

SCHENGEN receptor module drives localized ROS production and lignification in plant roots.

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ROS production by localized SCHENGEN receptor module drives lignification at 1 subcellular precision 2

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4 Authors: Satoshi Fujita^{1, *}, Damien De Bellis^{1,2}, Kai H. Edel³, Philipp Köster^{3, †}, Tonni Grube Andersen^{1, ‡}, Emanuel Schmid-Siegert⁴, Valérie Dénervaud Tendon¹, Alexandre Pfister¹, Peter 5

Marhavý^{1, ††}, Robertas Ursache¹, Verónica G. Doblas^{1, §}, Marie Barberon^{1, §§}, Jean Daraspe^{1,2}, 6

Audrey Creff⁵, Gwyneth Ingram⁵, Jörg Kudla³ and Niko Geldner^{1,*} 7

Affiliations: 8

- 9 1 Department of Plant Molecular Biology, Biophore, Campus UNIL-Sorge, University of Lausanne, CH-1015
- 10 Lausanne, Switzerland
- 11 2 Electron Microscopy Facility, University of Lausanne, 1015 Lausanne, Switzerland
- 12 3 Institut für Biologie und Biotechnologie der Pflanzen, Westfälische Wilhelms-Universität Münster, Schlossplatz 7, 13 Münster 48149, Germany
- 4 Vital-IT Competence Center, Swiss Institute of Bioinformatics, Lausanne, Switzerland 14
- 5 Laboratoire Reproduction et Développement des Plantes, Univ Lyon, ENS de Lyon, CNRS, INRAE, Lyon, France 15 16

Present address: National Institute of Genetics, 1111 Yata, Mishima, Shizuoka 411-8540, Japan

17 † Present address: Department of Plant and Microbial Biology, University of Zurich, Zollikerstrasse 107, CH-8008 18 19 Zurich, Switzerland

- ^{††}Present address: Umeå Plant Science Centre (UPSC), Department of Forest Genetics and Plant Physiology, 20
- Swedish University of Agricultural Sciences (SLU), 90183, Umeå, Sweden. 21
- 22 [±] Present address: Max Planck Institute for Plant Breeding Research, 50829 Cologne, Germany
- 23 § Present address: Institut Jean-Pierre Bourgin, INRAE, AgroParisTech, Université Paris-Saclay, 78000, Versailles, 24 France
- §§ Present address: Department of Botany and Plant Biology Quai Ernest-Ansermet 30 CH-1211 Geneva 25 Switzerland 26 27
- *Correspondence to: satoshi.fujita@nig.ac.jp (S.F.) and niko.geldner@unil.ch (N.G.) 28

Abstract: 29

Production of reactive-oxygen species (ROS) by NADPH oxidases (NOXs) impacts many 30 processes in animals and plants and many plant receptor pathways involve rapid, NOX-dependent 31 increases of ROS. Yet, their general reactivity has made it challenging to pinpoint the precise role 32 and immediate molecular action of ROS. A well-understood ROS action in plants is to provide the 33 co-substrate for lignin peroxidases in the cell wall. Lignin can be deposited with exquisite spatial 34 control, but the underlying mechanisms have remained elusive. Here we establish a kinase 35 signaling relay that exerts direct, spatial control over ROS production and lignification within the 36 cell wall. We show that polar localization of a single kinase component is crucial for pathway 37 function. Our data indicates that an intersection of more broadly localized components allows for 38 micrometer-scale precision of lignification and that this system is triggered through initiation of 39 ROS production as a critical peroxidase co-substrate. 40

As in animals, NADPH oxidase-produced ROS in plants is important for a multitude of processes and 41 42 the number of NADPH oxidase genes (10 in Arabidopsis, called RESPIRATORY BURST OXIDASE HOMOLOGs, RBOHs, A-J) suggests a high complexity of regulation of ROS production in plants. Among its 43 44 many roles, ROS-dependent regulation of plant cell wall structure and function is considered to be among its 45 most important (Kärkönen & Kuchitsu, 2015). The cell wall is the nano-structured, sugar-based, pressureresisting extracellular matrix of plants and NOXs are thought to be the predominant ROS source in this 46 compartment (also termed apoplast) (Kärkönen & Kuchitsu, 2015). 47

48 A staggering number of kinases have been shown to regulate plant NOXs and the activation mechanism 49 of NOX-dependent ROS production is well established, especially in response to microbial pattern-recognition by immune receptors (Zipfel, 2014). However, the specific role and direct molecular targets of ROS during 50 microbial pattern-recognition have remained elusive (Qi et al, 2017). The same applies to the central role of ROS 51 in tip growing cells, such as root hairs or pollen tubes, where ROS is thought to be part of an intricate oscillation 52 of cell wall stiffening and loosening, aimed at allowing cell wall expansion without catastrophic collapse 53 (Boisson-Dernier et al, 2013; Monshausen et al, 2007). In this case, ROS is proposed to be important for 54 counteracting cell wall loosening pH decreases, but it is again unclear what direct targets of ROS would mediate 55 56 cell wall stiffening. Cell wall lignification by apoplastic peroxidases, can therefore be considered as the most well-established role of ROS, where the peroxidases themselves are the direct "ROS targets", using it as a co-57 58 substrate for the oxidation of mono-lignols (Barbosa et al, 2019; Liu, 2012). In the case of lignification, however, a molecularly-defined signaling pathway that induces ROS production during lignification has not been defined. 59 A few years ago, our group identified a specific NADPH oxidase, RBOHF, to be required for the localized 60 formation of lignin in the root endodermis (Lee et al, 2013). Lignin is a poly-phenolic polymer that is generated 61 by the radical-coupling of mono-lignols, oxidized through the action of ROS-dependent peroxidases, as well as 62 laccases (Liu, 2012). The hydrophobic lignin polymer impregnates the cellulosic cell wall of plants, rendering it 63 unextendible and highly resistant to degradation. Lignin in the root endodermis is deposited in a central, 64 longitudinal band around every endodermal cell. Named Casparian strips (CS), these ring-like lignin structures 65 66 fuse into a supracellular network, establishing a tissue-wide, extracellular diffusion barrier (Fig. 1A), analogous to epithelial tight junctions in animals (Geldner, 2013; Barberon & Geldner, 2014). Functionality of this barrier 67 can be easily visualized by a block of penetration of a fluorescent cell wall dye, propidium iodide, into the 68 vasculature (Alassimone et al, 2010; Naseer et al, 2012). CS localization occurs through the action of 69 CASPARIAN STRIP DOMAIN PROTEINS (CASPs), 4-TM proteins, which form a highly scaffolded 70 71 transmembrane protein platform, assembling RBOHF and cell wall peroxidases and other proteins at the 72 Casparian strip domain (CSD) (Lee et al, 2013; Hosmani et al, 2013a; Roppolo et al, 2011). CSD formation and lignin deposition are coordinated such that the aligned rings of endodermal neighbors' fuse, leading to a 73 74 supracellular network that seals the extracellular space between endodermal cells, generating a tissue-wide 75 diffusion barrier.

76 More recently, we identified a pair of peptide ligands, a leucine-rich repeat receptor-like kinase (LRR-77 RLK) and a cytoplasmic kinase, whose phenotypes, genetic interaction and specific subcellular localizations led us to propose that they combine into a barrier surveillance pathway. Previous reports had shown that the CIF1/2 78 (CASPARIAN STRIP INTEGRITY FACTORs 1/2) peptides are SCHENGEN3 (SGN3) (also called GASSHO1 79 (GSO1)) ligands and that the SGN1 and SGN3 kinases govern Casparian strip integrity (Alassimone et al, 2016; 80 Doblas et al, 2017; Nakayama et al, 2017; Pfister et al, 2014). cifl cif2 and sgn1, sgn3 mutants have similar, 81 82 discontinuous CS, caused by a discontinuous CSD, as well as a conspicuous absence, or strong attenuation of, compensatory lignification and suberization observed in other CS mutants (Doblas et al. 2017; Hosmani et al. 83 84 2013a; Kalmbach et al, 2017; Kamiya et al, 2015; Li et al, 2017; Pfister et al, 2014). Their phenotypic similarities 85 suggested that these factors act in one pathway. CIF1 and 2 peptides do not express in the endodermis, where the CS is formed, but in the stele. In contrast, both SGN1 and SGN3 present specific localization on the 86 endodermal plasma membrane; SGN3 receptor-like kinase resides along both sides of the CS, while 87 palmitoylated SGN1 polarly localizes on cortex facing (outer) plasma membranes (Doblas et al, 2017). 88 Remarkably, their localization overlaps only in a small region next to the cortex-facing side of the CS 89 (Alassimone et al, 2016). This would require peptides from the stele to diffuse across the CS in order to access 90 91 the signaling complex. This is only possible while the CS is still permeable (Doblas et al, 2017) (Fig. 1E).

