid chromatography was coupled to an EVOQ Elite triple quadrupole mass spectrometer (Bruker) equipped with an electrospray ion source operated in positive and negative ionization mode. Instrument parameters were optimized by infusion experiments with pure standards. For analysis of GAs, the ion spray voltage was maintained at +4000 V and -4000 V in positive and negative ionization mode, respectively, and the heated probe temperature was set to 200 °C with probe gas flow at 50 psi. For ABA, the ion spray voltage was maintained at -3300 V in negative ionization mode, and heated probe temperature was set to 120 °C with probe gas flow at 40 psi. Remaining settings were identical for all analytical methods with cone temperature set to 350 °C and cone gas to 20 psi. Nebulizing gas was set to 60 psi and collision gas to 1.6

mTorr. Nitrogen was used as probe and nebulizing gas, and argon as collision gas. Active exhaust was constantly on. Multiple reaction monitoring was used to monitor analyte parent ion to product ion transitions for all analytes. Multiple reaction monitoring transitions and collision energies were optimized by direct infusion experiments. Detailed values for mass transitions can be found in **Supplemental Table 4**. Both Q1 and Q3 quadrupoles were maintained at unit resolution. Bruker MS Workstation software (Version 8.2.1) was used for data acquisition and processing. Linearity in ionization efficiencies were verified by analyzing dilution series of standard mixtures. Sinigrin glucosinolate was used as internal standard for normalization but not for quantification. Quantification of all compounds was achieved by external standard curves diluted with the same matrix as the actual samples. All GAs were analyzed together in a single method. GA<sub>12</sub> suffered from severe ion suppression when combined with the other GAs in the standard curve, thus quantification was not achieved for GA<sub>12</sub>.

### Root suberin monomer profiling by GC-MS

Suberin monomers were extracted from Col-0 and mutant roots according to the protocols previously described by <sup>71,72</sup>. A sample volume of 1 µL was injected in splitless mode on a GC-MS system (Agilent 7693A Liquid Auto injector, 8860 gas chromatograph, and 5977B mass spectrometer). GC was performed (HP-5MS UI column; 30 m length, 0.250 mm diameter, and 0.25 µm film thickness; Agilent J&W GC Columns) with injection temperature of 270°C, interface set to 250°C, and the ion source to 200°C. Helium was used as the carrier gas at a constant flow rate of 1.2 mL min<sup>-1</sup>. The temperature program was 0.5 min isothermal at 70°C, followed by a 30°C min<sup>-1</sup> oven temperature ramp to 210°C and a 5°C min<sup>-1</sup> ramp to 330°C, then kept constant during 21 min. Mass spectra were recorded with an *m/z* 40 to 850 scanning range. Chromatograms and mass spectra were evaluated using the MSD ChemStation software (Agilent). Integrated peaks of mass fragments were normalized for sample dry weight and the respective C32 alkane internal standard signal. For identification, the corresponding mass spectra and retention time indices were compared with the NIST20 library as well as in-house spectral libraries.

### *Xenopus* oocyte injection-based efflux transport assays and competition assays

For injection-based export assays, on the second day of gene expression, oocytes were injected with 23 nl 8.2 mM in 98 mM KCl, 1 mM CaCl<sub>2</sub>, 10 mM HEPES, pH 7.4. T1 oocytes were left 10 min to heal and were then transport was evaluated as described above. T2 oocytes were left for approximately 20 h in HEPES-based kulori at 16° C, followed by transport analysis.

### Quantification of nitrate from oocytes by HPLC

Nitrate concentration in the oocyte extracts was quantified using a Dionex ICS-2100 anion exchange chromatography system (Thermo Scientific). The separation was done on a Dionex IonPac AG11-HC analytical column coupled to the AS11-HC guard column (Thermo Scientific). The columns were connected to a Dionex AERS 500 anion suppressor (Thermo Scientific). The analyses were performed under the following conditions: sample injection volume 4.8 µl, column temperature 30 °C, flow rate of 0.38 ml/min, isocratic

eluent gradient using 30 mM KOH solution in QH<sub>2</sub>O, suppressor current of 29 mA, and runtime of 15 min. The nitrate detection was done at 220 nm using a Dionex UltiMate 3000 (Thermo Scientific). QH<sub>2</sub>O water dilutions of Dionex Combined Seven Anion Standard (Thermo Scientific) were used to create a standard calibration curve. Accuracy and precision of the quantification was checked by including samples of potassium nitrate throughout the sequence.

### **pH measurements of oocyte lumen**

The pH stabilization was performed as described previously<sup>34</sup>. pH-electrodes were pulled from borosilicate glass capillaries (KWIK-FIL TW F120-3 with filament) on a vertical puller (Narishige Scientific Instrument Lab), baked for 120 min at 220 °C and silanized for 60 min with dimethyldichlorosilane (Silanization Solution I, Sigma Aldrich). Electrodes were backfilled with a buffer containing 40 mM KH<sub>2</sub>PO<sub>4</sub>, 23 mM NaOH and 150 mM NaCl (pH 7.5). The electrode tip was filled with a proton-selective ionophore cocktail (hydrogen ionophore I cocktail A, Sigma-Aldrich) by dipping the tip into the cocktail. Oocytes, as described above, were placed in freshly made HEPES-based ekulori (2 mM LaCl<sub>3</sub>, 90 mM NaCl, 1 mM KCl, 1 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 5 mM HEPES pH 7.4) for at least 30 min prior to three-electrode voltage clamp experiments. Before each oocyte a pH calibration curve was made for each oocyte using 100 mM KCl pH 5.5, 100 mM KCl pH 6.5 and 100 mM KCl pH 7.5. Oocytes were clamped at 0 mV and perfused with HEPES-based ekulori pH 7.4, followed by MES-based ekulori (2 mM LaCl<sub>3</sub>, 90 mM NaCl, 1 mM KCl, 1 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 5 mM MES pH 5) and internal pH response was measured continuously as a function of external pH change.

### **Membrane potential measurements**

Membrane potentials of oocytes were measured using the automated two-electrode voltage clamp system, Roboocyte2 (Multi channel systems), in ekulori (90 mM NaCl, 1 mM KCl, 1 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 5 mM MES, 2 mM LaCl<sub>3</sub>, pH 5) with electrodes backfilled with 1 M KCl and 1.5 M potassium acetate. All oocytes were measured using the same electrodes with a resistance of 280-350 kΩ. The experiment was terminated when the resistance of one of the electrodes shifted to approximately 600 kΩ.

### **Two-electrode voltage clamp electrophysiology**

The electric signal elicited by GA treatment of oocytes was measured using the automated two-electrode voltage clamp system Roboocyte2 (Multi channel systems), in ekulori (90 mM NaCl, 1 mM KCl, 1 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 5 mM MES, 2 mM LaCl<sub>3</sub> pH 4.5 or pH 5) with electrodes (resistance 280-1000 kΩ) backfilled with 1 M KCl and 1.5 M potassium acetate. Oocytes were clamped at – 60 mV, and IV curves were obtained before and after substrate addition. Substrate dependent currents were calculated by subtracting currents before addition of substrate from currents after addition of substrate.

