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TOR dynamically regulates plant cell–cell transport

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Contributed by Patricia Zambryski, January 13, 2020 (sent for review November 4, 2019; reviewed by Mary Lou Guerinot and Robert Turgeon)

The coordinated redistribution of sugars from mature “source” leaves to developing “sink” leaves requires tight regulation of sugar transport between cells via plasmodesmata (PD). Although fundamental to plant physiology, the mechanisms that control PD transport and thereby support development of new leaves have remained elusive. From a forward genetic screen for altered PD transport, we discovered that the conserved eukaryotic glucose-TOR (TARGET OF RAPAMYCIN) metabolic signaling network restricts PD transport in leaves. Genetic approaches and chemical or physiological treatments to either promote or disrupt TOR activity demonstrate that glucose-activated TOR decreases PD transport in leaves. We further found that TOR is significantly more active in mature leaves photosynthesizing excess sugars than in young, growing leaves, and that this increase in TOR activity correlates with decreased rates of PD transport. We conclude that leaf cells regulate PD trafficking in response to changing carbohydrate availability monitored by the TOR pathway.

plasmodesmata | cell–cell signaling | rapamycin | Reptin | TARGET OF RAPAMYCIN

Plant cells are connected by plasmodesmata (PD), nanoscopic membrane-bound channels in the cell wall that permit molecules up to ~80 kDa to traffic between neighboring cytoplasts (1–3). PD are absolutely essential for plant development and physiology, and their importance is reflected in the repeated, parallel evolution of PD or analogous structures in every lineage of complex multicellular eukaryotes that have cellulosic cell walls (4, 5). The upstream signal transduction pathways that regulate PD transport in plants remain poorly understood, however. We and others discovered that chloroplasts can regulate the rate of PD transport and the biogenesis of PD in pathways that we named “organelle-nucleus-plasmodesmata signaling” (6–11). When plants undergo developmental transitions, such as during lateral root formation, PD transport is often restricted, which is proposed to isolate cells during differentiation and allow signaling molecules (such as phytohormones and transcription factors) to accumulate to high concentrations (3, 12–16). PD transport is also tightly regulated during various stresses, especially upon recognition of infection by a pathogen (5, 17–21).

During plant development, plants redistribute sugars photosynthesized in mature leaves (“sources”) to support the development of young leaves and roots (“sinks”). Tracheophytes traffic sugars via the phloem, specialized vascular tissue that concentrates high levels of sugars for long-distance transport through the plant body. As leaves develop, they switch from importing nutrients out from the phloem (sinks) to exporting sugars into the phloem (sources), a physiological shift that is called the “sink-to-source” transition. Among different plant species, there is considerable variation in how sugars are loaded from the photosynthetic mesophyll into the phloem; here, we will focus on herbaceous plants, which typically use an “apoplastic” loading mechanism. In source leaves of apoplastic loaders, sugars are loaded against a concentration gradient into the phloem companion cells by active transporters in the plasma membranes (22–25). To prevent passive backflow of sugars from the vasculature into the leaf, PD transport is tightly restricted in source

leaves (1). Phloem loading has a dual effect: The concentration of solutes in the phloem increases, creating hydrostatic pressure that can promote phloem transport, and perhaps more importantly, the concentration of free sugars in the source leaf is reduced (i.e., export of photo-assimilated carbohydrates from the source to growing sink tissues is maximized) (26). In contrast to source leaves, PD transport is unrestricted in young sink leaves, which allows rapid, passive transport of sugars out from the vasculature into the growing leaf (27–29). Although this change in PD transport as leaves age has been recognized for several decades, little is known about how PD transport between cells is coordinated with leaf development and physiology during the sink-to-source transition.

We conducted a screen to identify mutants defective in regulating PD transport, using *Arabidopsis thaliana* embryogenesis as a model system (30). Four of these mutants, *ise1* through *ise4*, do not restrict PD transport at the midtorpedo developmental stage of embryogenesis, which is when wild-type embryos become photosynthetic and PD transport rapidly decreases (30). We previously characterized *ise1* (*At1g12770*) and *ise2* (*At1g70070*), which disrupt chloroplast biogenesis and trigger a retrograde signaling pathway that controls intercellular trafficking (6). Here, we characterize *ise3* (*At2g25570*) and *ise4* (*reptin-1*, *At5g67630*), two mutants that disrupt the TARGET OF RAPAMYCIN (TOR) signaling network.

Significance

Plant cells are connected to each other by plasmodesmata (PD), membrane-bound cytosolic channels that cross the cell wall. Diverse molecules, ranging from metabolites to small RNAs and proteins, can traffic between plant cells via PD. During development, PD transport is regulated to control the redistribution of carbohydrates from mature leaves that photosynthesize excess sugars to young leaves that rapidly consume sugars. Here, we show that the ancient eukaryotic TARGET OF RAPAMYCIN (TOR) metabolic signaling network controls PD transport in plants. TOR is a protein kinase that is activated by nutrients, including sugars, and then coordinates growth with nutrient availability. Our study demonstrates that TOR has evolved new roles in plants by regulating cell–cell connectivity via PD.

Author contributions: J.O.B. and P.Z. designed research; J.O.B., M.X., M.R.S., S.C., E.A.S., H.M.G., and P.Z. performed research; J.O.B., M.X., M.R.S., S.C., E.A.S., H.M.G., and P.Z. analyzed data; and J.O.B. and P.Z. wrote the paper.

Reviewers: M.L.G., Dartmouth College; and R.T., Cornell University.

The authors declare no competing interest.

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Data deposition: The sequence reported in this paper has been deposited in the NCBI Sequence Read Archive, <https://www.ncbi.nlm.nih.gov/sra> (BioProject PRJNA600773).

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TOR is an essential, deeply conserved serine/threonine protein kinase that coordinates metabolism with nutrient availability in all eukaryotes (31–33). When nutrients are available, TOR promotes growth and anabolism, but when nutrients are scarce, TOR activity is suppressed and cells become quiescent, using catabolic pathways, such as autophagy, for sustenance (34). TOR is widely studied in biomedical fields, because mutations in TOR and its interacting partners can cause or contribute to a range of diseases, including many cancers (34, 35); much less is known about TOR signaling in plants. TOR is active in complex with at least two other conserved proteins, RAPTOR and LST8, to make the TOR complex 1 (TORC1). In its active form, TORC1 is a dimer; dimerization is promoted by a cochaperone complex called R2TP-TTT (Telo2-Tti1-Tti2) in association with the well-studied protein chaperone, HSP90. R2TP-TTT, named for its yeast components (Pontin/RuvBL1, Reptin/RuvBL2, Spaghetti/Tah1, and Pih1) compose the R2TP subcomplex; Telo2, Telo2-interacting protein 1, and Telo2-interacting protein 2 compose the TTT subcomplex, is proposed to act as an energy sensor upstream from TOR: Pontin and Reptin form an ATPase in R2TP-TTT that drives TORC1 dimerization when ATP is plentiful, but cannot dimerize TORC1 when ATP is scarce (36). When TORC1 is active, it can phosphorylate a number of downstream targets, including a phosphorylation cascade mediated by the TOR substrate S6K (ribosomal protein eS6 Kinase) that can promote cell division, cell growth, and protein translation, among diverse additional metabolic pathways.

Much less is known about TOR signaling in plants. TOR is essential for plant development: *tor* mutants are lethal at very early stages of embryogenesis (37) and strong inhibition of TOR through chemical inhibition or RNAi causes plant growth to arrest (38, 39). As in other eukaryotes, plant TOR directly phosphorylates S6K (there are two tandem S6K paralogues in *A. thaliana*, named S6K1 and S6K2) at a conserved serine (pT389 in humans, pT449 in *A. thaliana* S6K1), and S6K-pT449 levels are therefore a reliable marker for TOR activity in plants (38). A seminal study discovered that photosynthesis-stimulated TOR activity is required for seedling establishment in *A. thaliana* (40), and then took advantage of this physiological transition to clearly define a set of ~2,000 genes that are differentially expressed specifically in response to glucose activation of TOR in plants. Interactomics have demonstrated that TOR closely associates with many of the R2TP-TTT subunits (41), and in maize, loss of one of the TTT subunits causes embryo lethality and a drastic reduction in TOR levels, presumably due to protein instability when TOR is not chaperoned by R2TP-TTT to form the TORC1 dimer (42). The biological functions of plant TOR remain to be elucidated, however, and much less is known about the roles of TOR at later developmental stages (such as in mature leaves). Here, we show that TOR has evolved a role in plants to regulate PD transport and the sink-to-source transition in leaves.