This pathway would therefore provide a mechanism that allows to probe the tissue-wide integrity of an extracellular diffusion barrier and to respond to barrier defects through compensatory over-lignification (Doblas *et al*, 2017). Here we demonstrate that the receptor, cytoplasmic kinase and NADPH oxidase molecularly connect into one pathway, with great resemblance to plant immune signaling pathways, but whose direct action is to locally produce ROS for spatially restricted lignification. We establish the crucial importance of the restricted subcellular localization of its components and demonstrate that stimulation of this signaling pathway additionally leads to strong transcriptional activation of target genes that further drive and sustain endodermal lignification, as well as suberization and endodermal sub-domain formation and differentiation. We
 thus provide a molecular circuitry in which an endogenous peptide from the stele stimulates localized signaling
 kinases and NADPH oxidases in the endodermis, causing extracellular ROS production at micrometer-scale
 precision and a precisely localized lignification of the plant cell wall.

103 **RESULTS**

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We generated a new *cif1-2 cif2-2* double mutant allele by CRISPR-Cas9 in a pure Col background, because the previous *cif1-1 cif2-1* double mutant allele (Nakayama *et al*, 2017) was a mixture of Ws and Col alleles. The new CRISPR allele was complemented by CIF1 or CIF2 application, visualized by PI (propidium iodide) uptake assays or reconstitution of CASP1 membrane domain connectivity (Fig. EV1 D,E and Appendix Fig. S1A).

Apolar SGN1 kinase leads to constitutive barrier defect signaling

Central to the barrier surveillance model is the polar localization of SGN1, which is thought to limit 111 112 the potential for signal activation to the cortex side of the endodermis, requiring passage of CIF peptides across the CS region (Fig. 1A,E). Consistently, application of peptide ligand to the media, leading to stimulation from 113 the outside, causes overlignification at the cortex-facing endodermal edges (Fig. 1B). In an attempt to falsify the 114 model we had proposed and to interrogate the importance of SGN1 polar localization, we generated a SGN1 115 variant that localized in an apolar fashion, by adding a myristoylation (myr) and palmitoylation (palm) motifs 116 on the N terminus (Vermeer et al, 2004). This myrpalm SGN1-mCitrinerine (Citrine) variant was expressed 117 under the control of the endodermis-specific CASP1 promoter, which is strongly active during Casparian strip 118 119 formation and complemented the sgn1 barrier phenotype (Fig. EV 1A). In planta, the wild-type SGN1-Citrine 120 variant resides polarly on the cortex-facing side of endodermal cells, while myrpalm SGN1-Citrine was observed at both sides of the endodermal plasma membranes, even though preferentially accumulation at the cortex side 121 122 could still be observed (Fig. 1C, 1D). Both variants were excluded from the central position where the Casparian 123 strip domain is formed (Fig. EV 1B) and the localization patterns of the two variants did not change when introgressed into sgn1, sgn3 or cif1 cif2 mutants (Fig. 1C, 1D). An apolar SGN1 localization would allow SGN3 124 125 to encounter SGN1 also on the stele-facing side, not only on the cortex side (Fig. 1E). This should lead to 126 constitutive signal activation in the absence of barrier defects, because the CIF peptides would now be able to 127 access a SGN3/SGN1 signaling module on the stele-facing side without crossing the barrier. Indeed, we found that the apolar SGN1 variant caused both, ectopic lignin deposition and precocious suberization in endodermal 128 cells (Figs. 1D, EV 1F,G), as previously described for endodermal barrier mutants. Yet, no barrier defect was 129 observed in the lines complemented with apolar SGN1 (Fig. EV 1A) and, consistently, we found CASP1-130 mCherry distribution to be normal in these lines, forming a continuous band in the central position of the 131 endodermal cells, indistinguishable from wild-type (Fig. EV 1B). This indicates that presence of SGN1 at the 132 plasma membrane to the inside of the CS leads to signaling in the absence of barrier defects. CASP1 promoter-133 driven wild type SGN1 lines as a control, complemented the mutant (Fig. EV 1A) and did not cause any changes 134 135 in lignin accumulation pattern (Fig. 1C). Interestingly, the apolar SGN1 lines accumulated lignin mainly on the stele-facing edges of the endodermal cell walls, as opposed to the cortical lignin deposition observed by ectopic 136 ligand treatment (compare Fig. 1B with Fig. 1D, arrows). The ectopic lignin deposition in apolar SGN1 lines 137 was fully dependent on the presence of receptor and ligand, as both in sgn3 and cif1 cif2 mutants, no excess 138 lignification at the stele-facing side could be observed in apolar SGN1 lines (Fig. 1D). This strongly suggests 139 140 that mislocalized SGN1 does not become constitutively active, but leads to continuous, ectopic transduction of CIF1/2 signals through SGN3. An apolar, but kinase-dead variant of SGN1 was also unable to induce ectopic 141 lignification (Fig. EV 1C), suggesting that a phosphorylation relay downstream of SGN1 is necessary for 142 lignification. 143

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SGN1 is a downstream component of the CIF/SGN3 pathway

Previous data showed that *sgn1* is less sensitive to high doses of externally applied CIF peptide, consistent with a role of SGN1 downstream of the SGN3 receptor (Doblas *et al*, 2017), at least with respect to 148 CIF-induced excess lignification. In order to address whether SGN1 is indeed a generally required downstream 149 component of the CIF/SGN3 pathway during CS formation, we evaluated whether the sgn1 mutant is resistant to complementation by CIF2 peptide treatment. In contrast to the full complementation of the discontinuous CS 150 151 domain of *cif1 cif2* double mutants, the *sgn1* mutant did not show full restoration of domain integrity, neither on 10 nM nor 100 nM CIF2 medium (Fig. 2A, Appendix Fig. S1A). While some degree of rescue occurred, only 152 about 50% of discontinuities were rescued. This was corroborated by testing CS functionality using the 153 propidium iodide (PI) assay. Only a weak complementation of barrier formation was observed even when grown 154 on 100 nM CIF2 medium (Fig. 2B). Our results indicate that SGN1 functions downstream of CIFs/SGN3, but 155 156 suggest that additional factors can partially compensate for its absence, most probably homologs of the extended RLCKVII family to which SGN1 belongs. 157

159 We then tested for direct connectivity between SGN3 receptor kinase and SGN1 by carrying out an in vitro kinase assay. Some RLCKVII members are reported to be phosphorylated and activated by LRR receptor 160 kinases (Kim et al, 2011, 1; Lu et al, 2010, 1). A Glutathione S-transferase (GST)-fused SGN3 kinase domain 161 was incubated with a kinase-dead form of a trigger factor (TF)-SGN1 fusion protein - which does not have 162 autophosphorylation activity - in the presence of radioactive ATP. Kinase-dead TF-SGN1 was efficiently 163 phosphorylated by SGN3 kinase domain, but not by a kinase-dead form of SGN3 in vitro (Fig. 2C, Appendix 164 Fig. S1B). This further supports our model that SGN1 is a direct downstream component of the SGN3/CIF 165 166 pathway.

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Two NADPH oxidases are absolutely required for CIF-induced lignification in the endodermis

Our generation of an apolar SGN1 thus appears to have reconstituted a functional CIF/SGN3 pathway 170 171 at the stele-facing (inner) endodermal surface, causing ectopic lignification in the absence of barrier defects. Yet, 172 the second intriguing aspect of this manipulation is the observation that lignification occurs almost exclusively at the inner endodermal edges (Fig. 1D), the side where endogenous CIF peptide must be assumed to be present. 173 174 External treatment, by contrast, leads to predominant lignification at the outer endodermal edges (Fig. 1B). This surprising spatial correlation between the site of signal perception and localized lignification suggests a very 175 direct molecular connection between the two events that would allow to maintain spatial information. In a cell 176 primed for lignification, i.e. with mono-lignol substrates available and polymerizing enzymes expressed, 177 lignification could be simply "switched on" by activating ROS production through NADPH oxidases. Previously, 178 we had found one of the NOXs, RBOHF, to be crucial for lignification at the CS. Intriguingly, RBOHF is the 179 180 only transmembrane protein known to accumulate at the Casparian strip domain, safe the CASPs themselves (Lee et al, 2013) (Fig. 3A). Moreover, homologous NADPH oxidases, such as RBOHB or RBOHD, also present 181 182 in the endodermal plasma membrane, are excluded from this domain (Lee et al, 2013) (Fig. 3A). We therefore asked whether CIF peptides would induce lignification through activation of RBOHF as a downstream 183 component. To our surprise, CIF treatment still led to induced lignification in *rbohF*, despite the fact that RBOHF, 184 185 is strictly required for CS lignification in untreated conditions. Yet, when RBOHD was knocked-out in addition to RBOHF, a complete absence of lignification upon CIF treatment was observed (Fig. 3B). The fact that 186 RBOHD single mutant neither showed defects in developmental CS lignification, nor in CIF-induced 187 lignification, indicates that RBOHF is required for both processes, but that upon strong stimulation with 188 exogenous CIF peptide application, RBOHD can be additionally used. Indeed, using the general NADPH 189 oxidase inhibitor diphenyleneiodonium (DPI) in short-term co-treatment with CIF was also able to fully block 190 CIF-induced ectopic lignification in wild-type with intact Casparian strips, further supporting that NADPH 191 192 oxidases act downstream in the CIF/SGN3 pathway (Fig. EV 2A).