### **Root hormone quantification**

Hormone extraction and analysis was performed as described in Zhang et al., 2021. Standards (both labelled and non-labelled) were obtained from Olchemim Ltd. (Olomouc,

Czech Republic) and National Research Council (NRC-CNRC, Canada). Standard grade solvents were used for sample preparation, Methanol, Acetic acid (LiChrosolv, Sigma-Aldrich, USA), Acetonitrile (J.T.Baker, Avantor, PA, USA), Formic acid (Honeywell Fluka, Thermo Fisher Scientific, MA, USA) and de-ionized water (Milli-Q, Synergy-UV millipore system, USA). Briefly, root tissue frozen in liquid nitrogen was grounded using motor and pestle. Around 200 mg of root sample was measured from ground powder and extracted with ice cold methanol/water/formic acid (15/4/1 v/v/v) added with deuterium labelled internal standards (IS). Similar concentrations of IS of abscisic acid and gibberellin (GA<sub>4</sub>) were added into samples and calibration standards. The samples were purified using Oasis MCX SPE cartridges (Waters, USA) according to manufacturer's protocol. The samples were injected on Acquity UPLC BEH C18 column (1.7 µm, 2.1x100 mm, Waters; with gradients of 0.1% acetic acid in water or acetonitrile), connected to Acquity UPLC H class system (with Waters Acquity QSM, FNR sample manager and PDA) coupled with UPLC-ESI-MS/MS triple quadrupole mass spectrometer (Xevo TQ-S, Waters, equipped with ESI probe) for identification and quantification of hormones. The hormones were measured using MS detector, both in positive and negative mode, with two MRM transitions for each compound. External calibration curves were constructed with hormone standards added with IS, used for quantification, and calculated through Target Lynx (v4.1; Waters) software by comparing the ratios of MRM peak areas of analyte to that of internal standard.

### Phloem extract and hormone quantification

Rosette leaves of 5-week-old Col-0 and *npf2-12 npf2-13* mutant plants (before bolting) were cut with a razor blade at the base of the petiole, and each leaf was dipped in a tube containing 80 µL of exudation buffer (50 mM potassium phosphate buffer, pH7.6, 10 mM EDTA). Exudation was carried out for 3h in dark in high humidity to limit transpiration. Exudation of 75 leaves was regrouped and concentrated under vacuum centrifugation. Hormone contents in phloem exudates were determined by UPLC system-MS/MS (Waters Quattro Premier XE). Concentrated residue of phloem sap was resuspended with 80% methanol-1% acetic acid including 17-<sup>2</sup>H<sub>2</sub>-labeled GA internal standards (Olchemim), mixed and passed through an Oasis HLB column. The dried eluate was dissolved in 5% acetonitrile-1% acetic acid, and the GAs were separated by UPHL chromatography (Accucore RP-MS column 2.6 µm, 100 x 2.1 mm; ThermoFisher Scientific) with a 5 to 50% acetonitrile gradient containing 0.05% acetic acid, at 400 µL/min over 22 min. The concentrations of GAs in the extracts were analyzed with a Q-Exactive mass spectrometer (Orbitrap detector; ThermoFisher Scientific) by targeted SIM using embedded calibration curves and the Xcalibur 2.2 SP1 build 48 and TraceFinder programs.

### Grafting assays

Grafting was performed without collars on water imbibed 0.45 µM MCE membrane (Millipore) between hypocotyls of rootstocks and scions of 6-day-old seedlings grown on 1x MS agar plate. Grafted seedlings were then kept vertically to recover, for 5 days under constant humidity. Successful grafts were transferred onto ½x MS agar plates and grown under a 16h photoperiod at 22°C. Root growth was measured every day for 3 days with ImageJ (<https://imagej.nih.gov/ij/download.html>). Nile red suberin staining fluorescence

intensity was assessed as previously described, in roots of 13-day-old grafted seedlings, two days after transfer onto ½x MS agar plates.

### DELLA degradation assays

12-day-old seedlings were transferred to 1x MS agar modified medium without nitrogen (bioWORLD plant media) supplemented with 0.5 mM KNO<sub>3</sub> and 1 μM paclobutrazol (Sigma). 4 days after transfer, a drop of GA<sub>12</sub> (5 μl at 1 μM) was placed on one of the first two leaves formed. Roots were collected 6, 12 and 24h after adding GA<sub>12</sub>. Total proteins were extracted in 2x SDS-PAGE sample buffer and separated on 10% SDS-PAGE gel. After transfer onto membrane, immunoblots were performed using a 2000-fold dilution anti-RGA (Agriser) and a 10000-fold dilution of peroxidase-conjugated goat anti-rabbit (Thermo Fisher Scientific). Signals were detected with Fusion FX (Vilber) using Immobilon Forte Western HRP Substrate (Millipore). The blot was subsequently stained with Coomassie blue. Quantification of the signals was determined using ImageJ package.

### Mathematical model

Root templates were segmented from an experimental image using the CellSeT image analysis tool <sup>73</sup> (Sup. Fig. 20). We used CellSeT to manually assign a cell type to each cell and then read the geometrical and cell-type data into a tissue database (based on the OpenAlea tissue structure <sup>74</sup>), extending the data structure to incorporate vacuolar compartments within each cell. The geometrical, topological and transporter-distribution data were used to form a system of ordinary differential equations (ODEs) to describe the GA transport, synthesis and degradation within the multicellular root cross-section. Parameters associated with the passive and transporter-mediated transport components were estimated using the oocyte data (Fig. 1a, Fig. 3b) and the remaining parameter values were obtained from the literature (Sup. Table 5). These ODEs were simulated using the `solve_ivp` package in python 3.6.5. Full details of the model equations and assumptions are provided as Supplementary text.

### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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## Data and materials availability

All the data supporting the findings of this study are available within the article and the Supplementary Materials.