Results and Discussion

The R2TP Complex Regulates PD Transport. *ise4* is an embryo-lethal mutant that we identified in a screen for increased PD transport. For these experiments, PD transport is assayed by the movement of a 10-kDa fluorescent dextran tracer that does not readily move between cells at the midtorpedo stage in wild-type, but does move throughout the midtorpedo embryos in *ise* mutants (Fig. 1A) (30). In the siliques of *ise4/+* heterozygous parents, ~25% of embryos arrest at the midtorpedo stage and exhibit increased PD transport (Fig. 1A). *ise4* maps to a single nucleotide missense mutation in *At5g67630*, which encodes the *Arabidopsis* ortholog of Reptin (also called RuvBL2/RVB2) (*SI Appendix, Fig. S1A* and *Dataset S1*). Genetic complementation confirmed that this *At5g67630* allele causes the PD transport phenotype, because transformation with the native promoter and genomic sequence of *At5g67630* rescued the *ise4* mutants. We therefore renamed *ise4* for clarity,

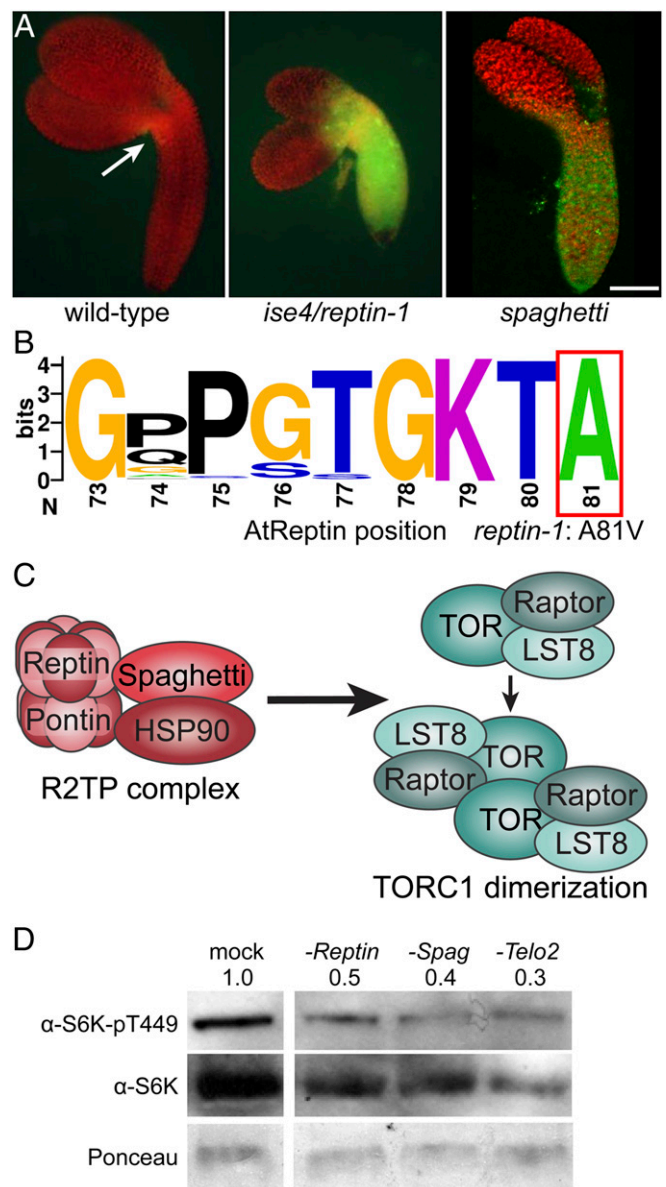


Fig. 1. R2TP cochaperone complex activity restricts cell-cell transport during embryogenesis. (A) Midtorpedo-stage wild-type, *ise4/reptin-1*, and *spaghetti Arabidopsis* embryos were removed from seed coats in a 10 kDa FITC-dextran solution, which loaded into plants at the hypocotyl-cotyledon boundaries (indicated in wild-type with a white arrow). After several washes to remove external FITC-dextran, embryos were observed using fluorescence microscopy (FITC, green; chlorophyll, red). The 10 kDa FITC-dextran rapidly moves through *ise4/reptin-1* (Center) and *spaghetti* (Right), but does not move between cells in their wild-type siblings (Left). (Scale bar, 100 μ m.) (B) Consensus Walker A motif across all eukaryotic Reptin orthologs. Amino acid positions follow *AtReptin*. *reptin-1* causes an A81V missense mutation. (C) R2TP is a cochaperone complex composed of Reptin, Pontin, and Spaghetti, which recruits HSP90. R2TP stabilizes the TORC1 (TORC1: dark teal, Raptor; teal, TOR; light teal, LST8) dimer. (D) Knocking down expression of R2TP-TTT cochaperone subunits *Reptin*, *Spaghetti*, or *Telo2* in *Arabidopsis* Col-0 with VIGS strikingly decreased levels of the TOR-dependent phosphopeptide S6K-pT449, reflecting significant inhibition of TOR activity. Total S6K levels and Ponceau staining were used for comparison. Note that steady-state levels of total S6K are also somewhat lower in *reptin*, *spaghetti*, and *telo2* relative to Col-0, as expected in plants with chronic low TOR activity.

and refer to this allele of *AtReptin* as *reptin-1* throughout. An apparent paralogue, previously named *AtTIP49b1* (*At3g49830*) (43), is a pseudogene: It is not expressed in available *A. thaliana*

RNA sequencing (RNA-seq) data (44), we could not amplify *AtTIP49b1* transcripts by RT-PCR, and we did not observe any phenotypes in a line carrying a T-DNA insertion in the *AtTIP49b1* coding sequence (SALK_036858). Therefore, we focused exclusively on the functional *AtReptin* gene, *At5g67630*.

Reptin is an AAA+ type ATPase found in all eukaryotes that acts as a chaperone for various multiprotein complexes, and has conserved roles in diverse processes, including chromatin remodeling, cell division, telomerase biogenesis, and R2TP-TTT-mediated phosphatidylinositol 3-kinase-related kinase (PIKK) complex assembly (45, 46). The mutation in *reptin-1* is a missense A81V mutation (*SI Appendix*, Fig. S1 A and D). A81 is a universally conserved residue in eukaryotic Reptin orthologs that is immediately C terminal to the nucleotide-binding Walker A motif (Fig. 1B), hinting that *reptin-1* is defective in ATP binding or ATPase activity. For most of its roles, Reptin assembles in a heteromeric complex with a closely related AAA+ ATPase, Pontin (also called RuvBL1/RVB1). As in other eukaryotes and recently confirmed in plants (46), *AtReptin*-GFP localizes to the nucleus and cytosol, and *AtReptin* physically associates with *AtPontin* (encoded by *At5g22330*) (43, 46) (*SI Appendix*, Fig. S1 B and E). A homozygous null allele of *AtReptin* with a T-DNA insertion in the coding sequence, which we named *reptin-2*, causes female gametophyte lethality (*SI Appendix*, Fig. S1 A and C), reflecting the essential roles this protein plays in basic cellular functions.

Reptin and Pontin are the only ATPases in R2TP, an HSP90 cochaperone complex (47) that recruits HSP90 via another R2TP subunit, Spaghetti (an ortholog of human RPAP3, a tetratricopeptide-repeat/carboxylate clamp protein with a C-terminal RPAP3-specific domain; for names of other orthologs, see *Dataset S1*) (Fig. 1C) (47, 48). Although Reptin interacts with several protein complexes, Spaghetti is a component specific to the R2TP complex. Using a null allele of *spaghetti* in *Arabidopsis* (encoded by *At1g56440*) (*SI Appendix*, Fig. S2) (49), we directly tested whether R2TP specifically regulates PD transport by assaying intercellular trafficking in *spaghetti* embryos. Strikingly, PD transport is also increased in R2TP-deficient *spaghetti* mutants at the midtorpedo stage of embryogenesis (Fig. 1A). Thus, we decided to focus further analysis on the roles of R2TP-dependent pathways in regulating PD transport.