CIF2 induces highly localized ROS production through RBOHF and RBOHD

196 The direct activity of NOX enzymes is not lignification, but production of superoxide (O_2^-) that becomes 197 dismutated to hydrogen peroxide (H_2O_2) in the apoplast (Kärkönen & Kuchitsu, 2015). We had previously 198 established that the different subcellular distribution of the SGN3 receptor and the SGN1 kinase only intersect 199 at a very restricted domain at the outer (cortex-facing) edge of the CS (Alassimone *et al*, 2016)(Fig. 4A). We 190 therefore attempted to visualize whether ROS might be produced locally in response to CIF treatment. Many 201 ways exist to visualize and quantify ROS, but only few allow for high spatial resolution and for discrimination 202 between extracellular and intracellular ROS. An older method, based on ROS-induced cerium precipitation that can be detected using transmission electron microcopy (TEM), has been used by us previously to demonstrate 203 204 that a highly localized ROS production indeed occurs at the CS and is dependent on NOX activity (Bestwick et al, 1997a; Lee et al, 2013) (Fig. 4B). Using this method, we could detect a strong ROS production in response to 205 206 CIF2, exclusively at the regions of endodermal-endodermal cell walls outside of the CS, but nowhere along the endodermis-cortex cell walls, which are equally reached by the cerium chloride and where NOX enzymes are 207 also present in the plasma membrane (Fig. 4B,C). Based on this striking spatial coincidence between the 208 209 SGN3/SGN1 overlap region (Fig. 4A) (Alassimone et al, 2016) and CIF2-induced ROS production, we developed a procedure to quantitatively assess cerium precipitates and checked whether this localized ROS 210 211 production is indeed dependent on the SGN3 pathway (Figs. 4D,E, EV 3A, see also Experimental Procedures section). For SGN3, we found that already steady-state ROS production was undetectable in the mutant and that 212 there was no increase upon CIF2-treatment (Fig. 4D,E). The sgn1 mutant showed lower, but still detectable 213 steady state ROS levels, but no significant increase upon CIF2-treatment (Fig. 4D,E). Thus, the highly localized 214 ROS accumulation induced by CIF2 is entirely dependent on the localized SGN3/SGN1 receptor module, 215 suggesting that the localization of the module determines the spatial extent of ROS production. As in the case 216 of lignification, we observed that CIF2-induced ROS could be produced by either RBOHF or RBOHD, as only 217 the double mutant caused a complete absence of ROS after CIF-stimulation. As expected, but not previously 218 219 demonstrated, the steady-state ROS production at the CS observed before stimulation was exclusively dependent 220 on RBOHF, but not RBOHD (Fig. 4D,E). 221

In contrast to lignin polymerization, suberin might not require NADPH oxidase activity. It was shown previously that CIF2 triggers excess suberization in WT (Fig. EV 2B,C)(Doblas *et al*, 2017). We found excess suberin deposition in *rbohD* and *rbohF* upon peptide treatment, although a slight enhancement is already observed in *rbohF*. In *rbohDF*, excess suberin deposition is very strong, even without treatment, likely due to a strong activation of the surveillance system due to complete absence of a CS (Fig. EV 2B,C). These data indicate that CIF2-triggered suberin accumulation is not affected by NADPH oxidases and suggests existence of a ROSindependent branch of the SCHENGEN pathway that regulates this process (see below).

SGN1 can directly activate RBOHF and RBOHD via phosphorylation

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232 The above data strongly suggest a direct connection between the SGN3/SGN1 kinase module and the 233 two NOX enzymes. We therefore conducted an *in vitro* kinase assay in order to ask whether SGN1 can directly phosphorylate the N-terminal cytoplasmic region of RBOHF and RBOHD. We found that recombinantly 234 235 expressed TF-SGN1 could phosphorylate both the recombinant N-terminal part of RBOHF and RBOHD (Fig. 5A). Plant NOX regulation has been intensively studied and shown to be highly complex, requiring possibly 236 237 interdependent activities of kinases, as well as small GTPases (Kadota et al. 2015). We therefore tested whether 238 SGN1 might be sufficient for activation of RBOHF activity in a cellular context. To do so, we made use of a heterologous reconstitution system in human HEK293T cells, which show very low endogenous ROS production 239 240 and for which it had been previously demonstrated that plant NADPH oxidases can be expressed and their activation mechanism be studied (Han et al, 2019). As a positive control the previously described calcium-241 242 dependent kinase complex of calcineurin B-like (CBL) interacting protein kinases 26 (CIPK26) and CBL1 was used and shown to be active (Fig. 5B). When expressing wild-type SGN1 in this cell line, we noticed that it did 243 not activate RBOHF or RBOHD, but that it also did not localize to the plasma membrane as in plant cells (Fig. 244 EV 4A). However, when we used the functional, constitutively plasma membrane-localized myrpalm-SGN1 245 version, a significant induction of ROS production was observed for RBOHF and to a lesser extent for RBOHD 246 247 (Figs. 5B, EV 4B).

CIF peptide-induced CASP domain growth requires new protein synthesis, but not ROS production

The direct phospho-relay from SGN3 receptor, to SGN1 kinase to RBOHF and RBOHD outlined above draws a direct molecular connection from perception of a peptide hormone stimulus to cell wall lignification. Moreover, it accounts for both, the highly localized ROS production that we observe upon CIF 254 stimulation, as well as the observation that localization of lignification is correlated with the site of active SGN3 255 signaling. Yet, the massive enhancement of lignification observed upon CIF stimulation (Fig. 3B), the increase of CASP accumulation and ectopic patch formation (Appendix Fig. S1A), as well as the non-localized formation 256 257 of precocious and enhanced suberin (Fig. EV 2B,C), are additional outcomes of SGN3 pathway stimulation and should be driven by transcriptional changes. Indeed, some degree of transcriptional upregulation of CASP genes 258 259 has been reported previously upon CIF1 treatment (Nakayama et al, 2017). CIF stimulation does not only lead to enhanced accumulation and ectopic patches of CASP1-GFP, absence of SGN3 signaling also leads to 260 discontinuous CASP1-GFP signals that could be explained by insufficient amount of CASPs and other factors 261 262 being produced during endodermal differentiation (Fig. 6A) (Doblas et al, 2017; Pfister et al, 2014). Neither the single, nor the double NADPH oxidase mutants display discontinuous CASP1-GFP signals, indicating that ROS 263 264 production is not required for this aspect of the CIF/SGN3 pathway (Fig. 6A). In order to directly demonstrate that formation of a continuous CASP domain requires newly formed gene products, we treated our cifl cif2 265 double mutant with CIF2 peptide in the presence or absence of protein synthesis inhibitor cycloheximide (CHX). 266 267 CASP1-GFP signal strongly increased during 8 hours of CIF treatment, during which the discontinuous CASP1-GFP domains became continuous. This effect was abrogated by CHX treatment (Fig. 6 B,C, Movie EV1). 268

The NADPH oxidase-independent branch of the CIF/SGN3 pathway is associated with MAP kinase stimulation and causes strong activation of gene expression.

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In pattern-triggered immune receptor signaling, gene activation is thought to depend in large parts on activation of Mitogen-activated protein (MAP) kinases (Dodds & Rathjen, 2010). We therefore tested whether CIF-treatment leads to MAP kinase phosphorylation and indeed found that CIFs can induce MAP kinase phosphorylation in a SGN3-dependent manner, further extending the molecular parallels between immune receptor signaling and the CIF/SGN3 pathway and suggesting that gene induction in the CIF pathway might equally depend on MAP kinase signaling (Fig. EV 5B).

