Cellulose synthase complexes act in a concerted fashion to synthesize highly aggregated cellulose in secondary cell walls of plants

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Cellulose, often touted as the most abundant biopolymer on Earth, is a critical component of the plant cell wall and is synthesized by plasma membrane-spanning cellulose synthase (CESA) enzymes, which in plants are organized into rosette-like CESA complexes (CSCs). Plants construct two types of cell walls, primary cell walls (PCWs) and secondary cell walls (SCWs), which differ in composition, structure, and purpose. Cellulose in PCWs and SCWs is chemically identical but has different physical characteristics. During PCW synthesis, multiple dispersed CSCs move along a shared linear track in opposing directions while synthesizing cellulose microfibrils with low aggregation. In contrast, during SCW synthesis, we observed swaths of densely arranged CSCs that moved in the same direction along tracks while synthesizing cellulose microfibrils that became highly aggregated. Our data support a model in which distinct spatiotemporal features of active CSCs during PCW and SCW synthesis contribute to the formation of cellulose with distinct structure and organization in PCWs and SCWs of Arabidopsis thaliana. This study provides a foundation for understanding differences in the formation, structure, and organization of cellulose in PCWs and SCWs.

Significance

Plant cell walls are important in plant development and for textiles, wood products, and bioenergy. Cellulose, the microfibrillar component of primary cell walls (PCWs) and secondary cell walls (SCWs), is formed by cellulose synthase complexes (CSCs) at the plasma membrane. Here, we show that CSCs behave differently during PCW and SCW synthesis and form microfibrils with different organization. During PCW synthesis, dispersed CSCs synthesize cellulose microfibrils with low aggregation, whereas during SCW synthesis, densely arranged groups of CSCs move coherently to synthesize highly aggregated microfibrils. Our study suggests that controlled alterations in CSC distribution and orchestrated movements contribute to the high density and bundling of cellulose microfibrils in SCWs.


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Results

Transdifferentiated Cells Synthesize SCWs with Highly Aggregated Cellulose Microfibrils. Similar to previous inducible xylem transdifferentiation systems (22, 23), we generated plants expressing the master regulator of xylem differentiation, Vascular-related NAC-domain 7, fused to the glucocorticoid receptor (35S::VND7-GR), and induced transdifferentiation by dexamethasone (DEX) treatment. Approximately 80% of epidermal cells of etiolated seedlings transdifferentiated after 60-h DEX treatment as indicated by lignin autofluorescence in SCW thickenings (Fig. 1A). SCW thickenings were also observed in epidermal peels from transdifferentiated seedlings using field emission scanning electron microscopy (FE-SEM) (Fig. 1B). Focused ion beam ablation was used to create windows through which the most recently deposited layer of cellulose microfibrils was observed (Fig. 1C). Cellulose microfibrils of SCWs in transdifferentiated cells were aggregated into bundles within the lignified cell wall thickenings and in the nonlignified regions between thickenings (Fig. 1D and G). In contrast, cellulose microfibrils of PCWs from epidermal peels of nontransdifferentiated control seedlings were much less aggregated (Fig. 1E and F). Sum-frequency generation (SFG) vibrational spectroscopy, which can selectively detect the orientation and lateral packing of cellulose microfibrils in intact plant cell walls, showed that transdifferentiated seedlings exhibited spectral features similar to native SCWs in Arabidopsis stems (24), suggesting that the cell walls of transdifferentiated seedlings were structurally similar to native SCWs (Fig. 1H). The 2,944 cm⁻¹ and 2,868 cm⁻¹ alkyl stretch peaks of transdifferentiated seedlings indicate highly ordered cellulose, perhaps due to high cellulose aggregation, and the enhancement of 3,320 cm⁻¹ and 3,440 cm⁻¹ hydroxyl stretch peaks suggest that transdifferentiated seedlings have an elevated crystalline cellulose content (Fig. 1H) (24, 25). Analogous spectral features were absent in nontransdifferentiated seedlings where the PCW-related 2,904 cm⁻¹ peak was dominant (Fig. 1H) (24).