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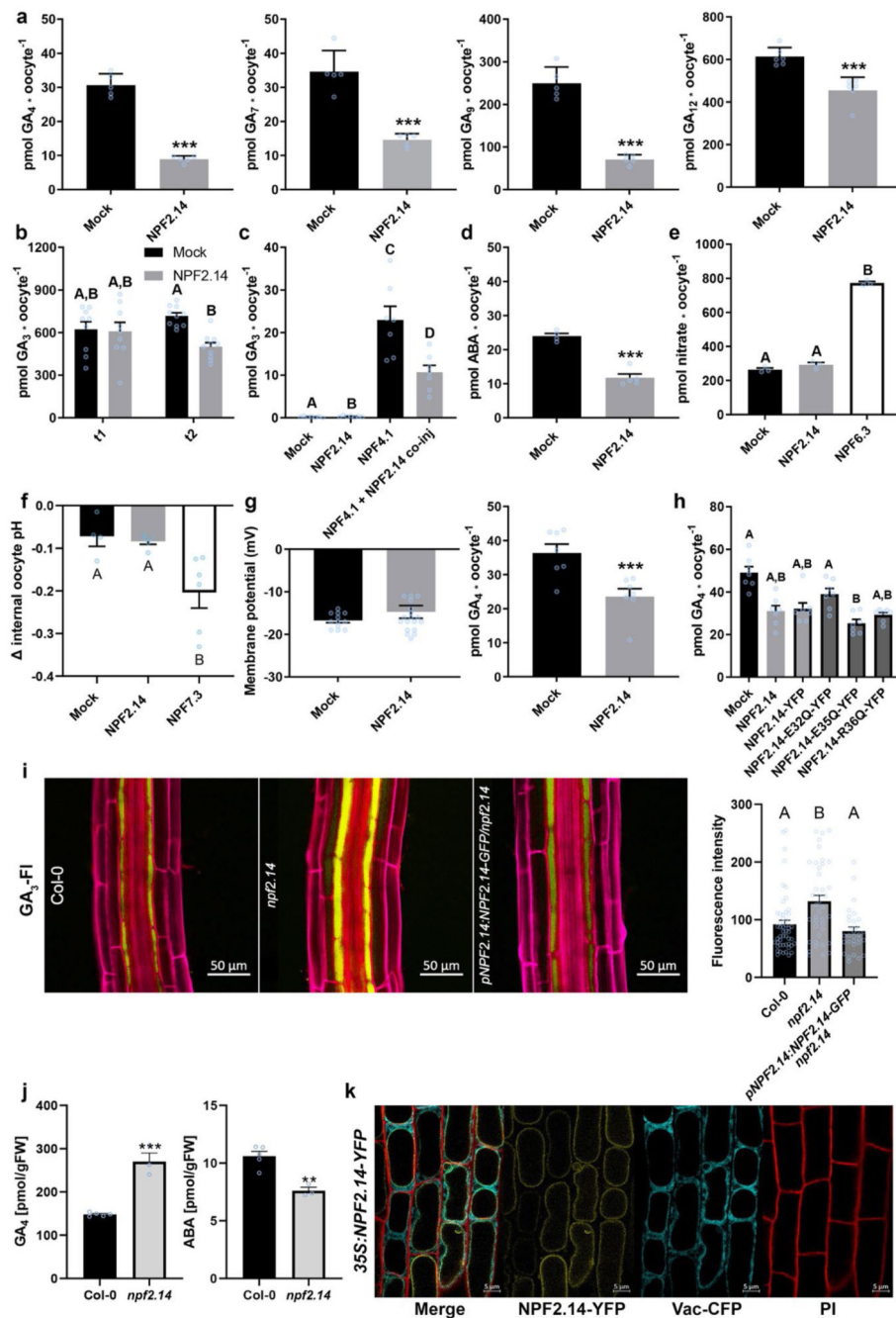
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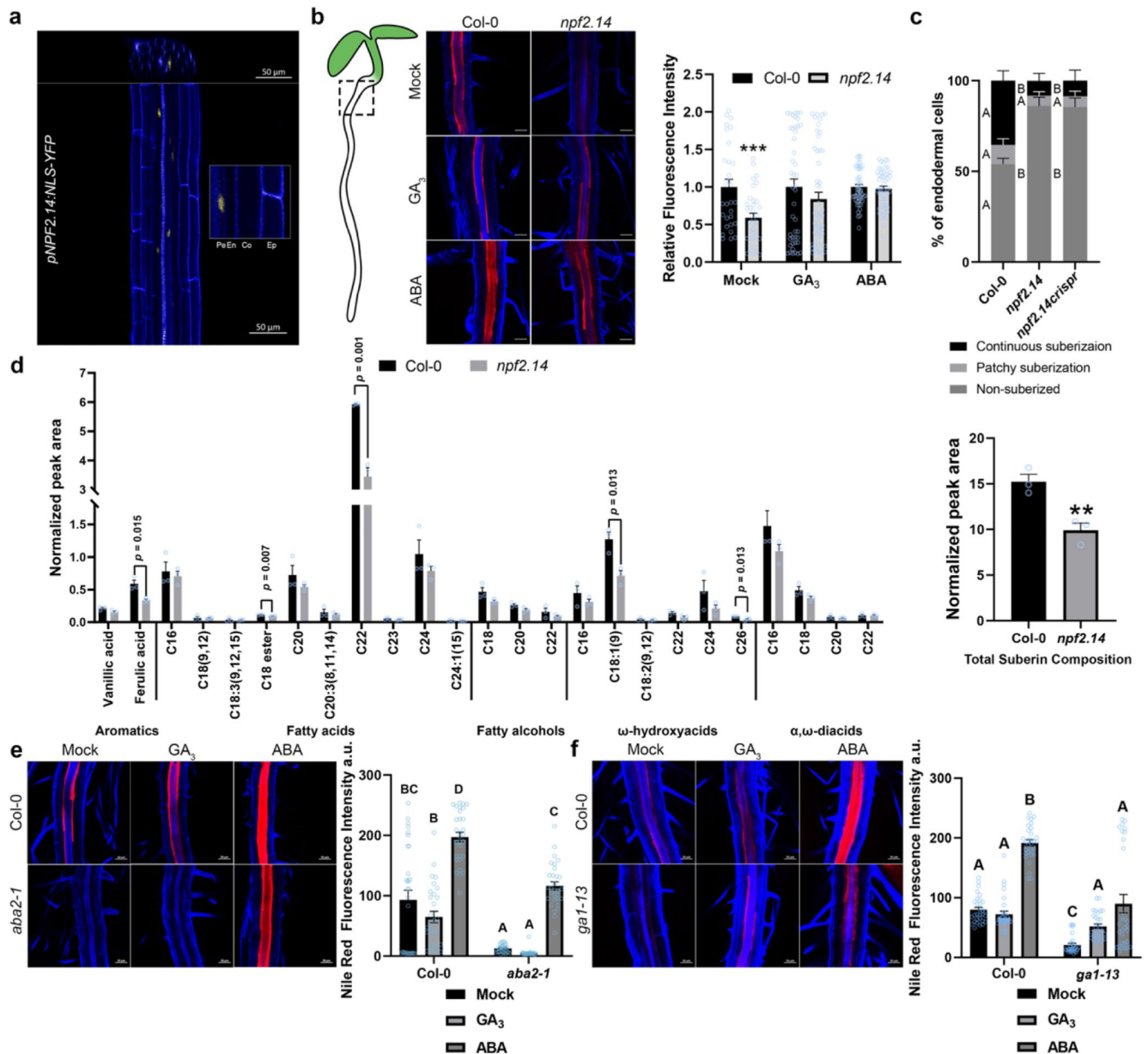
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**Fig. 1. NPF2.14 is a vacuolar GA and ABA transporter.**

(a) GA accumulation in NPF2.14-expressing and control oocytes exposed to the indicated GAs at 50 μM concentration for 60 min at room temperature at pH 5 for GA<sub>4</sub> ( $p = 0.0031$ ) and GA<sub>7</sub> ( $p = 0.001$ ) and at pH 6 for GA<sub>9</sub> ( $p < 0.0001$ ) and GA<sub>12</sub> ( $p = 0.0004$ ). Mean + S.E. ( $n = 5$  for GA<sub>4</sub>, GA<sub>7</sub>, and GA<sub>9</sub> and  $n = 6$  for GA<sub>12</sub>). Statistical significance was evaluated by two-tailed t-tests (b) GA accumulation at 20 h after direct injection of 23 nl of 8.2 mM membrane impermeable GA<sub>3</sub> at pH 7.4 at 16° C (t1) and after 60 min at room temperature at pH 5 (t2). Mean + S.E. ( $n = 6$  single oocytes). Statistical significance