We then undertook an RNA profiling of seedling roots at different time points (30, 120 and 480 min) 280 after CIF stimulation, using wild-type, *cif1 cif2* and *sgn3* mutants as genotypes. 930 genes were found to be 281 282 differentially expressed across any of the genotypes and time points combined, using a stringent cut-off (adj.pval. ≤ 0.05 ; logFC ≥ 1 or logFC ≤ -1) (Table EV1). After normalization of batch effects, the three replicates 283 clustered closely with a large degree of variance explained by the peptide treatment in wild-type and *cif1 cif2* 284 (Fig. EV 5A). Wild-type and *cif1 cif2* samples showed nearly identical responses (co-relation efficiency values 285 286 were 0.90, 0.89, 0.94 at 30, 120, 480 min respectively), with cifl cif2 displaying a slightly stronger overall amplitude (Fig. 6D). Importantly, sgn3 had virtually no differentially expressed genes across treatments, 287 288 indicating the specificity of the CIF responses and further corroborating that SGN3 is the single, relevant receptor for CIF responses in roots (Figs. 6D, EV 5A, Table EV1). Our data extends on the previous data of Nakayama 289 290 et al. (2017), by showing that all 5 CASP genes are differentially regulated upon CIF-treatment (Fig. EV 5C). 291 Moreover, we observed upregulation of MYB36, a central transcription factor for Casparian strip formation and CASP expression, thus potentially accounting for the increases in CASP1-4 expression (Kamiya et al, 2015; 292 293 Liberman et al, 2015)(Fig. EV 5C). The fact that CIF stimulates MYB36 expression also is consistent with the 294 recent report that CIF treatment can enhance ectopic endodermal differentiation, driven by overexpression of the 295 SHR transcription factor (Drapek et al, 2018).

297 Of the four response clusters defined by k-mean clustering, the two largest clusters (1 and 3) group early response genes (cluster 1) and later response genes (cluster 3). Cluster 2 groups a smaller fraction of genes 298 showing strong and largely sustained responses over the time scale of the experiment, while cluster 4 groups a 299 300 minor fraction of genes downregulated at later time points. GO term analysis indicates that many of the most significant, overrepresented terms in the "early response" (1) and "strong and sustained" (2) gene clusters are 301 302 related to immune and defense responses (response to chitin, bacterium, callose, etc.), as well as responses to oxidative stress (Table EV2). The top GO categories of the late response cluster (3) were suberin/cutin 303 biosynthesis, supporting our observation that suberin accumulation is a late effect of CIF stimulation (Figs. EV 304 305 5D, EV 2B,C)(Doblas et al, 2017). Additional, enriched terms were related to oxidative stress and cell wall 306 remodeling, nicely fitting the CIF2-induced ROS production we observed. Intriguingly, the cluster showing late

downregulation of gene expression contains many overrepresented GO terms related to a wide variety of
 transport processes, reaching from water, nitrate and ammonium transport, to primary and secondary metabolite
 or hormone transport (Fig. 6D, Table EV2). We interpret this as a consequence of CIF-induced lignification and
 suberization, which must profoundly impact endodermal ability for shuttling organic and inorganic compounds,
 as well as signaling molecules between stele and cortical root cell layers (Barberon & Geldner, 2014).

While GO terms related to lignification were also overrepresented, they were not as highly ranked as 313 expected from the strong induction of lignification upon CIF treatment. A number of reasons might account for 314 315 this: Firstly, GO terms for pathogen, defense and oxidative stress responses, are all bound to overlap with and contain genes that mediate lignification. Moreover, already elevated levels of developmental, lignin-related gene 316 317 expression in the endodermis might lead to less-pronounced fold-changes. Finally, the multiple roles for lignin biosynthetic genes in secondary metabolism further obscure clear categorization. Nevertheless, upregulation of 318 lignin biosynthesis by CIF-treatment became evident when clustering a set of laccases and peroxidases, the two 319 main families of enzymes implicated in lignin polymerization in the apoplast (Liu, 2012). A significant number 320 of both are strongly upregulated (Fig. EV 5E,F). Moreover, MYB15, a transcription factor shown to be involved 321 322 in stress and MAMP(microbe-associated molecular pattern)-induced lignification (Chezem et al, 2017, 15) is 323 about 6-fold induced after 30 min and about 17-fold after 120 min in both treated wild-type and *cif1 cif2*, nicely correlating with the upregulation of peroxidases and laccases (Fig. EV 5E,F, Table EV1). We then made use of 324 325 two of the most highly differentially expressed genes upon CIF2-treatment (PER15 and PER49) in order to 326 establish whether SGN1 and the NADPH oxidases are also required for gene regulation downstream of SGN3 327 activation. In qPCR analysis both peroxidase genes still showed a slight, but significant upregulation upon CIFtreatment in the sgn1 mutant (compared to the complete absence of response in sgn3) (Fig. 6E-H), clearly 328 indicating that, as for ROS production, SGN1 is an important, yet not absolutely required, downstream 329 330 component in CIF-regulated gene expression. When directly plotting peroxidase fold-inductions in *rbohF*, 331 rbohD and double mutant, a seemingly strong attenuation was observed (Fig. 6F,H). However, when independently plotting gene expression in treated and untreated conditions (Fig. 6E,G), this lower fold-induction 332 is largely explained by an already significant basal increase in marker gene expression in *rbohF* and *rbohD* 333 rbohF double mutants. Indeed, both mutants display an absence of Casparian strips, which should lead to an 334 335 endogenous stimulation of the SGN3 pathway, nicely supporting the barrier surveillance model. Absolute expression of both marker genes in the NADPH oxidase mutants is in effect higher than in wild-type after CIF 336 treatment, leading us to conclude that ROS production and gene activation in response to CIF are two 337 independent branches of this signaling pathway, with the branching occurring downstream of the SGN1 kinase. 338 339 We finally wanted to know whether the mislocalization of the SGN1 kinase reported initially is only affecting CIF/SGN3 signaling at the plasma membrane, or whether it also affects gene activation. We found that both 340 341 PER15 (Fig. 6I) and PER49 (Fig. 6J) are dramatically upregulated in the non-polar SGN1 lines, independent of exogenous peptide application, strongly corroborating our model whereby the SCHENGEN pathway function 342 343 crucially depends on the correct subcellular localization of its downstream kinase.

DISCUSSION

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The data presented here sketch out an entire signaling pathway. Previously, SGN3 had been established 347 348 as the receptor for CIF1 and 2, but its connection to SGN1 had exclusively been based on genetic evidence. Here we show that SGN3, SGN1 and RBOHD/F are, biochemically and functionally, part of a signal transduction 349 chain, leading directly from localized peptide perception to localized ROS production and lignification (Fig. 7A). 350 Moreover, the pathway branches downstream of SGN1, leading to MAP kinase activation and stimulation of 351 gene expression. Some of the most strongly induced genes being peroxidases and laccases that would further 352 353 enhance and sustain lignification. The SCHENGEN pathway therefore elegantly integrates fast, plasma membrane-based responses that maintain positional information (ROS and lignin are produced close to where 354 355 the ligand is perceived) and slower, gene expression-based responses, which have lost positional information, but would allow to enhance and maintain the ROS-burst-controlled activation of peroxidases (Fig. 7A). 356

The SCHENGEN pathway bears a striking overall resemblance to well-established signaling pathways for perception of MAMPs (Fig. 7B). In MAMP perception, structurally similar receptor kinases, such as FLS2 360 or EFR bind to microbial patterns and interact with the SERK family of co-receptors (Chinchilla et al, 2007), as is the case for SGN3 (Okuda et al, 2020). The signal is then transduced through kinases of the RLCKVII family, 361 such as BIK1 or PBLs, which are homologs of SGN1 (Liang & Zhou, 2018). BIK1 in turn was shown to 362 phosphorylate RBOHD, driving the well-described MAMP-induced ROS burst(Kadota et al, 2014, 1; Li et al, 363 2014, 1). Moreover, it has recently been shown that kinases of the RLCKVII family directly phosphorylate 364 365 MAPKKKs, which now mechanistically explains how MAMP perception induces MAP kinase phosphorylation (Bi et al, 2018). It is thus tempting to speculate that the SCHENGEN pathway represents an ancient neo-366 functionalization of an immune receptor pathway. Indeed, this has been proposed recently and it was pointed out 367 368 that the closest receptor homologs to SGN3 and GSO2 are PEPR1 and PEPR2 (Creff et al, 2019). The latter are receptors to an endogenous plant peptides (AtPEPs), whose activities resemble that of MAMPs and are best 369 370 thought of as "phytocytokines", i.e. agents able to induce an immune-like response in cells that have not yet encountered MAMPs (Gust et al, 2017). Such an original phytocytokine might then have been neo-371 functionalized to induce pre-formed defensive cell wall barriers in a developmental context, in the absence of 372 any actual biotic or abiotic stress. MAMP perception and MAMP-induced ROS production are found already in 373 mosses and clearly precede the wide-spread adoption of lignin as a major, cell-wall-reinforcing polymer 374 375 (Bressendorff et al, 2016; Weng & Chapple, 2010). Moreover, lignin was speculated to originate from phenylpropanoid-derived defense compounds, making it plausible that immune signaling pathways have been at 376 the origin of developmental regulation of lignification. The intriguing innovation of the SCHENGEN pathway 377 378 would then reside in the subcellular arrangement of its signaling components, especially that of SGN1. Indeed, 379 we demonstrate that polar localization of SGN1 to the outside of the Casparian strip domain is crucial for barrier function - since its simple mislocalization to the inside leads to ligand- and receptor-dependent overlignification 380 at inner cell corners. Thus, the only feature that arrests signaling in wild-type is the formation of a lignified 381 diffusion barrier in the cell wall, preventing access of the stele-produced peptides to SGN3 receptors at the outer 382 383 domain - as the only population able to stimulate the polar SGN1 kinase. Since we have shown here that a main 384 read-out of the SCHENGEN pathway is cell wall lignification itself, this designs a fascinating, spatial negativefeedback loop, in which SCHENGEN pathway-stimulated lignification feedback-regulates itself once enough 385 lignin has been produced to form a tight diffusion barrier and to fully prevent further CIF peptide penetration to 386 387 the outside.