TOR Restricts PD Transport. TOR is a serine/threonine protein kinase that is universally conserved in eukaryotes (31, 32). Nutrients and various hormonal signaling pathways converge to activate TOR, which then promotes growth and anabolism (33). R2TP and the TTT complex (for names of other orthologs, see *Dataset S1*) are required to stabilize the TOR complex, and R2TP-TTT is proposed to function as a metabolic sensor of intracellular ATP levels to prevent TOR complex assembly when ATP levels are very low (36). A recent report showed that *tti2* mutants in maize are embryo-lethal and accumulate drastically lower levels of TOR kinase, reflecting TOR complex instability (42). We directly tested whether R2TP-TTT is required to maintain TOR activity in plants by probing R2TP-TTT knockdowns for differential phosphorylation of the canonical TORC1 substrate, S6 kinase (S6K), which is phosphorylated by TORC1 at T449 in *Arabidopsis* (38). S6K-pT449 levels are lower in TRV:*reptin*, TRV:*spaghetti*, and TRV:*telo2* knockdown plants than in mock (TRV:*gus*) controls (Fig. 1D), demonstrating that R2TP-TTT is required for TOR kinase activity in plants. Furthermore, we used RNA-seq to define the transcriptome of *reptin-1* mutants, and found that the *reptin-1* and *tor* transcriptional profiles are significantly similar, as discussed in more detail below (see, for example, Fig. 3 E and G). Given the strong impact of *reptin* and *spaghetti* on TOR activity, we next focused on directly testing whether TOR regulates intercellular transport in plants.

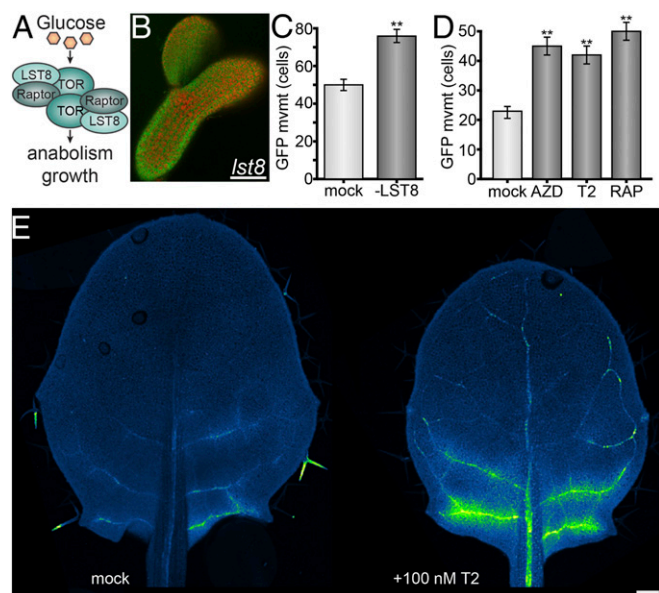


Fig. 2. TOR restricts cell-cell transport. (A) TORC1 model. TORC1 is a dimer composed of TOR, RAPTOR, and LST8. Glucose stimulates TORC1 to promote anabolism, cell division, and growth, depending on developmental context. (B) The 10 kDa FITC-dextran rapidly moves through *lst8* midtorpedo-stage *Arabidopsis* embryos (for wild-type, see Fig. 1A; FITC, green; chlorophyll, red). (Scale bar, 100 μ m.) (C) Silencing *LST8* expression with VIGS increases PD transport in *N. benthamiana* leaves ($n \geq 43$, $**P < 0.001$; error bars indicate SEM). Leaves equivalent to L8 (Fig. 5B) were used for this experiment. (D) Chemically inhibiting TOR activity with ATP-competitive kinase inhibitors [AZD-8055 or Torin 2 (39)] or rapamycin [steric inhibitor through interaction with FKBP12 (38, 75–78)] increases PD transport in *N. benthamiana* leaves ($n \geq 108$, $**P < 0.001$; error bars indicate SEM) L6 leaves (Fig. 5B) were used for this experiment. (E) Chemically inhibiting TOR activity with Torin 2 in young leaves increased import of GFP from source leaves via the phloem in *SUC2_{PRO}:GFP Arabidopsis* plants. Mock-treated leaves of the same age and size showed little to no GFP import. GFP fluorescence is shown in green; background chlorophyll autofluorescence is shown in blue; some trichome autofluorescence (not GFP) is also visible and shown in green. (Scale bar, 100 μ m.)

TOR forms a complex with two WD40-repeat-containing proteins, RAPTOR and LST8 (Fig. 2A) (37, 50, 51). We cannot directly assay PD transport in null *tor* and *raptor* mutants, because they cause complete growth arrest at early stages of embryogenesis under standard growing conditions (37, 50). *Arabidopsis lst8* is not lethal under standard growing conditions, however, although it does exhibit pleiotropic mutant phenotypes, including reduced growth and sensitivity to photoperiod (51). We found that *lst8* mutants are defective in the restriction of PD transport during embryogenesis (Fig. 2B). Unlike wild-type midtorpedo-stage embryos (Fig. 1A), 10-kDa FITC-dextran rapidly moves via PD throughout *lst8* midtorpedo-stage embryos, directly demonstrating that the TOR–RAPTOR–LST8 complex regulates cell-cell trafficking in plants.

Next, we assayed the rate of GFP movement through PD in mature *Nicotiana benthamiana* leaves after genetically or chemically disrupting TOR activity. Single epidermal cells were transformed by agroinfiltration to express GFP (27 kDa), and GFP movement from the transformed cell into neighboring cells was determined after 48 h using a quantitative assay (52, 53). For all experiments, we visualized GFP movement only in the proximal 25% of the leaf. Results of this movement assay are presented as the number of cells to which GFP spread. Silencing *LST8* significantly increased PD transport (Fig. 2C), confirming that *LST8* is required to limit intercellular transport in leaves. Treatment with rapamycin or two different ATP-competitive TOR kinase inhibitors, AZD-8055 and Torin 2, also significantly increased

PD transport of GFP (Fig. 2D). Collectively, these experiments demonstrate that active TOR restricts trafficking between plant cells.

TOR Promotes the Sink-To-Source Transition in Phloem Transport. We hypothesized that TOR-mediated restriction of PD transport corresponds to the sink-to-source transition, when tissues switch from phloem importers to phloem exporters. To directly measure phloem transport dynamics after manipulating TOR activity, we took a classic experimental approach using *SUC2_{PRO}:GFP* transgenic *A. thaliana* seedlings (28). In this line, GFP is transcribed and translated exclusively in the phloem of source leaves, but the GFP protein then rapidly moves through the phloem and accumulates in cells surrounding the veins in sink leaves. *SUC2* transcription has not been found to be regulated by TOR (40, 54). We applied 100 nM of the TOR inhibitor Torin 2 or mock treatment to the shoot apex of *SUC2_{PRO}:GFP* seedlings (with five emerged leaves), and assayed GFP accumulation in leaves transitioning from sink to source (leaf four [L4], ~1.5-mm long, which was in direct contact with the applied Torin 2) 1 d later. Across three experiments, we consistently observed GFP unloading from the phloem of leaves after treatment with Torin 2, but little or no GFP unloading from the phloem of leaves that received mock treatment (Fig. 2E). Therefore, inhibiting TOR interferes with the sink-to-source transition in leaves, such that the leaf continues to import resources from the phloem.

TOR Signaling Is Also Disrupted in *ise3*. Across eukaryotes, TOR activity is highly sensitive to mitochondrial bioenergetics and the production of ATP by oxidative phosphorylation (OXPHOS) (36, 55). In plants, glucose stimulates TOR through its metabolism via glycolysis and OXPHOS (40). We found that another mutant identified in our genetic screen for increased PD transport is also defective in TOR activity, likely due to disruption of OXPHOS in the glucose-glycolysis-OXPHOS-TOR pathway. *ise3* (Fig. 3A) is an embryo-lethal mutant identified in the same genetic screen for altered trafficking of 10 kDa FITC-dextran at the midtorpedo stage of embryogenesis that identified *ise1*, *ise2*, and *ise4* (30). We mapped *ise3* to a missense mutation in *At2g25570*, which encodes a SEL1-like repeat-containing (SLR) protein that localizes to mitochondria (SI Appendix, Fig. S3 A and D). *At2g25570* genomic DNA or *At2g25570* coding sequence expression driven by the native *At2g25570* promoter complemented the *ise3* phenotype, confirming that *ise3* is a loss-of-function allele of *At2g25570*. *ISE3* is highly conserved within land plants, but there are no clear orthologs of *ISE3* in other eukaryotes (SI Appendix, Fig. S3C). SLR proteins are typically involved in protein-protein interactions; for example, a structurally comparable mitochondrial SLR protein in humans has been implicated in the assembly of complex IV of the OXPHOS chain (56). In *Arabidopsis*, *At2g25570* (*ISE3*) cofractionates with mitochondrial complex V (ATP synthase), implying that it participates in ATP synthesis (57). The *ise3* allele is a G134E missense mutation; G134 is a consensus SLR residue, and G134E is predicted to destabilize the third SLR, presumably disrupting *ISE3*'s affinity for protein binding partners (Fig. 3 B and C and SI Appendix, Fig. S3 B and D).