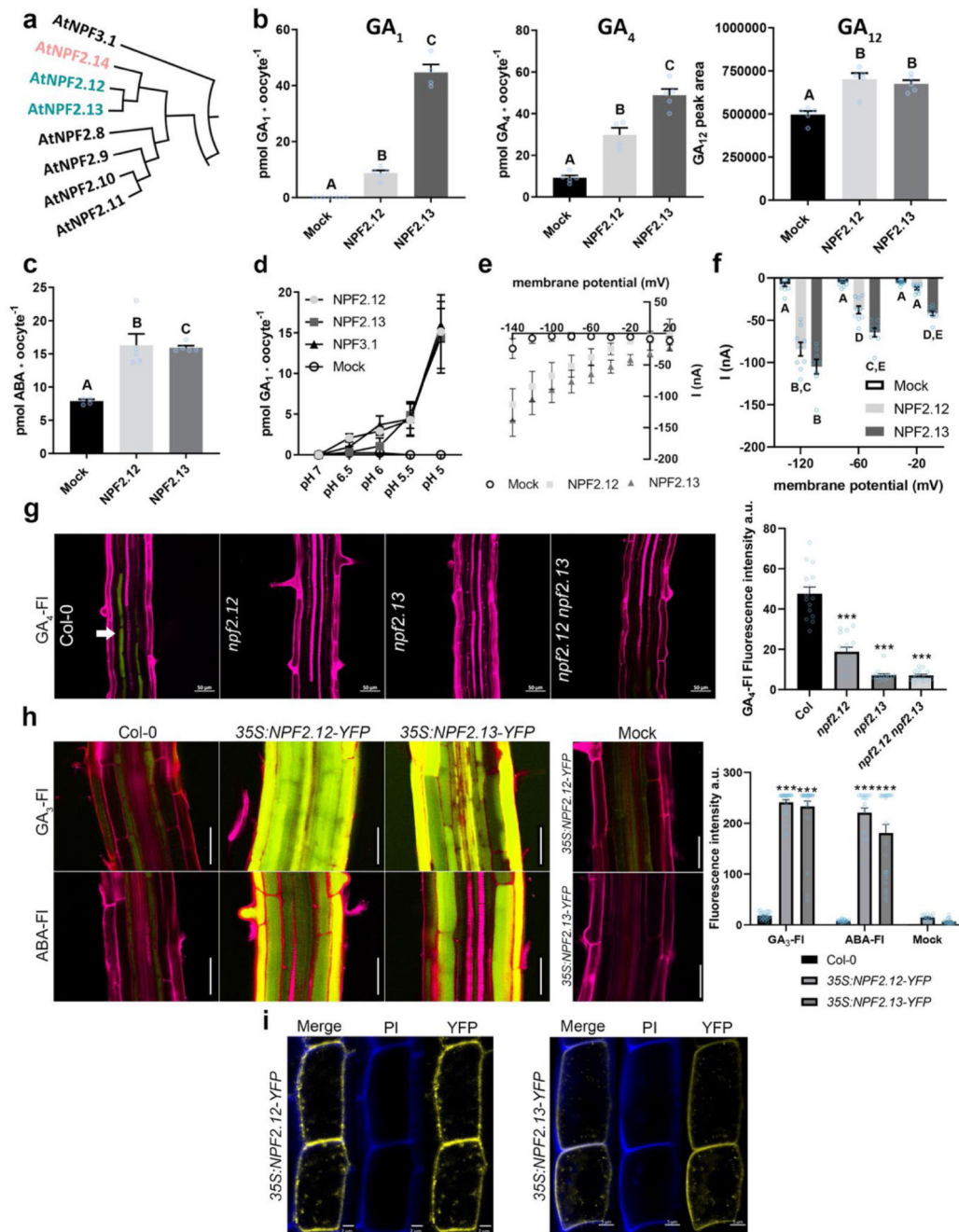
was determined by a two-way ANOVA with a Holm-Sidak post hoc test to adjust for multiple comparisons ( $p < 0.05$ ). (c) GA accumulation in control oocytes or oocytes that express NPF2.14, NPF4.1, or both proteins exposed to 50  $\mu\text{M}$  GA<sub>3</sub> at pH 5 for 60 min at room temperature and analyzed by LC-MS/MS. Mean + S.E. ( $n = 7$  single oocytes). Statistical significance was assessed using a Holm Sidak one-way ANOVA ( $p = 0.05$ ). (d) ABA accumulation in oocytes exposed to 50  $\mu\text{M}$  ABA at pH 5 for 60 min at room temperature ( $n = 5$  single oocytes). Statistical significance was determined by two-tailed t-test ( $p = 0.05$ ). (e) Nitrate accumulation in oocytes exposed to 5 mM nitrate at pH 5 at room temperature for 60 min. Mean + S.E. (three replicates of 5 oocytes analyzed by analytical anion chromatography,  $n = 3$ ). Statistical significance was assessed using a Holm Sidak one-way ANOVA ( $p = 0.05$ ). (f) Internal oocyte pH measured using three-electrode voltage clamp electrophysiology of control oocytes and NPF2.14- and NPF7.3-expressing oocytes. Oocytes were perfused at pH 7.4 for 5 min followed by perfusion at pH 5 for 5 min. Mean + S.E. ( $n = 4$  oocytes for Mock and NPF2.14, 6 for NPF7.3). Statistical significance was assessed using Holm Sidak one-way ANOVA ( $p = 0.05$ ). (g) Membrane potentials of control oocytes ( $n = 12$ ) and NPF2.14-expressing oocytes ( $n = 19$ ) measured at pH 5 using two-electrode voltage clamp electrophysiology. Oocytes with measured membrane potential were exposed to 50  $\mu\text{M}$  GA<sub>4</sub> at pH 5 ( $n = 7$  single oocytes) for 60 min at room temperature and analyzed by LC-MS/MS. Mean + S.E. Assessed using two-tailed t-tests ( $*** p < 0.001$ ). (h) GA accumulation in control oocytes or oocytes that express NPF2.14 with wild-type or mutant ExxE[K/R] motifs exposed to 50  $\mu\text{M}$  GA<sub>4</sub> at pH 5 for 60 min at room temperature and analyzed by LC-MS/MS ( $n = 7$  oocytes). Mean + S.E. Statistical significance was assessed using a Holm Sidak one-way ANOVA ( $p = 0.05$ ). (i) Left: Representative images of 6-day-old *npf2.14* mutant and *npf2.14* complementation lines. Roots were treated with 5  $\mu\text{M}$  GA<sub>3</sub>-Fl (Yellow); propidium iodide (PI, pink). Right: GA<sub>3</sub>-Fl fluorescence intensity in the endodermis, mean  $\pm$  S.E (5 endodermal cells sampled from each root,  $n = 50$  for Col-0, 40 for *npf2.14* and 30 for *pNPF2.14:NPF2.14-GFP/npf2.14*). (j) GA<sub>4</sub> ( $p = 0.0002$ ) and ABA content ( $p = 0.0024$ ) in 10-day-old roots of control and *npf2.14*-mutant plants measured using LC-MS. Each replicate is 200 mg of pooled *Arabidopsis* roots. Mean + S.E. ( $n = 5$ ). Significance was evaluated using a two-tailed Student's t-test. (k) Representative confocal image of 6-day-old root epidermis *35S:NPF2.14-YFP* cells stained with PI (red) and tonoplast marker Vac-CFP (cyan)<sup>1</sup>. Experiment was repeated 3 times with similar results.