Active CSCs Are Evenly Distributed Early in Transdifferentiation and Are Later Distributed Within Confining Hoops During the Synthesis of SCW Thickenings. Having established a system in which epidermal cells reliably transdifferentiate into xylem-like cells with genuine SCWs, we investigated whether differences in the spatiotemporal behavior of CSCs during PCW and SCW synthesis could explain the structural differences of cellulose microfibrils in PCWs and SCWs. We generated a line expressing green fluorescent protein (GFP) fused to CESA7 driven by the native CESA7 promoter (ProCESA7::GFP-CESA7) that was able to rescue the growth phenotype of the null cesa7-2 irx3-1 mutant (Fig. S1A) and crossed it with 35S::VND7-GR. When 3-d-old etiolated seedlings were treated with DEX for 24 h, GFP-CESA7 was observed in cells undergoing transdifferentiation (Fig. S1C and D). Like CSC markers during PCW formation, such as GFP-CESA3, GFP-CESA7 labeled active CSCs at the PM and intracellular Golgi bodies (Fig. S1 B–D) (12, 15, 16). In addition to localizing to distinct puncta, PM-localized GFP-CESA7 localized to swaths of motile signal, which might represent stretches of densely arranged CSCs (Fig. 2D, F, G, I, J, L, and Fig. S1C and D, and Movie S1). Early in transdifferentiation PM-localized GFP-CESA7 particles were positioned along tracks that were distributed evenly across the cell surface (Fig. 2D and Fig. S1C). Later, GFP-CESA7 became focused within defined hoop-like regions of the PM (Fig. 2G and Fig. S1D), which is consistent with previous observations (22). These observations suggest that a uniform layer of SCW is synthesized over the entire cell surface before hoop formation, which is probably the source of highly aggregated cellulose in the nonlignified regions between cell wall thickenings (Fig. 1D), and that SCW thickenings are subsequently formed when CSC distribution is constrained within hoop regions.

Active CSCs Exhibit Bidirectional Movement During PCW Synthesis and Directionally Coherent Movement During SCW Synthesis. The striking changes in CSC distribution during transdifferentiation caused us to investigate whether CSCs also developed unique dynamic features. Both during PCW and SCW synthesis, PM-localized CSCs moved steadily along linear trajectories (Fig. 2 A, D, and G and Movie S1), which occurs during active cellulose synthesis (11, 12). The velocities of several CSC markers have been recorded during PCW synthesis (6, 7, 12, 13, 16, 17, 19, 22, 26, 27), each exhibiting a slight variation in velocity, which makes the direct comparison of the velocities of two dissimilar CSC markers difficult. To account for the marker-to-marker variation in CSC velocity, the velocities of several CSC markers were measured during PCW synthesis and compared with the velocities of GFP-CESA7–labeled CSCs during SCW synthesis (Fig. S2). The velocities of GFP-CESA7–labeled CSCs during SCW synthesis in this study were similar to recent measurements of YFP-CESA7 particles during SCW synthesis (22). However, these velocities of CSCs during SCW synthesis fall within the range of velocities exhibited by CSC markers during PCW synthesis, making it uncertain whether CSC velocities are meaningfully different during PCW and SCW synthesis (Fig. S2).

However, other aspects of CSC dynamics were significantly different during PCW and SCW synthesis. In contrast to the bidirectional movement of CSCs along shared tracks during PCW synthesis (12), a majority of CSCs moved coherently in a favored direction along shared tracks during SCW synthesis. The
Fig. 2. Active CSCs exhibit bidirectional movement along tracks during PCW synthesis and directionally coherent movement along tracks during SCW synthesis. (A–L) GFP-CESA3 cesa3+ and GFP-CESA7 cesa7+ were imaged in nontransdifferentiating cells as a CSC marker during PCW synthesis (A–C), and GFP-CESA7 cesa7+/− 35S::VND7-GR was imaged during two stages of xylem cell transdifferentiation, before hoop formation (BHF) and during hoop formation (DHF), as a CSC marker during SCW synthesis (D–L). (J–L) The influence of CMTs on GFP-CESA7 particle behavior was assessed via 8–12 h treatments with 25 μM oryzalin. (A, D, G, and J) Representative single-frame images and 7-min projection images show the distribution and trajectories of GFP-CESA particles, respectively. Arrows indicate the apical direction. (Scale bars: 10 μm) (B, E, H, and K) Kymographs were derived from the indicated tracks in each 7-min projection image and the directions of particle movements along the track were color-coded in schematic kymographs. (Scale bars: 5 μm) (C, F, I, and L) Images from 5 s and 5 min after photobleaching the lateral sides of cells are shown to depict the direction of particle movement along tracks of interest. Upper images are raw images, and lower images are highlighted to show the bleached regions (gray boxes), the particles within tracks of interest (white boxes), and the favored direction of particle movements (arrows). (Scale bars: 10 μm.) The frequency of particle direction along individual tracks was quantified (M). Error bars are standard errors of the mean; n ≥ 48 tracks and 6 seedlings per data point. *P < 0.0001.

Cortical Microtubules Are Not Required to Maintain the Directionally Coherent Movement of Active CSCs During SCW Synthesis. We next asked whether the directional coherence of CSC movement was controlled by the CMTs along which the CSCs travel (12, 22). Because End Binding Protein 1B (EB1b) follows the plus-ends of polymerizing microtubules, Red Fluorescent Protein (RFP)-conjugated EB1b (RFP-EB1b) can be used as a marker for the position and polarity of newly formed microtubules. Like GFP-CESA7 particles, RFP-EB1b particles were distributed evenly across the cell early in transdifferentiation and became restricted to hoop-like regions at later stages of transdifferentiation during the synthesis of SCW thickenings. The trajectories of GFP-CESA7 and RFP-EB1b were often colocalized (Fig. S3A). Kymographs were created from colocalized GFP-CESA7 and RFP-EB1b trajectories, and the direction of coherent CSC movement was compared with the CMT polarity of each track. The direction of coherent CSC movement had a similar tendency to travel toward the plus-end of CMT tracks and toward the minus-end of CMT tracks (Fig. S3 B–G), suggesting that there is no correlation between the polarities of newly formed CMTs and the direction of coherent CSC movement.