It will be important to further describe the extent to which the SCHENGEN pathway uses components 389 of MAMP signaling (as for example the requirement for SERK-family co-receptors, which has not been 390 demonstrated yet for SGN3), but we propose that our findings on the SCHENGEN pathway function can already 391 392 be of considerable interest for MAMP receptor signaling. The particular, restricted spatial overlap of receptor and downstream kinase has allowed us to visualize that, even if NADPH oxidases are non-localized (as in the 393 394 case of RBOHD), ROS can be locally produced in a micrometer-scale region at the plasma membrane, through localized receptor stimulation. In MAMP receptor signaling, the non-localized nature of both MAMP receptor 395 and NADPH oxidase would not have allowed to visualize this unexpected degree of spatial control. Yet, during 396 397 actual microbial infections, highly-localized receptor stimulation and ROS production might indeed occur and be relevant for the outcome of the immune response. In addition, while it is established that MAMP-stimulated 398 399 ROS production is an important part of the immune response, its direct molecular downstream action is not well understood. Lignin production has long been associated with immune responses, as well as responses to cell wall 400 401 damage, yet a direct molecular connection from MAMP perception to lignification has rarely been drawn (Chezem et al, 2017, 15). It will be intriguing to investigate whether and how much MAMP-induced ROS 402 production is actually used for lignification during defense and to which degree this explains the importance of 403 ROS in plant defense responses. 404 405

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The SCHENGEN pathway might not be limited to the regulation of lignified diffusion barriers, as it has been shown that it is also important in the formation of the embryonic cuticle. The peptide ligand used in this context has not been identified and it remains unclear whether an equally precise barrier surveillance mechanism is also acting to ensure separation of endosperm and embryo during embryonic cuticle formation (Creff *et al*, 2019). In this context, the SCHENGEN pathway might be used to drive production and deposition of cutin instead of lignin and suberin as in the case of the endodermis. In the future, it will be fascinating to investigate whether the SCHENGEN pathway is of even broader developmental significance and to understand 413 414 the molecular basis that enables its distinct, organ and cell-type specific activities.

415 MATERIAL AND METHODS

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417 **Plant material and growth conditions**.

For all experiments, *Arabidopsis thaliana* (ecotype Columbia) was used. The T-DNA tagged lines, *sgn1-2*(SALK_055095C) and *sgn3-3* (SALK_043282) were obtained from NASC (Alassimone *et al*, 2016; Pfister *et al*,
2014). *rbohD dSpm*, *rbohF dSpm* and *rbohD dSpm rbohF dSpm* are described in (Tissier *et al*, 1999; Torres *et al*,
2002). *esb1-1* was kindly shared by Prof. David Salt's group (Baxter *et al*, 2009; Hosmani *et al*, 2013b). The *cif1-2 cif2-2* double mutant was generated by CRISPR-Cas9 method (Fauser *et al*, 2014) using 5'- gctttggtttaggactggag -3'
as a protospacer sequence to target both CIF1 and CIF2 loci.

pCASP1::CASP1-GFP marker line was crossed into *sgn1-2*, *sgn3-3* or *cif1-2 cif2-2*. The *pCASP1::myrpalm SGN1*construct was independently transformed into each mutant background using the floral dip method(Clough & Bent,
1998). For observations and histological analysis, seeds were kept for 2 days at 4°C in the dark for stratification, then
grown for 5 days at 22°C under 16 h light/8 h dark vertically on solid half-strength Murashige–Skoog (MS) medium.
Cycloheximide (Sigma) and Diphenyleneiodonium chloride (Sigma) were prepared as 50 mM water stock solution
and as 20 mM DMSO stock solution, respectively.

432 Plasmid construction

433 To generate myrpalm SGN1, GGCFSKK (5'- GGAGGATGCTTCTCTAAGAAG -3') (Vermeer et al, 2004) was 434 added to SGN1 cDNA right after the start codon by inverse PCR. This SGN1 variant was combined with pCASP1 435 and mCitrine coding sequence by LR clonase (Thermo Fisher Scientific). Coding sequences of RBOHF 2-383 a.a. 436 and RBOHD 1-376 a.a. were inserted between BamHI and XhoI sites of a modified pET24(+) vector, in which GST 437 (glutathione S-transferase) was added to the N-terminus. For constructing a GST-SGN3 kinase domain vector, a 438 cDNA fragment coding for 899 a.a. to 1249 a.a. of SGN3 was fused into pDEST15 by Gateway LR reaction. TF-439 SGN1 was as described previously (Alassimone et al, 2016). Kinase-dead mutations in SGN1 (K134E) and SGN3 440 (K979E) were introduced by site-directed mutagenesis. 441

442 **Protein expression in bacteria and purification**

443 GST or Trigger factor (TF) proteins were expressed from pGex6p-1 or pCold-TF vectors in BL21 (DE3) CodonPlus 444 RIL and purified in a glutathione sepharose column (Thermo Fischer Scientific) or a Co-Sepharose column (Clontech) according to the manufacturer's protocols. GST-RBOHD N-ter and GST-RBOHF N-ter were produced 445 and purified as described in (Drerup et al, 2013). Bacterially expressed GST-SGN3 kinase domain (WT or kinase-446 dead) in BL21 (DE3) CodonPlus RIL was purified with a glutathione sepharose column following the manufacture's 447 protocol. For TF-SGN1 (WT or kinase-dead), we followed the method in the previous report (Alassimone et al, 448 449 2016). The buffer of all proteins was changed to 50 mM HEPES-KOH pH7.5 and 1 mM DTT with PD-10 or NAP-450 5. Purified kinase protein solutions were mixed with glycerol to 30% [v/v] final concentration and kept at -20°C until 451 use. 452

453 In vitro kinase assay

In vitro kinase assay was done as described previously (Alassimone *et al*, 2016) with small modifications. Purified 250 ng GST-SGN3 kinase domain was incubated with 250 ng TF-SGN1 or TF proteins in reaction buffer (50 mM HEPES-KOH pH7.5, 1 mM MnCl2. 1 mM DTT, 1 mM ATP, 185,000 Bq [γ -³²P] ATP) at 30°C for 30 min. 250 ng TF-SGN1 and 1 \Box g GST or 500 ng GST-N-terminal cytoplasmic domains of RBOHD or F were treated in the same way as above. The reaction was stopped by adding 4 x LDS sample buffer (Invitrogen) and heating at 75°C for 10 min. The samples were separated on 4-12 % gradient Nu-PAGE gels or 10 % Nu-PAGE gels (Invtirogen). After drying the gels, signal was detected using Typhoon FLA7000 (GE Healthcare).