**Fig. 2. *NPF2.14* expression in the differentiated pericycle is required for endodermal root suberization.**

(a) Confocal image of 6-day-old *pNPF2.14:NLS-YFP* roots stained with PI (blue) and imaged for YFP (yellow). Inset is magnification of cells with *pNPF2.14:NLS-YFP* signal. Pe – Pericycle, En – Endodermis, Co – Cortex, Ep – Epidermis. Experiment was repeated 3 times with similar results. (b) Left: Images of 5-day-old Col-0 and *npf2.14* mutant roots supplemented with mock solution, 5  $\mu$ M GA<sub>3</sub>, or 1  $\mu$ M ABA and stained with Nile red (red) and calcofluor (blue). Right: Quantification of Nile red fluorescence intensity, normalized to Col-0. Mean + S.E. Illustration indicates imaged area. Fluorescence intensity quantified from 5 endodermal cells per root, minimum of 6 roots per treatment (n = 30 for Mock treated Col-0, 35 for Mock treated *npf2.14* and GA<sub>3</sub> treated Col-0. Rest is 50). Statistical

significance was assessed using a 2-way ANOVA followed by a Sidaks's ad-hoc multiple comparisons test (\*\*\*)  $p = 0.003$ . (c) Quantifications of suberin pattern along the root using Nile red fluorescence intensity. Mean + S.E.  $n = 10$  individual plants. Statistical significance determined using a 2-way ANOVA with a Dunnett's multiple comparisons ad hoc test. Different letters indicate significant differences ( $p < 0.0001$ ). (d) Root suberin profile for 10-day-old plants, measured by GC-MS. The  $y$ -axis represents relative peak areas following normalization to a C32-alkane internal standard. Right graph indicates total suberin composition. Mean  $\pm$  S.E. ( $n = 3$ ). Statistical significance determined by a two-tailed Student's  $t$ -test. Total suberin composition calculated by summing up all monomers ( $p = 0.0092$ ) (e) Left: Images of 5-day-old Col-0 and *aba2-1* roots from plants grown on mock MS or MS with 1  $\mu$ M ABA. Roots were stained with Nile red (red) and calcofluor (blue). Right: Fluorescent intensity was quantified from 6 roots per treatment, 5 endodermal cells per root. Mean + S.E.  $n = 30$  individual cells. Significance was determined using a 2-way ANOVA with a Tukey's multiple comparisons ad-hoc test. Different letters mark significant difference ( $p < 0.001$ ). (f) Left: Images of 5-day-old Col-0 and *gal-13* mutant roots grown on mock MS or MS with 5  $\mu$ M GA<sub>3</sub>. Roots were stained with Nile red (red) and calcofluor (blue). Right: Fluorescent intensity was quantified from 6 roots per treatment, 5 endodermal cells per root. Mean + S.E.  $n = 30$  individual cells. Significance was determined using a 2-way ANOVA with a Tukey's multiple comparisons ad-hoc test. Different letters mark significant difference ( $p < 0.05$ ).

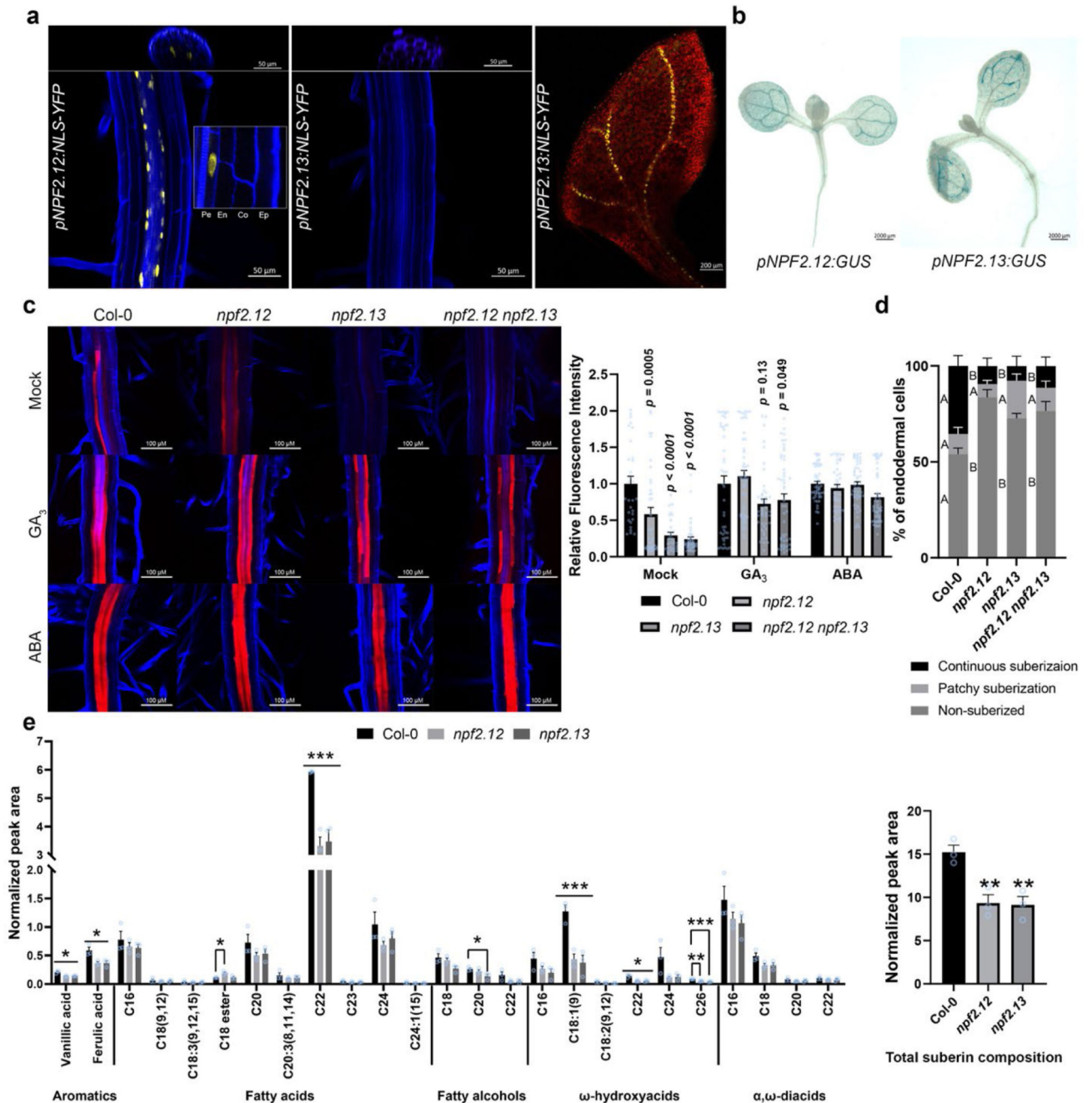


**Fig. 3. NPF2.12 and NPF2.13 are plasma membrane-localized GA and ABA importers that facilitate endodermal hormone accumulation.**