Pharmacological disruption of CMTs was used to further probe the role of CMTs during SCW synthesis. After 8–12 h of treatment with 25 μM oryzalin, a microtubule-depolymerizing drug, RFP-EB1b particles were abolished in transdifferentiating cells, indicating that the polymerization of new CMTs was arrested (Fig. S4 A and B). Under the same oryzalin treatment conditions, the microtubule marker, GFP-Microtubule Associated Protein4 (GFP-MAP4), no longer localized to CMTs, but rather in the cytosol, suggesting that CMT arrays were abolished (Fig. S4 C and D). In transdifferentiating cells, oryzalin treatment occasionally caused GFP-CESA7 trajectories to appear wavy but did not influence the crosshatched pattern of trajectories in kymographs from GFP-CESA3-labeled CSCs during PCW synthesis is indicative of bidirectional CSC movement (Fig. 2 B and M), whereas trajectories on kymographs from GFP-CESA7-labeled CSCs mostly run in a common direction during SCW synthesis both before and during hoop formation (Fig. 2 E, H, and M). The directionally coherent movement of GFP-CESA7 existed on a track-by-track basis such that distinct tracks could exhibit coherent particle movement in opposite directions, resulting in no net directional bias on a cell-wide scale and an antiparallel packing of cellulose microfibrils in SCWs at the mesoscale (25). Adjacent CSC tracks became condensed within the hoop regions, which sometimes obscured the ability to resolve individual tracks and potentially caused an underrepresentation of directional coherence in the quantification during hoop formation (Fig. 2M). Likewise, in a recent study, a kymograph of motile YFP-CESA7 particles displayed a cross-hatched pattern, which would seem to suggest that CSCs moved bidirectionally during SCW synthesis (22). However, it is likely that the kymograph in the previous study encompassed particles from multiple adjacent tracks due to the close packing of adjacent tracks during hoop formation and due to technical differences in the kymograph generation methods of the two studies (ref. 22 and Materials and Methods).

To more clearly visualize the directional trends of CSC movement, a lateral photobleaching experiment was designed in which the lateral regions of the cell were bleached during time-lapse imaging and the direction of the movement of the remaining unbleached particles within a central strip of the cell was monitored after photobleaching (Fig. 2 C, F, and I and Movie S2). To better analyze the direction of the movement of the CSC from an unbleached region to bleached region without influence of newly delivered CSCs, we selected photobleaching regions with minimal underlying CESA-associated intracellular compartments. The bidirectional movement of GFP-CESA3 during PCW synthesis resulted in an equal number of particles on each track moving into the left and right bleached regions (Fig. 2C). The directionally coherent movement of GFP-CESA7 during SCW synthesis resulted in an uneven distribution of signal from each track moving into the left and right bleached regions before and during hoop formation (Fig. 2 F and I). In many cases, swaths of GFP-CESA7 signal, rather than distinct particles, moved coherently during SCW synthesis, which might represent groups of CSCs that are too crowded to be resolved as individual particles by light microscopy.
of underlying intracellular signal (Fig. 3A). During SCW synthesis, GFP-CESA7 occupied a slightly higher percentage of the PM area before hoop formation than GFP-CESA3 during PCW synthesis (Fig. 3A and C). During hoop formation, a high percentage of the PM was occupied by GFP-CESA7 signal within hoop regions, but the area between hoops was nearly devoid of PM-localized signal (Fig. 3A and C). Under the presumption that the maintenance of crowded populations of CSCs during SCW synthesis would require high rates of CSC delivery to the PM, the delivery rate of CSCs to the PM was measured via photobleaching experiments and compared during PCW and SCW synthesis (Fig. 3B and Movie S3). Both before and during hoop formation, GFP-CESA7-labeled particles were delivered to the PM at more than triple the rate of GFP-CESA3 particles during PCW synthesis (Fig. 3B and D). During hoop formation, delivery of new PM-localized GFP-CESA7 particles was confined to the hoop regions. The elevated rates of CSC delivery during SCW synthesis and the more extensive coverage of PM area that we observed, along with previous electron microscopy measurements of high CSC density under developing SCW thickenings, suggest that CSCs are arranged in crowded groups during SCW synthesis (29–32).

