Metal substituents can be used to tune the physical properties of these Li-rich phases because they affect the (O₂)ₓ⁻³ stability against oxygen recombination or voltage fade, as previously demonstrated for Li₂Ru₁₋ₓMₓO₃ (M = Sn, Ti, Mn). The benefits of Sn, in that it limits both O₂ release and voltage fade, are preserved in the Li₂IrₓSnₓO₃ system but are mitigated by the emergence of a capacity fade mechanism that is linked to the emergence and accumulation of stacking faults. This finding emphasizes that the origins of voltage and capacity fading in these Li-rich layered phases are different, a point that has previously been a source of confusion.

In summary, combined TEM, neutron diffraction, and ab initio studies on high-capacity Li-rich LiₓIrₓSnₓO₃ layered phases permitted the atomic-scale visualization of the (O₂)ₓ⁻³ peroxo-like dimers responsible for the capacity gain in Li-rich layered electrode materials. These observations lead to a better understanding of peroxo formation and localization, O₂ recombination, and the effect of the transition metal substituents. Additionally, these findings provide a chemical handle for tuning the performances of Li-rich layered materials.

PLANT SYMBIOSES

Rice perception of symbiotic arbuscular mycorrhizal fungi requires the karrikin receptor complex

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In terrestrial ecosystems, plants take up phosphate predominantly via association with arbuscular mycorrhizal fungi (AMF). We identified loss of responsiveness to AMF in the rice (Oryza sativa) mutant hebbia, reflected by the absence of physical contact and of characteristic transcriptional responses to fungal signals. Among the 26 genes deleted in hebbia, DWARF 14 LIKE is, the one responsible for loss of symbiosis. It encodes an alpha/beta-fold hydrolase, that is a component of an intracellular receptor complex involved in the detection of the smoke compound karrikin. Our finding reveals an unexpected plant recognition strategy for AMF and a previously unknown signaling link between symbiosis and plant development.

REFERENCES AND NOTES

22. Materials and methods are available as supplementary materials on Science Online.

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SUPPLEMENTARY MATERIALS

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Most land plants establish symbioses with arbuscular mycorrhizal fungi (AMF) of the phylum Glomeromycota (1). These symbioses contribute to global carbon and mineral nutrient cycles, because AMF provide mineral nutrients to the plant and receive carbohydrates in return. Colonization of plant roots by AMF requires reciprocal recognition initiated by diffusible molecules before fungal attachment to the root surface and root penetration via hyphopodia (2). Diffusible recognition signals include strigolactones, released from plant roots that activate the fungus before physical interaction (3), and fungal (lipo)chitooligosaccharides and chitotetraose, secreted by AMF that trigger plant calcium signaling, gene expression, and lateral root formation (4, 5). Plant LysM receptor-like kinases (6) are required for perception of chitinous microbial molecules that trigger either symbiosis or defense signaling (7–9). Plant signaling mutants impaired in root colonization by both AMF and nitrogen-fixing bacteria still exhibit transcriptional responses to fungal signaling molecules (10–12). Therefore, additional signaling modules have been postulated (12). We identified the rice receptor for karrikin, a plant growth regulator first identified in smoke (13–16), as a necessary signaling component for establishment of arbuscular mycorrhizal (AM) symbiosis. We found that the jasmonate-deficient rice (Oryza sativa) mutant hebbia was unable to
establish symbiosis with either of two AM—
*Rhizophagus irregularis* and *Gigaspora rosea*—as reflected by the absence of hyphopodia, intraradical colonization, and induction of colonization marker genes (Fig. 1, A to C) (10). The lack of fungal interaction persisted upon increased inoculum strength imposed by growing *hebiba* alongside colonized wild-type plants (Fig. 1D). This suggested that the mutant is compromised at a very early stage of the interaction, during presymbiotic signaling.

The *hebiba* mutant is due to a genomic deletion of 169 kb, which contains 26 annotated genes (17, 18). One of the genes encodes allene oxide cyclase (AOC), part of the jasmonate synthetic pathway, loss of which leads to jasmonate deficiency (17). However, transgenic complementation of *hebiba* with *AOC* (*hebibaAOC*) did not restore AM symbiosis (fig. S1) (17, 19). Therefore, another gene contained within the deleted interval must be required for AM development.

