Characterization of Cellulose Synthesis Complexes in *Physcomitrella patens*

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CHARACTERIZATION OF CELLULOSE SYNTHESIS COMPLEXES
IN PHYSCOMITRELLA PATENS

BY

XINGXING LI

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OF

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ABSTRACT

Due to the enormous economic value and significance of cellulose in human consumption and plant cell walls, production of cellulose microfibrils is considered to be one of the most critical biochemical processes in plant biology. In the past decades, cellulose biosynthesis has been extensively studied in vascular plants. More and more fundamental questions related to this key process are being answered. One such question is: What are the protein components of the enzymatic complex for cellulose synthesis? In seed plants, membrane-embedded rosette Cellulose Synthesis Complexes (CSCs) producing cellulose microfibrils are obligate hetero-oligomeric, being assembled from three functionally distinct and non-interchangeable cellulose synthase (CESA) isoforms. For instance, Arabidopsis has two types of CSCs. One contains AtCESA1, AtCESA3, and AtCESA6, involved in cellulose synthesis in primary cell walls; the other consists of AtCESA4, AtCESA7, and AtCESA8, specialized for secondary cell wall deposition. Recently, the stoichiometry for the three Arabidopsis CESAs forming a CSC was determined to be a 1:1:1 molecular ratio. The constructive neutral evolution hypothesis has been proposed as a mechanism for evolution of these hetero-oligomeric complexes.

Physcomitrella patens, a non-vascular plant, is one of the most popular models for genetics studies. A relatively small genome, dominant haploid phase, and high rate of homologous recombination make P. patens a simple and efficient system for genetic manipulation. Seven CESA genes (PpCESA3, PpCESA4, PpCESA5, PpCESA6, PpCESA7, PpCESA8, and PpCESA10) were identified in the P. patens genome, but proteins encoded by these genes are not orthologs of functionally distinct seed plant
CESAs according to phylogenetic studies. The similar rosette-type of CSCs were observed in *P. patens* by freeze-fracture electron microscopy. It is not yet known whether the *P. patens* CSCs are homo-oligomeric complexes consisting of only a single type of CESA, or hetero-oligomeric complexes assembled by different CESAs like those in seed plants. Knowing this information would be helpful for understanding the roles of different CESAs that compose seed plant CSCs. Furthermore, answers to this question potentially will be useful for testing the constructive neutral evolution hypothesis, since moss CESAs diversified independently from seed plant CESAs.

In this study, I generated PpCESA knock out (KO) mutants. Morphological analyses were carried out to identify mutant phenotypes of these KOs together with several previously made KO mutants. Cellulose defects in these mutants were also analyzed using quantitative methods. Reverse transcriptase PCR (RT-qPCR) was performed to examine the expression of all seven PpCESAs in KO lines to identify co-expressed PpCESAs that potentially reside within the same CSCs as the deleted PpCESA. Immunoblot analysis using specific monoclonal antibodies was used as an additional method to detect co-expression based on the accumulation of the protein products of these *PpCESA* genes. Finally, I carried out Co-immunoprecipitation (Co-IP) assays to identify potential physical interactions between different PpCESA isoforms. The results show that functionally distinct CESA isoforms have evolved in the moss *P. patens* independently from seed plants, and CSCs synthesizing cellulose microfibrils in secondary cell walls of *P. patens* gametophore leaves are obligate hetero-oligomeric complexes. Meanwhile, our research also suggests that PpCESA5 alone is able to form
homo-oligomeric CSCs, making *P. patens* an intriguing model in which to study the evolution of cellulose synthase.
ACKNOWLEDGMENTS

Special thanks to my major advisor, Dr. Alison Roberts, and co-advisor, Dr. Joanna Norris, for giving me an opportunity to join their lab and guiding me on all of my projects. During the four and half years of my graduate school, not only I learned lab techniques from Dr. Roberts and Dr. Norris, but also I learned scientific thinking from them which will be important for my future career. As an international student whose native language is not English, I really appreciate the kind assistance of my advisors in writing of my manuscripts. I would like to thank my committee members, Dr. Gongqin Sun and Dr. Steve Irvine, for their valuable time and comment for my projects and dissertation. I would like to thank the lab members, Mai Tran, Arielle Chaves, and Tess Scavuzzo-Duggan, for teaching me lab skills, helping me with trouble shooting, and providing insights into my projects. I especially appreciate Dr. Jodi Camberg, Dr. Niall Howlett, Dr. Gongqin Sun, Dr. Steve Irvine and their students for allowing me to use their equipment, sharing their protocols, and helping me setup experiments. Finally, I would like to thank my parents for being supportive financially and spiritually throughout these years of my life in the United States.
PREFACE

Manuscript format is used in this dissertation.

Chapter 2 is a manuscript published in *Plant Physiology* 175:210-222 on August 2, 2017 in collaboration with Joanna H. Norris, Mai L. Tran, Bailey Mallon, Danielle Mercure, and Arielle M. Chaves at the University of Rhode Island; Shixin Huang and Seong H. Kim from the Pennsylvania State University; Ashley Tan and Rachel A. Burton from University of Adelaide, Australia; and Allison M.L. Van de Meene and Monika L. Doblin from University of Melbourne, Australia. In this manuscript, I did the phenotypic analysis for some *PpCESA* knockout (KO) mutants and quantitative PCR analysis for expression of the *PpCESA* genes in the KO mutants. Results of these experiments lead to a hypothesis: obligate hetero-oligomeric Cellulose Synthesis Complexes (CSCs) are involved in the *P. patens* secondary cell wall deposition.

Chapter 3 is a manuscript that describes work done in collaboration with Mai L. Tran and Joanna H. Norris at the University of Rhode Island. This manuscript includes morphological analysis for several *PpCESA* KO mutants that identified roles for *PpCESA4* and *PpCESA10* in tip-growing protonema. In this manuscript, I generated the quadruple *ppcesa4/6/7/10* KO mutants. I also did most of the morphological assays, except the assays for *ppcesa6/7* KOs and *ppcesa3/8* KOs.

Chapter 4 is a manuscript that includes work done in collaboration with John McManus from the University of Pennsylvania State University. In this manuscript, the hypothesis proposed in Chapter 1 is directly tested by reverse transcriptase PCR and Co-immunoprecipitation. In this manuscript, the majority of the work was done by me, except the creation of antibodies. The manuscript is currently being prepared for
submission. Mass spectrometry (M.S.) data will be added to the manuscript before submission. Currently, the M.S. data will be collected and analyzed with our collaboration with Ian S. Wallace and Tori Speicher at University of Nevada, Reno.
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CHAPTER 1

Introduction

Cellulose synthesis complex (CSC)

Cellulose is a biopolymer of β(1,4)-linked glucose that forms the microfibrils essential in most plant cell walls. It is extensively used for a variety of commercial and industrial purposes including lumber and textiles. The synthesis of cellulose in plants is catalyzed by enzymatic complexes called cellulose synthesis complexes (CSCs) located in plasma membrane (Delmer et al., 1999; Somerville et al., 2006; McFarlane et al., 2014). The membrane-bound CSCs were first observed to have a "rosette" structure and to be associated with the ends of microfibrils in freeze-fracture electron microscopy studies on maize (Mueller & Brown, 1980). By searching a cotton fiber EST library for sequences similar to a bacterial cellulose synthase gene, the first putative plant gene encoding a cellulose synthase catalytic subunit (CESA) was identified (Pear et al., 1996). Antibodies against cotton CESAs were later produced to label the rosettes in freeze-fractured bean hypocotyls indicating CESAs are components of the multi-protein complexes inserted into the plasma membrane (Kimura et al., 1999).

Cellulose synthase catalytic subunits (CESAs)

Among currently identified protein components in CSCs, CESAs are implicated by all sorts of evidence (Delmer et al., 1999; Somerville et al., 2006; McFarlane et al., 2014) to be the only functional subunits that produce individual glucan chains. Recently, a
heterologously-expressed CESA isoform, PttCESA8, from *Populus* tremula x tremuloides (hybrid aspen), was reconstituted in liposomes and shown to be functional for cellulose microfibril formation in vitro (Purushotham et al., 2016) for the first time.

The CESA family is contained within the glycosyltransferase-2 (GT-2) superfamily characterized by an eight-transmembrane-helix topology and conserved cytosolic substrate binding and catalytic site (McFarlane et al., 2014). The site for substrate binding and catalysis consists of a D, DxD, D, QxxRW motif and is predicted to be in the loop bounded by transmembrane helix 2 and 3 (Pear et al., 1996). In this motif, the first two conserved aspartic acid residues are predicted to bind the substrate, UDP-glucose. This was supported by the results of mutational analysis (Pear et al., 1996). The functions of these residues have been confirmed by x-ray crystallography of bacterial cellulose synthase (Morgan et al., 2013). The third aspartic acid is thought to be involved in the addition of UDP-glucose to the existing glucan, and the QxxRW region is predicted to be a binding site for the growing glucan chain (Morgan et al., 2013). Compared with bacterial cellulose synthase, plant CESAs are larger. That is because the plant CESA also contains an extended N-terminal Zn-binding RING finger domain, a plant-conserved domain within the N-terminal cytoplasmic loop, and a class-specific domain within the central cytoplasmic loop in addition to the conserved catalytic region (Pear et al., 1996). These domains are specific to plants, hence they are thought to be important for the interactions between the CESA subunits and presumably involved in the formation of the rosette CSCs (McFarlane et al., 2014).
Interactions between CESAs

*CESA* genes are members of multigene families in plants. For example, *Arabidopsis* has 10 *CESA* genes from which distinct combinations are required for primary and secondary cell wall synthesis (McFarlane et al., 2014). The *AtCESA4*, *AtCESA7*, and *AtCESA8* genes were first shown to be specifically involved in secondary cell wall deposition (Turner & Somerville, 1997; Taylor et al., 1999; Taylor et al., 2000; Taylor et al., 2003). The mRNAs of the three genes are found to be coregulated in microarray analysis (Brown et al., 2005; Persson et al., 2005a). Proteins encoded by the three genes physically interact and are exclusively required for assembly of CSCs in cells with thickened secondary walls (Taylor et al., 2000; Taylor et al., 2003).

Mutations in *AtCESA1*, *AtCESA3*, and *AtCESA6* cause primary cell wall defects (Arioli et al., 1998; Fagard et al., 2000; Burn et al., 2002; Robert et al., 2004).

*AtCESA3* and *AtCESA6* interact with each other according to results of in vitro pull-down assays, and Bimolecular Fluorescence Complementation (BiFC) experiments show that *AtCESA1*, *AtCESA3*, and *AtCESA6* can interact in vivo (Desprez et al., 2007). *AtCESA2* and *AtCESA5* were shown to be closely related and partially functionally redundant with *AtCESA6* (Desprez et al., 2007). In *Arabidopsis*, therefore, a primary wall CSC might consist of *AtCESA1*, *AtCESA3*, and one or perhaps several *AtCESA6* like *AtCESAs* (McFarlane et al., 2014).

Characterization of CSCs has also been carried out in another vascular plant model, *Populus trichocarpa*. Two types of CSCs are identified in the xylem of *Populus* by co-immunoprecipitation (Co-IP) experiments: one type contains PdxtCESA7A and PdxtCESA8B; the other one contains PdxtCESA1A and PdxtCESA3 (Song et al.,
2010a). Altogether, current evidence suggests that vascular plant CSCs are obligately hetero-oligomeric. A theory known as constructive neutral evolution addresses how homo-oligomer complexes are driven towards hetero-oligomeric by neutral processes during evolution. According to this theory, in the initial complex assembled from multiple copies of the same subunit, additional obligate subunits could be evolved by gene duplication followed by relatively high frequency degenerative mutations causing specific interaction sites among them to be lost (Doolittle, 2012; Finnigan et al., 2012). A study showed that the extant Vo complex of the fungi V-ATPase proton pump which is composed of three obligate subunits, evolved from an ancient two-subunit complex by a gene duplication and subsequent complimentary loss of specific interfaces on each daughter isoforms on which they rely to interact with other subunits in the complex (Finnigan et al., 2012). So far, this is the only study that provided convincible experimental evidence. Hence, the generality of this hypothesis needs to be further tested. Plant CSCs are similar to the fungal Vo complex, which are also composed by paralogous CESA isoforms sharing a considerable amount of similarities. Thus, characterizing the CESAs in plant CSCs will be helpful for continuing testing this theory.

Other components of the CSC

Other than CESAs, several other protein components (Endo et al., 2009; Gu et al., 2010) of seed plant CSCs have been identified successively by Co-IP and BiFC. For instance, a putative endo-1,4-β-D-glucanase, KORRIGAN1 (KOR1), was identified to be a part of the primary cell wall CSCs in Arabidopsis (Vain et al., 2014). A microtubule-binding protein, Cellulose synthase interactive protein 1 (CSI1), was
discovered to associate with CSCs and serve as a linker protein between CSCs and microtubule (Li et al., 2012). Genetic evidence and the observed size of the cytosolic portion of the rosette demonstrated in electron micrographs (Bowling & Brown, 2008) imply that more other proteins related to cellulose synthesis might also participate in assembly of CSCs.

The moss *Physcomitrella patens*

*Physcomitrella patens*, a moss species, has also been shown to have rosette CSCs, but not members of the CESAs clades that contain the functionally distinct isoforms of the hetero-oligomeric CSCs in seed plants (Roberts et al., 2012). The *PpCESA* family includes seven members that cluster in two clades (Roberts & Bushoven, 2007). The A-clade contains *PpCESA3*, *PpCESA5*, and *PpCESA8*. The B-clade contains *PpCESA4*, *PpCESA6*, *PpCESA7*, and *PpCESA10*. Currently, the functions of these *P. patens* CESAs are still under investigation. It is also not yet known if *P. patens* and other mosses have homo-oligomeric or hetero-oligomeric CSCs. Understanding whether or not the PpCESAs serve distinct functions from those of the seed plant CESAs and determining the organization of *P. patens* CSCs will provide insight into the roles of the different CESA isoforms forming CSCs in seed plants, and possibly allow us to test the constructive neutral evolution hypothesis. *Physcomitrella patens* has many advantages as a research model for studying genetics, including a relatively small and fully sequenced genome and predominate haploid phase (Rensing et al., 2008; Zimmer et al., 2013). More importantly, *P. patens* is capable of being genetically manipulated as a result of its high rate of homologous recombination (Reski & Frank, 2005; D. G. Schaefer & Z., 1997). Taking advantage of this
unique property, functions of genes of interest can be identified by knockout (KO) mutations (Schaefer, 2002).

**Thesis outline**

*ppcesa5*KO*Os have cellulose defects in primary cell walls affecting gametophore bud development and resulting in a "no leafy gametophore" phenotype (Goss et al., 2012). However, other single *PpCESA* KOs do not show obvious phenotypic changes. To investigate functions for the other *PpCESA*s, double *PpCESA*s KO mutants (*ppcesa3/8KO, ppcesa6/7KO, and ppcesa4/10KO*) were generated. In manuscript 1, we show that *ppcesa3/8KO* has defects in secondary cell wall deposition in gametophore leaf midribs (Norris et al., 2017). I used reverse transcription quantitative PCR (RT-qPCR) analysis to measure the expression of *PpCESA3* and *PpCESA8* in corresponding *PpCESA* KO mutants. The results show that *PpCESA8* is up-regulated for the loss of *PpCESA3*, suggesting these two *PpCESA*s are partially functionally redundant. My phenotypic analysis of other double KO mutants revealed that *ppcesa6/7KO* also have significantly decreased cellulose deposition in the midribs of gametophore leaves. This indicates that *PpCESA3, PpCESA8, PpCESA6*, and *PpCESA7* are all involved in cellulose deposition during secondary cell wall formation in gametophore leaves and may be members of the same CSCs. *Ppcesa4/10KO* showed slightly but significantly decreased cellulose content in the midribs of gametophore leaves, suggesting a minor role of these two *PpCESA*s during secondary cell wall deposition. Together, this study provided important clues for characterization of composition and function of *P. patens* CSCs.
On the protein sequence level, PpCESAs within the same clade are highly similar, compared to lower similarity between the clade A and B PpCESAs. For instance, the clade B PpCESAs are 90-99% similar to each other and PpCESA6 and PpCESA7 differ by only three amino acids in protein sequence (Wise et al., 2010; Norris et al., 2017). This indicates that PpCESAs from this clade may have overlapping functions, which can mask potential phenotypic defects when carrying out mutational analysis.