(a) Phylogenetic clade of *NPF2.12*, *NPF2.13*, *NPF2.14*, and their close paralogues. (b and c) Hormone uptake in control and *NPF2.12*- and *NPF2.13*-expressing *Xenopus* oocytes. Mean + S.E. (b) Oocytes were exposed to 50 μM GA<sub>1</sub> at pH 5 (n = 5 single oocytes), 50 μM GA<sub>4</sub> at pH 5.5 (n = 5), or 50 μM GA<sub>12</sub> at pH 6 (n = 5). (c) Oocytes were exposed to 50 μM ABA at pH 5 (n = 5). Hormone uptake analyzed by LC-MS/MS. Statistical significance assessed using Holm Sidak one-way ANOVA ( $p = 0.05$ ). (d) Oocytes were

exposed to 50  $\mu\text{M}$  GA1 at pH ranging from 5 to 7. Mean + S.E. (n = 5). **(e and f)** Mean + S.E. of control oocytes (n = 10), oocytes expressing NPF2.12 (n = 9), and oocytes expressing NPF2.13 (n = 8) exposed to 500  $\mu\text{M}$  GA<sub>3</sub> at pH 5, currents were measured using two-electrode voltage clamp electrophysiology over a range of membrane potentials from +20 to -140 mV. Statistical significance assessed using Holm Sidak two-way ANOVA ( $p = 0.01$ ). **(g)** Left: Representative images of roots of 6-day-old Col-0 and single and double *npf2.12* and *npf2.13* knockout plants treated with 5  $\mu\text{M}$  GA<sub>4</sub>-Fl (Yellow) overnight. Pink indicates PI. White arrow indicates GA<sub>4</sub>-Fl signal. Right: Quantification of GA<sub>4</sub>-Fl fluorescence intensity in the endodermis, mean  $\pm$  S.E. (5 endodermal cells per 3 biological repeats, n = 15). Significance was determined by 2-way ANOVA with an ad-hoc Dunnett's multiple comparisons test (\*\* $p < 0.0001$ ). **(h)** Left: Representative images of roots of 6-day-old *35S:NPF2.12-YFP* and *35S:NPF2.13-YFP* plants treated with 5  $\mu\text{M}$  GA<sub>3</sub>-Fl and ABA-Fl for 2 hours. Pink indicates PI, yellow indicates GA<sub>3</sub>-Fl or ABA-Fl. Right: GA<sub>3</sub>-Fl and ABA-Fl fluorescence intensity quantified in the epidermal cells, mean  $\pm$  S.E. (5 epidermal cells sampled from at least 4 biological repeats, n > 20). *35S:NPF2.12-YFP* and *35S:NPF2.13-YFP* mock do not show fluorescence under these confocal gain settings. Statistical significance assessed using a 2-way ANOVA followed by a Dunnett's multiple comparisons test (\*\* $p < 0.0001$ ). **(i)** Confocal imaging of 6-day old root meristem epidermis cells expressing *35S:NPF2.12-YFP* and *35S:NPF2.13-YFP*. Blue is PI, yellow is YFP. Experiment was repeated 3 times with similar results.





**Fig. 4. *NPF2.12* and *NPF2.13* regulate root endodermis suberization.**

(a) Confocal imaging of 6-day-old *pNPF2.12:NLS-YFP* and *pNPF2.13:NLS-YFP* seedlings.

Inset is magnification of cells with *pNPF2.12:NLS-YFP* signal. Pe – Pericycle, En –

Endodermis, Co – Cortex, Ep – Epidermis. Blue indicates either calcofluor, yellow

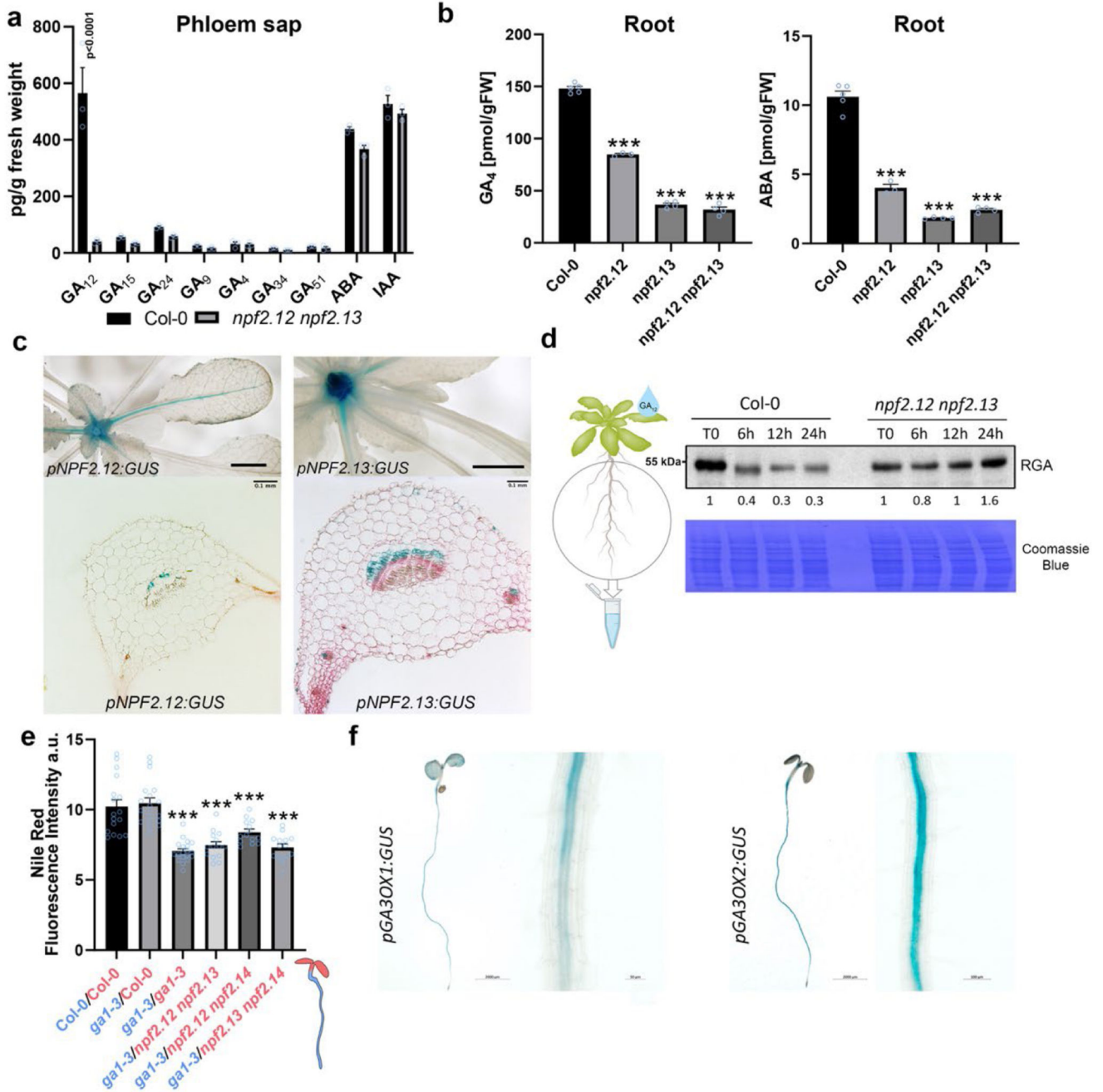
indicates YFP. Right image is *pNPF2.13:NLS-YFP* cotyledon (red indicates chlorophyll).