We directly tested whether *ISE3* is required for TOR activity by silencing *ISE3* expression and assaying S6K-T449 phosphorylation, and found that S6K-pT449 levels are significantly lower in TRV::*ise3* knockdowns than in mock (TRV::*gus*) controls (Fig. 3D). Moreover, silencing *ISE3* significantly increases PD transport of GFP in leaves (Fig. 3F). Next, we took a comparative transcriptomic approach by comparing significantly differentially expressed genes (DEGs) of *ise3*, *reptin-1*, and *tor* (Datasets S2–S4). The *tor* transcriptome is the set of DEGs that respond 2 h after glucose activation of TOR in glucose-deprived quiescent *Arabidopsis* seedlings; these DEGs are specifically sensitive to TOR, because glucose supply has no transcriptomic effect in

these seedlings when TOR is genetically or chemically inhibited (40). As mentioned above, the *reptin-1* and *tor* transcriptomes overlapped significantly in DEGs ($P < 10^{-27}$) (Fig. 3E and Dataset S3), and the vast majority of overlapping DEGs were both induced or both repressed in the two datasets (75.6% coregulated) (Fig. 3E). Similarly, the set of DEGs in *ise3* significantly overlaps with the *tor* transcriptome ($P < 10^{-60}$, 74.2% coregulated) (Fig. 3E and Dataset S4).

More broadly, both the *reptin-1* and *ise3* transcriptomes exhibit lowered TOR activity, with high expression of genes related to catabolic recycling and quiescence, as expected for a mutant deficient in TOR activity, instead of gene-expression profiles reflecting the rapid anabolic growth typically associated with embryogenesis (Fig. 3G, SI Appendix, Fig. S4, and Dataset S2). Analysis of significantly enriched categories of DEGs in *reptin-1* revealed widespread repression of nuclear-encoded photosynthesis-associated genes and cytosolic ribosomal protein genes, alongside significant induction of genes associated with late embryogenesis and seed maturation (SI Appendix, Fig. S4). Cytosolic ribosomal protein gene expression, in particular, is a well-established target promoted by TOR activity in eukaryotes (58, 59). The *ise3* transcriptome shows comparable patterns to *reptin-1*, with repression of genes encoding ribosomal proteins, histones, cyclin-dependent protein kinases (and related proteins that drive cell cycle progression), components of both photosystems, and Calvin-Benson cycle enzymes (SI Appendix, Fig. S4). Additionally, genes encoding autophagosome components and E3 ubiquitin ligases that target proteins for proteasomal degradation are strongly induced in *ise3* (Fig. 3G). Many of these patterns in the *ise3* transcriptome are canonical signatures of TOR repression, such as the induction of autophagy-related genes (60–62) and suppression of ribosomal protein genes (58, 59) and cell cycle genes (40, 63). We conclude that the glucose-TOR transcriptional program is inactivated in *ise3* and *reptin-1*, demonstrating their critical roles in promoting TOR activity.

We probed the *ise3*, *reptin-1*, and *tor* transcriptomes for DEGs related to PD, but we did not identify any changes in gene expression that could readily explain the altered PD transport phenotype. For example, *PDLP5*, which encodes a PD-LOCALIZED PROTEIN that decreases PD transport in leaves (17, 18), is induced 6.7-fold in *reptin-1* and 11.9-fold in *ise3* (Dataset S2); induction of *PDLP5* would not be predicted to increase PD transport in these mutants. If the effect of these mutants on PD transport is not due to transcriptional changes in gene expression, an alternative hypothesis is that TOR could posttranslationally regulate PD transport by phosphorylating proteins that localize to PD or in other ways impact PD trafficking. We expect that ongoing efforts to map the TOR signaling network in plants will illuminate how TOR regulates PD transport at a mechanistic level.

The Glucose-TOR Signaling Axis Restricts PD Transport. Building on our discovery that mitochondrial *ISE3* is critical for maintaining TOR activity, we next interrogated whether the glucose-glycolysis-OXPHOS-TOR signaling axis regulates PD transport in plants using chemical inhibitors (Fig. 4A). First, we treated young leaves (L8) with 15 mM glucose, providing the leaves with a physiologically relevant increase in sugar availability, and assayed GFP movement 48 h later. Glucose treatment significantly decreased PD transport (Fig. 4C). We then inhibited glycolysis in middle-aged leaves (L7) with 15 mM 2-deoxy-D-glucose (2-DG), a glucose analog that cannot undergo glycolysis and competitively inhibits glucose-6-phosphate isomerase, and assayed GFP movement 48 h later. The 2-DG inhibition of glycolysis had the opposite effect of glucose, increasing PD transport (Fig. 4C). Finally, we focused on inhibiting OXPHOS. We previously showed that disrupting mitochondrial energetics with salicylhydroxamic acid increases PD transport (8) (Fig. 4B). Antimycin A, a direct OXPHOS inhibitor that disrupts complex III activity