We identified the gene responsible for AM symbiosis by transforming *hebibaAOC* with genomic clones of individual genes from the deleted interval (table S1) (17, 18). Reintroduction of the *LOC_Os03g32270* gene restored fungal colonization of *hebibaAOC* roots in independent rice transformants (Fig. 2, A and B, and table S1). Quantitative measurements of colonization correlated ($R^2 = 0.84$) with the amount of transcript accumulation from the *LOC_Os03g32270* transgene (Fig. 2C). Transgenic lines such as C10 (Fig. 2B), with transgene mRNA levels below the detection limit, retained the *hebiba* mutant phenotype. *LOC_Os03g32270* encodes the alpha/beta-fold hydrolase DWF4/HLIKE (D14L), homologous to *Arabidopsis thaliana* KARRIKIN INSUSCEPTIVE2/HYPOSENSITIVE TO LIGHT (KAI2/HTL). This hydrolase acts together with the F-box protein DWF9/MORE AXILLARY GROWTH2 (D3/MAX2) in the perception of karrikins, a group of butenolide compounds found in smoke that induce seed germination in fire-chasing plants (13–16). The structurally related strigolactones are perceived by a receptor complex involving D3 and the alpha/beta-fold hydrolase DWF14 (D14), the paralog of D14L (D3/MAX2) (17–19). However, the strigolactone-insensitive rice mutant *d14* is not perturbed in AM symbiosis (23) (Fig. 3A); thus, the strigolactone receptor gene D14 is not required for establishment of the interaction. A rice d3 mutant was also severely impaired in AM colonization and marker gene induction (Fig. 3, A and B) (23), revealing the importance of the karrikin receptor complex for the earliest stages of AM development. We further confirmed the requirement of D14L in AM development by using a set of RNA interference (RNAi) lines generated in the *Oryza sativa* cv. Nipponbare background. The RNAi lines displayed diverse levels of AM suppression that correlated ($R^2 = 0.69$) with the degree of down-regulation of endogenous *LOC_Os03g32270* (fig. S2, A to C). The D14L RNAi line R43 supports AMF colonization (23); however, we found a decrease ($P = 0.047$) in total fungal colonization relative to wild type in this line. The phenotypic diversity among the D14L RNAi lines suggests a low transcript threshold for AM symbiosis establishment.

In *Arabidopsis*, KAI2/HTL controls hypocotyl elongation in response to light and karrikin (33, 24). Overexpression of rice D14L in an *Arabidopsis* htl-2 mutant restored wild-type hypocotyl length in two independent F$_2$ populations homozygous for htl-2 (fig. S3A). Mesocotyl elongation assays in rice demonstrated that *hebibaAOC* is insensitive to karrikin but responds to the synthetic strigolactone GR24 (fig. S3B). In contrast, mutations of D14 specifically compromised strigolactone but not karrikin responses in rice, whereas mutation of the F-box protein encoding D3 resulted in insensitivity to both (fig. S3B). Thus, in rice, D14L and D14 mediate perception specificity to karrikin versus strigolactone in an overall similar manner to *Arabidopsis* (13). However, the partial response of *Arabidopsis* d14 to racemic...
GR24 (J3, 25) was not reproduced in rice di4 mutants (fig. S3B) (26), suggesting that D14L has less redundant activity in rice. Fluorescently tagged D14L in Arabidopsis (24) and rice localized to both nucleus and cytoplasm (fig. 2E). D14L in rice (fig. 2D) as in Arabidopsis (24) is expressed in all rice organs, and transcript accumulation in roots is not altered during AM colonization.

We asked whether D14L is required for suppression of defense responses against AMF. We found no evidence for increased activation of selected defense marker genes (27) during the early stages of mycorrhizal colonization (fig. S4, A and B). Moreover, hebibaAOC was susceptible to colonization by the root endophyte Piriformospora indica and the pathogen Magnaporthe oryzae (fig. S4, C and D), implicating D14L in symbiotic compatibility. On the basis of the early and pronounced complementation lines. Values represent means and SEs from two to five replicate plants. (C) D14L transcript levels were assessed by real-time RT-PCR in the independent transgenic complementation lines. The averages for the WT, hebibaAOC, and the complementation hebibaAOC,D14L lines were plotted against the corresponding averages for total RLC. The Spearman rank correlation was calculated and squared to give the proportion of the variation accounted for by the correlation. (D) Real-time RT-PCR—based expression of D14L in control root (C), mycorrhizal roots (M), stem, leaf, panicle, and embryophy of Nipponbare rice. Expression values were normalized to those of the constitutively expressed gene Cyclophilin2 (LOC_Os02g02890). Means and SDs of three technical replicates are shown. (E) Subcellular localization of D14L. A plasmid containing a D14L overexpression construct (i) maize ubiquitin promoter:D14L cDNA:green fluorescent protein was co-transfected with the plasmid containing a genomic clone of D14L driven by its native promoter (ii) pD14LgD14L:RFP in rice root protoplasts. (iii) DAPI staining. (v) Overlay of all channels, including bright field (iv). Scale bar, 10 μm.

representative genes was validated by quantitative reverse transcription polymerase chain reaction (RT-PCR) on the same RNA used for the RNA-seq experiment (fig. S5A) and on RNA from two additional biological replicates, which included the complemented line C11 (fig. S5B). Thus, D14L is required to support initial colonization events by AMF. Despite its effect on mesocotyl elongation, treatment with karrikin did not induce significant gene expression changes in roots of wild-type rice (table S4). Also, the exogenous application of karrikin did not stimulate colonization of wild-type roots by R. irregularis (fig. S6).