**Discussion**

Cellulose is synthesized by diverse organisms, including plants, algae, tunicates, protists, fungi, and bacteria (28, 31). CESAs of these divergent organisms are organized in a variety of spatial arrangements—including rows of CESAs, linear arrays of CESAs, and rosette-shaped complexes of CESAs—which dictate the size, shape, and aggregation of the cellulose that is produced by each organism (33, 34). It has also been shown that some organisms can alter the spatial arrangement of CESAs to alter the structure of the cellulose being produced. For example, freeze–fracture electron microscopy studies of the green algae, *Microstodium denticulatum*, showed that *Microstodium* uses a sparse population of disperse rosette CSCs to synthesize randomly oriented arrays of cellulose during PCW synthesis and uses large, clustered arrays of rosette CSCs to synthesize highly aggregated bands of cellulose during SCW synthesis (35). Freeze–fracture electron microscopy has also showed that individual rosette CSCs are capable of synthesizing cellulose in higher plants, but some investigators have speculated that clusters and groups of rosette CSCs that are positioned along shared cellulose microfibril impressions might cooperate to synthesize bundled cellulose microfibrils in SCWs (29, 30, 32). Now—with dynamic, spatiotemporal details of CSC behavior during PCW and SCW synthesis in living plants—we can propose a hypothetical model to explain how plants synthesize PCWs with low cellulose microfibril aggregation and SCWs with highly aggregated and bundled cellulose microfibrils by altering the behavior of active CSCs at the PM (Fig. 4).

Our data are consistent with the following hypothetical model. Cellulose of low aggregation is synthesized by disperse CSCs that exhibit bidirectional movement during PCW synthesis (Fig. 4 A and B). During SCW synthesis, elevated rates of CSC delivery to the PM produce crowded populations of closely arranged CSCs, which move in a coherent direction during cellulose synthesis. The concerted activity of coherently moving and densely arranged CSCs during SCW synthesis results in the synchronous extrusion of many closely spaced cellulose microfibrils, promoting the aggregation and bundling of adjacent microfibrils at the time of synthesis (Fig. 4 C–E).

As of yet, it is unclear whether the coherent movement of CSCs occurs passively or through an unknown regulated process. In either case, the observations that the coherent CSC movement occurs on a track-by-track basis and often involves swaths of CSC signal suggest that coherent CSC movement is a spatially confined process, potentially caused by the crowding or physical association of groups of CSCs at the PM (31, 32). Although the role of CMTs in the maintenance of the coherent movement of CSCs has been ruled out, the role of CMTs in the initial
establishment of the coherent movement of CSCs and in other aspects of SCW synthesis remains to be tested.

In addition to the observed differences in CSC distribution and behavior during PCW and SCW synthesis, other differences between PCWs and SCWs might also contribute to the differences in cellulose microfibril aggregation. For example, it is unclear what role, if any, other components of the cell wall play in the aggregation status of cellulose microfibrils in PCWs and SCWs. Several models for the interaction of hemicelluloses—xyloglucan in PCWs and xylan in SCWs—with cellulose have been described (36, 37), suggesting that hemicellulose–cellulose interactions might influence cellulose aggregation. Another possibility could be that lignin contributes to the high aggregation of cellulose in SCWs. Immunofluorescence experiments showed that LM10-labeled xylan was localized to discrete regions of the cell wall defined by SCW thickenings in transdifferentiated xylem-like cells of Arabidopsis (38). Similarly, lignin and the laccase enzymes that catalyze lignin polymerization were confined to SCW thickenings in transdifferentiated xylem-like cells (Fig. 1D) (38). Similar confinement of xylan/lignin within hoop regions may support their role in assisting cellulose aggregation during SCW synthesis. Interestingly, we observed highly aggregated cellulose microfibrils not only within the SCW thickenings of transdifferentiated xylem-like cells but also in the nonlignified regions spanning between SCW thickenings that lack detectable LM10-labeled xylan (Fig. 1D) (38), suggesting that increased levels of cellulose aggregation might be possible without substantial contributions from xylan or lignin. Nevertheless, future studies should more thoroughly address the contribution of xylan and/or lignin in cellulose aggregation of SCWs, perhaps by combining the tools used in the current study with xylan- or lignin-deficient mutants. It is our hope that this study can serve as a foundation to help stimulate future investigations into how plants control and alter cell wall structure and synthesis.

**Materials and Methods**

**Plasmid Construction.** proCESA7::GFP–CESA7, 35S::VND7–GR, and proEB1b:: mCherry–EB1b were constructed by gateway cloning. See SI Materials and Methods for details on the construction.

**Plant Transformation and Generation of Lines.** CESAs2 transfer DNA (SALK_029040c, inx3-4) was obtained from the Arabidopsis Biological Resource Center and confirmed with the following primers: left genomic primer, 5′-AGAGAAAGC-TTAAGAAACCCGC3′; right genomic primer, 5′-GAAACAAACAGGACAAGGC3′; and left T-DNA border primer, 5′-ATTIGCGGTATGGCCAGAC3′. To obtain the GFP–CESA7 transgenic line, proCESA7::GFP–CESA7 was transformed into inx3-4 using the floral dip method of Agrobacterium-mediated transformation. Both 35S::VND7–GR and proEB1b::mCherry–EB1b were transformed into WT (Col-0) Arabidopsis. A cross was made to obtain a line expressing both GFP–CESA7 and 35S::VND7–GR. To obtain a line expressing GFP–CESA7, 35S::VND7–GR, and mCherry–EB1b, a cross was made between a homozygous GFP–CESA7 line and a homozygous line containing VND7–GR and mCherry–EB1b.