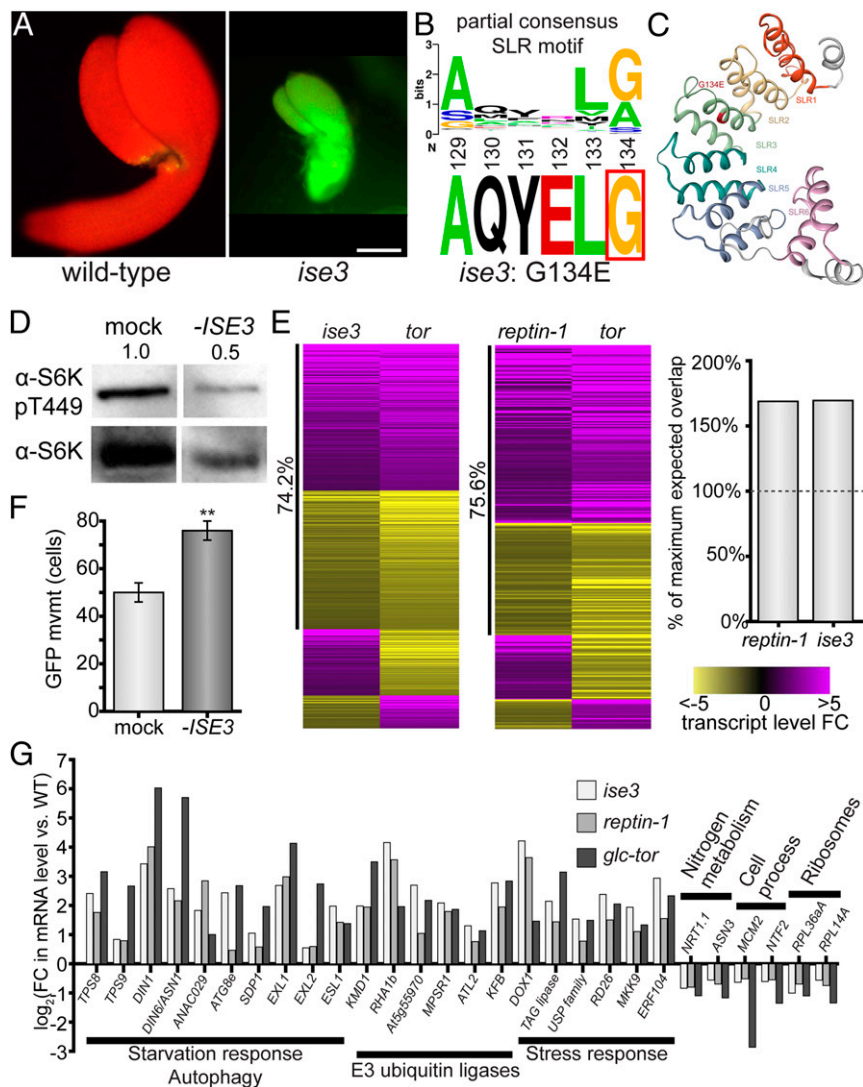


Fig. 3. Dysfunctional mitochondria inhibit glucose-TOR signaling and increase cell–cell transport. (A) The 10 kDa FITC-dextran rapidly moves through *ise3* midtorpedo-stage *Arabidopsis* embryos (Right) compared to wild-type siblings (Left; FITC, green; chlorophyll, red). (Scale bar, 100 μ m.) (B) *ise3* is a missense mutation, G134E, at a conserved G/A residue in SLR proteins. (C) Predicted structure of ISE3 protein, after cleavage of the mitochondrial transit peptide. ISE3 is composed of 6 Sel1-like repeats (SLR1–SLR6), of which SLR5 and SLR6 are relatively unstructured. *ise3* is a missense G134E mutation at the consensus SLR glycine/alanine residue in SLR3 (indicated in red). (D) Silencing *ISE3* with VIGS strongly reduces S6K-pT449 levels in *Arabidopsis* leaves, demonstrating that *ISE3* promotes TOR activity. Total S6K levels are also shown. Numbers indicate S6K-pT449/S6K ratio relative to mock-infected leaves, across three replicates. (E) The glucose-TOR transcriptional program is disrupted in *ise3* and *reptin-1*, reflecting TOR inactivity. Heatmaps show all overlapping significant DEGs between either the *ise3* or *reptin-1* mutant transcriptome and the *tor* transcriptome (41), with purple indicating increased transcript abundance and yellow indicating decreased transcript abundance, as indicated by the legend (Right). The number of overlapping genes in each pairwise comparison is significantly more than expected, as shown in the bar chart (Right; “100% maximum expected value” is the number of genes that could overlap with $P > 0.05$, as determined by a hypergeometric test; specifically $P < 10^{-27}$ for *reptin-1* and $P < 10^{-62}$ for *ise3*). *reptin-1* and *ise3* transcriptomes show strong coregulation with the *tor* transcriptome (75.6% and 74.2% genes coregulated, respectively) (Datasets S2–S4). (F) Silencing *ISE3* expression with VIGS increases PD transport in *N. benthamiana* leaves ($n \geq 59$, $**P < 0.001$; error bars indicate SEM). (G) Representative DEGs affected in *ise3*, *reptin-1*, and *tor* compared to wild-type. Genes involved in starvation response/autophagy, proteasomal degradation, and stress response are induced in each mutant. Cytosolic ribosomal protein genes, cell process/cell cycle genes, and some nitrogen metabolism genes are repressed (SI Appendix, Fig. S4A and Datasets S2–S4).

(40), also increases PD transport in leaves (Fig. 4B). Supporting the conclusion that PD transport increases when ATP synthesis is inhibited, we also recently showed that silencing expression of a subunit of the chloroplast ATP synthase, *AtpC*, increases PD transport in leaves (53). To summarize, stimulating glucose-TOR signaling with glucose decreases PD transport, whereas inhibiting the glucose-glycolysis-OXPHOS signaling axis upstream of TOR increases PD transport.

We extended this hypothesis to test whether modulating sink/source relations in shoots could impact PD transport. We defoliated plants to remove either sink or source leaves, followed immediately

by agroinfiltrating a middle-aged leaf that had not fully expanded (L7) and assayed PD transport 48 h later. Without source leaves to provide sugars, GFP transport through PD was significantly higher in L7; without sink leaves consuming sugars, GFP transport through PD was significantly lower in L7 (Fig. 4D). These results are consistent with the hypothesis that sugar metabolism and transport dynamically regulate PD transport in leaves.

TOR Acts as a Rheostat to Coordinate Leaf Maturation. As a working model, we propose that the nutrient sensor TOR regulates PD transport in response to the balance of sugar synthesis and catabolism

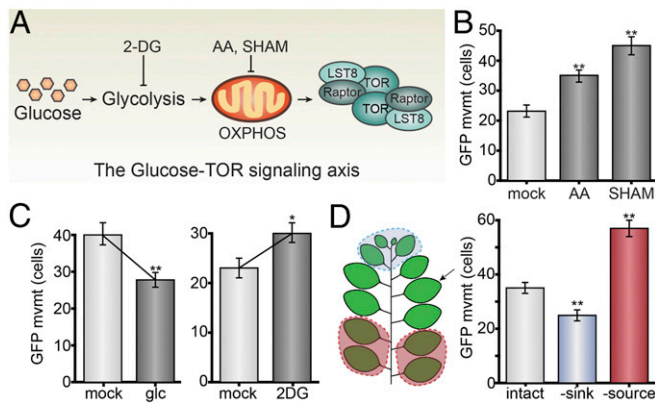


Fig. 4. Glucose-TOR signaling restricts PD transport in leaves. (A) Summary of the glucose-glycolysis-OXPPOS-TORC1 signaling axis (40) interrogated in this figure, with inhibitors of glycolysis and OXPPOS indicated. (B) Chemically inhibiting OXPPOS with antimycin A (AA) increases PD transport ($n \geq 135$, $P < 0.001$; error bars indicate SEM), similar to the previously reported effect of salicylhydroxamic acid (SHAM) (8). L7 leaves were used for these experiments. (C) 15 mM D-glucose (glc) activates TOR and decreases PD transport in young leaves ($n \geq 95$, $**P < 0.001$; error bars indicate SEM); 15 mM 2-DG inhibits glycolysis and increases PD transport. L7 leaves were used for these experiments. (D) GFP movement was assayed in the proximal 25% of a leaf (L7, indicated with an arrow) that has not yet fully expanded. PD transport increases after limiting sugar availability by source removal (L1 through L4 defoliated for 48 h, red), and PD transport decreases after increasing sugar availability by sink removal (L10 through L13 defoliated for 48 h, blue).

in leaves. In young leaves that are actively growing, we hypothesize that sugar is metabolically limiting and TOR is relatively inactive, increasing PD transport and phloem import to promote sugar import and limit sugar export (Fig. 5A). In mature leaves, we hypothesize that sugar is synthesized in excess and TOR is strongly active, decreasing PD transport and promoting active sugar export (Fig. 5A). To test this model, we first used the quantitative GFP movement assay (52, 53) to measure rates of PD transport in young to old leaves of mature plants, and found that PD transport steadily declines in a linear relationship with leaf age (Fig. 5B and C). Next, we directly assayed TOR activity by measuring S6K-pT449 and total S6K levels in each leaf of ~4-wk-old plants (Fig. 5D), using a transgenic line, *35S_{PRO}::AtS6K1-HA*, that drives consistent expression of S6K throughout the plant (38). S6K-pT449 levels gradually and steadily increase as the leaves mature (Fig. 5G), such that the S6K-pT449 to S6K ratio increases ~3.5-fold (Fig. 5H). These results are further confirmed by analysis of the set of significantly DEGs as a leaf ages. The set of DEGs that gradually change during leaf maturation (L11 to L5) and the set of DEGs between a young sink leaf (L11) and mature source leaf (L1) are significantly coregulated with the glucose-TOR transcriptional signaling network (40) (Fig. 5D–F and Datasets S7 and S8). Many of these DEGs are canonical TOR targets, including, notably, over half of the cytosolic ribosomal protein genes, which are induced alongside TOR activity in more mature leaves (Dataset S9). Therefore, we propose that TOR acts as a metabolic rheostat (33), coordinating multiple physiological pathways, including the sink-to-source transition, during vegetative development.