In manuscript 2, I show that quadruple *ppcesa4/6/7/10* KOs in which all B-clade PpCESAs are knocked out have dramatically reduced cellulose deposition in the midribs of gametophores as expected. However, overall morphology of leafy gametophores is normal in these quadruple KOs indicating the clade B PpCESAs are not required for gametophore morphogenesis. Since *ppcesa3/8* KOs also produce normal looking gametophores (Norris et al., 2017), current results of mutational analyses are consistent with the hypothesis that PpCESA5 forms homo-oligomeric CSCs responsible for cellulose deposition in primary cell walls during gametophore bud development.

In manuscript 2, I used quantitative analysis of colony morphology, to show that quadruple KOs are defective in tip-growth of protonemal filaments indicated by significantly increased circularity and solidity of protonema colonies regenerated from single protoplasts. Later, I found that knocking out *PpCESA4* and *PpCESA10* together is enough to cause this phenotype. *Ppcesa6/7* KOs are not different from wild-type in protonema colony morphology. Different phenotypes of *ppcesa6/7* KOs and *ppcesa4/10* KOs might be related to different gene expression patterns. As shown by previous studies, *PpCESA6* and *PpCESA7* are expressed in leafy gametophores at
higher levels, while \textit{PpCESA4} and \textit{PpCESA10} have higher expression in protonema filaments (Hiss et al., 2014; Tran & Roberts, 2016). The mutant phenotype of \textit{ppcesa4/10}KOs indicates the \textit{PpCESA4} and \textit{PpCESA10} play some roles in tip-growing protonema cells, supporting the idea that cellulose is an essential cell wall component in cells undergoing tip growth (Newcomb & Bonnett, 1965; Emons & Wolters-Arts, 1983; Emons, 1994; Cosgrove, 2005; Park et al., 2011).

Results of mutational analyses suggest that CSCs involved in cellulose deposition in \textit{P. patens} secondary cell walls might be hetero-oligomeric, consisting of \textit{PpCESAs} from both A-clade and B-clade and I tested this hypothesis in manuscript 3. First, I measured expression of all seven \textit{PpCESAs} by RT-qPCR analysis in knockout mutants in order to identify which \textit{PpCESAs} are downregulated, as predicted for those that reside within the same CSC as the deleted \textit{PpCESA}. Results show that gene expression of \textit{PpCESA3}, \textit{PpCESA8}, and \textit{PpCESA7} are co-regulated. Western blot analysis of the microsomal proteins isolated from wild-type \textit{P. patens} showed that \textit{PpCESA3}, \textit{PpCESA8}, and \textit{PpCESA6/7} are highly expressed in gametophores which is consistent with cellulose defects in secondary cell walls of corresponding KO mutants. The Co-immunoprecipitation (Co-IP) experiments show that \textit{PpCESA3} and \textit{PpCESA8} can both interact with \textit{PpCESA6/7} \textit{in planta}. Taken together, these results are consistent with the hypothesis that \textit{PpCESA3}, \textit{PpCESA8}, \textit{PpCESA6}, and \textit{PpCESA7} form obligate hetero-oligomeric CSCs that produce cellulose microfibrils during secondary cell wall deposition in \textit{P. patens} gametophore leaves.

To summarize, my work reveals: 1) In the moss \textit{P. patens}, CSCs that synthesize cellulose in secondary cell walls are obligate hetero-oligomeric, with members from
clade A and clade B; 2) PpCESA4 and PpCESA10 function in elongating protonemal implying important role of cellulose in tip growth; 3) Clade B PpCESAs are not required for gametophore morphogenesis, which also means PpCESA5 possibly can form homo-oligomeric CSCs. Taken together, these discoveries indicate that functional specialization of CESAs occurred independently in mosses and seed plants through both subfunctionalization and neofunctionalization, which are consistent with the theory of constructive neutral evolution providing a possible mechanism for the convergent evolution of plant CSCs.
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Short title: CESA functional specialization in *Physcomitrella*

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Title: Functional specialization of cellulose synthase isoforms in a moss shows parallels with seed plants

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One sentence summary: Regulatory uncoupling of primary and secondary cellulose synthases occurred independently in mosses and seed plants, and is associated with convergent evolution of secondary wall structure.

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Abstract

The secondary cell walls of tracheary elements and fibers are rich in cellulose microfibrils that are helically oriented and laterally aggregated. Support cells within the leaf midribs of mosses deposit cellulose-rich secondary cell walls, but their biosynthesis and microfibril organization have not been examined. Although the Cellulose Synthase (CESA) gene families of mosses and seed plants diversified independently, CESA knockout analysis in the moss Physcomitrella patens revealed parallels in CESA functional specialization of Arabidopsis and P. patens, with roles for both sub-functionalization and neo-functionalization. The similarities include regulatory uncoupling of the CESAs that synthesize primary and secondary cell walls, a requirement for two or more functionally distinct CESA isoforms for secondary cell wall synthesis, interchangeability of some primary and secondary CESAs, and some CESA redundancy. The cellulose-deficient midribs of ppcesa3/8 knockouts provided negative controls for structural characterization of stereid secondary cell walls in wild type P. patens. Sum frequency generation spectra collected from midribs were consistent with cellulose microfibril aggregation, and polarization microscopy revealed helical microfibril orientation only in wild type leaves. Thus, stereid secondary walls are structurally distinct from primary cell walls, and they share structural characteristics with the secondary walls of tracheary elements and fibers. We propose a mechanism for convergent evolution of secondary walls in which deposition of aggregated and helically oriented microfibrils is coupled to rapid and highly localized cellulose synthesis enabled by regulatory uncoupling from primary wall synthesis.
Introduction

In vascular plants, cellulose is a major component of both primary cell walls that are deposited during cell expansion and secondary cell walls that are deposited after expansion has ceased (Carpita and McCann 2000). Secondary cell walls of water-conducting tracheary elements and supportive fibers are rich in cellulose with microfibrils arranged in helices that vary in angle according to developmental stage and environmental conditions (Barnett and Bonham 2004). Secondary cell wall microfibrils are also more aggregated than those of primary cell walls (Donaldson 2007; Fernandes et al. 2011; Thomas et al. 2014). Recently, Sum Frequency Generation (SFG) spectroscopy has been used to compare the mesoscale structure of cellulose microfibrils in primary and secondary cell walls. Both high cellulose content and microfibril aggregation contribute to a strong secondary cell wall signature in SFG spectra of mature angiosperm tissues (Barnette et al. 2012; Lee et al. 2014; Park et al. 2013).

Cellulose microfibrils are synthesized by cellulose synthase (CESA) proteins that function together as cellulose synthesis complexes (CSCs) in the plasma membrane (Delmer 1999; Kimura et al. 1999). Recent analyses of CSC and microfibril structure indicate that the rosette CSCs of land plants most likely contain 18 CESA subunits (Fernandes et al. 2011; Jarvis 2013; Newman et al. 2013; Nixon et al. 2016; Oehme et al. 2015; Thomas et al. 2014; Vandavasi et al. 2016) in a 1:1:1 ratio (Gonneau et al. 2014; Hill et al. 2014). Seed plants have six phylogenetic and functional classes of CESA proteins, three required for primary cell wall synthesis (Desprez et al. 2007; Persson et al. 2007) and three required for synthesis of the lignified secondary cell
walls of tracheary elements and fibers (Taylor et al. 2003). Mutation of any of the secondary CESAs results in a distinctive irregular xylem phenotype characterized by collapsed xylem tracheary elements and weak stems (Taylor et al. 2004). The secondary cell wall CESAs of Arabidopsis are regulated by master regulator NAC domain transcription factors that also activate genes required for the synthesis of other secondary cell wall components, such as xylan and lignin (Schuetz et al. 2013; Yang and Wang 2016; Zhong and Ye 2015).

The moss *Physcomitrella patens* (Hedw.) B. S. G. has seven CESA genes (Goss et al. 2012; Roberts and Bushoven 2007). Phylogenetic analysis has revealed that the *P. patens* CESAs do not cluster with the six CESA clades shared by seed plants (Roberts and Bushoven 2007). Like other mosses, *P. patens* lacks the lignified secondary cell walls that are characteristic of vascular plant tracheary elements and fibers. However, mosses do have support cells (stereids) with thick unlignified cell walls (Kenrick and Crane 1997) and water-conducting cells (hydroids) that have thin cell walls and undergo programmed cell death like tracheary elements (Hebant 1977). Although the stereid cell walls of *P. patens* are known to contain cellulose (Berry et al. 2016), the mesoscale structure has not been examined. Only one of the seven *P. patens* CESAs has been characterized functionally. When *PpCESA5* was disrupted, gametophore buds failed to develop into leafy gametophores, instead forming irregular cell clumps. The associated disruption of cell expansion and cell division are consistent with an underlying defect in primary cell wall deposition (Goss et al. 2012). Recently it was shown that *PpCESA3* expression is regulated by the NAC transcription factor *PpVNS7*, along with thickening of stereid cell walls (Xu et al. 2014).
Here we show that PpCESA3 and PpCESA8 function in the deposition of stereid cell walls in the gametophore leaf midribs of *P. patens* and are sub-functionalized with respect to PpCESA5. We also used polarization microscopy and SFG to reveal similarities in the mesoscale organization of the microfibrils synthesized by PpCESA3 and PpCESA8 and those in the secondary cell walls of vascular plants. Finally, we propose a mechanism through which uncoupling of primary and secondary CESA regulation played a role in independent evolution of secondary cell walls with aggregated, helically arranged cellulose microfibrils in the moss and seed plant lineages.

**Results**

**PpCESA3 and PpCESA8 function in secondary cell wall deposition**

Cellulose synthase genes *PpCESA3* and *PpCESA8* were independently knocked out by homologous recombination in an effort to examine their roles in development and cell wall biosynthesis in *P. patens*. Stable antibiotic resistant lines generated by transforming wild type *P. patens* with CESA3KO or CESA8KO vectors were tested for integration of the vector and deletion of the target gene by PCR (Fig. S1). Integration was verified for five *ppcesa8KO* lines recovered from two different transformations, line 8KO5B from a transformation of the GD06 wild type line and lines 8KO4C, 8KO5C, 8KO7C and 8KO10C from a transformation of the GD11 wild type line (Fig. S1). Integration was verified for three *ppcesa3KO* lines recovered from a single transformation of GD11 and three double *ppcesa3/8KO* lines recovered from a single transformation of the *ppcesa8KO5B* line with the CESA3KO vector (Fig. S1).
The GD06 and GD11 lines are from independent selfings of the same haploid wild type line, as described in Materials and Methods.

The colonies that developed from wild type and KOs consisted of protonemal filaments and leafy gametophores (Fig. 1). Whereas wild type, ppcesa3KO, and ppcesa8KO gametophores grew vertically, the gametophores on ppcesa3/8KO colonies were unable to support themselves and adopted a horizontal orientation.

Superficially ppcesa3/8KO colonies appeared to produce fewer gametophores (Fig. 1), but dissection revealed similar numbers of horizontal gametophores that had been overgrown by protonemal filaments. Thus, PpCESA3 and PpCESA8 are not required for gametophore initiation or morphogenesis, but they appear to contribute to structural support.

When examined with polarized light microscopy, the wild type gametophore leaves exhibited strong cell wall birefringence in the midribs and margins (Fig. 1). In contrast, the leaves produced by ppcesa3/8KOs lacked strong birefringence in these cells, consistent with reduced crystalline cellulose content. The ppcesa3KO leaves appeared similar to wild type leaves (Fig. 1) and ppcesa8KO leaves had an intermediate phenotype. Staining with the fluorescent cellulose binding dye Pontamine Fast Scarlet (S4B) (Anderson et al. 2010) produced similar results with strong fluorescence in the midribs of wild type and ppcesa3KO leaves, weak fluorescence in ppcesa3/8KO leaves, and intermediate fluorescence in ppcesa8KO leaves (Fig. 1).

Cellulose Binding Module (CBM) 3a provides a third method for detecting cellulose and can be used to probe thin sections (Blake et al. 2006). In sections from fully expanded wild type leaves, the walls of the lamina cells were labeled relatively weakly
with CBM3a, whereas the thickened cell walls of the central midrib and bundle sheath cells were strongly labeled (Fig. 1). The same was true for ppcesa3KO leaves. However, midrib and bundle sheath cell labeling was nearly absent in ppcesa3/8KO and diminished in ppcesa8KO (Fig. 1) compared to wild type and ppcesa3KO. Differential interference contrast microscopy of the same sections showed enhanced contrast in wild type and ppcesa3KO midribs (Fig. 1). Partial cell collapse occurred during embedding in ppcesa3/8KO leaves (Fig. 1).

The cellulose content of the leaf midribs in wild type and single and double ppcesaKO mutants was quantified by measuring the intensity of S4B fluorescence. Statistical analysis confirmed that the S4B fluorescence was significantly reduced in double KOs, but not in ppcesa3KOs (Fig. 2). The intermediate phenotype of the ppcesa8KOs was confirmed and shown to be significantly different from both wild type and the double KOs (Fig. 2). Updegraff analysis showed that cellulose content of cell walls from whole ppcesa3/8KO gametophores (mean±S.E. of three genetic lines = 33.8±0.034%) was reduced significantly \( p = 0.004 \) compared to wild type (GD06, mean±S.E. of three independent cultures = 60.1±0.030%).

To confirm that the observed ppcesa3/8KO phenotype was due to the absence of PpCESA3 and PpCESA8, the selection cassette was removed from ppcesa3/8KO-86 by Cre-mediated recombination of flanking \( \text{lox-p} \) sites (Vidali et al. 2010) to allow transformation with vectors that drive expression of PpCESA3 or PpCESA8 with their native promoters (Fig. S2). Stable antibiotic resistant lines selected for the presence of numerous erect gametophores were examined with polarization microscopy (Fig. S2). For the transformation with \( \text{proCESA8::CESA8} \), 13 lines were examined, 6 of these
had strong midrib birefringence, and the first 3 were used for further analysis. For the transformation with \textit{proCESA3::CESA3}, the first three lines examined had strong midrib birefringence and were used for further analysis. S4B staining confirmed that expression of PpCESA8 or PpCESA3 rescued the defects in cellulose deposition in the leaf midribs of the double \textit{ppcesa3/8KO} (Fig. 2). Lines from the transformation with \textit{proCESA8::CESA8} were expected to be restored to the wild type phenotype because \textit{ppcesa3KO}, which also expresses \textit{PpCESA8} under control of the \textit{PpCESA8} promoter, showed no defects in cellulose deposition in the leaf midrib. All three \textit{proCESA8::CESA8} lines had significantly stronger S4B fluorescence than \textit{ppcesa8KO}. This demonstrates substantial restoration of the phenotype, although fluorescence was still significantly weaker than the wild type (Fig. 2). Two lines from a transformation with \textit{proCESA3::CESA3} (3R29 and 3R52) were not significantly different from \textit{ppcesa8KO}-5B, which is expected since they both lack \textit{PpCESA8} and express \textit{PpCESA3} under control of the \textit{PpCESA3} promoter. In the third line (3R45) fluorescence was restored to wild type levels (Fig. 2). Y-axis scales differ between experiments due to the use of different exposure time settings.