**Transgenic lines with markers of CSCs during PCW synthesis were previously described as follows:** GFP–CESA3 cesa3+++ (6), GFP–CESA3 cesa3+++–RFP–TUAS (16), mCherry–CESA3 cesa3+++ (19), GFP–CESA5 (26), GFP–CESA6 cesa6−/− (6), YFP– CESA6 cesa6−/−–RFP–TUAS (16), mCherry–CESA6 cesa6−/−–RFP–TUAS (13), GFP–CS1 cs1−/− (17), GFP–CS1 cs1−/− (17), RFP–CS1 cs1−/− (27), and GFP–KOR1 kor1−/− (19).

**Induction of Transdifferentiation and Drug Treatment.** Seeds were surface sterilized with 30% (wt/vol) bleach for 15 min, washed three times with autoclaved double-distilled H2O (ddH2O), and stored at 4 °C for 3–7 d. Seeds were grown for 3 d at 21 °C in the dark on vertical half-strength Murashige and Skoog (MS) plates containing 1% sucrose. Three-day-old seedlings were placed in open microfuge tubes containing half-strength MS 35 °C (wt/vol) sucrose liquid medium that was supplemented with either 20 μM DEX for induced samples or an equivalent volume of the solvent, dimethyl sulfoxide (DMSO), for control seedlings. Seedlings that were subjected to live-cell fluorescence imaging were treated for 60 h with DEX or DMSO. After DEX or DMSO treatment, samples for SFG analysis were frozen on MS plates containing 1% sucrose at −80 °C overnight. After thawing, groups of 20–30 seedlings were aligned parallel to close to one another in a single layer on glass slides and air dried before SFG analysis. All other analyses were conducted on fresh, never-frozen, never-dried samples.

**SFG Vibrational Spectroscopy Analysis.** The broadband SFG spectroscopy system and common SFG spectra collecting procedures were previously described (19, 39). SFG spectroscopic probing was performed in the reflection geometry and s-polarized sum frequency output, s-polarized visible input, and p-polarized IR input (SFG, 800 nm, and IR, respectively) polarization combination. To account for inherent heterogeneity within each sample, SFG data collection was performed at four different locations on each sample and averaged to generate a representative SFG spectrum for each sample.

**FE-SEM.** Epidermal peels from DEX or DMSO-treated hypocotyls were gently shaken in buffer containing 20 mM Hepes and 0.1% Tween-20 for 2 h to remove membrane material and subjected to five 3-min washes with ddH2O. Peels were transferred to 200-mesh grids with the inner part of the peel facing up and treated with 100% (wt/vol) ethanol (USP KOPTEC) for 30 min. Samples were dried with a Leica EM CPD300 automated critical point drier and coated with iridium using a Emitech K575X sputter coater. Images were obtained using a Zeiss Sigma VP-FESEM.
Cells of epidermal peels of transdifferentiated seedlings remained intact and required focused ion beam ablation to open the cells with an FEI Quanta 3D FEG focused ion beam. After focused ion beam ablation, samples were sputter-coated once again to coat the newly exposed regions and imaged.

**Live-Cell Imaging and Analysis.** The imaging system consisted of a Leica DMi6000 microscope with a Leica 100×/1.4 NA oil objective, a Yokogawa CSUX1 spinning disk system, and a Photometrics QuantEM:512SC CCD camera as described previously (20). Aciousto-opaque tunable filter-controlled excitation lasers (491 nm for GFP and 561 nm for RFP and lignin autofluorescence) and emission filters (520/50 nm for GFP and 620/60 nm for RFP and lignin autofluorescence) were used for fluorescence imaging. An integrated iLas photobleaching assembly (Roper Scientific) with a 405-nm laser was used for photobleaching experiments. Image acquisition was controlled using Metamorph (Molecular Devices) software. See SI Materials and Methods for details on image acquisition and data analysis.

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To construct the GFP and 5′ (and Movie S2), the two lateral pCR8/GW/TOPO pGWB2(35S) were digested with EB1b CTCTCTGAAACGAA−3′ EB1b′ GGAGCAATGGCGACGAA−linker and ′F was scored to both the pCR8−direction was cloned into " and 5′ encoding fragment); ′F′ and 5′ GFP-CESA7 and 5′ proEB1b:mCherry-EB1b was created by using Gateway LR Clonase to perform a ′F′ and the coding sequence of VND7-GR were cut with KpnI, ligated together to create the was amplified with primers 5′-CCGGTACCATGGAAGCTAGCGCCGT-3′ and 5′-TCAGCACTTGGATGCACACTGGGATG-3′. The coding sequence of GFP was amplified with primers 5′-ATGGTGACCAAGGCGAGGAGCTTTC-3′ and 5′-CCG- GTACCCCTGTACAGCTCTCCATGC-3′. Both CESA7 and GFP were cut with KpnI, ligated together to create the GFP-CESA7 sequence, and PCR-amplified. GFP-CESA7 was cloned into pCR8/GW/TOPO to produce pCR8-GFP-CESA7 and confirmed by sequencing. To generate proCESA7::GFP-CESA7, a recombination reaction was preformed between pGW2-proCESA7 and pCR8-GFP-CESA7 using Gateway LR Clonase (catalog no. 12558-120; Invitrogen).