Experiments were repeated 3 times with similar results. (b) GUS staining of 7-day-old

seedlings expressing *pNPF2.12:GUS* and *pNPF2.13:GUS*. (c) Left: Nile red-stained 5-day-old

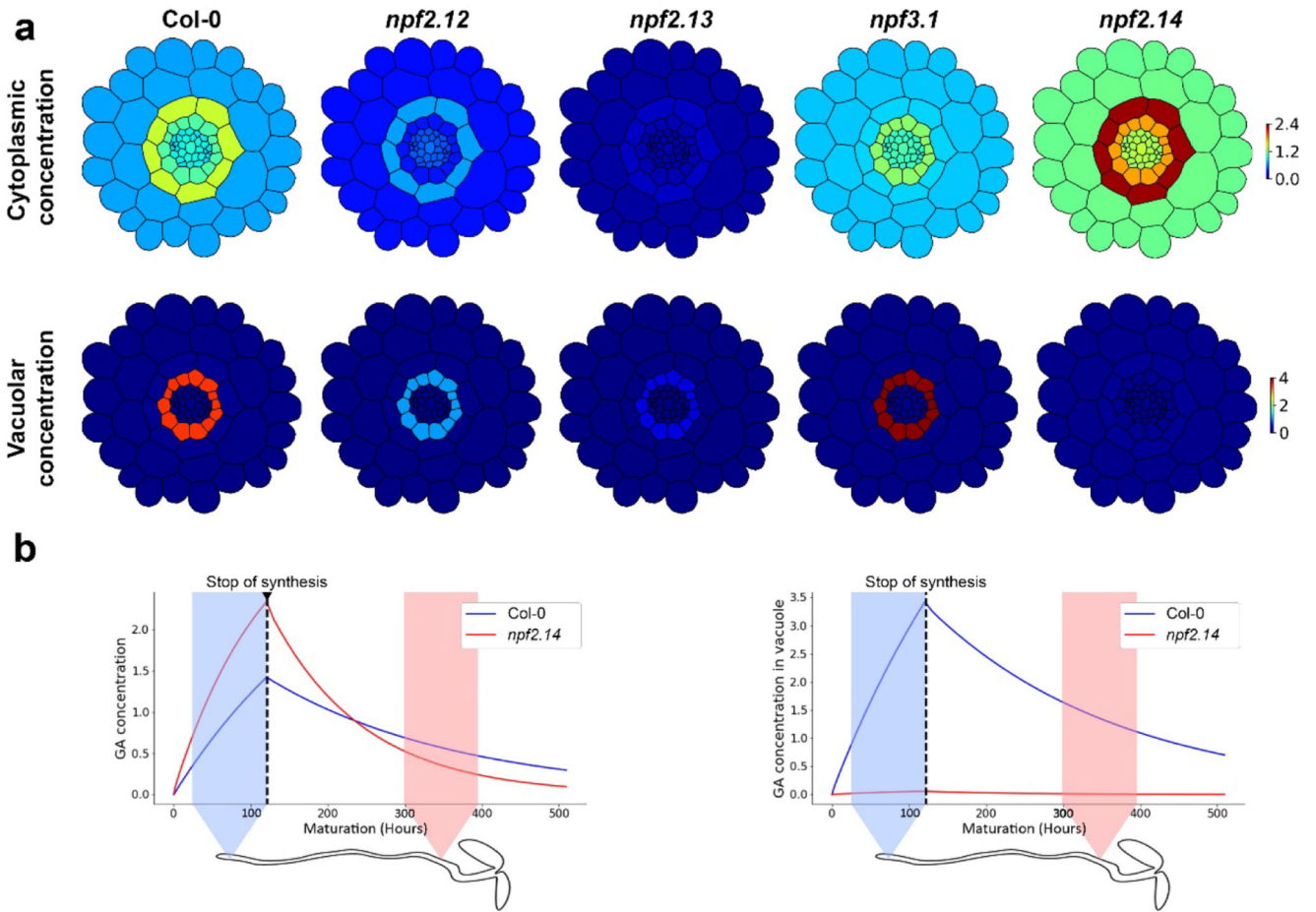
*npf* mutant roots mock treated or treated with 5  $\mu$ M GA<sub>3</sub> or 1  $\mu$ M ABA. Right:

Fluorescent intensity relative to Col-0 quantified from a minimum of 6 roots per treatment, 5 endodermal cells per root ( $n = 30$  for Mock treated Col-0 and *npf2.13*, 40 For Mock and ABA treated *npf2.12* 45 for GA<sub>3</sub> treated Col-0 and *npf2.13*, rest are 50). Statistical significance was assessed using a 2-way ANOVA followed by a Dunnett's multiple comparisons test. **(d)** Quantifications of suberin pattern along the root using Nile red fluorescence intensity. Mean + S.E.  $n = 10$  individual roots. Statistical significance determined using a 2-way ANOVA with a Tukey's multiple comparisons ad hoc test. Different letters indicate significant differences ( $p < 0.005$ ). Different letters indicate significant differences between conditions ( $p < 0.05$ ). **(e)** Root suberin profile for 10-day-old plants, measured by gas chromatography-mass spectrometry. The  $x$ -axis represents relative peak areas following normalization to a C32-alkane internal standard. Right graph indicates for total suberin composition. Data in bars represent the means  $\pm$  SE of three biological replicates. \* indicates statistically significant differences compared with Col-0 at  $p = 0.015$ , \*\*  $p = 0.01$  and \*\*\*  $p = 0.001$  by a two-tailed Student's t-test. Total suberin composition significance assessed using a one-way ANOVA with a Dunnett's multiple comparisons test ( $p = 0.0072$  for *npf2.12*,  $p = 0.006$  for *npf2.13*).