We should note that the sink-to-source transition occurs within a leaf during development: For a number of leaves undergoing the transition (in Fig. 5D, approximately L7 through L10) (28), the distal portion of the leaf stops importing nutrients from the phloem while the proximal portion of the leaf continues to grow, undergo cell division, and import nutrients from the phloem. Therefore, while the youngest leaves are exclusively sinks and the more mature leaves are exclusively sources, some leaves represent

a composite of sink and source tissue. By conducting all leaf PD transport assays in only the proximal 25% of the leaf, however, we minimized this confounding factor in our analysis. In fact, the results shown here suggest that the reduction in PD transport rates in the leaf epidermis as leaves mature is not a binary switch between “sink” and “source,” but instead a gradual, approximately linear decline with leaf age (Fig. 5C). Similarly, TOR activity gradually increases with leaf age, even after all leaves have completed the transition to phloem exporters (approximately L6 through L1). Thus, rather than simply switching from “low” activity in sink tissues to “high” activity in sources, TOR activity progressively increases as leaves mature and begin to photosynthesize more sugars than they consume. This supports our hypothesis that TOR acts as a metabolic rheostat in leaves, with steady changes in TOR activity causing gradual shifts in PD transport and contributing to the coordination of the discrete sink-to-source transition.

Conclusions. We define a pathway regulating intercellular transport through PD in plants that coordinates growth with sugar availability. Analysis of the *reptin-1* mutant led us to postulate that TOR decreases PD transport during plant development, a hypothesis that we then tested directly using multiple genetic, chemical, and physiological approaches (Fig. 6), and elaborated upon this hypothesis by testing how TOR impacts phloem transport. We propose a dynamic, metabolic model of PD and phloem transport, in which relative differences in TOR activity stimulate sugar transport from mature, self-sufficient leaves to younger, rapidly growing leaves. Beyond their roles in nutrient partitioning, PD also traffic a variety of signaling molecules, including transcription factors, cytosolic proteins, and hormones, and are the primary route for viral spread. Dynamic regulation of PD transport by TOR may therefore contribute to a variety of plant biological processes by controlling the mobility of viruses and endogenous micro- and macromolecular signaling molecules. We conclude that TOR, an ancient, conserved mechanism for sensing and responding to metabolic cues, has been coopted in plants to coordinate redistribution of carbohydrates during plant vegetative development.

Materials and Methods

Plant Materials and Growth Conditions. Unless otherwise stated, plants were grown under standard conditions with 16-h day/8-h night at ~120 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$ of photosynthetically active radiation. *ise4/reptin-1* and *ise3* were identified in the Landsberg *erecta* (*Ler*) background and then mapped in crosses to Col-0. SALK_020547 (*spaghetti*) (49), GK543F01-021955 (*reptin*), SALK_036848 (*pseudoreptin*), and SALK_002459 (*lst8-1-1* or *lst8a-1*) (51) were obtained from the *Arabidopsis* Biological Resources Center and their genotypes were confirmed by PCR (Dataset S10). Unless otherwise stated, all *Arabidopsis* experiments used the standard Col-0 ecotype as wild-type. For all *N. benthamiana* experiments, the reference Nb-1 genotype obtained from the Boyce Thompson Institute was used.

Molecular Cloning. Virus-induced gene silencing (VIGS) vectors were prepared as previously described (52), using oligonucleotides listed in Dataset S10. RNA was isolated from Nb-1 or Col-0 shoots with the Spectrum Plant Total RNA (Sigma-Aldrich) with on-column DNase I digestion (New England Biolabs). cDNA was synthesized from RNA using random hexamers and SuperScript III reverse transcriptase (Fisher Scientific). Silencing triggers were amplified with Phusion DNA polymerase (New England Biolabs), digested alongside pYL156 with restriction enzymes as listed in Dataset S10 (enzymes from New England Biolabs), and ligated with Promega T4 DNA ligase (Fisher Scientific). Ligations were transformed into house-made chemically competent XL1-Blue *Escherichia coli*. Kanamycin-resistant bacterial colonies were screened by colony PCR for positive clones. Plasmids were then miniprep (Bioneer) and Sanger-sequenced to confirm insert sequences. Manufacturer’s protocols were followed throughout. All constructs were then transformed into *Agrobacterium* GV3101 for plant transformation. For *NbLST8*, we tested two silencing triggers that had the same effects on *NbLST8* expression and PD transport; primers for each silencing trigger are described in Dataset S10.

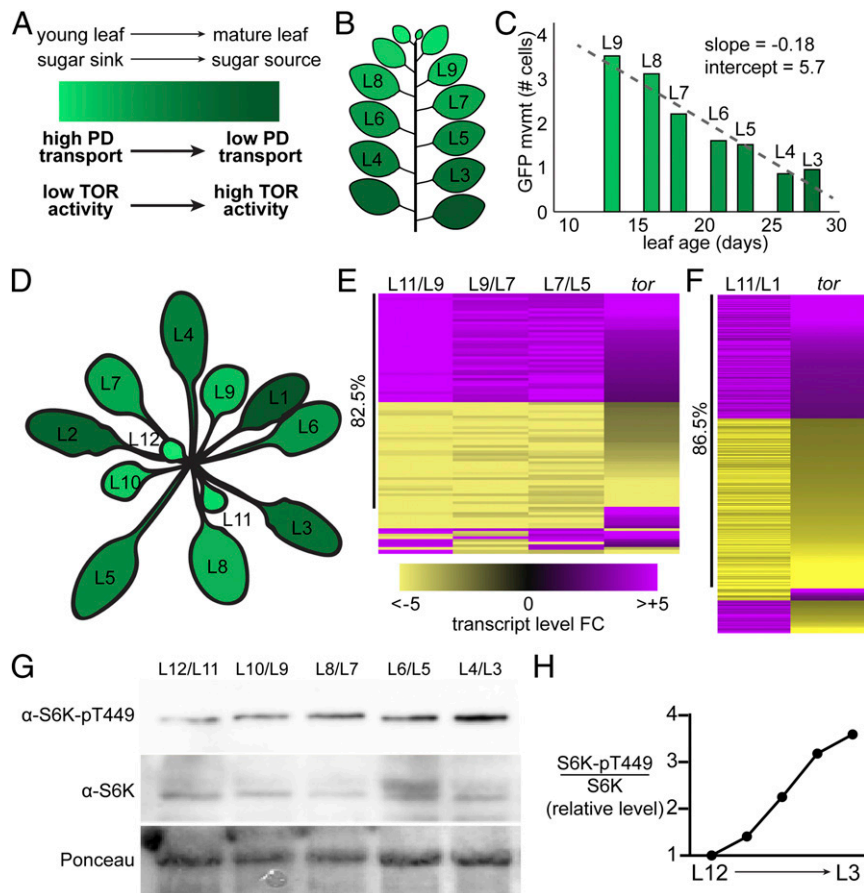


Fig. 5. TOR activity coordinates leaf metabolism with leaf age. (A) As leaves age, they transition from young sinks (light green) that rapidly consume sugars to mature sources (dark green) that photosynthesize excess sugars for export to the rest of the plant. Here, we show that TOR activity steadily increases and PD transport steadily decreases during this transition. (B) Leaf emergence of *N. benthamiana* plants was tracked for 6 wk, during which time 13 leaves emerged. The first true leaf is labeled leaf 1 (L1), and subsequent leaves are numbered by order of emergence. (C) Six weeks after germination, PD transport in L9 (a relative sink leaf) through L3 (a relative source leaf) was quantified using the GFP movement assay. PD transport was assayed in the proximal 25% of every leaf. PD transport decreases linearly with leaf age (as in ref. 53) in *N. benthamiana*. (D) Schematic of an *Arabidopsis* vegetative rosette used for transcriptomes and the Western blot analyses in this figure. (E–H) (E) The sets of genes that are differentially expressed in *Arabidopsis* leaves of different ages significantly overlap with the *tor* transcriptome ($P < 10^{-26}$), and are highly coregulated (82.5%), indicating that the TOR-promoted transcriptional program is activated as leaves mature (Dataset S7). The *tor* transcriptome is the set of gene-expression changes that respond to glucose only in the presence of TOR (40), and are therefore transcriptional targets of the TOR signaling pathway. Pairwise analyses of DEGs for L11 and L9, L9 and L7, and L7 and L5 are shown, respectively, from left to right. As an example, a gene is considered “induced” in the L11/L9 dataset if it is significantly more highly expressed in L11 than in L9. (F) Over 30% of the glucose-TOR target genes (732 of 2,350, $P < 10^{-138}$) are significantly differentially expressed between a young sink leaf (L11) and a mature source leaf (L1), and the vast majority of these DEGs (86.5%) are coregulated, suggesting that the glucose-TOR signaling network is more active in source leaves than in sink leaves (Dataset S8). (G) S6K-Thr449 phosphorylation levels, a proxy for TOR activity in plant cells, were assayed by Western blot with a phospho-specific antibody. A *35S_{PRO}:S6K1* transgenic *Arabidopsis* line was used for this experiment, because it maintains consistent S6K protein levels in leaves of different ages. S6K-Thr449 levels steadily increase with leaf age in an *Arabidopsis* rosette. Ponceau staining was used to show loading of total protein (Lower). (H) The quantified ratio of S6K-pT449 levels to total S6K levels in leaves of different ages confirms that TOR activity increases with leaf age in a mature *Arabidopsis* rosette.