To produce the proEB1b:mCherry-EB1b construct, the promotor of EB1b (At5G62500) was amplified with primers 5′-ATGGA-TAATATAATGCAATCTGTAATGCAATCC-3′ and 5′-CCGGAATCCCTCAAAGCTACATCAAG-3′ (containing the overhang of a mCherry fragment); mCherry was amplified with primers 5′-GGTGAAGGTCTAAAAATGTGGAGAAAGGGCGAGGAGGATATA-3′ (containing the overhang of the EB1b promoter fragment) and 5′-AATGTTGCGTGGCATCCTGCTGTCCTGCTGACCTGTCCTGC-3′ (containing overhangs of the "glycine-alanine" linker encoding sequence and the EB1b encoding fragment); EB1b genomic sequence was amplified with primers 5′-GACG-GGCGCTGTAAGGGAGGAGGAGCAATGGCCGAGGAAATTGGAATCAA-3′ and 5′-ATACCATATTCTTGGATATTTC-3′. The above-mentioned three PCR fragments were further linked together by fusion PCR. The resulting PCR fragment was cloned into the pENTR/D-TOPO (catalog no. K2500-20; Invitrogen) vector to produce pENTR-proEB1b:mCherry-EB1b, confirmed with sequencing, and delivered into pEarlyGate302 by recombination to get the final proEB1b:mCherry-EB1b construct.

To analyze the direction of CSC particle movement along a track was performed through kymograph analysis on a track-by-track basis. Kymographs were generated using the multiple kymograph tool in ImageJ with a one-pixel line-width setting. Care was taken to select a single track for each kymograph and to minimize the possibility of capturing particles from multiple adjacent tracks within a single kymograph. For each kymograph, the number of particles that traveled from left-to-right and from right-to-left was recorded. The percentage of particles that moved in that direction was then calculated for each track. If a majority of particles moved in the same direction along a given track, that direction was assigned as the favored direction and the alternative direction was assigned as the opposed direction. If an equal number of particles moved in each direction along a track, “50%” was scored to both the “favored” and “opposed” direction dataset. A minimum of 48 kymographs and 6 seedlings from each experimental condition was used to calculate the average percentage of particles moving in the favored and opposed direction.

Lateral photobleaching experiments were conducted such that the two lateral sides of a cell were bleached, leaving a strip of unbleached GFP-CESA particles down the center of the cell. The movement of PM-localized GFP-CESA particles was then tracked for 5 min after photobleaching to discern whether the particles moved from the center strip to the left and/or right bleached region.

To analyze the correlation between directionally coherent GFP-CESA7 movement and CMT polarity, kymographs of both GFP-CESA7 and RFP-EB1b were generated from shared tracks. If 65% or more of EB1b particles were moving in a common direction, that direction was assigned as the microtubule plus-end of the track. If less than 65% of EB1b particles were moving in a common direction, the track was deemed not to have a well-defined microtubule polarity. Likewise, if 65% or more GFP-CESA7 particles were moving in a common direction, the track was considered to exhibit directionally coherent CSC movement either toward the apical hook of 3-d-old dark grown seedlings, a common model tissue for PCW synthesis. Seedlings used for imaging during PCW synthesis were grown on half-strength MS plates with 0% sucrose, and slides were prepared with half-strength MS liquid medium with 0% sucrose.

Images were acquired every 5 s for periods of 5–8.5 min to generate time-lapse movies to assess the localization, dynamics, and distribution of fluorescent protein-tagged proteins of interest. Photobleaching was caused by three rapid pulses with a 405-nm laser in designed ROIs. In lateral photobleaching experiments designed to visualize the directional movement of GFP-CESA particles (Fig. 2 C, F, I, and L and Movie S2), the two lateral regions of cells were bleached during time-lapse imaging. In these experiments, the bleach event occurred at the 2-min time point of 7-min time-lapse acquisition. In photobleaching experiments designed to measure the delivery rate of CSCs (Fig. 3 B and Movie S3), a region that was ~20 μm long and spanned the entire width of the cell was bleached at the 30-s time point of 8.5-min time-lapse acquisitions.

Image Analysis. The velocities of CSC particles were measured using the automated particle-tracking feature of Imaris software (Bitplane). Particle tracking parameters were set to exclude Golgi compartments and remove noise from the analysis. These parameters constrained the particle diameter to 250 nm and required each particle to be tracked continuously for more than 60 s (six frames). The accuracy of automated velocity measurements was confirmed by conducting manual kymograph-based particle velocity measurements in ImageJ (NIH).