\textbf{Secondary cell wall microfibrils are helically oriented and laterally aggregated}

A first order retardation plate was used with polarized light microscopy to determine the optical sign, and thus the cellulose microfibril orientation, of wild type and \textit{ppcesa3/8KO} midrib cell walls (Fig. 3). In mature wild type leaves, the larger bundle sheath-like cells that surround the central stereids showed blue addition colors when oriented parallel to the major axis of the plate and yellow subtraction colors when oriented perpendicular to the major axis (Fig. 3), indicating that the net orientation of
positively birefringent cellulose microfibrils is longitudinal. In contrast, the walls of
the smaller central stereids were colorless when oriented parallel or perpendicular to
the major axis (Fig. 3). However, when oriented at 45° to the retardation plate, these
cells showed alternating bands of blue and yellow (Fig. 3), indicating that the
microfibrils in their walls are helical with an angle near 45°. The central midrib cells
of developing wild type leaves showed a transition from colorless to blue to yellow
along the apical to basal developmental gradient when the midrib was oriented parallel
to the major axis of the plate (Fig. 3). This indicates that the microfibril orientation
changes from transverse to longitudinal and then to helical as the cells mature. In
contrast, the central midrib stereids of mature ppcesa3/8KO leaves had blue addition
colors when oriented parallel to the major axis, yellow subtraction colors when
oriented perpendicular to the major axis, and no interference color when oriented at
45° to the retardation plate indicating that microfibrils are longitudinal, rather than
helical. Developing ppcesa3/8KO leaves had no longitudinal gradient in interference
colors (Fig. 3).

The walls of midrib cells were examined by transmission electron microscopy in
ultrathin sections of chemically fixed gametophore leaves. Despite the reduced
cellulose content detected by other means, the walls of midrib cells were thickened
compared to walls of adjacent lamina cells in all ppcesaKOs, as well as wild type
leaves (Fig. 4). When we attempted to prepare specimens by high pressure freezing
and freeze-substitution, the leaves fractured in a plane parallel to the midrib. This
resulted in a loss of midrib cells and precluded examination of midrib cell walls in
these specimens. We were able to examine the lamina and margin cells of freeze-
substituted leaves in wild type and two lines of each mutant. The walls of these cells appeared similar between wild type, and single and double ppcesaKOs (Fig. S3). However, measurements revealed that lamina cell external walls, i.e. those facing the external environment, were thinner in ppcesaKOs (Fig. S4).

The mesoscale organization of cellulose in the midribs of wild type, ppcesa3/8KO, and ppcesa8KO leaves was examined using a broadband SFG microscope (Lee et al. 2016). Because it detects only non-centrosymmetric ordering of functional groups, SFG provides a means of analyzing cellulose in intact cell walls with relatively little interference from matrix components (Barnette et al. 2011). For each genotype, full SFG spectra collected from three different locations along the midribs of each of three different leaves were averaged (Fig. 5). The sampling depth of the SFG microscope for cellulosic samples is 20-25 μm (Lee et al. 2016). Given that the thickness of turgid leaves is about 50-60 μm at the midrib and that they likely collapse to less than half their thickness when dried, we conclude that most of the leaf thickness contributes to the SFG signal. In spectra collected from the wild type, a strong peak at 2944 cm\(^{-1}\), which is characteristic of secondary cell walls, was observed in the CH/CH\(_2\) stretch region along with a 3320 cm\(^{-1}\) peak in the OH stretch region. In contrast, the spectra collected from ppcesa3/8KO and ppcesa8KO midribs had weaker peak intensity overall with a broad CH/CH\(_2\) stretch peak centered around 2910 cm\(^{-1}\). Compared to ppcesa3/8KO, the spectra from ppcesa8KO midribs had a weak signal at 2963 cm\(^{-1}\) that was absent in spectra collected from ppcesa3/8KO midribs. A scan across a wild type leaf shows that the 2944 cm\(^{-1}\) signal is associated with the midrib and was not observed in the cells of the lamina (Fig. 5). Equivalent scans of ppcesa3/8KO and
ppcesa8KO leaves confirm the absence of a strong 2944 cm⁻¹ peak from the midribs of these mutants (Fig. 5).

**PpCESA proteins are functionally specialized**

Based on the *ppcesa3KO, ppcesa8KO, and ppcesa3/8KO* phenotypes, PpCESA3 and PpCESA8 appear to be partially redundant. To determine whether the relative strengths of these phenotypes are related to gene expression levels, we used reverse transcription quantitative PCR to measure the expression of *PpCESA3* and *PpCESA8* in the wild type and mutants. In the *ppcesa3KOs, PpCESA8* was significantly upregulated compared to wild type (Fig. 6), providing a possible explanation for the lack of a mutant phenotype in these lines. In contrast, *PpCESA3* was not significantly upregulated in the *ppcesa8KOs* compared to wild type, potentially explaining the intermediate phenotype in these mutants.

*ppcesa3KOs, ppcesa8KOs and ppcesa3/8KOs* were tested for changes in rhizoid and caulonema development to determine whether developmental defects were restricted to the gametophores. When cultured on medium containing auxin, all lines produced the expected leafless gametophores with numerous rhizoids (Fig. S5), indicating no defects in rhizoid development in any of the KOs. Caulonema produced by colonies grown in the dark on vertically oriented plates were all negatively gravitropic (Fig. S6). Although appearance of the caulonema varied among experiments, those produced by KOs were always similar to control wild type within the same experiment. Caulonemal length was not significantly different between *ppcesa3/8KOs* and wild type (Table 1).
To determine whether other PpCESAs are functionally interchangeable with PpCESA3 and PpCESA8, we tested for rescue of ppcesa3/8KO-86lox by various PpCESAs driven by the PpCESA8 promoter. Polarization microscopy screening of at least 21 and up to 27 stably transformed lines for each vector revealed little or no midrib birefringence for the proCESA8::CESA4, proCESA8::CESA7 and proCESA8::CESA10 lines and moderate to strong midrib birefringence for 92% and 78% of the proCESA8::CESA3 and proCESA8::CESA5 lines, respectively. Quantitative analysis of S4B staining (Fig. 7) confirmed that the ppcesa3/8KO phenotype was partially rescued by proCESA8::CESA3 (3 out of 3 lines) and proCESA8::CESA5 (2 out of 3 lines) as we observed for proCESA8::CESA8 (Fig. 2). However, the proCESA8::CESA4, proCESA8::CESA7 and proCESA8::CESA10 vectors showed no rescue (Fig. 7). Western blot analysis confirmed that PpCESA proteins were expressed in all lines except proCESA8::CESA4-11 and proCESA8::CESA5-7 (Fig. S7). PpCESA6 differs from PpCESA7 by only 2 amino acids and was not tested. Although expressed with the same promoter, protein accumulation varies among the different transgenic lines (Fig. S7). Similar differences in protein accumulation may also explain variation in the extent of rescue by the proCESA3::CESA3 and proCESA8::CESA8 vector (Fig. 2).

Finally, we examined ppcesa4/10KOs and ppcesa6/7KOs produced for another study to determine whether they phenocopy the ppcesa3/8KO phenotype. Genotype verification for these lines is presented in Fig. S8 and Fig. S9. The ppcesa4/10KOs showed slight, but significant reduction in midrib S4B fluorescence. However, for ppcesa6/7KOs the reduction was substantial and significant (Fig. 7), showing the
PpCESA6/7 and PpCESA3/8 have non-redundant roles in secondary cell wall deposition in leaf midrib cells.

**Discussion**

**PpCESA3 and PpCESA8 function redundantly in cellulose deposition in steroid secondary cell walls.**

Targeted knockout of *PpCESA3* and *PpCESA8* blocked deposition of cellulose in the thick walls of steroid cells as indicated by 1) reduction of the strong birefringence associated with the midribs in *ppcesa3/8* KOs, 2) reduction in the midrib fluorescence of *ppcesa3/8* KO leaves stained with S4B, 3) lack of CBM3a labeling of sections from *ppcesa3/8* KO leaf midribs (Fig. 1), and 4) reduction in *ppcesa3/8* KO gametophore cell wall cellulose content as measured by Updegraff assay. Evidence that knockout of *PpCESA3* and *PpCESA8* is responsible for the observed phenotype includes consistency of the phenotype in three independent KOs and restoration of cellulose deposition in the midribs by transformation of *ppcesa3/8* KO with vectors driving expression of PpCESA3 or PpCESA8 (Fig. 2). Whereas we detected no reduction in midrib cellulose in *ppcesa3* KO, the phenotypes of *ppcesa8* KOs were intermediate between wild type and *ppcesa3/8* KO (Fig. 2). This, combined with the observations that only *PpCESA8* is up-regulated to compensate for loss of its paralog (Fig. 6) and expression of PpCESA3 under control of its native promoter only partially restores the wild type phenotype (Fig. 2), are consistent with the hypothesis that the PpCESA3 and PpCESA8 proteins are functionally interchangeable and that a dosage effect is responsible for the *ppcesa8* KO phenotype. The formation of morphologically normal
gametophores in ppcesa3/8KO s (Fig. 1) indicates that PpCESA3 and PpCESA8 serve a different role in development than PpCESA5, which supports normal cell division and cell expansion required for gametophore development (Goss et al. 2012). It is possible that PpCESA3 and PpCESA8 contribute to primary cell wall deposition since ppcesa3/8KO lamina cells had thinner external walls (Fig. S4) and tended to collapse during embedding (Fig. 1). Alternatively, PpCESA3 and PpCESA8 may contribute to secondary thickening of lamina cell walls after they stop expanding.

**CESA evolution in both P. patens and Arabidopsis involve sub-functionalization and neo-functionalization.**

There are many parallels in the evolution of the P. patens and Arabidopsis CESA families. In both species, different CESAs are responsible for primary and secondary cell wall deposition. In Arabidopsis, the secondary CESAs are AtCESA4, -7 and -8 (Taylor et al. 2003) and primary CESAs are AtCESA1, -3, and members of the 6-like group (Desprez et al. 2007; Persson et al. 2007). In P. patens, midrib secondary cell wall synthesis involves PpCESA3, -6, -7 and -8, whereas gametophore primary cell wall synthesis requires PpCESA5 (Goss et al. 2012). At least some primary CESAs can substitute for secondary CESAs and vice versa in both species. In Arabidopsis, AtCESA3pro::AtCESA7 partially rescues atcesa3, and AtCESA8pro::AtCESA1 partially rescues atcesa8 (Carroll et al. 2012). In P. patens, PpCESA8pro::PpCESA5 rescues ppcesa3/8KO. This indicates that the CESA division of labor for primary and secondary cell wall deposition in vascular plants and mosses is due at least in part to sub-functionalization. However, neo-functionalization has also occurred in both species, resulting in the requirement for two or more non-interchangeable CESA
isoforms for secondary cell wall biosynthesis. In Arabidopsis, \textit{atcesa4}, \textit{atcesa7}, and \textit{atcesa8} null mutants share a phenotype (Taylor et al. 2000) that cannot be complemented by expressing one of the other secondary AtCESAs with the promoter for the missing isoform (Kumar et al. 2016). Likewise in \textit{P. patens}, \textit{ppcesa3/8KO} and \textit{ppcesa6/7KO} share the same phenotype and \textit{ppcesa3/8KO} is not complemented by \textit{PpCESA8pro::PpCESA7}. Studies are ongoing to determine whether the secondary PpCESAs physically interact to form a CSC, as has been shown for the secondary AtCESAs (Taylor et al. 2003; Timmers et al. 2009). Finally, the CESA families of both species show some redundancy. In Arabidopsis the 6-like CESAs (\textit{AtCESA2}, -5, -6 and -9) are partially redundant (Persson et al. 2007), as are \textit{PpCESA3} and -8 in \textit{P. patens}. PpCESA6 and -7 differ by only three amino acids and the genes that encode them appear to be redundant (Wise et al. 2011).

A recent study has shown that secondary cell wall deposition, including CESA expression, is regulated by NAC transcription factors in both \textit{P. patens} and Arabidopsis (Xu et al. 2014). Three \textit{P. patens} NAC genes, \textit{PpVNS1}, \textit{PpVNS6}, and \textit{PpVNS7}, were preferentially expressed in leaf midribs and \textit{ppvns1/ppvns6/ppvns7KO}s were defective in stereid development. Overexpression of \textit{PpVNS7} activated PpCESA3 (Xu et al. 2014). Phylogenetic analyses of NACs place eight \textit{PpVNS} proteins within the clade that has variously been named subfamily NAC-c (Shen et al. 2009), subfamily Ic (Zhu et al. 2012), or the VNS group (Xu et al. 2014), and also includes the Arabidopsis vascular-related NACs VND6 (ANAC101), VND7 (ANA030), NST1 (ANAC043), NST2 (ANAC066) and NST3/SND1 (ANAC012). The three \textit{PpVNS} proteins that regulate stereid development form a single sister clade.
with five other PpVNS proteins implicated also in other processes (Xu et al. 2014). Based on this phylogenetic analysis, the common ancestor of the mosses and seed plants had a single VNS gene, and it also had a single CESA gene (Kumar et al. 2016; Roberts and Bushoven 2007; Yin et al. 2009). Both lineages now include secondary CESAs that are regulated by VNSs and primary CESAs that are not, indicating that CESA subfunctionalization occurred independently in mosses and seed plants.

**Secondary cell wall microfibrillar texture is similar in mosses and vascular plants.**

In vascular plants, both water conducting tracheary elements and supportive fibers are characterized by helical (Barnett and Bonham 2004) and aggregated (Donaldson 2007; Fernandes et al. 2011; Thomas et al. 2014) cellulose microfibrils. The midribs of *P. patens* leaves include hydroid cells that transport water and stereid cells that provide support, but only the stereids have thick cell walls (Xu et al. 2014). With highly reduced cellulose in their stereid secondary cell walls, *ppcesa3/8KOs* provided a negative control for structural characterization of secondary cell walls in wild type *P. patens*. A sharp SFG CH/CH$_2$ stretch peak at 2944 cm$^{-1}$ is characteristic of angiosperm secondary cell walls (Park et al. 2013) and extensive empirical testing has shown that this spectral feature is attributable to lateral microfibril aggregation (Lee et al. 2014). The 2944 cm$^{-1}$ peak was also present in SFG spectra of wild type *P. patens* midribs. In contrast, the spectra of *ppcesa3/8KO* leaf midribs lacked the 2944 cm$^{-1}$ peak and instead had a broad peak between 2800 and 3000 cm$^{-1}$, which is characteristic of primary cell walls and other samples lacking aggregated microfibrils (Lee et al. 2014;
This suggests that lateral aggregation of microfibrils is a common feature of the secondary cell walls of moss stereids and vascular plant tracheary elements and fibers. Polarization microscopy with a first order retardation plate revealed that the microfibrils in the stereid cell walls are deposited in a helical pattern, as observed in secondary cell walls of tracheary elements and fibers (Barnett and Bonham 2004). Although deficient in cellulose, the stereid cell walls of ppcesa3/8KOs were thickened, indicating that secondary cell wall synthesis involves deposition of non-cellulosic components, which proceeded in the absence of cellulose deposition. This has also been observed in developing tracheary elements treated with cellulose synthesis inhibitors (Taylor et al. 1992). Thus, stereid cell walls share structural characteristics with the cell walls of tracheary elements and fibers.

**Mosses and vascular plants have acquired similar secondary cell walls through convergent evolution.**

Thick, cellulose-rich secondary cell walls provide added support for aerial organs of mosses and vascular plants alike. Within these cell walls, the lateral aggregation and helical orientation of the microfibrils contributes to their strength and resiliency. Although cortical microtubules play an important role in cellulose microfibril orientation, oriented cellulose deposition can occur in the absence of cortical microtubules, and it has previously been suggested that aggregation and helical orientation of microfibrils in secondary walls is a consequence of high CSC density during rapid cellulose deposition (Emons and Mulder 2000; Lindeboom et al. 2008). Regulation at the level of CSC secretion was emphasized in this model (Emons and
Mulder 2000), but CSC density can potentially be regulated at the level of transcription.

Rapid cellulose synthesis during secondary cell wall deposition in specific cell types requires precise temporal and spatial regulation of CESA expression that is distinct from the regulatory requirements for primary cell wall synthesis. We suggest that these distinct regulatory needs were met through the evolution of independent regulatory control of primary and secondary CESAs by sub-functionalization in both mosses and seed plants. In seed plants, phylogenetic analysis shows that the first divergence of the CESA family separated the genes that encode the primary and secondary CESAs and was followed by independent diversification within each group (Roberts et al. 2012). This, along with evidence that some primary CESAs are interchangeable with secondary CESAs (Carroll et al. 2012), indicates that sub-functionalization was an early event in the evolution of the seed plant CESA family. In *P. patens*, the genes that encode secondary PpCESA3 and PpCESA8 and primary PpCESA5 are also sub-functionalized and therefore specialized, although they encode interchangeable proteins.