Next-Generation Mapping of *ise3* and *ise4*. *ise3* and *ise4* were identified from a population of embryo-defective ethyl methane sulfonate-mutagenized *A. thaliana* Ler lines, as previously described (30, 64, 65). F2 mapping populations were constructed by crossing heterozygotes carrying the *ise* allele with Col-0, and then selfing heterozygous *ise*+ F1 plants to generate the F2. Using indel and SNP markers (oligonucleotides for PCR listed in Dataset S10), the mutated *ise3* locus mapped to a 112-kb region on chromosome 2 between SNPs SGC5NP606 at *At2g25470* and PERL0363471 at *At2g25710*, and *ise4* locus was mapped to a 120-kb region between the SNP PERL1125350 at *At5g67310* and the end of chromosome 5. Due to low recombination rates in the *ise4* region, we used next-generation mapping to identify the causal mutations for both *ise3* and *ise4*. Eighteen overlapping genomic fragments between 4- and 8-kb long were amplified with iProof high-fidelity DNA polymerase (Bio-Rad Laboratories) from genomic DNA isolated from homozygous *ise3* or *ise4* mutants. The fragments were purified by gel extraction, and pooled in equal amounts. DNA-seq libraries of bulked DNA were constructed using the Nextera DNA Flex Library Prep Kit

(Illumina) following the manufacturer’s protocol, and sequenced on the Illumina HiSeq 2000 platform. Data were mapped to the unmasked *Arabidopsis* genome and compared using Sequencher (66).

ise3 was complemented by three different constructs (cloned with oligonucleotides in Dataset S10). A full genomic sequence was constructed by cloning from ~1.8-kb upstream of the *At2g25570* start codon through 858 bp downstream from the *At2g25570* stop codon. The same genomic ~1.8-kb promoter was also used to drive expression of the *At2g25570* coding sequence or the coding sequence C-terminally tagged with GFP. All constructs rescued *ise3* lethality. *ise4* was complemented with three different constructs (cloned with oligonucleotides in Dataset S10). A long genomic sequence was composed of ~2.4-kb upstream of *At5g67630*, the *At5g67630* gene, and ~1-kb downstream of *At5g67630*. A gene of unknown function, *At5g67640*, is within the ~2.4-kb region upstream of *At5g67630*; no SNPs were identified in *At5g67640* between wild-type and *ise4*, but to be sure that expression of *At5g67640* did not affect the *ise4* phenotype, a short genomic sequence lacking the *At5g67640* translation start site was also

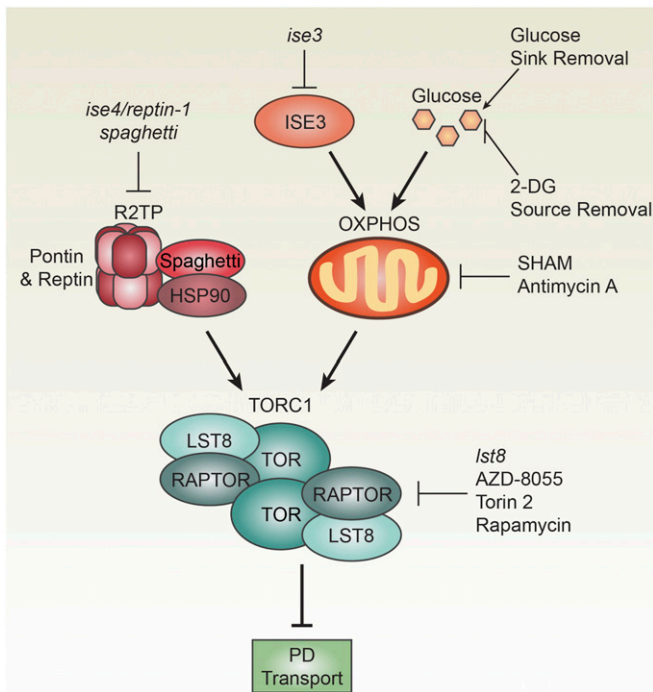


Fig. 6. Summary of the glucose-TOR-PD signaling network in plants. A forward genetic screen uncovered two mutants, *ise3* and *ise4/reptin-1*, that decrease TOR activity and increase PD transport. As a component of R2TP, Reptin promotes assembly of the active TORC1 dimer (Fig. 1). *spaghetti*, another R2TP mutant, also decreases TOR activity and increases PD transport (Fig. 1). OXPHOS promotes TOR activity and is disrupted by loss of ISE3 (Fig. 3) or inhibition of the glucose-glycolysis-OXPHOS-TOR signaling axis (Fig. 4). Direct assays reported here demonstrate that TOR restricts PD transport (Fig. 2), and TOR activity correlates with decreased PD transport during leaf maturation (Fig. 5). Chemical, genetic, and physiological treatments that impact each step in this pathway are indicated; each of these conditions affects PD transport, as described in the text.

constructed, and this sequence complemented *ise4*. Finally, the ~2.4-kb upstream sequence was used to drive expression of the coding sequence of *At5g67630* cloned from *Arabidopsis* cDNA; this construct also complemented *ise4*. Complementation was performed by transforming *ise4/+* heterozygotes with the floral dip method (67), screening for transformants with Basta (0.3% [vol/vol]), and testing for rescue of the embryo-defective phenotype.

PD Movement Assays. Embryo dye-loading assays were performed as previously reported (30, 68). Briefly, 10 kDa FITC-dextran (Sigma-Aldrich) was prepared at 5 mg/mL in half-strength liquid Murashige and Skoog (MS) (pH = 5.5) media, with any dye less than 10 kDa removed by centrifugal filtration (Amicon Ultra 10K membranes, Millipore Sigma). Embryos were removed from seeds and immediately treated with FITC-dextran for 5 min before several washes with MS media to remove excess dye. Col-0 and *ise2* (30) were used as negative and positive controls for dye loading, respectively, alongside each experiment. Experiments were replicated three times. For newly identified *ise* mutants (including *spaghetti* and *Ist8*), dye movement throughout the hypocotyl of the embryo was observed in >80% of mutant embryos loaded with 10 kDa FITC-dextran.

GFP movement assays in *N. benthamiana* were performed as previously reported (52, 53). Briefly, *Agrobacterium* was grown overnight in LB with antibiotics at 28 °C. Cultures were spun down at 700 × g for 10 min, washed, and resuspended to OD_{600nm} = 1.0 in 10 mM MgCl₂, 10 mM 2-(N-morpholino) ethanesulfonic acid (pH = 5.5, adjusted with KOH), and 200 μM acetosyringone from a recently prepared 100-mM stock solution frozen in DMSO. *Agrobacterium* were left to induce virulence at room temperature for 2 to 4 h with gentle shaking before infiltration. For GFP movement assays, immediately before infiltrating, cells carrying 35S_{PRO}:GFP T-DNA were diluted to OD_{600nm} = 10⁻⁴ in the same virulence-inducing media. GFP movement was scored 48 h after infiltration. For VIGS, equal parts of pYL192 (which expressed *TRV1*) and pYL156

(which expresses *TRV2* with silencing triggers) were mixed and infiltrated into leaves. A mock *TRV2* vector, pYC1, which is pYL156::GUS_{fragment} (64), was used as a negative control for all VIGS experiments.

Unless otherwise noted, all *N. benthamiana* movement assays used the fourth-youngest emerged leaf (as counted from the shoot apex basipetally). GFP movement was only visualized in the proximal 25% of the leaf. For chemical treatments and overexpression, 5- to 6-wk-old plants were used. For VIGS experiments, 3-wk-old plants were infiltrated to express *TRV*, and movement assays were conducted 2 wk later. Heteroscedastic unpaired *t* tests were used to compare GFP movement between samples throughout.