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To analyze the correlation between directionally coherent GFP-CESA7 movement and CMT polarity, kymographs of both GFP-CESA7 and RFP-EB1b were generated from shared tracks. If 65% or more of EB1b particles were moving in a common direction, that direction was assigned as the microtubule plus-end of the track. If less than 65% of EB1b particles were moving in a common direction, the track was deemed not to have a well-defined microtubule polarity. Likewise, if 65% or more GFP-CESA7 particles were moving in a common direction, the track was considered to exhibit directionally coherent CSC movement either toward the apical hook of 3-d-old dark grown seedlings, a common model tissue for PCW synthesis. Seedlings used for imaging during PCW synthesis were grown on half-strength MS plates with 0% sucrose, and slides were prepared with half-strength MS liquid medium with 0% sucrose.
plus-end or toward the minus-end of the track. If less than 65% of GFP-CESA7 particles moved in a common direction, the track was deemed not to have well-defined coherent CSC movement.

To compare the percentage of PM area occupied by GFP-CESA signal, it was first necessary to select ROIs that lacked intracellular GFP signal. Selected ROIs were confirmed to lack interference from intracellular compartments by visualizing the dynamics of the GFP-CESA signal in time-lapse movies corresponding to the frame used for the analysis. The threshold tool was used to convert each ROI to a binary image such that pixels brighter than the threshold gray value were converted to white and pixels dimmer than the threshold gray value were converted to black. The percentage of white pixels in each binary ROI was used to represent the percentage of the ROI occupied by signal. At least four small ROIs were analyzed from each cell and averaged together to generate a representative data point for each cell. A minimum of 28 ROIs and 7 cells from 7 different seedlings were analyzed for each experimental condition. The selection of many small ROIs was favored over the selection of one large ROI because there was heterogeneity in the brightness of PM-localized GFP-CESA particles across a single cell in a single image. This heterogeneity prevented a single threshold gray value from being applied across a large ROI. Conversely, the use of small ROIs allowed for the proper tuning of the threshold gray value for each ROI.

The delivery rate of CSCs was measured by manually counting the number of bona fide PM-localized CSCs delivered within a photobleached ROI in the 5 min after bleaching. The bleached region for each delivery rate experiment was ∼20 μm long and encompassed the entire width of the cell. To be classified as a bona fide PM-localized CSC delivery event, the following criteria were required: the particle had to appear within the bleached region and not migrate in from the edge of the bleached region, the particle had to appear within the first 5 min after photobleaching, the particle had to establish linear lateral movement that is characteristic of active PM-localized CSCs during the 8 min of time-lapse imaging after bleaching. Each bleached region was dissected into smaller, more manageable ROIs for delivery rate analysis and the average delivery rate of multiple ROIs was used to represent each cell. A minimum of 14 ROIs from 7 cells from 7 different seedlings was measured. Because CSC delivery was restricted to hoop regions during hoop formation, only ROIs within hoop regions were analyzed during hoop formation. The delivery rate was expressed as delivery events per square micron per minute by dividing the number of delivery events by the area of each ROI and by 5 min.

The movies were assembled in ImageJ. Contrast enhancement (0.4% saturated pixels), background subtraction (rolling ball radius of 50 pixels), and walking average (4 frames) were applied to time-lapse movies for improved visualization of CSC particles. The draw tool was used to accentuate features within the movies and to display a scale bar. The frame rate of movies was adjusted to appropriately view the features of each movie and time stamps were added to movies to comprehend the playback speed of each movie.