Western blotting

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Plants were grown in liquid MS medium supplemented with 0.5% sucrose with cell strainer for 5 days under long
day (18 h light /6 h dark) condition after 2-day vernalization at 4°C. Hydroponically grown seedlings were
transferred into fresh liquid MS medium containing 0.5% sucrose with or without 1 □M CIF2 peptide and incubated
for 15 min. The seedlings were immediately frozen in liquid nitrogen and ground by TissueLyser II (Qiagen).
Extraction buffer (20 mM Na phosphate pH 7.4, 150 mM NaCl, 1 mM EDTA, 0.1% Tween, 50 mM β

468 glycerophosphate, 0.5 mM PMSF, 100 µM sodium orthovanadate, 5 mM Na fluoride, Complete cocktail) was added 469 to the frozen samples. The samples were briefly mixed by vortex and centrifuged at 15,000 rpm for 15 min at 4°C. 470 The supernatant was transferred in new tubes and measured the concentration by Bradford method (Thermo Fischer Scientific). Equal amount of protein was loaded ($20 \Box g / lane$) and separated in 10% acrylamide gel (Eurogentec). 471 472 After the electrophoresis, the separated proteins were transferred onto nitrocellulose membrane (Amersham, GE 473 Healthcare) by XCell SureLock (Invitrogen) and stained with Ponceau S to show the loading control. The blot was 474 incubated in blocking buffer (3% skim milk in TBS) for 1 hour and probed with anti-phospho-p42, p44 antibody 475 (1:1000 dilution, Cell signaling technologies #9101) in 0.5% skim milk in TBS-0.1% Tween for overnight at 4°C. After washing the blot three times with TBS-0.1% Tween, the blot was treated with anti-rabbit secondary antibody 476 (1:30,000 dilution, Agrisera) in 0.5% skim milk in TBS-0.1% Tween for 1 hour and washed three times with TBS-477 0.1% Tween. Signals were detected on X-ray film with SuperSignal West Femto Kit (Thermoscientific). For the 478 MPK6 detection, the membrane was stripped (Pierce), incubated in blocking buffer for 1 hour and probed with anti-479 MPK6 antibody (1:10000 dilution, Sigma) in blocking buffer for 1 hour at room temperature. After washing the blot 480 481 three times with TBS-0.1% Tween, the membrane was treated with anti-rabbit secondary antibody (1:30000, Agrisera) in blocking buffer and washed three times. Signal were detected with SuperSignal West Femto kit 482 483 (Thermo Scientific).

Confocal microscopes

Confocal pictures were obtained using Leica SP8 or Zeiss LSM 880 confocal microscopes. The excitation and
detection window settings to obtain signal as follows: When using Leica SP8 (excitation, detection window), GFP
(488 nm, 500–550 nm), mCitrine, (514 nm, 518–560 nm), mCitrine/mCherry, (514 nm and 594 nm, 518–560 nm,
600–650 nm, sequential scan), when using Zeiss LSM880 (excitation, detection window), GFP (488 nm, 500–550 nm), Calcofluor White (405 nm, 425-475 nm), Basic Fuchsin (561 nm, 570-650 nm) and Fluorol Yellow (488 nm,
500-550 nm).

PI assay

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PI assay was done as described previously (Lee *et al*, 2013) with small changes. Seedlings were incubated in water
containing 10 µg /mL PI for 10 min and transferred into fresh water. The number of endodermal cells were scored
using a Zeiss LSM 700 confocal microscope (excitation 488 nm, SP640, split at 570 nm) from the onset of cell
elongation (defined as endodermal cell length being more than two times than width in the median, longitudinal
section) until PI could not penetrate into the stele.

500 Scoring discontinuities in CS domain

501 CASP1-GFP signal was obtained by Leica SP8 as 1 □m step z-stack images from 5-day old seedlings. After the
 502 images were projected as maximum projections, total lengths of CS domains were measured in maximum
 503 projections and the number of discontinuities in the domain was counted manually.

Lignin and cell wall staining

506 ClearSee adapted cell wall staining was performed as described in recent publications (Kurihara et al, 2015; Ursache 507 et al, 2018). Briefly, 7-8 five-day-old seedlings were fixed in 3 mL 1 x PBS containing 4% para-formaldehyde for 1 hour at room temperature in 12-well plates and washed twice with 3 mL 1 x PBS. Following fixation, the seedlings 508 were cleared in 3 mL ClearSee solution under gentle shaking. After overnight clearing, the solution was exchanged 509 to new ClearSee solution containing 0.2% Fuchsin and 0.1% Calcofluor White for lignin and cell wall staining 510 511 respectively. The dye solution was removed after overnight staining and rinsed once with fresh ClearSee solution. 512 The samples were washed in new ClearSee solution for 30 min with gentle shaking and washed again in another 513 fresh ClearSee solution for at least one overnight before observation.

515 Methanol-based Fluorol Yellow staining of Arabidopsis root suberin

516 Vertically grown 5-day old seedlings were incubated in methanol for three days at room temperature. The cleared 517 seedlings were transferred to a freshly prepared solution of Fluorol Yellow 088 (0.01%, in methanol) and incubated 518 for 1 hour. The stained seedlings were rinsed shortly in methanol and transferred to a freshly prepared solution of 519 aniline blue (0.5%, in methanol) for counterstaining. Finally, the seedlings were washed for 2-3min in water and 520 transferred to a chambered coverglass (Thermo Scientific), covered with a piece of 1% half-strength MS agar and 521 imaged using a Zeiss LSM 880 confocal microscope as described above.

522 523 Detection of H₂O₂ production *in situ* using transmission electron microscopy

524 Visualization of H₂O₂ production around Casparian strip was done by cerium chloride method as described 525 previously (Bestwick et al. 1997b; Lee et al. 2013) with some modifications, 4-day-old Arabidopsis seedlings were 526 transferred onto fresh 1/2 MS solid medium with or without 1 µM CIF2 and incubated for 24 hours. After the 527 peptide treatment, the seedlings were incubated in 50 mM MOPS pH7.2 containing 10 mM CeCl₃ for 30 min. After incubation with CeCl₃, seedlings were washed twice in MOPS buffer for 5 min and fixed in glutaraldehyde 528 solution (EMS, Hatfield, PA) 2.5% in 100 mM phosphate buffer (pH 7.4) for 1 hour at room temperature. Then, they 529 were post-fixed in osmium tetroxide 1% (EMS) with 1.5% of potassium ferrocyanide (Sigma, St. Louis, MO) in 530 531 phosphate buffer for 1 hour at room temperature. Following that, the plants were rinsed twice in distilled water and 532 dehydrated in ethanol solution (Sigma) at gradient concentrations (30% 40 min; 50% 40 min; 70% 40 min; two times (100% 1 hour). This was followed by infiltration in Spurr resin (EMS) at gradient concentrations (Spurr 33% 533 534 in ethanol, 4 hours; Spurr 66% in ethanol, 4 hours; Spurr two times (100% 8 hours) and finally polymerized for 48 535 hours at 60°C in an oven. Ultrathin sections 50 nm thick were cut transversally at 1.3 ± 0.1 mm from the root tip, on a Leica Ultracut (Leica Mikrosysteme GmbH, Vienna, Austria) and picked up on a copper slot grid 2x1 mm (EMS) 536 537 coated with a polystyrene film (Sigma). Micrographs were taken with a transmission electron microscope FEI CM100 (FEI, Eindhoven, The Netherlands) at an acceleration voltage of 80kV and 11000x magnification (pixel size 538 539 of 1.851nm, panoramic of 17x17 pictures), exposure time of 800ms, with a TVIPS TemCamF416 digital camera 540 (TVIPS GmbH, Gauting, Germany) using the software EM-MENU 4.0 (TVIPS GmbH, Gauting, Germany). All the 541 pictures were taken using the same beam intensity, and panoramic aligned with the software IMOD (Kremer et al, 542 1996). For quantification, thresholding of cerium precipitate on pictures was scored using IMOD software (Kremer 543 et al, 1996). Briefly, all pictures subjected to the quantification were normalized to one picture, a section from nontreated wild-type seedlings, by comparing the gray value of plastids in pericycle cells. After normalization, cell wall 544 545 spaces from the beginning of the Casparian strip at the pericycle side until the cortex corner between endodermal 546 cells were selected and one identical threshold setting was applied to all pictures in order to highlight signals around Casparian strips. The values were shown as a percentage of thresholded pixels to selected area. 547

ROS production assay in HEK cells

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550 Measurements of RBOHD and RBOHF activity were performed as described in (Han et al, 2019) with some modifications. RBOHD or RBOHF were transiently expressed in HEK293T cells with or without coexpression of 551 SGN1 or myrpalm SGN1. HEK293T cells were cultivated at 5 % CO2 and 37 °C in Dulbecco's modified Eagle's 552 medium (DMEM) mixture F-12HAM (Sigma) enriched with 10% fetal bovine serum. Before transfection, lysine 553 coated white 96-well plates were inoculated with HEK293T cells and incubated for 24 h. For the transient 554 555 transfection of the cells, GeneJuice transfection reagent (Novagen) was used according to the manufacturer's 556 guidelines. The transfected expression cassettes were cloned into modified pEF1 vectors (Drerup et al. 2013). Each 557 well was transfected with 110 ng of plasmid mixture (50 ng pEF1-RBOH; 30 ng of each effector; empty pEF1 558 vector to ensure equal loading). Transfected cells were incubated for 48 h and subsequently measured in a buffer consisting of Hank's Balanced Salt Solution (Gibco) with 62 µM L-012 and 60 µg / mL HRP. To stimulate SGN1, 559 560 calyculin A, a phosphatase inhibitor, was added at the final concentration of 0.1 µM directly before the start of the measurements. ROS production was measured in a LB 943 Mithras² (Berthold) microplate reader and is presented as 561 relative luminescence units s⁻¹ (RLU s⁻¹) in graphs. For all experiments, the values were gained from 6 wells 562 563 measured in parallel and average values were plotted with S.D. Experiments have been repeated three times.