SUC2_{PRO}:GFP Phloem Transport Assays. Transgenic *A. thaliana* SUC2_{PRO}:GFP seeds were grown as described above for 3 wk (at this stage, five leaves had emerged). Five microliters of 100 nM Torin 2 or a mock control was applied by pipette to the shoot apex of seedlings, such that the Torin 2 was in direct contact with L4 and L5. Care was taken not to apply any Torin 2 to the mature source leaves, to prevent any unintended effects of TOR inhibition on translation of GFP in cells with active SUC2 promoters. One day later, young leaves (L4, ~1.5-mm long) were excised and immediately observed by confocal scanning laser microscopy, collecting z stacks and tiling several fields of view to visualize the entire leaf. At least three mock-treated seedlings and three Torin 2-treated seedlings were observed for each experiment, and the entire experiment was repeated three separate times.

Confirming Gene Silencing with RT-qPCR. To confirm gene silencing, reverse transcription-quantitative polymerase chain reaction (RT-qPCR) was performed on VIGS tissue in parallel with GFP movement assays, with mock *TRV*::GUS infected plants as controls. Three biological replicates were included for each experiment. RNA was isolated with the Spectrum Plant Total RNA (Sigma-Aldrich) with on-column DNase I digestion (New England Biolabs). cDNA was synthesized from RNA using oligo (dT)18 and SuperScript III reverse transcriptase (Fisher Scientific). qPCR was performed in parallel for all samples using a validated reference gene (*EF1α*) and gene-specific oligonucleotides (Dataset S10) for *N. benthamiana* VIGS experiments (69). Specificity of primers for target genes was validated by resolving RT-qPCR products on 3% agarose gels. VIGS constructs were included in this study only if they reduced gene expression to ≤20% of wild-type levels.

Chemical Inhibitors. TOR kinase inhibitors used in this report have previously been demonstrated to suppress TOR signaling (38, 39). Stock solutions of rapamycin (10 mM), Torin 2 (10 mM), AZD-8055 (10 mM), cantharidin (100 μM), okadaic acid (100 μM), and antimycin A (5 mM) were prepared within days of use in DMSO (except okadaic acid, prepared in ethanol) and frozen until used. All chemicals were obtained from Cayman Chemicals, except antimycin A (Sigma-Aldrich). Stocks were serially diluted in infiltration media to ensure accurate reporting of inhibitor concentrations. Dextros or 2-DG (Fisher Scientific) was dissolved directly in virulence-inducing infiltration media at 150 mM, and then diluted to 15 mM just prior to infiltration. Concentrations for inhibitor treatments were determined based on previous reports of minimum effective concentrations for rapamycin (38), AZD-8055 (39), Torin 2 (39), antimycin A (40), dextrose (40), and 2-DG (40).

Western Blot Assays. Leaves from Col-0 or 35S_{PRO}:S6K1-HA plants were snap-frozen in liquid nitrogen. Protein was extracted from leaves in 100 mM Mops (pH 7.6), 100 mM NaCl, 5% SDS, 0.5% β-mercaptoethanol, 10% glycerol, 2 mM PMSF, and 1× PhosSTOP phosphatase inhibitor (Sigma-Aldrich). Specific protein levels were assayed by Western blot protein using primary antibodies against S6K (Santuz Cruz Biotechnology, sc-230) and S6K-pThr449 (Abcam no. 207399) along with an HRP-conjugated goat anti-rabbit IgG secondary antibody (Jackson Immuno Research, no. 111-035-003). Total protein was visualized after transfer using Ponceau S red staining. All Western blot experiments were repeated at least three times, with representative results shown in figures.

RNA-seq and Transcriptome Analysis. Homozygous *ise3* or *ise4/reptin-1* seeds at the midtorpedo stage of embryogenesis and their wild-type or heterozygous midtorpedo-stage siblings were collected as previously described and stored at -80 °C in RNAlater (6). For each pooled RNA sample, ~100 to 150 seeds were collected, and four replicates were collected (three replicates were used for RNA-seq, the fourth replicate was stored as back-up). RNA was isolated with the Spectrum Plant Total RNA (Sigma-Aldrich) with on-column DNase I digestion (New England Biolabs). RNA was treated with RiboMinus (Life Technologies) to deplete 16S, 18S, 23S, and 25S rRNA, and then used to prepare Ion Torrent libraries with the Ion Total RNA-Seq Kit v2 (Life Technologies).

Libraries were sequenced with an Ion Proton I Chip. Torrent Suite Software was used to filter polyclonal reads, adapter dimers, reads of very low quality, and to trim 5' adapter and repetitive sequences. The remaining computational steps were completed using an instance of Galaxy. FASTQ Quality Trimmer was then used to remove low quality 3' ends (window size 10, step size 1, quality score ≥ 20) from reads, then reads were filtered with Filter FASTQ to remove reads less than 25-bp long.

To determine differences in gene expression between *ise4/reptin-1* and wild-type seeds, RNA-seq results were compared using the Tuxedo Suite. Reads were mapped with TopHat2 to the *Ler* genome and assembled into transcripts with Cufflinks. Cuffmerge was used to assemble the transcriptome with a *Ler* reference transcriptome. Cuffdiff2 was used to generate a list of differentially expressed genes comparing three replicates of *ise3* or *ise4/reptin-1* seeds with three replicates of wild-type seeds ($P < 0.05$, $q < 0.05$).

The *ise3*, and *ise4/reptin-1* transcriptomes were analyzed using MapMan software, which assigns genes to various categories based on their biological function and uses a Wilcoxon rank sum test, corrected with the Benjamini-Hochberg procedure to reduce the false-positive rate, to identify categories that are significantly altered in gene expression.

DEG lists from different transcriptomes were compared using a hypergeometric test for significant overlap using R software (phyper function). Within the subset of overlapping genes, any genes that were up-regulated or down-regulated (regardless of the measured fold-change in expression) in all transcriptomes were considered "coregulated." Gene lists were only considered significantly overlapping if the hypergeometric test revealed $P < 10^{-5}$, and the maximum expected number of overlapping genes was calculated by determining the number of overlapping genes that would yield $P = 0.05$.

Tables were generated in Excel using TAIR10 and MapMan annotations, including a large table with all transcriptomic data included in this study, and individual tables showing data underlying heatmaps in all figures. In tables, as in the heatmap figures, purple is used to indicate transcript induction, and yellow is used to indicate transcript repression.

Computational Analysis. ISE3 structure was modeled by Phyre2, which used structures of *E. coli* EsiB (70) and *Pseudomonas fluorescens* AlgK (71) as templates. The resulting model was visualized with NGL. SLRs were identified by TPRpred (72) and by comparison to the structure of *Helicobacter pylori* HcpC (73). ISE3 orthologs were identified using a BLASTp search with the *Arabidopsis* amino acid sequence as a query against the reference protein sequence database for eukaryotes. BLASTp results were then filtered to remove paralogues within a species and to highlight representative species from different clades. No proteins with worse scores than the closest-matching *Homo sapiens* protein were included, except for the next-closest match within *A. thaliana* to root the trees.

Reptin consensus sequences were determined by aligning all Reptin orthologs available in Ensembl databases with MAFFT, trimming the sequence to include only the Walker A motif and surrounding residues, and then generating a consensus sequence logo with WebLogo (74). SLR consensus sequences were determined by aligning the curated seed alignment of SLRs from PFAM, trimming the sequence to remove uncommon insertions sometimes present between residues 134 and 135, and then generating a consensus sequence logo with WebLogo (74).

N. benthamiana orthologs of *LST8* and *ISE3* were identified by reciprocal BLASTp searches starting with the *A. thaliana* protein sequences as queries against the Sol Genomics Network *N. benthamiana* genome v1.0.1 predicted protein database. The resulting aligned protein sequences are available in Dataset S11. There are two homeologues (paralogues derived from an ancestral polyploidization event) of each of these genes in *N. benthamiana*; therefore, in selecting silencing triggers, we chose regions of the transcripts that are nearly perfectly homologous in nucleic acid sequence between the two homeologues (paralogues), but had no sufficient homology to any other transcripts in *N. benthamiana* to cause off-target effects. In the case of *NbLST8*, we made two separate silencing triggers, which had the same effects on *NbLST8* expression and PD transport.

Data Availability. RNA-seq data described in this article have been deposited in National Center for Biotechnology Information Sequence Read Archive as BioProject PRJNA600773.

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