Several lines of evidence indicate that the capacity to deposit a secondary cell wall evolved independently in mosses and seed plants. Structural and paleobotanical evidence suggests that the support and water-conducting cells of bryophytes and vascular plants are not homologous (Carafa et al. 2005; Ligrone et al. 2002). Phylogenetic evidence indicates that the primary and secondary CESAs diversified independently in mosses and seed plants (Kumar et al. 2016; Roberts and Bushoven 2007; Yin et al. 2009) and, as explained above, so did the NAC transcription factors.
that regulate the secondary CESAs. There are even examples of convergent evolution of secondary cell walls within the angiosperm lineage. Cotton fiber secondary cell walls are synthesized by the same CESAs that are responsible for secondary cell wall deposition in tracheary elements and fibers (Haigler et al. 2012), whereas the secondary cell walls of epidermal trichomes are synthesized by the primary CESAs (Betancur et al. 2011). These observations are consistent with independent evolutionary origins for secondary cell walls in different land plant lineages and different cell types within angiosperm lineages.

Taken together, these data indicate that CESA duplication, followed by adoption of regulatory elements within the secondary CESA promoters that enable control by NAC transcription factors, occurred independently in mosses and vascular plants. The resulting uncoupling of the secondary CESAs from the regulatory constraints associated with primary cell wall deposition, along with a mechanistic linkage between CESA expression and microfibril texture as well as selection for strength and resiliency, may have contributed to the capacity of different plants to synthesize cellulose-rich secondary cell walls with similar microfibrillar textures.

Materials and methods

Vector construction

All primer pairs are shown in Table S1, along with annealing temperatures used for PCR. Amplification programs for Taq Polymerase (New England Biolabs, Ipswich, MA, USA) consisted of a 3 min denaturation at 94°C; 35 cycles of 15 s at 94°C, 30 s at the annealing temperature, and 1 min/kbp at 72°C. Amplification programs for
Phusion Polymerase (New England Biolabs) consisted of a 30 s denaturation at 98°C; 35 cycles of 7 s at 98°C, 7 s at the annealing temperature, and 30 s/kbp at 72°C.

To construct the CESA8KO vector, a 3’ homologous region was amplified from *P. patens* genomic DNA with primers 174JB and 193JB using Taq DNA polymerase, cut with SalI and BspD1, and cloned into the SalI/BstBI site of pBHSNR (gift of Didier Schaefer, University of Neuchâtel). The resulting plasmid was cut with KasI and NsiI to accept the KasI/NsiI fragment of a 5’ homologous region amplified from *P. patens* genomic DNA with primers 203JB and 185JB (Table S1). The CESA8KO vector was cut with EcoRI and NsiI for transformation into wild type *P. patens*. The CESA3KO, CESA4KO, CESA6/7KO, and CESA10KO vectors were constructed using Gateway Multisite Pro cloning (Invitrogen, Grand Island, NY, USA) as described previously (Roberts et al. 2011). Flanking sequences 5’ and 3’ of the coding regions were amplified with appropriate primer pairs (Table S1) using Phusion DNA polymerase (New England Biolabs) and cloned into pDONR 221 P1-P4 and pDONR 221 P3-P2, respectively, using BP Clonase II (Invitrogen). Similarly, an *nph* selection cassette was amplified from pMBL6 (gift of Jesse Machuka, University of Leeds) cloned into pDONR 221 P3r-P4r. All entry clones were sequence-verified. For vectors conferring hygromycin resistance, entry clones with flanking sequences in pDONR 221 P1-P4 and pDONR 221 P3-P2 were inserted into BHSNRG (Roberts et al. 2011). For vectors conferring G418 resistance, entry clones with flanking sequences in pDONR 221 P1-P4 and pDONR 221 P3-P2 were linked with the entry clone containing the *nph* selection cassette and inserted into pGEM-gate (Vidali et al. 2009) using LR Clonase.
II Plus (Invitrogen). The vectors in BHSNRG or pGEM-gate were cut with BsrGI for transformation into wild type or mutant *P. patens* lines.

Expression vectors for HA-tagged PpCESAs under control of *PpCESA* promoters were constructed using Gateway Multisite Pro cloning (Invitrogen). The *PpCESA4* (DQ902545), *PpCESA5* (DQ902546), *PpCESA7* (DQ160224) and *PpCESA8* (DQ902549) coding sequences were amplified from cDNA clones pdp21409, pdp24095, pdp38142 and pdp39044 (RIKEN BioResource Center, Tsukuba, Ibaraki JP), respectively, using forward primers containing a single hemagglutinin (HA) tag and appropriate reverse primers (Table S1) and cloned into pDONR 221 P5-P2 using BP Clonase II (Invitrogen). The *PpCESA3* (XP_001753310) and *PpCESA10* (XP_001776974) coding sequences were similarly amplified from expression vectors. pDONR 221 P1-P5r entry clones containing approximately 2 kB of sequence upstream of the *PpCESA3* or *PpCESA8* start codon (Tran and Roberts 2016), were linked to the sequence verified entry clones containing the HA-*PpCESA* coding sequences and inserted into pSi3(TH)GW (Tran and Roberts 2016) using LR Clonase II Plus (Invitrogen). These vectors target the expression cassettes to the intergenic 108 locus, which can be disrupted with no effect on phenotype (Schaefer and Zryd 1997). Rescue vectors were cut with SwaI for transformation into a *P. patens ppcesa3/8KO* line from which the *hph* resistance cassette had been removed (see below).

**Culture and transformation of *P. patens***

Wild type *P. patens* lines (haploid) derived from the sequenced Gransden strain (Rensing et al. 2008) by selfing and propagation from a single spore in 2006 (GD06) or 2011 (GD11) were gifts of Pierre-Francois Perroud, Washington University. Wild
type and transformed *P. patens* lines were cultured on basal medium supplemented with ammonium tartrate (BCDAT) as described previously (Roberts et al. 2011). Protoplasts were prepared and transformed as described previously (Roberts et al. 2011). Stable transformants were selected with 50 μg mL⁻¹ G418 (CESA3KO vector) or 15 μg mL⁻¹ hygromycin (CESA8KO and complementation vectors). The *hph* selection cassette was removed from *ppces3/ppcesa8KO* by transforming protoplasts with NLS-Cre-Zeo (Vidali et al. 2010) selecting for 7 d on BCDAT plates containing 50 μg mL⁻¹ zeocin, replica plating zeocin resistant colonies on BCDAT with and without 15 μg mL⁻¹ hygromycin, and recovering hygromycin-sensitive colonies.

Protein expression was tested by western blot analysis as described previously (Scavuzzo-Duggan et al. 2015) in selected lines transformed with HA-PpCESA expression vectors.

**Genotype analysis**

For PCR screening, DNA was extracted as described previously (Roberts et al. 2011) and 2.5 μL samples were subjected to 35 cycles of amplification (45 s at 94°C, 45 s at the annealing temperature shown in Table S1, 1 min/kbp at 72°C) with PAQ5000 DNA polymerase (Agilent Technologies, http://www.home.agilent.com/) in 25 μL reactions. Primers used to test for target integration, target-gene disruption, and selection cassette excision are listed in Table S1.

**Phenotype analysis**

Cell wall birefringence of unfixed leaves mounted in water was examined using an Olympus BHS compound microscope with D Plan-Apo UV 10X/0.4, 20X/0.7, and 40X/0.85 objectives, and polarizer and circular-polarizing analyzer, with and without a
first order retardation plate (Olympus, Center Valley, PA, USA). Images were captured with a Leica DFC310FX digital camera with Leica Application Suite software, version 4.2.0 (Leica Microsystems Inc., Buffalo Grove, IL, USA) with manual exposure under identical conditions.

For direct fluorescent labeling of cellulose, whole gametophores (3 per line) dissected from colonies grown for four weeks on solid BCDAT medium were dipped in 100% acetone for 5 sec to permeabilize the cuticle, rinsed in phosphate buffered saline (PBS), incubated in PBS containing 0.1 mg/ml S4B (Anderson et al. 2010) for 30 min, and rinsed in PBS. All fully expanded leaves (12-20) were cut from each gametophore and mounted in PBS. Fluorescence images of each leaf, centered on the brightest part of the midrib, were captured using a Zeiss Axio Imager M2 with 43HE DsRed filter set, Plan-Neofluar 20X/0.5 objective, AxioCam MR R3 camera, and Zen Blue software, version 1.1.2.0 (Carl Zeiss Microscopy, Jena, Germany) under identical conditions using manual exposure. The midrib in each image was selected manually (Fig. S10) and average pixel intensity was measured using ImageJ, Fiji version (Schindelin et al. 2012). For comparison of KOs to the wild type, three independent lines of each KO genotype (n=3) and two independent wild type lines (GD06 and GD11, n=2) were sampled in triplicate. For analysis of rescue lines, three independent explants were sampled for each genetic line (n=3).

For affinity cytochemistry of cellulose, gametophores dissected from colonies grown for two weeks on BCDAT medium were fixed and embedded in LR White resin (Polysciences, Inc., Warrington, PA, USA) as described previously (Kulkarni et al. 2012). Sections (1 μm) were mounted and labeled with CBM3a as described
previously (Berry et al. 2016). Images were captured with a Zeiss Axio Imager M2 with 38 Green Fluorescent Protein filter set, EC Plan-Neofluar 40X/0.75 objective, AxioCam MR R3 camera, and Zen Blue software, version 1.1.2.0 (Carl Zeiss Microscopy) under identical conditions using manual exposure. Fluorescence and polarization images were not altered after capture. Bright field and differential interference contrast images were captured using automatic exposure and some images used for illustrative purposes were adjusted for uniformity using the color balance and exposure functions in Photoshop, version CS6 (Adobe Systems, San Jose CA, USA). 

ppcesa3KOs, ppcesa8KOs, and ppcesa3/8KOs were tested for changes in caulonema gravitropism and rhizoid development as described previously (Roberts et al. 2011). Images were captured using a Leica M165FC stereomicroscope with Leica DFC310FX camera and Leica Application Suite software, version 4.2.0 (Leica Microsystems Inc.). Caulonema length for each colony was measured as the distance from the edge of the colony to tip of the longest caulonema filament using Leica Application Suite software.

Cell wall analysis

Alcohol insoluble residue (AIR) was prepared from gametophores dissected from 8-10 4-week-old explants of P. patens wild type (three samples from independent cultures) and ppcesa3/8KO (samples from three independent lines) cultured on BCDAT medium. Tissue was ground in liquid nitrogen and extracted three times, 30 min each, with 70% (v/v) ethanol and once with 100% ethanol and the residue was dried under vacuum. The AIR (~1 mg) was weighed to 0.001 mg and mixed with 1 mL of acetic acid:water:nitric acid (8:2:1, v/v) in screw-cap vials and the suspension was heated in
a boiling water bath for 30 min (Updegraff 1969). After cooling, the tubes were centrifuged at 16,900 x g for 5 min and the supernatant discarded. The pellet was resuspended in 2 mL of deionized water, centrifuged, and the supernatant was discarded. The washing step was repeated at least 10 more times until the supernatant was neutralized and the pellet was resuspended in 1 mL of water. The amount of cellulose remaining after hydrolysis was quantified by sulfuric acid assay (Albalasmeh et al. 2013) with glucose as the standard. Briefly, 100 µL of hydrolysate (six technical replicates per sample) was diluted to 1 mL with water in a glass tube, 3 mL of concentrated sulfuric acid was added, and samples were vortexed for 30 s and chilled on ice for 2 min. Reactions were measured at 315 nm against a reagent blank.

**High pressure freezing-freeze substitution and transmission electron microscopy**

Gametophytes of *P. patens* GD06 and PpCESAKOs were high pressure-frozen using a Leica EMPACT2 high pressure freezer (Leica Microsystems, Inc.) followed by freeze-substitution in 0.1% uranyl acetate in acetone for 48 h at -90°C before the temperature was ramped up slowly to -50°C (Wilson and Bacic 2012). The samples were rinsed with acetone twice at -50°C before the acetone was replaced with ethanol and the samples were subsequently infiltrated with LR White resin (ProSciTech Pty. Ltd., Thuringowa Central QLD Australia) in a series of ethanol/resin dilutions. The samples were rinsed three times in 100% resin before polymerization with UV light at -20°C for 48 h. Thin sections (70 nm) were cut using a Leica Ultracut R (Leica Microsystems, Inc.) and post-stained with uranyl acetate and lead citrate (Wilson and Bacic 2012). Images were taken using a Tecnai G2 Spirit transmission electron
microscope (FEI, Hillsboro, OR USA). Cell wall thickness was measured using
ImageJ, Fiji version (Schindelin et al. 2012).

Ultrathin sections (70 nm) were also cut from blocks prepared for affinity
cytochemistry (see above), mounted on Formvar coated copper grids, and stained with
uranyl acetate and lead citrate (Wilson and Bacic 2012). Sections were imaged using a
FEI/Phillips CM-200 transmission electron microscope (FEI).

**Sum Frequency Generation spectroscopy**

Leaves of wild type GD06, 8KO-5B, and 3/8KO-86 lines were mounted abaxial side
down in water on glass slides and allowed to air-dry overnight. SFG spectra were
collected 5 µm intervals along a 200 µm line scan perpendicular to the midrib at its
thickest point using an SFG microscope system described previously (Lee et al. 2016).
The SFG spectra were collected with the following polarization combination: SFG
signal = s-, 800 nm = s-, and broadband mid-IR = p-polarized with the laser incidence
plane and the laser incidence plane aligned along the axis of midrib.

**Reverse transcription quantitative PCR**

RNA was extracted from gametophores from two independent wild type and three
independent lines each of *ppcesa3KO* and *ppcesa8KO* as described previously (Tran
and Roberts 2016). cDNA samples were tested in duplicate as described previously
using primer pairs for amplification of *PpCESA3* and *PpCESA8*. The primers have
been previously tested for specificity and efficiency (Tran and Roberts 2016). Primers
for actin and v-Type H⁺translocating pyrophosphatase reference genes were described
previously (Le Bail et al. 2013). Target/average reference cross point ratios were
calculated for each sample and standard errors were calculated for independent genetic lines.

**Statistical analysis**

For statistical analysis, one-way Analysis of Variance (ANOVA) with post-hoc Tukey Honest Significant Difference (HSD) test was performed at astatsa.com/OneWay_Anova_with_TukeyHSD/.

**Supplemental Materials**

Table S1. Primers used for vector construction and genotype analysis.

Fig. S1. Genotype analysis of ppcesa8, ppcesa3 and ppcesa3/8 KO lines.

Fig. S2: Phenotype analysis of a ppcesa3/8 double KO line transformed with vectors driving expression of PpCESA3 or PpCESA8 with their native promoters.

Fig. S3. Transmission electron microscopy images of leaf cell walls from wild type and cesaKO lines of *P. patens*.

Fig. S4. Thickness of outer cell walls measured from transmission electron microscopy images.

Fig. S5: *P. patens* wild type and KO lines cultured on medium containing 1 μM naphthalene acetic acid (auxin) to induce rhizoid initiation and inhibit leaf initiation.

Fig. S6: *P. patens* wild type and KO lines cultured in the dark on vertically oriented plates containing medium supplemented with 35 mM sucrose to test for caulonema gravitropism.
Fig. S7. Western blot analysis of protein expression for *P. patens* lines derived from transformation of *ppcesa3/8KO-86lox* with vectors driving expression of PpCESAs under control of the *PpCESA8* promoter.