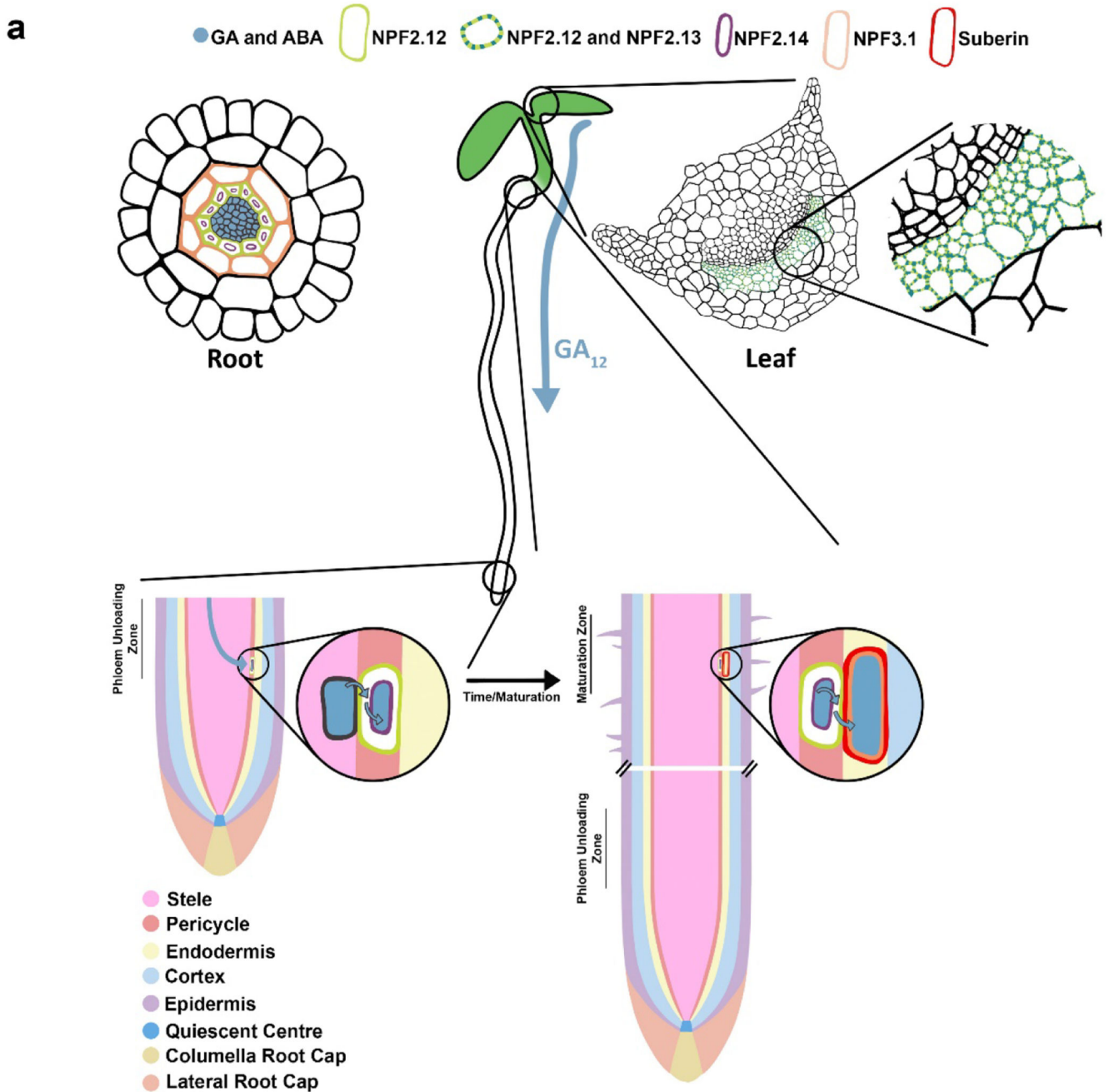
**Fig. 5. NPF2.12 and NPF2.13 facilitate long-distance shoot-to-root GA transport.** (a) Quantification of GA, ABA and IAA contents (in  $\text{pg}\cdot\text{g}^{-1}$  fresh weight) in phloem exudates collected from leaf petiole of 5-week-old Col-0 and *npf2.12 npf2.13* mutant plants. Mean + S.E. Exudate was collected from 75 leaves, pooled into 3 separate biological repetitions ( $n = 3$ ). Statistical significance evaluated by a one-way ANOVA followed by a Dunnett's multiple comparisons test, comparing to Col as control. ( $p = 0.0009$  when comparing Col-0/Col-0 to *ga1-3/npf2.12 npf2.14*,  $p < 0.0001$  for the rest of \*\*\* marked lines) (b) GA<sub>4</sub> and ABA quantification (in  $\text{pmol}\cdot\text{g}^{-1}$  fresh weight) in 10-day-old roots of

Col-0 and *npf2.12 npf2.13* doublemutant plants. Mean + S.E. (n = 5). Significance was assessed using a 2-way ANOVA followed by a Dunnett's multiple comparisons test (\*\*\*)  $p < 0.0001$ ). (c) GUS staining patterns in rosette leaves of 4 week-old *pNFP2.12:GUS* and *pNPF2.13:GUS* plants. Top images are whole plants; scale bars represent 0.25 cm. Bottom images are cross-sections of GUS-stained leaf petioles; scale bars represent 0.1 mm. (d) Immunodetection of RGA protein accumulation in the root of 16-day-old Col-0 and *npf2.12 npf2.13* mutant plants. The plants were grown *in vitro* for 12 days and then moved to MS plates containing a low concentration of nitrogen (0.5 mM KNO<sub>3</sub>) and paclobutratol (PAC, 1 μM). 4 days after transfer, a drop of GA<sub>12</sub> (5 μl at 1 μM) was added to one of the first two leaves formed. Proteins were extracted from the root at 4-time points following GA<sub>12</sub> application. Similar results were obtained in two independent experiments. (e) Quantification of Nile red fluorescence intensity (mean ± SEM) of various combinations of grafted seedlings. Blue font indicates for rootstock genotype, red indicates for grafted scion. Fluorescence intensity was quantified from a 5 roots per genotype/grafting, at least 3 endodermal cells per root, n = 17 for Col-0/Col-0, 18 *gal-3*/Col-0, 19 for *gal-13/gal-13* and 14 for the rest). Statistical significance was assessed using a 2-way ANOVA followed by a Dunnett's multiple comparisons test (\*\*\*)  $p < 0.001$ . (f) Images of 6-day-old plants that express GUS driven by the *GA<sub>3</sub>OXI-2* promoters. Experiment was repeated 3 times with similar results.



**Fig. 6. NPF transporters mediate pericycle-specific hormone uptake into the vacuoles at the phloem unloading zone to facilitate a hormone slow-release mechanism that allows suberization at the maturation zone.**

(a) Spatial distributions of cytoplasmic and vacuolar GA concentration in the root cross-section, predicted by the multicellular mathematical model, for the wild type, and the *npf2.12*, *npf2.13*, *npf3.1*, and *npf2.14* mutants. The time of the simulations is 5 days (120 hours), corresponding to the age of the plants used in the experiments. (b) Predicted dynamics of the endodermal cytoplasmic and pericycle vacuolar GA concentration for the wild type and the *npf2.14* mutant. The model is initially simulated with a constant GA source in the stele (pale blue region), then synthesis is set to zero to simulate the GA redistribution after cells leave the phloem unloading zone. The model predicts that NPF2.14 leads to higher endodermal cytoplasmic concentrations where suberin forms (pale red region).



**Fig. 7. Proposed model illustrating NPF2.12, NPF2.13, NPF2.14 and NPF3.1 function in regulating endodermal suberin formation.**

Proposed model incorporating experimentally observed distributions of root GA transporters, hormone accumulation, endodermal suberin formation and mathematical model predictions. NPF2.12 and NPF2.13, which are localized to the shoot phloem, are both required for  $GA_{12}$  long-distance shoot-to-root translocation. NPF2.12 is expressed in the root pericycle cell membranes and promotes the movement of ABA and GA from the vasculature to the pericycle. Once in the pericycle cytoplasm, NPF2.14 imports the hormones into the vacuole to form a reservoir which will be available in later stages. When

the root elongates over time and the cells that accumulated high levels of GA and ABA in the vacuoles mature, the hormones are exported out of the pericycle vacuole and imported into the endodermis by NPF3.1 to induce suberization.