Sample preparation for RNA-seq experiments

566 Wild type, *sgn3-3* and *cif1-2 cif2-2* were grown on solid half-MS medium with mesh for 5 days and transferred onto 567 fresh half MS medium with or without 100 nM CIF2. After 30, 120, or 480 min incubation, aerial parts were cut off 568 and whole roots were collected. Samples were immediately frozen in liquid nitrogen and RNA was extracted using a 569 Trizol-adapted Reliaprep RNA extraction Kit (Promega).

RNA-seq library preparation and sequencing

RNA-seq libraries were prepared as described in (Jan *et al.* 2019) in using 1000 ng of total RNA. Libraries were
 sequenced using the Illumina TruSeq SR Cluster Kit v4 reagents on the Illumina HiSeq 2500 and processed using
 the Illumina Pipeline Software version 2.20.

576 **RNA-seq data processing and analysis**

578 Data processing was performed by the Lausanne Genomic Technologies Facility using their in-house RNA-seq 579 pipeline. It includes purity-filtered read trimming for adapters and low quality sequences with Cutadapt (v. 1.8)

- (Martin, 2011). Removal of reads matching ribosomal RNA sequences with fastq_screen (v. 0.11.1), followed by
 low complexity read filtering with reaper (v. 15-065) (Davis et al, 2013). Cleaned reads were aligned against
 Arabidopsis thaliana TAIR10 genome using STAR (v. 2.5.3a) (Dobin et al, 2013). Read counts were obtained per
 gene locus with htseq-count (v. 0.9.1) (Anders et al, 2015) using A. thaliana TAIR10 Ensembl 39 gene annotation.
 The quality of the data alignment was evaluated with RSeQC (v. 2.3.7) (Wang et al, 2012).
- 584 The quality of the data 585

Gene-level statistical analysis was performed in R (R version 3.4.3). Filtering of genes with low counts was
achieved based on the rule of 1 count per million (cpm) in at least 1 sample. TMM normalization was applied for
library sizes scaling followed by log-transformation into counts per million or CPM (EdgeR package version 3.20.8)
(Robinson et al, 2010). PCA was computed using normalized values corrected for batch effect using limma function
removeBatchEffect.

- 591 Differential expression was computed with limma-trend approach (Ritchie *et al*, 2015) by fitting all samples into one 592 linear model. The batch factor was added to model matrix.
- 593 Pairwise comparisons treated vs untreated per time point were assessed using moderated t-tests. The adjusted p-
- value is computed by the Benjamini-Hochberg method, controlling for false discovery rate (FDR or adj.P.Val).
- Differential expression of untreated mutant vs wt per time point was assessed using moderated F-test and Post-Hoc
 classification. The adjusted p-value is computed by the Benjamini-Hochberg method, controlling for false discovery
 rate (FDR or adj.P.Val).
- Differential expression of treated vs untreated over time paired by genotype was assessed using moderated F-test
 and Post-Hoc classification. The adjusted p-value is computed by the Benjamini-Hochberg method, controlling for
 false discovery rate (FDR or adj.P.Val).
- Interaction between time and treatment (paired by genotype, excluding SGN3 of the model) was assessed using
 moderated F-test. The adjusted p-value is computed by the Benjamini-Hochberg method, controlling for false
 discovery rate (FDR or adj.P.Val).
- Time effect in untreated conditions (paired by genotype, excluding SGN3 of the model) was assessed using
 moderated F-test. The adjusted p-value is computed by the Benjamini-Hochberg method, controlling for false
 discovery rate (FDR or adj.P.Val).
- 608Genes were considered significant in further analysis if the adjusted p-value was equal or below 0.05 and the log2-609fold change was >= 1. Heatmaps were constructed using the package ComplexHeatmap (v1.99.4, pearson distance610for row clustering) (Gu *et al*, 2016). Genes were clustered using kmeans (factoextra v1.0.5,
- https://github.com/kassambara/factoextra) using a non-supervised approach resulting in 3 suggested clusters. We
 found that 4 inferred clusters split one gene cluster in a more sensible way and adjusted the number of clusters
 accordingly to four. GO analysis were conducted using the package topGO (v. 2.34.0, weight01 algorithm) (Alexa &
 Rahnenfuhrer, 2019). GO annotations were obtained through Biomart (version 2.40.0, Ensembl Plants release 43 -
- April 2019) (Durinck *et al*, 2009). Genes for pathway specific heatmaps were obtained from the corresponding GO
 term through the Ensembl Plant database.

qPCR analysis

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619 620 For the CIF2 peptide treatment, 5-days-old seedlings grown on half MS were moved to fresh 1/2 MS medium in the 621 absence or presence of 100 nM CIF2 and incubated for 30 or 120 min. Otherwise, the plants were grown with mesh. Only root parts (around 100 mg) were collected at each time point and total RNA was extracted using a Trizol-622 adapted ReliaPrep RNA Tissue Miniprep Kit (Promega). Reverse transcription was carried out with PrimeScript RT 623 Master Mix (Takara). All steps were done as indicated in the manufacturer's protocols. The qPCR reaction was 624 performed on an Applied Biosystems QuantStudio3 thermocycler using a MESA BLUE SYBR Green kit 625 626 (Eurogentech). All transcripts are normalized to *Clathrin adaptor complexes medium subunit family protein* 627 (AT4G24550) expression. All primer sets are indicated in star methods.

629 Statistical analysis for experiments

All statistic-related analysis was done with R software(R Core Team, 2013)(<u>https://www.r-project.org/</u>). For multi-

631 comparison analysis, one-way ANOVA was carried out and Tukey's test was subsequently performed.

632 Data and materials availability

- All data to support the conclusions of this manuscript are included in the main text and supplementary materials.
- The full RNA seq dataset was deposited in GEO (accession GSE144182,
- 635 <u>https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE144182</u>).

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648	experiments. S.F., D.D.B., K.H.E, P.K, T.G.A, V.D.T, A.P., P.M., R.U., V.G.D., M.B. and A.C.
649	performed the experimental work. D.D.B., T.G.A., E.S.S. and J.D. performed image quantification
650	and RNA-seq analysis. S.F., and N.G. wrote the manuscript. G.I., J.K. and all other authors revised
651	the manuscript and were involved in the discussion of the work.
652	
653	Conflict of interest: The authors declare no competing interests.
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655	REFERENCES
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659	
660	Supporting information
661	Appendix file
662	Tables EV1-2
663 664	Movie EV1



Figure1:

666 Fig. 1 Apolar SGN1 leads to ectopic lignin accumulation in endodermal cells

(A) Schematic of Casparian strip development (magenta). Casparian strips start to appear as centrally aligned
 discontinuous dots in the endodermal cell layer, progressing into a network of fused rings functioning as a root
 apoplastic barrier.

672 (B) Lignin accumulation patterns at endodermal surface or median positions with or without the 100 nM CIF2 673 (Casparian strip integrity factor 2) ligand treatment. Lignin and cellulosic (unmodified) cell walls are stained 674 with Basic Fuchsin and Calcofluor White, shown in magenta and white respectively. Schematics are indicating 675 the position of optical sections in a 3D illustration. For each condition, at least 10 roots were tested and showed 676 similar results in two independent experiments. White arrows indicate sites of excess lignification on the cortex 677 facing (outer) side. Scale Bar = 5 μ m.

679 (C,D) Localization of SGN1-Citrine and lignin deposition patterns in pCASP1::SGN1-Citrine lines in wild-type 680 (Col) and different mutant backgrounds (*sgn1, sgn3, cif1 cif2*) (C). myrpalm-SGN1-Citrine localization and 681 lignin deposition patterns in pCASP1::myrpalm-SGN1-Citrine lines (D). Lignin (Basic Fuchsin) and cell walls 682 (Calcoflour White) are shown in magenta and white respectively. For this experiment, two or three independent 683 lines were tested. From each transgenic line, 2 positions from 12 roots were observed and representative pictures 684 are shown in the figure. Schematics are indicating the position of optical sections in a 3D illustration. White 685 arrows in (D) highlight excess lignification on the pericycle-facing (inner) side. Scale bars = 5 μ m.

(E) Schematic illustrating how signal activation can be governed by SGN1 localization and peptide liganddiffusion from the stele.

Data information: "inner" designates the stele-facing endodermal surface, "outer", the cortex-facing surface.