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Table 1. Caulonema length for wild type and *ppcesaA3/8*KO s grown on vertical plates in the dark. Data are from two independent experiments (n=2). ANOVA analysis showed no significant differences between genetic lines.

| Genetic line     | Caulonema length (mm) | Standard Error |
|------------------|------------------------|----------------|
| WT GD06          | 4.69                   | 0.50           |
| *ppcesaA3/8KO-43*| 5.70                   | 0.87           |
| *ppcesaA3/8KO-57*| 4.51                   | 1.14           |
| *ppcesaA3/8KO-86*| 5.69                   | 0.47           |
Figure 1: Phenotypes of *ppcesa3/8KO*, *ppcesa3KO* and *ppcesa8KO* compared to wild type *Physcomitrella patens*. (A-D) Colony morphology is similar in wild type, *ppcesa3KOs* and *ppcesa8KOs*: horizontal growth is typical of gametophores produced by *ppcesa3/8KO* (arrowheads). (E-H) Polarized light microscopy of leaves shows that the midribs of wild type and *ppcesa3KO* are highly birefringent. The midribs of *ppcesa3/8KO* leaves have low birefringence and *ppcesa8KO* leaves have moderate birefringence. (I-L) Fluorescence microscopy of leaves stained with S4B shows strong fluorescence in the midribs of wild type and *ppcesa3KO*, low fluorescence in the midribs of *ppcesa3/8KO* leaves and intermediate fluorescence in the midribs of *ppcesa8KO* leaves. (M-P) Differential interference contrast microscopy of sections through the midribs of maturing leaves (L=lamina cell, *=bundle sheath cell). In wild type and *ppcesa3KO*, the walls of bundle sheath cells and the stereid cells they surround show enhanced contrast due to higher refractive index. (Q-T) Fluorescence microscopy of the same sections shown in M-P labeled with CBM3a. The bundle sheath and stereid cells of wild type and *ppcesa3KO* leaves are strongly labeled, whereas labeling is weak in *ppcesa3/8KO* and intermediate in *ppcesa8KO* leaves.
Figure 2: Quantitative analysis of S4B fluorescence intensity in leaf midribs of *P. patens* wild type, *ppcesa*KO, and rescue lines. (A) Fluorescence was significantly weaker in *ppcesa*3/8KOs compared to wild type (WT). *ppcesa*3KOs were not significantly different from wild type, whereas *ppcesa*8KOs were intermediate between the wild type and *ppcesa*3/8KOs and significantly different from both. For each mutant genotype, three independent genetic lines were sampled in triplicate. Two independent wild type lines (GD06 and GD11) were sampled in triplicate. Bars indicate the standard error of the mean for three mutant (n=3) or two wild type (n=2) lines. Genotypes with different letters are significantly different. (B) Lines derived from transformation of *ppcesa*3/8KO-86lox with *proCESA8::CESA8* (8R) had significantly higher fluorescence compared to the parent double KO line and *ppcesa*8KO, but significantly less than WT. (C) Lines derived from transformation of *ppcesa*3/8-86lox with *proCESA3::CESA3* (3R) had significantly higher fluorescence compared to the parent double KO line (except 3R29) and were not significantly different from either *ppcesa*8KO lines (3R29 and 3R52) or WT (3R45). For B and C, three independent explants were sampled for each genetic line. Bars indicate the standard error of the mean for three explants from the same line (n=3 or n=2 (WT, 3/8KO, 8KO in C)).
Figure 3: Polarized light microscopy with first order retardation plate. Double pointed arrow indicates the vibration direction of the major axis. (A-C) Midrib of a mature wild type leaf oriented parallel, perpendicular, and at 45° to the major axis of the retardation plate. Bundle sheath cells (*) flank the central midrib. (D) Midrib of a developing wild type leaf oriented parallel to the major axis of the retardation plate showing change in microfibril orientations through the basal (b), medial (m), and apical (a) regions of the midrib. (E-G) Midrib of a mature ppcesa3/8KO leaf oriented parallel, perpendicular, and at 45° to the major axis of the retardation plate. (H) Midrib of a developing ppcesa3/8KO leaf oriented parallel to the major axis of the retardation plate showing no change in microfibril orientation through the basal, medial, and apical regions of the leaf. Bar in A is also for B-C and E-G and bar in D is also for H.
Figure 4: Transmission electron microscopy images of leaf midribs of *P. patens* showing adjacent cells with primary cell walls (PW) and secondary cell walls (SW) in (A) wild type, and (B-D) mutant leaves.
Figure 5: Sum Frequency Generation (SFG) spectroscopy of *P. patens* leaves. (A) Full SFG spectra collected from leaf midribs (each is the average of nine spectra, from three different positions on each of three different leaves). A strong peak in the C-H stretch region (2944 cm$^{-1}$) is present in spectra from wild type (WT), greatly diminished in spectra from *ppcesa*8KO (8KO), and absent in spectra from *ppcesa*3/8KO (3/8KO). (B) *P. patens* wild type, *ppcesa*8KO, and *ppcesa*3/8KO leaves with SFG scan trajectories traversing the midribs. Step size was 5 μm/step. SFG spectra were collected from 2850 to 3150 cm$^{-1}$, covering the entire CH region. (C) 2D projection image of SFG spectra collected across the midribs of each leaf shown in B. Each column in each image is an entire spectrum collected from one point plotted against displacement along the scan trajectory. Colors indicate SFG intensity as shown in the legend.
Figure 6: RT-qPCR analysis of *PpCESA3* and *PpCESA8* expression in wild type, *ppcesa3* KOs and *ppcesa8* KOs. Target/average reference cross point ratios (using actin and ν-Type H’ translocating pyrophosphatase reference genes) were determined for three independent lines of each mutant (3KO-5, -35, -126; 8KO-5B, -4C, -10C; and 3/8KO-43, -57, -86) and two independent wild type lines (GD06 and GD11) with two technical replicates each. Bars indicate the standard error of the mean for the three mutant (*n*=3) or two wild type (*n*=2) lines.
Figure 7: Quantitative analysis of S4B fluorescence intensity in leaf midribs. (A,B) Wild type (WT), ppcesa3/8KO-86lox, and ppcesa3/8KO-86lox transformed with proCESA8::CESA expression vectors. For each rescue genotype, three independent genetic lines were sampled in triplicate and measured with 6 samples of wild type (GD06) and 8 samples of ppcesa3/8KO-86lox. (A) For lines derived from transformation of ppcesa3/8KO-86lox with proCESA8::CESA3 (8pro:3R), proCESA8::CESA7 (pro8:7R), and proCESA8::CESA10 (pro8:10R) genotypes, the three independent lines did not differ significantly and were combined. proCESA8::CESA7 and proCESA8::CESA10 lines did not differ significantly from the parent double KO line (p > 0.05), whereas proCESA8::CESA3 lines had significantly higher fluorescence compared to the parent double KO line, but significantly less than WT (p < 0.05). Bars indicate the standard error of the mean for three independent lines. Genotypes with different letters are significantly different. (B) For lines derived from transformation of ppcesa3/8KO-86lox with proCESA8::CESA5 (pro8:5R) and proCESA8::CESA4 (pro8:4R), the three independent lines were significantly different and were analyzed separately. proCESA8::CESA5 (5R) lines were not significantly different from the wild type (p > 0.05), except for 5R7, which was not significantly different from ppcesa3/8KO-86lox (p > 0.05). proCESA8::CESA5 lines did not differ significantly from ppcesa3/8KO-86lox (p > 0.05). Bars indicate the standard error of the mean for three gametophores from the same line (n=3). Lines with different letters are significantly different (p < 0.05). (C) Mid rib fluorescence was slightly, but significantly reduced in cesa4/10KO compared to wild type (p = 0.037). Reduction in midrib fluorescence in cesa6/7KO was substantial and highly significant (p = 0.0011). Bars indicate the standard error of the mean for three independent mutant lines or 3 replicates of wild type (n=3).
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### Supplemental Materials

Table S1. Primers used for vector construction and genotype analysis.

| Primer pair        | Sequences                                                                 | Annealing temp. | Amplicon size | Amplified region                  |
|--------------------|---------------------------------------------------------------------------|-----------------|---------------|-----------------------------------|
| 174JB 193JB        | **TACGGCAGGATGTATGAGCA**<br>**TACTTCCACGGCTTCTTGT**                          | 57°C            | 2003          | **5'** targeting region **PpCESA8** |
| 203JB 185JB        | **ATCAACAACAGCAAGGCCAT**<br>**AGCACCTGGTCAACCCATC**                          | 57°C            | 1041          | **3’** targeting region **PpCESA3** |
| 3KOattB1           | **GGGGACAACATTGGCTATATTAAGTGGGA**<br>**CGGAAGCTTGGATTCTCTAC**                 | 66°C            | 894           | **5'** targeting region **PpCESA3** |
| 3KOattB2           | **GGGGACCACTTGGCTATAGAAAAGCTGGG**<br>**TGCAAGCTTATGACATTTTTGCTG**            | 66°C            | 898           | **3'** targeting region **PpCESA3** |
| 4KOattB1           | **GGGGACAACATTGGCTATATTAAGTGGGA**<br>**CGGAAGCTTGGATTCTCTAC**                 | 68°C            | 1108          | **5'** targeting region **PpCESA4** |
| 4KOattB2           | **GGGGACCACTTGGCTATAGAAAAGCTGGG**<br>**TGCAAGCTTATGACATTTTTGCTG**            | 68°C            | 1148          | **3'** targeting region **PpCESA4** |
| 10KOattB1          | **GGGGACAACATTGGCTATATTAAGTGGGA**<br>**CGGAAGCTTGGATTCTCTAC**                 | 68°C            | 1121          | **5'** targeting region **PpCESA10** |
| 10KOattB4          | **GGGGACCAACTTGGCTATAGAAAAGCTGGG**<br>**TGCAAGCTTATGACATTTTTTCTG**           | 68°C            | 637           | **3'** targeting region **PpCESA10** |
| 8KOFlankF VectorR-hph | **CTGGACAGACTTTTCTCCTGGTTAT**<br>**TCCGAGGCAAAGAATAAGA**                     | 57°C            | 1121          | **5'** targeting region **PpCESA8** |
| 8KOFlankR          | **TCGAGTACGATTGGCTGCAATG**<br>**TCCGAGGCAAAGAATAAGA**                         | 57°C            | 637           | **3'** integration **PpCESA8**    |
| pMBL6attB4r        | **GGGGACAACTTGGCTATATTAAGTGGGA**<br>**CGGAAGCTTGGATTCTCTAC**                 | 68°C            | 2014          | Selection cassette from pMBL6     |
| pMBL6attB3r        | **GGGGACAACTTGGCTATATTAAGTGGGA**<br>**CGGAAGCTTGGATTCTCTAC**                 | 68°C            | 2014          | Selection cassette from pMBL6     |
| 8KOFlankF          | **See above**                                                             | 57°C            | 731           | **PpCESA8 KO** cassette excision  |
| 3KOFlankF VectorR2-npt | GTTTCGTTTTGGTTTCGCTGT TGCTTTGGGTAAGGAAGGGAG TGGAAGGATGTTGTAAGCAG | 57°C | 1362 | 5’ integration PpCESA3 |
|------------------------|-------------------------------------------------|------|------|----------------------|
| VectorF2-npt 3KOFlankR | AAGTGGCCGAGGAGGGAGGAG TTGAAGGCCCATGTTGTCAG | 57°C | 1259 | 3’ integration PpCESA3 |
| 4KOFlankF VectorR-hph | TGTCAGTGTCTAGCCATCCA TCTATTCTTTCGCCCTCAGGA | 59°C | 1520 | 5’ integration PpCESA4 hph cassette |
| VectorF-hph 4KOFlank-R2 | TGACACAGAGCTGGGCAATG GCAATGGTGGTGGTATC | 58°C | 1832 | 3’ integration PpCESA4 hph cassette |
| 4KOFlankF VectorR-npt | See above CCCGAAATTACCCCTTGTG | 57°C | 1263 | 5’ integration PpCESA4 npt cassette |
| VectorF-npt 4KOFlank-R2 | GCCCTGTGCAAGGTAAGAAG See above | 57°C | 1839 | 3’ integration PpCESA4 w/npt cassette |
| 6KOF2 VectorR-hph | GCTTCAATGCTGTACCACAAAGAACT TCCGAGGGGAAGAAGATAGA | 57°C | 1647 | 5’ integration PpCESA6 |
| VectorF-hph CESA7FlankR | TGACAGATAGCTGGGCAATG AAACCTTACTCTCCAGCACC | 57°C | 833 | 3’ integration PpCESA7 |
| 10KOFlankF VectorR-hph | TTCGCCCTGTGTAAGCTCGT See above | 57°C | 1461 | 5’ integration PpCESA10 hph cassette |
| VectorF-hph 10KOflankR2 | See above CATCCATCTATTTCTGATGC | 57°C | 1136 | 3’ integration PpCESA10 hph cassette |
| 10KOFlankF VectorR-npt | See above See above | 57°C | 1274 | 5’ integration PpCESA10 npt cassette |
| VectorF-npt 10KOflankR2 | See above See above | 57°C | 1132 | 3’ integration PpCESA10 npt cassette |
| CESA8TargetF CESA8TargetR | GTCTTTCTGCATGTCACAGCAGC TACTCTCAACGGCTCTTGTCT | 57°C | 339 | PpCESA8 deletion test |
| CESA3TargetF5 CESA3TargetR5 | CGTGGTCTCAAATCTGAGTG CTTTAATTCTGGCCAGCCTG | 64°C | 1266 | PpCESA3 deletion test |
| CESA6TargetF CESA6TargetR | GTTGGAGTAGAAGAAGAGAAG TTTTAAATCTGGCCAGCCTG | 60°C | 142 | PpCESA6 deletion test |
| CESA7TargetF CESA7TargetR | CTGGTACGAGAGGAGAAGAAG TTCTTAAATCTGGCCAGCCTG | 60°C | 1254 | PpCESA7 deletion test |
| CESA4TargetF CESA4TargetR2 | AGGTGAGTGCTGTGGGGAA GCCTGAGCTGACATCTCAG | 58°C | 1731 | PpCESA4 t deletion test |
| CESA10TargetF CESA10TargetR | TGGGATTTGACATCTGAGAG CACCCGCGGAAAACTATAGA | 57°C | 973 | PpCESA10 deletion test |
| 4KOattB1 4KOattB2 | See above See above | 68°C | 2321 | PpCESA4KO cassette excision |
| HACESA3attB5 | GGGGACAACTTTGTATACAAAGAGTTTGCG ATGGAGTACCCATAGGATTTTCCGATT ACGCTAGGGCAGAATGCTGGCCGGCTTGGT GGGGACACCTTTGTACAAAGAGTTTGCG TATACAAAGAGGAGGCGGACGCG | 68°C | 3370 | PpCESA8 coding sequence |
| CESA8attB2 | GGGGACAACTTTGTATACAAAGAGTTTGCG ATGGAGTACCCATAGGATTTTCCGATT ACGCTAGGGCAGAATGCTGGCCGGCTTGGT GGGGACACCTTTGTACAAAGAGTTTGCG TATACAAAGAGGAGGCGGACGCG | 68°C | 3370 | PpCESA3 coding |
| HACESA3attB5 CESA3CDSattB2 | See above See above | 68°C | 3370 | PpCESA3 coding |
| HACESA5attB5   | GGGGACAACCTTTGTATACAAAAAGTTGCG ATGGGCTACCCTCTACGATGTGCCCCGATT | 68°C | 3337 | PpCESA5 coding sequence |
| CESA5attB2     | ATGGCCTACCCTCCTACGATGTGCCCCGATT ATGGCTATGGAGGCTAATGCAGGCCTTTAT GGGGACCCTTTGTACAAAGAAGCTGGG |   |   |   |
|                | TACTAACAGCTAGCTAACGGCCGACACTGAC |   |   |   |
| HACESA4attB5   | GGGGACAACCTTTGTATACAAAAAGTTGCG ATGGAGTAACACCATACGATGTGCCCCGATT | 68°C | 3391 | PpCESA4 coding sequence |
| CESA4CDSatB2   | ATGGAGTAACACCATACGATGTGCCCCGATT ATGGCTATGGAGGCTAATGCAGGCCTTTAT GGGGACCCTTTGTACAAAGAAGCTGGG |   |   |   |
|                | TACTATCGACAGTTGATCCCACACTG |   |   |   |
| HACESA7attB5   | GGGGACAACCTTTGTATACAAAAAGTTGCG ATGGAGTAACACCATACGATGTGCCCCGATT | 68°C | 3382 | PpCESA7 coding sequence |
| CESA6_7attB2   | ATGGAGTAACACCATACGATGTGCCCCGATT ATGGCTATGGAGGCTAATGCAGGCCTTTAT GGGGACCCTTTGTACAAAGAAGCTGGG |   |   |   |
|                | TATCAACAGTTTATCCCGCACTGCGA |   |   |   |
| HACESA10attB5  | GGGGACAACCTTTGTATACAAAAAGTTGCG ATGGAGTACCCATACGATGTGCACACT | 68°C | 3379 | PpCESA10 coding sequence |
| CESA10CDSatB2  | ATGGAGTACCCATACGATGTGCACACT |   |   |   |
|                | ATGGCTATGGAGGCTAATGCAGGCCTTTAT GGGGACCCTTTGTACAAAGAAGCTGGG |   |   |   |
|                | TACTATCGACAGTTGATCCCACACTC |   |   |   |
Fig. S1. Genotype analysis of *ppcesa8*, *ppcesa3* and *ppcesa3/8* KO lines. (A) Genotyping strategy and results for *ppcesa8* KO lines. 5’ integration tested by PCR with primer pair 8KOFlankF/VectorR-hph produced the expected 1121 bp fragment in lines 8KO5B, 5KO4C, 5KO5C, 5KO7C and 8KO10C. 3’ integration tested by PCR with primer pair VectorF-hph/8KOFlankR produced the expected 637 bp fragment in the same 5 lines. Target deletion was verified in the 3 KO lines by the absence of a product from primer pair CESA8TargetF/CESA8TargetR, which anneal within the *PpCESA8* coding sequence and amplify a 339 bp fragment in the wild type. (B) Genotyping strategy and results for *ppcesa3* and *ppcesa3/8* KO lines. 5’ integration tested by PCR with primer pair 3KOFlankF/VectorR-npt produced the expected 1362 bp fragment in lines 3KO5, 3KO35, 3KO126, 3/8KO43, 3/8KO57, and 3/8KO86. 3’ integration tested by PCR with primer pair VectorF-npt/3KOFlankR produced the expected 1259 bp fragment in the same 6 lines. Target deletion was verified in the 6 KO lines by the absence of a product from primer pair CESA3TargetF5/CESA3TargetR5, which anneal within the *PpCESA3* coding sequence and amplify a 1266 bp fragment in the wild type.
Fig. S2: Phenotype analysis of a ppces3/8 double KO line transformed with vectors driving expression of PpCESA3 or PpCESA8 with their native promoters. Bright field images captured with a stereomicroscope show colony morphology (A-C, G-I, M-O) and polarization images show cell wall birefringence (D-F, J-L, P-R). (A-F) Wild type with erect gametophores (A) and strong cell wall birefringence (D), ppcesa8 KO with erect gametophores (B) and intermediate cell wall birefringence (E) and ppces3/8 KO with horizontal gametophores (C) and weak birefringence (F) are shown for comparison to complemented lines. (G-R) Complemented lines have erect gametophores (G-I, M-O) and strong cell wall birefringence (J-L, P-R).
Fig. S3. Transmission electron microscopy images of leaf cell walls from wild type and *cesa*KO lines of *P. patens*. In lamina cells, outer walls face the external environment, inner walls are between cells, and middle lamellae are from cell junctions. Margin cells are from leaf edges.
Fig. S4. Thickness of outer cell walls measured from transmission electron microscopy images. Error bars represent standard error of the mean (n=2 lines per genotype).
Fig. S5: *P. patens* wild type and KO lines cultured on medium containing 1 μM naphthalene acetic acid (auxin) to induce rhizoid initiation and inhibit leaf initiation. (A) A wild type colony with leafless gametophores (arrows). (B,C) Dark field images of wild type leafless gametophores with multiple rhizoids (arrowheads). (D-I) Dark field images of *ppcesa3* KO and *ppcesa3/8* KO leafless gametophores with multiple rhizoids. (J-L) Bright field images of *ppcesa8* KO leafless gametophores with multiple rhizoids. No defects in rhizoid initiation or growth were noted in any of the KO line
Fig. S6: *P. patens* wild type and KO lines cultured in the dark on vertically oriented plates containing medium supplemented with 35 mM sucrose to test for caulonema gravitropism. KO lines in columns 2-4 of each row are compared to their background wild type line from the same experiment in column 1. No significant differences in caulonema length or gravitropic behavior were detected.
Fig. S7. Western blot analysis of protein expression for *P. patens* lines derived from transformation of *ppcesa3/8KO-86lox* with vectors driving expression of PpCESAs under control of the *PpCESA8* promoter. Western blot probed with anti-HA is shown above the same blot stained with Ponceau S as a loading control. Protein loading per lane was 3.6 μg. Asterisks indicate lines that rescued the mutant phenotype.
Fig. S8: PCR-based genotyping of ppeesa6/7 KO lines. Primers used for amplification of 5' and 3' ends are indicated as black arrows on the diagram showing the PpCESA6/7KO vector integrated so as to delete PpCESA6 and PpCESA7, which occur as a tandem repeat. The products confirming 5' (1647 bp) and 3' (833 bp) integration amplified in three KO lines (6A, 7A and 1D) selected from two transformations. Products from amplification of the target genes PpCESA6 (142 bp) and PpCESA7 (1254 bp) were observed in wild type (WT), but not in KO line.
Fig. S9: PCR-based genotyping of *ppcesa4/10* KO lines. (A) Six *ppcesa10*KO lines recovered from one transformation with the PpCESA10KO vector conferring hygromycin resistance were verified by amplification the 5’ integration site (1461 bp) and 3’ integration site (1136 bp) and lack of amplification of the target gene (973 bp). (B) *ppcesa4*KO-13A recovered from a transformation with the CESA4KO vector conferring hygromycin resistance was verified by amplification the 5’ integration site (1521 bp) and 3’ integration site (1832 bp) and lack of amplification of the target gene (1731 bp) and *cre*-mediated deletion of the selection cassette was verified by amplification across the deletion site (2321 bp). (C) A double *ppcesa4/10*KO line from transformation of *ppcesa10*KO-5 with the CESA4KO vector conferring G418 resistance was verified by amplification the 5’ integration site (1263 bp) and 3’ integration site (1839 bp) and lack of amplification of the target gene (1731 bp). (D) Double *ppcesa4/10*KO lines from transformation of *ppcesa4*KO-lox with the CESA10KO vector conferring G418 resistance was verified by amplification the 5’ integration site (1274 bp) and 3’ integration site (1132 bp) and lack of amplification of the target gene (973 bp).
Fig. S10: Quantification method for S4B fluorescence. Representative paired DIC (A) and fluorescence (B,C) micrographs of a <i>P. patens</i> leaf stained with S4B. Insets show the central midrib and surrounding bundle sheath at higher magnification. The central midrib was selected manually using the polygon selection tool in ImageJ (Fiji version) as shown by the red lines in C.
Short title: Identify functions of CESAs in *Physcomitrella*