We found that a total of 104 transcripts differed significantly between untreated hebibaAOC and wild-type roots (table S4) derived from genes with borderline GO-term enrichment for metabolic processes (table S6C). Whereas mRNA levels of known genes essential for AM symbiosis accumulated independently of functional D14L, transcript levels of the rice homolog of DWARF14 LIKE 2 (35), LOC_Os05g51240, depended on D14L, as earlier observed in Arabidopsis (J3) (table S4). In the wild-type relied on functional D14L. Time-resolved gene ontology (GO) analyses of genes differentially regulated in response to GSEs in wild type but not in hebibaAOC demonstrated an overrepresentation of terms associated with responses to extracellular and biotic stimuli. Genes were induced or repressed at the earliest time points, 3 and 6 hpt, and in a D14L-dependent fashion, consistent with D14L playing a role in early signaling activation (fig. 4B and table S6, A and B). The expression pattern of (RNA-seq) to monitor the transcriptional changes of hebibaAOC and wild-type roots in response to GSEs over the first 24 hours post treatment (hpt) (supplementary materials, tables S2 and S3). Overall 140 genes showed statistically significant differences in average expression upon GSE treatment in wild-type plants (Fig. 4A and tables S4 and S5). In hebibaAOC plants, six genes responded significantly to GSEs, of which two genes (predicted to encode an expressed and a hypothetical protein) overlapped with the genes responding in wild type (fig. 4A and table S4), suggesting that the transcriptional response observed in the wild-type relied on functional D14L. 

Fig. 2. D14L is required for AM development. (A to C) AM phenotype of transgenic hebibaAOC,D14L complementation lines. (A) Trypan blue—stained roots at 6 wpi with R. irregularis: micrographs refer to (from left to right) WT Nihonmasari, hebibaAOC mutant, and two independent transgenically complemented hebibaAOC,D14L lines (C4 and C11). Scale bar, 50 μm. (B) RLC expressed as % of WT colonization at 6 wpi for independent hebibaAOC,D14L complementation lines. Values represent means ± SEs from two to five replicate plants. (C) D14L transcript levels were assessed by real-time RT-PCR in the independent transgenic complementation lines. The averages for the WT, hebibaAOC, and the complementation hebibaAOC,D14L lines were plotted against the corresponding averages for total RLC. The Spearman rank correlation was calculated and squared to give the proportion of the variation accounted for by the correlation. (D) Real-time RT-PCR—based expression of D14L in control root (C), mycorrhizal roots (M), stem, leaf, panicle, and embryophy of Nipponbare rice. Expression values were normalized to those of the constitutively expressed gene Cyclophilin2 (LOC_Os02g02890). Means and SDs of three technical replicates are shown. (E) Subcellular localization of D14L. A plasmid containing a D14L overexpression construct (i) maize ubiquitin promoter:D14L cDNA:green fluorescent protein was co-transfected with the plasmid containing a genomic clone of D14L driven by its native promoter (ii) pD14LgD14L:RFP in rice root protoplasts. (iii) DAPI staining. (v) Overlay of all channels, including bright field (iv). Scale bar, 10 μm.
Fig. 3. AM phenotype of d3 relative to hebibaAOC, d4, and corresponding WT cultivars. (A) Percentage of RLC and (B) induction of AM early marker genes at 7 wpi with R. irregularis of d3, d4, hebibaAOC, hebibaAOC-d4Al complementation line C2, and corresponding WT background Nihonmasari (Nih0) and Shiokari (Shi0), respectively. Expression values were normalized to those of the constitutively expressed gene Cyclophilin2 (LOC_Os02g02890). Values represent means and SEs from three biological replicates (A and B). The mutants and complementation line were compared with the appropriate WT by using the Kruskal-Wallis test. Symbols above the bars indicate statistical significance with respect to the WT (\(P < 0.10\); * \(P < 0.05\)).

Fig. 4. GSE-induced transcriptional responses of WT and hebibaAOC. (A) Venn diagram depicting the number of transcripts induced (black) and repressed (white) in WT and hebibaAOC plants treated with GSEs in comparison to plants receiving a mock treatment. (B) Time-resolved GO-term enrichment analysis (P ≤ 0.001) for genes differentially regulated in response to GSEs in WT but not in hebibaAOC. The color code represents odds ratios.

contrast to Arabidopsis, karrikin treatment of rice roots did not induce this gene. Because D14L is found in genomes of plants that germinate without fire stimulation and because Arabidopsis mutants lacking D14L show developmental phenotypes, we hypothesize that an endogenous ligand exists and is required for wild-type seedling development (29). In rice, the differences in transcripts between GSEs and mock or karrikintreated wild-type plants indicate either that this ligand is not karrikin or that D14L acts upstream of the GSE response, thereby possibly creating a condition permissive for AM symbiosis.

We show that the karrikin receptor complex is central to the everyday interaction of plants with AMF, involving more than 80% of all plant species as opposed to 1200 smoke-responsive plant species (30). Conservation of D14L in early land plants, such as liverworts (31), suggests that it has served this purpose since AMF started supporting terrestrial plant life. On poor natural supply and need to coordinate AMF development with their physiological and developmental needs. The karrikin receptor complex may represent a node in the cross-talk between plant development and AM signaling.

REFERENCES AND NOTES

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