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Figure2

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Fig. 2 SGN1 acts as a transducer of CIF2 signaling and is phosphorylated by the SGN3 receptor

695 (A) Quantification of defects in CSD formation as number of holes per 100 μ m in the CASP1-GFP domain at 696 around 10 cells after onset of CASP1-GFP expression in 5-day-old seedlings. In the box plot, boxes indicate 697 ranges from first to third quartiles, bold central lines display median. Upper and lower whiskers extend to 698 maximum or minimum values no further than 1.5 times IQR (interquartile range, the distance between the first 699 and third quartiles). One-way ANOVA was performed followed by Tukey's test. Different letters show significant 691 statistical differences. (p<0.05, One-way ANOVA and Tukey's test, 12 roots in total were observed for each 692 condition in two independent assays).

703(B) Propidium iodide (PI) penetration assay in the presence or the absence of CIF2. CS barrier function was704scored as exclusion of PI signal from the inner side of endodermal cells. In the box plot, boxes indicate ranges705from first to third quartiles, bold central lines display median. Upper and lower whiskers extend to the maximum706or minimum values no further than 1.5 times IQR. Different letters show significant statistical differences.707(p<0.05, One-way ANOVA and Tukey's test. During two independent experiments, 10 roots in total were tested</td>708for each condition).

710(C) $[\gamma^{-32}P]$ ATP radioactive *in vitro* kinase assay of SGN3 kinase domain against SGN1. Autoradiograph is shown711on top. Coomassie stained-gel below illustrates presence and equal loading of recombinant proteins. Note that a712kinase-dead SGN1 variant was used to avoid auto-phosphorylation activity of SGN1. Also note that trigger factor713represents a very big tag protein, accounting for the high migration of TF-SGN1. Representative result of three714independent experiments is shown.



Figure3

Fig. 3 Both RBOHD and F are required for CIF2-induced excess lignin accumulation

718 (A) Localization of Citrine-RBOHD (left) or Citrine-RBOHF (right) in endodermal (en) cells. Both proteins 719 were expressed under the control of pCASP1, an endodermis-specific promoter. Representative pictures are 720 shown, 2 positions from 10 roots for each transgenic line were inspected. Scale bar = $5 \mu m$.

722(B) Lignin accumulation in WT and *rbohD and rbohF* single mutants and a double mutant with or without 2-723hour 100 nM CIF2 peptide treatment. Arrowheads indicate excess lignification. Pictures are shown as overviews724(maximum projection) or median sections. Lignin and cell walls are shown with magenta (stained with Basic725Fuchsin) and gray (stained with Calcofluor White) respectively. Representative pictures are shown, 12 roots726(overview) and 2 positions in 12 roots (median section) were inspected. Scale bars = 20 μ m (lignin overviews),7275 μ m (median sections). "inner" designates the stele-facing endodermal surface, "outer", the cortex-facing728surface.

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Fig. 4 ROS production is enhanced by SGN3/CIFs and requires RBOHD and F

(A) Co-visualization of pCASP1::SGN1-mCherry and pCASP1::SGN3-GFP. Note that their respective localization at the PM has only restricted overlap at the cortical side of the CS domain (Arrow). Representative picture is shown, 3-6 roots in 4 different transgenic lines were inspected. Scale bar = 10 µm. "inner" designates the stele-facing endodermal surface, "outer", the cortex-facing surface.

(B,C) Overview of endodermal cells after with or without 1 µM CIF2 treatment for 24 hours. Red arrowheads
are indicating ROS production sites and boxes in dotted lines are corresponding to the regions in (sky blue and
creme boxes in C and black boxes in D). Similar patterns were obtained in 39 or 41 cells from 5 roots with or
without the peptide treatment. Scale bars = 500 nm.

(D) In situ H₂O₂ detection at Casparian strips in WT, sgn3, sgn1, rbohF, rbohD and rbohDF with or without 24hour treatment of 1 μ M CIF2. Brackets and arrowheads indicate Casparian strips (seen as uniformly whitish cell wall areas) and H₂O₂ production sites (black area) respectively. Scale bar = 1000 nm.

(E) Quantification of ROS production as a number of dark pixels area (n = 23 - 44 sites from 5 roots of each condition in one experiment.). For quantification methods, see fig. EV 3. In the box plot, boxes indicate ranges from first to third quartiles, bold central lines display median. Upper and lower whiskers extend to the maximum or minimum values no further than 1.5 times IQR. Different letters mean significant statistical differences. (p<0.01, One-way ANOVA and Tukey's test.)

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7 Fig. 5 SGN1 directly activates NADPH oxidases in a cellular context

759(A) $[\gamma^{-32}P]$ ATP radioactive *in vitro* kinase assay of TF-SGN1 against GST-N-terminal cytoplasmic domains of760RBOHD or F. Autoradiograph is shown on top. Coomassie stained-gel below illustrates presence and equal761loading of recombinant proteins. Experiments were done independently three times with similar results.

(B) HEK293Tcell based NOX activation assay. Cells were transfected with the indicated plasmid combinations.
The phosphatase inhibitor Calyculin A was added directly before the start of the measurements. Each data point represents the mean of six wells analyzed in parallel, bars indicate S.D.. Experiments were repeated three times (another set of results is shown in Fig. EV 4B).



Figure 6

771 Fig. 6 CIF2 induces large-scale transcriptional changes for cell wall remodeling

(A) CASP1-GFP and lignin deposition in WT, *sgn3*, *rbohD*, *rbohF* and *rbohDF*. CASP1-GFP and lignin (fuchsin) are presented in green and magenta respectively. Pictures were obtained from more than 10 roots from each background with similar results. Scale bar = $10 \mu m$.

777(B) Time lapse imaging of single- or cotreatment of 10 nM CIF2 with 25 μ M cycloheximide (CHX) on CASP1-778GFP in *cif1 cif2*. Seedlings were pretreated with or without CHX for 30 min and transferred onto each medium.779Scale bar = 10 μ m (See also movie EV1).

(C) Quantification of (B). Relative numbers of holes in CASP1-GFP domain after single- or co-treatment with
 CIF2 or CHX from the pictures in (B). Bars are S.D. * indicates statistical significance from all other conditions
 (p<0.01) after one-way ANOVA and Tukey test. 6 roots in total for each condition were observed during two
 independent tests.

786(D) Fold-change of 930 genes (p<0.05 and $log_2(Fold change) \ge 1$ or ≤ -1 at least one time point in one genotype)787after CIF2 treatment at indicated time points in WT, *cif1,2* and *sgn3*. Degree of the fold changes are shown in788color code as indicated.

790(E) Relative expression levels of *PER15* to *CLATHRIN* control in each genotype with or without 2-hour CIF2791treatment. Bars are S.D. (n=3). Different characters indicate statistically significant differences (p < 0.01,792ANOVA and Tukey test).

- (F)Fold changes of *PER15* in each genotype with or without 2-hour CIF2 treatment. Bars are S.D. (n=3). Different characters indicate statistical significance differences (p < 0.01, ANOVA and Tukey test).
- 797(G) Relative expression levels of *PER49* to *CLATHRIN* control in each genotype with or without 2-hour CIF2798treatment. Bars are S.D. (n=3). Different characters indicate statistically significant differences (p < 0.01,799ANOVA and Tukey test).

801 (H) Fold changes of *PER49* in each genotype with or without 2-hour CIF2 treatment. Bars are S.D. (n=3). 802 Different characters indicate statistically significant differences (p < 0.01, ANOVA and Tukey test). 803

804 (I)Relative fold changes of *PER15* in *pCASP1::SGN1-Citrine* and *pCASP1::myrpalm-SGN1-Citrine* compared 805 to the expression level in WT. Bars are S.D. (n=3). Different characters indicate statistically significant 806 differences (p < 0.01, ANOVA and Tukey test).

808 (J)Relative fold changes of *PER49* in *pCASP1::SGN1-Citrine* and *pCASP1::myrpalm-SGN1-Citrine* compared 809 to the expression level in WT. Bars are S.D. (n=3). Different characters indicate statistically significant 810 differences (p < 0.01, ANOVA and Tukey test).

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Figure 7 Overview of the plasma membrane-based and nuclear branches of the SGN3 pathway

Fig. 7 Overview of the plasma membrane-based and nuclear branches of the SGN3 pathway

A, B (A) Schematic of the spatially-restricted activation of ROS production at the plasma membrane after CIF stimulation (B) Comparison of the components of the cytoplasmic/nuclear signaling cascades induced by the flg22 bacterial pattern peptide (left) and the CIF peptides (right). Note that all of the signaling components in the two pathways belong to the same gene families, with the exception of the transcription factors. Components for which there is currently no direct experimental evidence are marked in gray, as are arrows indicating activation events that have not yet been experimentally established.

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822 **Table EV1.**

- ⁸²³ Differentially expressed genes from heatmap presented in Fig.6D
- 824 **Table EV2.**
- A list of enriched GO terms of clusters presented in Fig.6D

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