Title: Morphological analysis of cellulose synthase (CESA) mutants in *Physcomitrella patens*

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One sentence summary: The *Physcomitrella* clade-B PpCESAs (PpCESA4, PpCESA6, PpCESA7, and PpCESA10) are not required for leafy gametophore morphogenesis, indicating PpCESA5 forms homo-oligomeric cellulose synthesis complexes; meanwhile PpCESA4 and PpCESA10 are involved in the tip growth of protonema cells.

List of author contributions: A.W.R. conceived the project, and supervised experiments; X.L. designed and performed experiments, and analyzed the data; M.L.T and J.H.N performed experiments; X.L. and A.W.R. wrote the manuscript. All authors read and approved the manuscript.

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Abstract

Cellulose produced by plasma membrane rosette Cellulose Synthesis Complexes (CSCs) is an essential component of plant cell walls, providing vital mechanical strength. The catalytic subunits of CSCs, called cellulose synthase (CESA) proteins, are encoded by gene families that vary in size among different plant species. *Arabidopsis* has 10 functionally non-redundant *CESA* genes, and assembly of its CSCs requires the participation of at least three members from this gene family, which means these CSCs are obligate hetero-oligomeric. The moss *Physcomitrella patens* has rosette CSCs and seven CESA genes that have not been fully characterized functionally. According to phylogenetic studies, the PpCESAs are not members of the clades comprising the different subunits of the hetero-oligomeric seed plant CSCs. Hence, it is unknown whether *P. patens* CSCs are also hetero-oligomeric. Previous functional analyses showed that *ppcesa5* knockout (KO) mutants are unable to produce gametophores. Double *ppcesa3/8KOs* were shown to be defective in secondary cell wall deposition in gametophore leaf midribs. Here, we continue investigating functions of *PpCESAs* through morphological analysis of *ppcesa* KO mutants to gain clues about the composition of *P. patens* CSCs. Our results show that B-clade PpCESAs (PpCESA4, 6, 7, and 10) are not required for gametophore morphogenesis. However, PpCESA4 and 10 are found to serve a function in the tip growth of protonema filaments, indicating the potential roles of cellulose in the cells undergoing tip growth.
Introduction

Cellulose is a key component in plant cell walls. In the primary cell wall (deposited during cell expansion), the oriented deposition of cellulose microfibrils serves the vital load-bearing role important in determining the orientation of cell expansion and thus overall plant morphology (Taylor, 2008). After cell expansion has stopped, certain cells, such as collenchyma cells, sclerenchyma cells, and xylem cells, can deposit thickened secondary cell walls (inside the primary wall) that mechanically support plants to stand upright and efficiently conduct water and minerals (Mauseth, 2012). Cellulose is highly abundant in the secondary walls (Taylor, 2008). Cellulose microfibrils, in higher plants, are synthesized by rosette cellulose synthesis complexes (CSCs) embedded in the plasma membrane. The catalytic core of these complexes is assembled from cellulose synthase (CESA) subunits (Delmer, 1999; Kimura et al., 1999; McFarlane et al., 2014). In seed plants, the CSCs for cellulose deposition in both primary and secondary cell wall requires three types of functional distinct CESAs for function (McFarlane et al., 2014). In Arabidopsis, mutants for CESA1, CESA3, and CESA6 have cellulose defects in primary cells wall causing developmental retardation and phenotypic changes in hypocotyls and roots (Arioli et al., 1998; Fagard et al., 2000; Williamson et al., 2001; Burn, et al., 2002; Robert et al., 2004). Mutations in any of the three secondary cell wall CESAs (CESA4, 7, and 8) result in severe defects in secondary cell wall cellulose deposition leading to collapsed xylem cells in Arabidopsis (Turner & Somerville, 1997; Taylor et al., 1999; Taylor et al., 2000; Taylor et al., 2003). The moss Physcomitrella patens is an intriguing model bryophyte that is commonly used in genetics studies and mutational analysis because of its ability
to be genetically manipulated due to the naturally occurring high rate of homologous recombination. Gene knockin and knockout transformations can be accomplished within one month and phenotyped in a few weeks in *P. patens* (Kamisugi, Cuming, & Cove, 2005). This is rapid compared to transformation and phenotypic analysis in *Arabidopsis*, which takes about three months (Clough & Bent, 1998). Rapidly elongating protonema cells in *P. patens* can be used as an alternative model to examine tip-growth related mechanisms (Rounds & Bezanilla, 2013). Leafy gametophores consist of several distinguishable cell types including support cells (stereids) and water-conducting cells (hydroids), but they develop from single-celled shoot apical meristems, making *P. patens* a less complicated model to study plant organ morphogenesis (Harrison et al., 2009). *Physcomitrella patens* has seven CESA genes which can be divided into two sub-clades (A-clade: *PpCESA*3, 5, and 8; B-clade: *PpCESA*4, 6, 7, and 10), but are not orthologs of seed plants CESAs (Goss et al., 2012; Roberts & Bushoven, 2007).

We carried out morphological analysis of CESA knockout (KO) mutants in order to investigate functions of CESAs in *P. patens*. So far, PpCESA5 is known to be required in gametophore development based on the "no leafy gametophore" phenotype of *ppcesa5*KO mutant (Goss et al., 2012). Both of double *ppcesa3/8*KO and *ppcesa6/7*KO mutants show significantly reduced cellulose deposition in secondary cell walls in midribs of gametophore leaves, indicating PpCESA3, 8, 6, and 7 are involved in secondary cell wall thickening of stereids (Norris et al., 2017). Here, we show that the quadruple *ppcesa4/6/7/10*KOs are able to produce morphologically normal leafy gametophores, indicating that the B-clade PpCESAs are not required in...
gametophore morphogenesis. Since ppcesa3/8KO also produces morphologically normal gametophores (Norris et al., 2017), together the current results suggest that PpCESA5 might be able to form homo-oligomeric CSCs, solely functioning in gametophore development. In addition, knocking out PpCESA4 and PpCESA10 causes morphological changes in protonemal colonies, suggesting the importance of cellulose in the tip-growing P. patens protonema cells.

Results

Genotyping and morphological analysis (rhizoid, caulonema, and gametophore) of ppcesa4/6/7/10KO

Three verified quadruple ppcesa4/6/7/10KO lines were recovered from three different transformations of ppcesa4/10KO-4B with the CESA6/7KO vector (Norris et al., 2017) and tested for 5’ and 3’ integration of the vector and deletion of the target gene (Figure 1). All of the quadruple KO lines were able to produce leafy gametophores that were morphologically similar to wild type (Figure 2 A-H) indicating that the B-clade PpCESAs are not required for gametophore morphogenesis. The quadruple KOs were also tested for developmental defects in rhizoid and caulonema development. All of the three quadruple KOs produced leafless gametophores with several rhizoids similar to wild-type after growing on medium supplied with auxin for two weeks (Figure S1), indicating that PpCESA4, PpCESA6, PpCESA7, and PpCESA10 are not required for normal rhizoid development. When explants of quadruple KOs were cultured vertically in the dark, caulonemal filaments produced by the resulting colonies grew upright against gravity and were similar in appearance to wild-type
controls (Figure S2 A-F). Caulonemal length was also not significantly different
between the mutant lines and wild-type lines (Figure S2 G).

**Cellulose deposition of the secondary cell wall in ppcesa4/6/7/10KO**

By polarization microscopy and S4B staining, Norris et al. (2017) showed a large and
significant reduction in cellulose deposition in the midribs of ppcesa6/7KO
gametophore leaves, whereas the gametophore leaves of ppcesa4/10KOs showed a
small, but significant reduction compared to wild-type. To clarify the roles of the clade
B PpCESAs in secondary cell wall deposition, we used polarization microscopy to
examine midrib birefringence in ppcesa4/6/7/10KO compared to wild-type (Gd11).
We found that gametophore leaves of three ppcesa4/6/7/10KO lines all had
substantially reduced midrib birefringence (Figure 2 J-L), similar to the phenotypes
of previously described ppcesa6/7KOs and ppcesa3/8KOs, and more dramatic than
ppcesa4/10KOs (Norris et al., 2017). To quantify the defect in secondary cell wall
deposition relative to ppcesa4/10KOs and ppcesa6/7KOs, we stained mutant
gametophore leaves with cellulose-specific fluorescent dye S4B and used fluorescence
microscopy to measure the cellulose content in midribs of the mutant leaves. All
mutants showed a significant reduction in brightness compared with the midribs of
wild-type gametophore leaves (Figure 3), consistent with previous results. The
quadruple KO midribs had significantly reduced brightness compared to
ppcesa4/10KOs but were not significantly different from ppcesa6/7KOs (Figure 3).
The phenotype similarity of ppcesa4/6/7/10KO compared to ppcesa6/7KO, but not
ppcesa4/10KO (Norris et al., 2017) indicates a major role for PpCESA6 and
PpCESA7 and a minor role for PpCESA4 and PpCESA10 in secondary cell wall deposition in gametophore leaf midribs.