Fig. S1. ProCESA7::GFP-CESA7 rescues the *cesa2*/*cesa3* growth phenotype and GFP-CESA7 signal localizes to similar populations as previously visualized PCW CESA markers such as GFP-CESA3. (A) Adult WT (Col-0 ecotype), ProCESA7::GFP-CESA7 *cesa2*/*cesa3*, and *cesa2*/*cesa3* plants revealed that ProCESA7::GFP-CESA7 rescued the *cesa2*/*cesa3* growth phenotype. (Scale bar: 5 cm.) (B–D) ProCESA3::GFP-CESA3 *cesa3*/*je5* was imaged in nontransdifferentiating cells as a CSC marker during PCW synthesis (B), and ProCESA7::GFP-CESA7 *cesa2*/*cesa3* 35S::VND7-GR was imaged during two stages of xylem cell transdifferentiation, before hoop formation and during hoop formation, as a CSC marker during SCW synthesis (C and D). Single-frame images and 5-min projection images show the distribution and trajectories of GFP-CESA particles, respectively (B–D). Three arrowheads denote diffraction-limited PM-localized CSC particles and octagons denote GFP-CESA signal in globular, intracellular Golgi bodies (B–D). PM-localized GFP-CESA7 often localized to swaths of signal in addition to well-defined puncta that are commonly seen in PCW CSCs such as GFP-CESA3. Arrows indicate the apical direction. (Scale bars: 10 μm.)
Fig. S2. The distribution of GFP-CESA7 particle velocities in transdifferentiating cells during SCW synthesis is not drastically different from that of markers of CSCs during PCW synthesis. The velocities of CSC particles were quantified for the 11 CSC markers during PCW synthesis including GFP-CESA3 cea3je5, GFP-CESA3 cesa3je5 RFP-TUA5, mCherry-CESA3 cesa3je5, GFP-CESA5, GFP-CESA6 cea6prc1-1, YFP-CESA6 cea6prc1-1 RFP-TUA5, mCherry-CESA6 cea6prc1-1, GFP-CSEI1 csi1-6, GFP-CSEI3 csi3-1, RFP-CSEI1 csi1-6, and GFP-KOR1 kor1-3 (gray). The velocities of GFP-CESA7–labeled CSCs were quantified during SCW synthesis before hoop formation (BHF), during hoop formation (DHF), and under oryzalin treatment (8–12 h; 25 μM) before hoop formation (BHF + Ory) (white). The line within each box represents the mean. The bottom and top of each box represents the first and third quartile, respectively. The error bars represent the SD of the mean; n > 5,700 particles from ≥5 seedlings for each data point.
Fig. S3. The direction of coherent CSC movement is not correlated with the polarity of the CMTs along which the CSCs are moving. (A) In transdifferentiating cells, GFP-CESA7–labeled PM-localized CSCs traveled along trajectories that coincided with the trajectories of RFP-EB1b particles, which label the plus-ends of polymerizing microtubules, both before and during hoop formation. (Scale bars: 10 μm.) (B–E) Kymograph analysis displays the direction of EB1b particle movement, from which the polarity of new microtubules can be deduced, and the direction of CSC particle movements from four representative tracks. (Scale bars: 5 μm.) The direction of coherent CSC movement was occasionally toward the plus-ends of newly synthesized CMTs (B) or toward the minus-ends of newly synthesized CMTs (C). Some tracks did not exhibit a dominant CMT polarity (D and E) or coherent CSC movement (E). The relationship between the direction of coherent CSC movement and the polarity of newly synthesized CMTs was quantified before and during hoop formation (F and G). N is 56 tracks from 7 seedlings for each pie chart.
Fig. S4. Treatment with 25-μM oryzalin for 8–12 h disrupts CMTs. (A) Motile RFP-EB1b particles demarcate newly polymerized CMTs in mock-treated seedlings. (B) RFP-EB1b no longer localizes to particles at the cell cortex in oryzalin-treated seedlings but rather displays a faint cytosolic signal, which suggests that CMT polymerization is abolished. (C and D) GFP-MAP4 localizes to CMTs in mock-treated seedlings (C) but is diffusely distributed throughout the cytosol in oryzalin-treated seedlings, which suggests that CMTs have depolymerized (D). (Scale bars: 10 μm.)

Movie S1. The dynamic characteristics of GFP-CESA3 during PCW synthesis and GFP-CESA7 during SCW synthesis at various stages of transdifferentiation are displayed by time-lapse imaging. CSC dynamics during PCW synthesis are represented by GFP-CESA3 particles in nontransdifferentiated (NT) seedlings. CSC dynamics during SCW synthesis is represented by GFP-CESA7 particles before hoop formation (BHF), during hoop formation (DHF), and under oryzalin treatment (BHF + ORY). The frame rate is 60 frames per second. (Scale bar: 10 μm.)

Movie S1
Movie S2. Lateral photobleaching experiments display the bidirectional movement of GFP-CESA3 particles during PCW synthesis and the directionally coherent movement of GFP-CESA7 particles during SCW synthesis. Time-lapse imaging was conducted for 2 min before photobleaching the lateral edges of the cells. In PCW-synthesizing nontransdifferentiated (NT) seedlings, GFP-CESA3 particles on a shared track move equally into the right and left bleached areas during PCW synthesis. In contrast, GFP-CESA7 particles on a shared track move either into the left or right bleached area during SCW synthesis both before (BHF) and during (DHF) hoop formation. Oryzalin treatment (BHF + ORY) did not disrupt the directionally coherent movement of GFP-CESA7 particles during SCW synthesis. The frame rate is 15 frames per second. (Scale bar: 10 μm.)

Movie S3. The delivery of new CSCs to the PM is visualized after photobleaching during PCW synthesis and during SCW synthesis. The movie displays a single frame before photobleaching, followed by a single frame immediately after photobleaching and a 5-min period after photobleaching during which GFP-CESA particles repopulated the bleached region through the delivery of new CSCs to the PM. GFP-CESA3 particles in nontransdifferentiated (NT) seedlings represent CESA delivery during PCW synthesis, and GFP-CESA7 particles before hoop formation (BHF) and during hoop formation (DHF) represent CSC delivery during SCW synthesis. The frame rate is 15 per second. (Scale bar: 10 μm.)