**Morphological analysis of protonema colonies**

Protonemal filaments of *P. patens* extend by apical cell division and tip growth, branching to form colonies (Cove, 2005). To test whether clade B PpCESAs are required for protonemal tip growth, Chlorophyll autofluorescence images of colonies were analyzed for area, solidity, and circularity. **Figure 4** summarizes the results of this assay for ppcesa4/6/7/10KOs, ppcesa6/7KOs, and ppcesa4/10KOs. Circularity is the ratio of colony area to colony perimeter and indicates the degree of polarized extension. A score of 1 represents a perfect circle, while scores approaching 0 represent a more linear shape. Solidity quantifies the presence of concavities in the colony and reflects the degree of polarization and branching of the protonema filaments. The lowest solidity with the highest branching of the filaments was scored 0 and the highest solidity possible with less branching of filaments was scored 1 (Vidali et al., 2007). Graphs in **Figure 5** show that when compared with wild-type control, colonies of *ppcesa4/6/7/10KO* showed increased solidity and circularity. *pcesa6/7KOs* showed no difference in area, solidity or circularity compared to wild-type (*P* > 0.05, ANOVA). All *ppcesa4/10KOs* showed significantly increased solidity and circularity compared to wild-type similar to *ppcesa4/6/7/10KO*, consistent with defects in protonemal tip growth. We further analyzed single *ppcesa4KOs* and *ppcesa10KOs*. Only two of the three *ppcesa4KO* lines, *ppcesa4KO-13A* and *ppcesa4KO-14B*, had significantly increased solidity and circularity compared to wild-type. There was no significant difference observed among the three *ppcesa10KOs*. We also tested
ppcesa5KO and ppcesa3/8KO to test the roles of the clade A PpCESAs in protonema tip growth. One of the three ppcesa5KOs, ppcesa5KO-20, showed the significantly increased solidity and circularity compared to both wild-type and the other two ppcesa5KOs. None of ppcesa3/8KOs showed any significant difference compared with wild-type in the three parameters.

**Discussion**

Mutation analysis for the B-clade *PpCESA*S revealed that they are not required for gametophore morphogenesis. This is evident from the fact that quadruple ppcesa4/6/7/10KOs are still able to produce normal leafy gametophores (Figure 2), unlike ppcesa5KO. Gametophore buds of ppcesa5KOs are defective in cell expansion, cytokinesis, and leaf initiation, resulting in failure of leafy shoot formation (Goss et al., 2012). None of these phenomena were observed in ppcesa4/6/7/10KOs. The ppcesa3/8KOs also produce morphologically normal gametophores (Norris et al., 2017). Thus ppcesa5KOs are the only mutants that are defective in gametophore morphogenesis. It has also been shown that constitutively expressing PpCESA3 and PpCESA8 can rescue ppcesa5KO indicating A-clade PpCESA5s are functionally interchangeable (Norris et al., 2017). Thus, the unique mutant phenotype of ppcesa5KOs might be attributable to PpCESA5 having non-overlapping expression with PpCESA3 and PpCESA8 and the non-interchangeable functions with the B-clade PpCESAs (Scavuzzo-Duggan et al., unpublished). According to this, PpCESA5 might be able to form homo-oligomeric CSCs in order to properly deposit cellulose microfibrils into the cell walls of newly emerged gametophore buds. The
interchangeable functions of different CESA members are seen only in limited cases in seed plants. Promoter-swap assays in *Arabidopsis* showed that the defective phenotype of *atcesa3* mutants can be partially rescued by driving expression of *AtCESA7* using the *AtCESA3* promoter and *atcesa8* mutants can be partially rescued by driving expression of *AtCESA1* using the *AtCESA8* promoter (Carroll et al., 2012).

Results of S4B staining (Figure 3) showed that: 1) there is no significant difference between *ppcesa4/6/7/10*KO and *ppcesa6/7*KO in cellulose content in the midrib secondary cell walls of the mutant leaves; 2) there is a slight but significant reduction in *ppcesa4/10*KO compared to the wild-type. These results suggest that compared with PpCESA6 and PpCESA7, PpCESA4 and PpCESA10 only have a minor role in secondary cell wall deposition. This is consistent with previous gene expression data showing that *PpCESA4* and *PpCESA10* have lower expression in gametophores than in protonema (Hiss et al., 2014; Tran & Roberts, 2016). The fact that *ppcesa3/8*KO and *ppcesa6/7*KO are similar in phenotype showing cellulose defects in secondary cell walls provides a clue that CSCs involved in cellulose deposition in *P. patens* secondary cell walls might be hetero-oligomeric consisting of PpCESA3, PpCESA8, PpCESA6, and PpCESA7 (Norris et al., 2017).

Tip growth in certain types of cells, such as root hairs and pollen tubes, is regulated by highly coordinated mechanisms which guide deposition of new cell wall materials strictly proceeding in a limited area of the cell surface (Carol & Dolan, 2002; Cosgrove, 2005; Cheung & Wu, 2008; Lee & Yang, 2008; Nielsen, 2009; Gu & Nielsen, 2013). Several studies pointed out that cellulose is an essential cell wall component in cells undergoing tip growth (Newcomb & Bonnett, 1965; Emons &
Wolters-Arts, 1983; Emons, 1994; Galway et al., 2011; Park et al., 2011). Mutational analyses in Arabidopsis showed that some atcesa mutants are severely defective in germinating pollen and elongating pollen tube, indicating important roles of cellulose in the tip-growing cells (Persson et al., 2007). Elongating P. patens protonemal filaments are another ideal model to investigate the role of cell wall deposition in tip-growth related mechanisms (Roberts et al., 2012). Crystalline cellulose has been detected by affinity cytochemistry with Cellulose Binding Module 3A (CBM3A) in primary cell walls of subapical cells and the very tip region of the apical cells in expanding P. patens protonema filaments (Berry et al., 2016), indicating the potential roles of cellulose during tip growth of protonema. Here, our study shows that P. patens CESAs (PpCESA4 and 10) have roles in tip growing protonema, supporting the point of view that cellulose is significant for cell tip growth. This is evident from the abnormal protonema colony morphology of ppcesa4/10KOs, which show significantly increased circularity and solidity (Figure 5). Increased circularity and solidity are caused by slower elongation and less branches of the protonema filaments (Vidali et al., 2007). Quantitative affinity cytochemistry of cellulose content using S4B or CBM3A will be needed to prove that the mutant phenotypes were caused by the decreased cellulose in cell walls of tip-growing protonema cells. Based on available evidence, PpCESA6, PpCESA7, and the A-clade PpCESAs do not seem to contribute to protonemal tip growth, since no obvious phenotypic changes were observed in corresponding KO mutants. Although colony circularity and solidity of ppcesa5KO-20 was shown to be significantly increased in our analysis, this is likely due to other genetic effects since the other ppcesa5KO lines were not different from
wild-type. It remains possible that PpCESA5, PpCESA3 and PpCESA8 function redundantly in tip growth. This can be tested by producing and analyzing a ppcesa3/5/8 triple KO mutant. Tip growth in our ppcesa4/10KO and ppcesa4/6/7/10KO was not abolished, suggesting the deposition of cellulose in cell walls of tip-growing protonema involves proteins other than the PpCESAs. Several members from one of the Cellulose Synthase-like (CSL) gene family, CSLD, were shown to be required for tip growth of root hairs and pollen tubes in Arabidopsis (Favery et al., 2001; Wang et al., 2001; Bernal et al., 2008; Park et al., 2011). P. patens also has the CSLD gene family, and expression of these genes have enhanced expression in cultures containing only protonema (Roberts & Bushoven, 2007). Thus, it will be interesting to carry out mutational analysis to investigate functions of CSLD genes in P. patens protonema.

Materials and methods

Transformation and genotyping

Except ppcesa4/6/7/10KOs, the ppcesaKO lines used in this study were created previously and described in Norris et al. (2017).

To create the quadruple ppcesa4/6/7/10KO lines, the hygromycin sensitive ppcesa4/10KO-4B line (Norris et al., 2017) was transformed with the CESA6/7KO vector conferring hygromycin resistance and stably transformed colonies were genotyped as described for primary ppcesa6/7KO lines in Norris et al. (2017). Primers used for genotyping are listed in supplemental table 1.
Polarization microscopy of cell wall birefringence

Cell wall birefringence of leaf midribs was analyzed as described in Norris et al., (2017). Three independent lines of each knockout mutant and three biological replicates of wild-type were cultured for 15 days on BCDAT medium. The first fully expanded leaf of each gametophore was cut off with a pair of micro-dissecting scissors (Electron Microscopy Sciences, Hatfield, PA, USA) and mounted in the water on a glass slide. An Olympus BHS compound microscope equipped with a polarizer and circular-polarizing analyzer (Olympus Corp., Shinjuku, Tokyo, Japan) was used to visualize the gametophore leaves. Images were captured with a Leica M165FC digital camera (Leica Microsystems Inc., Buffalo Grove, IL, USA) using identical settings for the knockouts and the wild-type control.

Pontamine fast scarlet 4B (S4B) fluorescence histochemistry

S4B staining of leaf midribs was performed as describe (Norris et al., 2017). Three independent lines of each knockout along with three biological replicates of wild-type were cultivated on BCDAT medium for 15 days. For each genotype, three gametophores with 10-12 leaves were collected, permeabilized in acetone for 5 seconds, rinsed in PBS, and stained for 30 min in PBS containing 0.01% S4B. All leaves were rinsed in PBS after staining, cut off with a sharp razor blade, and mounted in PBS on a glass slide. Images were taken using the same microscope and conditions described previously in Norris et al (2017). For data analysis, the midrib of each leaf was outlined by hand and intensity was quantified using ImageJ as described previously (Norris et al., 2017).
Analysis of caulonema and rhizoid development

Caulonema and rhizoid assays were carried out as previously described (Roberts et al., 2011) to test ppcesa4/10KO, ppcesa6/7KO, ppcesa4/6/7/10 KO lines for phenotypic changes. For the caulonema and rhizoid assays, samples were analyzed using a Leica M165FC stereomicroscope, and images were recorded using a Leica DFC310FX camera (Leica). The length of caulonema was measured as described in Norris et al. (2017). Three independent experiments (n=3) were done. For each experiment, caulonema colonies were cultured on seven replicate plates containing solid BCDAT? medium. Four explants were placed along the equator of each plate, with each explant representing a unique genotype.

Protonema colony morphology assay

Colony morphology was analyzed as described previously (Bibeau & Vidali, 2014). Protoplasts were isolated from three independent lines for each genotype along with three biological replicates of wild-type using the method described previously (Roberts et al., 2011). However, it was necessary to add 21 units/mL of cellulase from *Trichoderma reesei* (Worthington Biochemical Corporation, Lakewood, NJ, USA) to the digestion mixture when using driselase lot # SLBP0654V (Sigma-Aldrich, St. Louis, MO, USA) for effective digestion. Five thousand protoplasts suspended in 1 mL of PRML were spread on each of three plates containing PRMB medium overlain with cellophane. The plates were incubated at 25°C with constant illumination at 50-80 μmol/m^2/s for 4 d and cellophane membranes were then transferred to BCDAT plates for an additional 2 d. Colony morphology was documented by capturing chlorophyll autofluorescence images of approximately 50 regenerated protoplasts per
plate at 63X magnification using an M165FC stereo microscope with 10447407 GFP filter and DFC310FX camera (Leica). Images were analyzed for area, solidity, and circularity with ImageJ (National Institutes of Health, USA) using a macro developed by Vidali et al. (2007).

**Statistical analysis**

One-way Analysis of Variance (ANOVA) followed by post-hoc Tukey Honest Significant Difference (HSD) test was performed using "R" programming (Vienna, Austria; http://www.R-project.org/) to identify the potential significant difference in caulonema assay and tip growth assay.

**Supplemental Materials**

Table. S1. Primers designed for knockout vector construction and genotyping.

Table. S2. Data of morphological analysis of protonema colonies.

Fig. S1: B clade PpCESAs are not required for rhizoid development.

Fig. S2. B-clade PpCESAs are not required for caulonema development and gravitropism.

**Acknowledgments**

This work was supported by National Science Foundation Award IOS-1257047. DNA sequencing and qPCR were conducted using the Rhode Island Genomics and Sequencing Center, a Rhode Island NSF EPSCoR research facility, supported in part by the National Science Foundation EPSCoR Cooperative Agreement EPS-1004057. We also thank Bowen Jiang for assistance with statistics.
Figure 1: PCR-based genotyping of ppcesa4/6/7/10KO lines. Genomic DNA from wild-type *P. patens* (WT) was used as positive control. The expected band for the target gene of 1178 bp (*PpCESA6*) and 141 bp (*PpCESA7*) was observed in WT, but not in KO lines. The expected 5' integration band of 833 bp was present in the KO lines created with PpCESA6/7 KO vector but was not seen in WT. The expected 3' integration band of 1647 bp was observed in the same KO lines above but was also not present in WT. Primer sets used for 5' and 3' ends amplification are indicated as black arrows on the graph showing each gene's locus.
Figure 2: B clade PpCESAs are not required for gametophore development. (A-D) Colony morphology is similar in wild-type (A) and three *ppcesa4/6/7/10KO* lines (B-D). (E-H) Normal looking leafy stalks were observed in all of the lines above. (I-L) Polarized light microscopy of detached leaves from gametophores cultured for 15 days on BCDAT medium for wild-type (I) and three *ppcesa4/6/7/10KO* lines (J-L). The leaf midribs of three knockout lines (J-L) were shown to have lower birefringence compared with wild-type (I) leaf midribs.
Figure 3: Quantitative analysis of S4B fluorescence intensity in leaf midribs of wild-type *P. patens* and B clade *PpCESA* knockout (KO) mutants. *Ppcesa4/10*KO leaf midribs have a moderate yet significant reduction in fluorescence intensity compared to wild-type. Fluorescence is significantly weaker in *ppcesa6/7*KO than it is wild-type and *ppcesa4/10*KO. *Ppcesa4/6/7/10*KO lines also have significantly decreased fluorescence intensity in leaf midribs compared to wild-type and *ppcesa4/10*KO, but there is no significant difference between *ppcesa6/7*KO and *ppcesa4/6/7/10*KO lines. Three independent genetic lines were tested in triplicate for each mutant genotype. The Gd11 line was used as the wild-type control and sampled in triplicate. Error bars indicate standard error of the mean (*ppcesa4/10*KO, n=3; *ppcesa6/7*KO, n=3; *ppcesa4/6/7/10*KO, n=3; wild-type, n=3).
Figure 4: Representative micrographs showing morphologies of B clade PpCESA knockout (KO) mutants protonemal colonies undergoing tip growth. Micrograph show the morphology of colonies with median (add parameter that you used to select the median) as determined by imaging chlorophyll autofluorescence.
Figure 5: Protonemal colonies of knockout (KO) mutant lines and wild-type *P. patens* line were analyzed for area, circularity, and solidity. Protonemal colonies of *P. patens* KO mutant lines and wild-type line were analyzed for three parameters, area, circularity, and solidity. For circularity, plants with values approaching one are more circular. Solidity scale of 1 represents the highest solidity with 0 as the lowest solidity with the highest branching. Here, the bar graphs show changes of the three parameters in KO mutants. The height of the bar represents the ratio (KO mutant/wild-type). A ratio larger than 1 (indicated by a dotted line on each graph) indicates an increase of that parameter in KO lines compared to wild type and a ratio less than 1 indicates a decrease for KO lines. Error bars display standard error of the mean between each data set (n=3 for each data set). Statistical significant difference between KO mutant and wild-type is indicated by the "▲" sign. The statistically significant difference among KO mutants is indicated by the "●" sign. Raw measurements are reported in Table S4.
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## Supplemental Materials

Table S2. Primers designed for knockout vector construction and genotyping.

| Name             | Sequence                                                                 | Tm (°C) | Amplicon Size (bp) | Description                                               |
|------------------|--------------------------------------------------------------------------|---------|--------------------|-----------------------------------------------------------|
| 6KOF2 VectorR-hph| GCTTCAATGCTGTACCACAAACCA C TCCAGGGCAAGAAATAGA                            | 57°C    | 1647               | 5' integration testPpCESA6 with hph cassette             |
| VectorF-hph      | TGACAGATAGCTGGGCAATG AAGCCCTAACTTCCAGCACC                                 | 57°C    | 833                | 3' integration testPpCESA7 with hph cassette             |
| CESA7FlankR      | GCGAATGCAGGGCTGCTG ACATTACTCAACGGCCTCGG                                  | 60°C    | 1178               | PpCESA7 deletion                                         |
| CESA6TargetF     | GTGAGGTGCGAGGAAAGAAG                                                    | 60°C    | 141                | PpCESA6 deletion                                         |
| CESA6TargetR     | GGCGAATGCAGGGCTGCTG                                                      |         |                    |                                                           |
| CESA7TargetF     | ACATTACTCAACGGCCTCGG                                                    | 60°C    | 1178               | PpCESA7 deletion                                         |
| CESA7TargetR     | ACATTACTCAACGGCCTCGG                                                    |         |                    |                                                           |
Table S2. Data of morphological analysis of protonema colonies. For each knockout mutant analyzed, three independent lines of each genotype were used. Three biological replicates of wild-type control (Gd11) were included in each assay.

| Ppcesa4/6/7/10KO | Gd11 | Area | Circularity | Solidity |
|------------------|------|------|-------------|----------|
|                  |      | 40399 | 0.04500 | 0.32044 |
| Ppcesa4/6/7/10KO | O1   | 20285.5 | 0.0742 | 0.45040 |
|                  |      | 28965.2 | 0.6975 | 0.40170 |
|                  |      | 27604.7 | 0.8108 | 0.44066 |
|                  | O2   | 18547.2 | 0.08993 | 0.46961 |
|                  |      | 30581.3 | 0.7216 | 0.41858 |
|                  |      | 26905.7 | 0.7559 | 0.41858 |
|                  | O3   | 18319.3 | 0.09851 | 0.47605 |
|                  |      | 33573.7 | 0.8378 | 0.45982 |
|                  |      | 35148.9 | 0.9548 | 0.46436 |
|                  |      |        |          |          |
| Ppcesa6/7KO1D    | Gd11 | 17151.1 | 0.12093 | 0.46761 |
|                  |      | 15971.6 | 0.16288 | 0.53438 |
|                  |      | 18077.3 | 0.12687 | 0.45907 |
|                  | Ppcesa6/7KO6A | 18683.8 | 0.09826 | 0.42167 |
|                  |      | 17356.7 | 0.10113 | 0.42660 |
|                  |      | 27964.1 | 0.10592 | 0.45876 |
|                  | Ppcesa6/7KO7B | 21736.6 | 0.11249 | 0.46012 |
|                  |      | 17140.6 | 0.12051 | 0.46605 |
|                  |      | 16779  | 0.18276 | 0.53886 |
|                  |      |        |          |          |
|                  | Ppcesa4/10KO1A | 21237.3 | 0.04925 | 0.33708 |
|                  |      | 23356.3 | 0.05085 | 0.33751 |
|                  |      | 23087.8 | 0.04897 | 0.34933 |
|                  | Ppcesa4/10KO4B | 18132.0 | 0.07399 | 0.39773 |
|                  |      | 19258.2 | 0.07932 | 0.40354 |
|                  |      | 22136.1 | 0.07762 | 0.40859 |
|                  | Ppcesa4/10KO7B | 18396.8 | 0.06135 | 0.38732 |
|                  |      | 17776.9 | 0.08310 | 0.41631 |
|                  |      | 17604.1 | 0.08334 | 0.40156 |
|                  |      |        |          |          |
|                  | Ppcesa10KO5    | 32858.3 | 0.03546 | 0.30819 |
|                  |      | 29972.5 | 0.03504 | 0.31219 |
|                  |      | 31806.7 | 0.03282 | 0.27408 |
|                  | Ppcesa10KO6    | 33235.3 | 0.04286 | 0.33303 |
|                  |      | 34680.9 | 0.04573 | 0.31052 |
|                  |      | 39200.3 | 0.03999 | 0.29706 |
|                  | Ppcesa10KO13   | 33246.8 | 0.03633 | 0.30547 |
|                  |      | 43001.8 | 0.03352 | 0.28189 |
|                  |      | 35311.7 | 0.03450 | 0.27599 |
|                  |      |        |          |          |
|                  | Ppcesa4KO12B   | 24418.7 | 0.04464 | 0.32436 |
|                  |      | 18226.8 | 0.04547 | 0.31683 |
|                  |      | 20659.8 | 0.04641 | 0.34460 |
|                  | Ppcesa4KO13A   | 24471.9 | 0.05777 | 0.38431 |
|                  |      | 19759.3 | 0.06349 | 0.36331 |
|                  |      | 24988.6 | 0.05506 | 0.33547 |
|                  | Ppcesa4KO14B   | 25267.9 | 0.05307 | 0.37208 |
|                  |      | 19759.0 | 0.06925 | 0.39133 |
|                  |      | 23972.9 | 0.07203 | 0.39576 |
|                  |      |        |          |          |
|                  | Ppcesa5KO      | 22012.5 | 0.04333 | 0.31432 |
|                  |      | 24818.3 | 0.03759 | 0.28210 |
|                  |      | 25844.6 | 0.03926 | 0.29756 |
|                  | Ppcesa5KO      | 19099.7 | 0.05255 | 0.35224 |
|                  |      | 24562.6 | 0.04243 | 0.31946 |
|                  |      | 23829.9 | 0.04049 | 0.30255 |
|                  | Ppcesa5KO      | 18242.9 | 0.04172 | 0.29151 |
|                  |      | 24602.5 | 0.03816 | 0.29486 |
|                  |      | 24648.7 | 0.04150 | 0.29269 |
|                  | Ppcesa5KO      | 19725.5 | 0.06487 | 0.38877 |
|                  |      | 26649.7 | 0.05430 | 0.35500 |
|                  |      | 22635.9 | 0.05432 | 0.36718 |
Figure S1: B clade PpCESAs are not required for rhizoid development. *P. patens* wild-type (Gd11) and three independent *ppcesa4/6/7/10KO* lines were cultured on medium supplemented with 1 μM naphthalene acetic acid (auxin) to stimulate rhizoid initiation, and inhibit leaf initiation. (A-C) Dark field images of wild-type leafless gametophores with numerous rhizoids. (D-F) Dark field images of *ppcesa4/6/7/10KO1*, *ppcesa4/6/7/10KO2*, and *ppcesa4/6/7/10KO3* leafless gametophores with numerous rhizoids. None of the three *ppcesa4/6/7/10KO* lines showed defects in rhizoid initiation or growth.
Figure S2: B-clade PpCESAs are not required for caulonema development and gravitropism. *P. patens* wild-type (Gd11) and three independent ppcesa4/6/7/10KO lines were cultured on medium containing sucrose to test for caulonema gravitropism. The plates were vertically oriented and kept in the dark. (A-C) Dark field images of wild-type (Gd11) colonies with upright growing caulonema filaments. (D-F) Dark field images of ppcesa4/6/7/10KO1, ppcesa4/6/7/10KO2, and ppcesa4/6/7/10KO3 colonies with upright growing caulonema filaments. Wild-type line and knockout lines were from the same experiment. (G) None of the three ppcesa4/6/7/10KO lines showed significant differences in caulonema length (ANOVA, n=3, P>0.05) or gravitropic behavior.
Manuscript 3

Manuscript formatted for the publication in *New Phytologist*

Short title: *Physcomitrella* has hetero-oligomeric Cellulose Synthesis Complexes

Title: A hetero-oligomeric Cellulose Synthesis Complex (CSC) involved in the secondary cell wall deposition in *Physcomitrella patens*

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One sentence summary: Similar to seed plants, the moss *Physcomitrella patens* contains obligate hetero-oligomeric Cellulose Synthesis Complexes (CSCs) involved in cellulose synthesis in secondary cell walls, indicating that the structure and function of the cellulose production machinery in moss and seed plants are undergoing convergent evolution.

List of author contributions: A.W.R. conceived the project, and supervised experiments; X.L. designed and performed experiments, and analyzed the data; J. M. performed experiments; X.L. wrote the manuscript.

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Abstract

Cellulose synthesis is catalyzed by plasma membrane Cellulose Synthesis Complexes (CSCs) that have been visualized by freeze-fracture electron microscopy as rosette structures with 6-fold symmetry. In seed plants, CSCs are obligate hetero-oligomeric, consisting of three functionally distinct and non-interchangeable cellulose synthase (CESA) isoforms. *Physcomitrella patens* has rosette CSCs, but its seven CESAs are not members of the clades that comprise the functionally distinct subunits of the hetero-oligomeric seed plant CSCs. Double *ppcesa3/8KOs* and *ppcesa6/7KOs* have defects in secondary cell wall deposition in gametophore leaf midribs, which suggests that PpCESA3, PpCESA8, PpCESA6, and PpCESA7 are required for forming hetero-oligomeric CSCs in gametophores. PpCESA5 is required for primary cell wall deposition in gametophore buds, but it is not known whether other PpCESAs are required for this processes. Here, Real-Time quantitative PCR (RT-qPCR) analysis shows that expression of *PpCESA3, PpCESA8*, and *PpCESA7* are co-regulated. Based on western blot analysis of isolated proteins, PpCESA3, PpCESA8, and PpCESA7 are all highly expressed in gametophores. Co-immunoprecipitation (Co-IP) shows that PpCESA3 and PpCESA8 can both interact with PpCESA6/7 *in planta*. These results support the hypothesis that cellulose microfibrils in the secondary cell walls of *P. patens* leaf midribs are synthesized by obligate hetero-oligomeric CSCs.
Introduction

Cellulose is a biopolymer of β-(1,4)-glucose that forms microfibrils essential in land plant cell walls. It is synthesized by cellulose synthase complexes (CSCs) located on the plasma membrane (Delmer et al., 1999; Somerville et al., 2006; McFarlane et al., 2014). The CSCs of vascular plants were first observed to have a "rosette" structure and associate with the ends of microfibrils in freeze-fracture electron microscopy studies (Mueller & Brown, 1980). Within the CSC, cellulose synthase catalytic subunits (CESAs) catalyze the synthesis of individual glucan chains and are currently the only verified functional subunits (Delmer, 1999; Somerville, 2006; McFarlane et al., 2014; Purushotham et al., 2016).

CESA genes are members of multigene families in vascular plants. Arabidopsis has 10 CESA genes (McFarlane et al., 2014). AtCESA4, AtCESA7, and AtCESA8 were first identified to be specifically involved in secondary cell wall deposition (Turner & Somerville, 1997; Taylor et al., 1999; Scheible et al., 2001; Taylor et al., 2000; Taylor et al., 2003). Proteins encoded by these three genes physically interact and are exclusively required for assembly of CSCs in cells having thickened secondary walls (Taylor et al., 2000; Taylor et al., 2003). Mutations in AtCESA1, AtCESA3, and AtCESA6 cause primary cell wall defects (Arioli et al. 1998; Fagard et al., 2000; Burn et al., 2002; Robert et al., 2004). AtCESA3 and AtCESA6 interact with each other according to results of in vitro pull-down assays, and BiFC experiments show that AtCESA1, AtCESA3, and AtCESA6 can interact in vivo (Desprez et al., 2007). AtCESA2 and AtCESA5 are shown to be closely related and partially functionally redundant with AtCESA6 (Desprez et al. 2007; Timmers et al. 2009; Carroll et al.
2012; Li et al., 2013). In *Arabidopsis*, therefore, a primary wall CSC likely consists of AtCESA1, AtCESA3, and one of the AtCESA6 like AtCESAs (McFarlane et al., 2014). The stoichiometry of *Arabidopsis* CSCs has been recently been determined to be a 1:1:1 molecular ratio (Gonneau et al., 2014; Hill et al., 2014). CSCs have also been characterized in other vascular plant models, such as *Populus trichocarpa*. Two types of CSCs (one type contains PdxtCESA7A and PdxtCESA8B; the other one contains PdxtCESA1A and PdxtCESA3) were identified in the xylem of *Populus* by coimmunoprecipitation (Co-IP, Song et al., 2010a). Altogether, current evidence suggests that vascular plant CSCs are hetero-oligomeric and can assembly only when three specific functionally distinct CESA isoforms are present.

Rosette-type CSCs have also been observed in the moss *Physcomitrella patens*, a model nonvascular plant, by freeze-fracture electron microscopy (Roberts et al., 2012; Nixon et al., 2016). The *PpCESA* gene family has seven members clustered in two clades, A-clade (*PpCESA3*, *PpCESA5*, and *PpCESA8*) and B-clade (*PpCESA4*, *PpCESA6*, *PpCESA7*, and *PpCESA10*). Mosses and seed plants are derived from a common ancestor that had a single *CESA*, and *PpCESAs* are not orthologs of vascular plant *CESAs* according to phylogenetic analyses (Roberts & Bushoven, 2007). It is not yet known if the CSCs of *P. patens* are homo-oligomeric or if they have evolved a hetero-oligomeric state independently from seed plants.

In addition to *PpCESA* knockout (KO) and phenotypic analysis, examination of co-expression and protein-protein interaction can provide insight into the composition of the *P. patens* CSCs. Previous mutational analyses showed that *ppcesa5*KO is required for the development of the gametophore (Goss et al., 2012). *Ppcesa3/8*KOs were
recently shown to have substantially reduced cellulose levels in midrib secondary cell walls of the gametophore leaf (Norris et al., 2017). A double B-clade PpCESAs KO, ppcesa6/7KO, phenocopies the ppcesa3/8KO midrib phenotype (Norris et al., 2017). These results are consistent with the hypothesis that cellulose in the secondary cell walls is produced by hetero-oligomeric CSCs consisting of PpCESA3, PpCESA6, PpCESA7 and PpCESA8. Other double B-clade PpCESAs KOs, ppcesa4/10KOs, were shown to have morphological changes in colonies formed by tip growing protonema filaments (unpublished data). This result indicates PpCESA4 and PpCESA10 are involved in synthesis of cellulose in primary cell walls. Here, our RT-qPCR analysis revealed a co-regulated expression of three PpCESA genes (PpCESA3, 7, and 8), which is similar to the gene expression pattern of Arabidopsis CESAs that reside in the same hetero-oligomeric CSCs. Moreover, coordinated accumulation of the corresponding PpCESAs proteins and physical interactions identified among them further support the hypothesis that a type of obligate hetero-oligomeric CSC consisting of PpCESA3, PpCESA8, and PpCESA6/7 is involved in the secondary cell wall deposition of P. patens gametophore leaves.

**Results**

**Expression levels of PpCESAs in the knockout mutants by RT-qPCR**

Correlated expression is expected for PpCESAs that reside in the same CSC. To test this, we performed RT-qPCR on RNA extracted from leafy gametophores collected from Gd11, ppcesa3KO, ppcesa8KO, ppcesa3/8KO, ppcesa4/10KO, ppcesa6/7KO, and ppcesa4/6/7/10KO plants. PpCESA8 was significantly upregulated in ppcesa3KO
(Figure 1) consistent with the results of the previous RT-qPCR analysis (Norris et al., 2017). Both \( PpCESA3 \) and \( PpCESA8 \) were significantly downregulated in \( ppcesa6/7\text{KO} \) and \( ppcesa4/6/7/10\text{KO} \) when compared with their expression in the wild-type controls. In reciprocal experiments, \( PpCESA7 \) was significantly downregulated in \( ppcesa3/8\text{KO} \). Expression of \( PpCESA4 \) and \( PpCESA10 \) were not significantly downregulated in \( ppcesa3/8\text{KO} \). In addition, no significant change in expression of \( PpCESA4 \) or \( PpCESA10 \) was detected in \( ppcesa6/7\text{KO} \). Expression of \( PpCESA7 \) was also not changed in \( ppcesa4/10\text{KO} \). No significant change of \( PpCESA5 \) expression was seen in any of the KO mutants. Expression of the other \( PpCESAs \) in the gametophores of \( ppcesa5\text{KO} \) could not be examined since \( ppcesa5\text{KOs} \) do not produce normal leafy gametophores (Goss et al., 2012). The results here are consistent with the correlated expression of \( PpCESA3, PpCESA8, PpCESA6, \) and \( PpCESA7 \).

Characterization of the PpCESA antibodies

Monoclonal antibodies were generated for detection of \( PpCESA3, PpCESA8, \) and \( PpCESA6/7, \) and specificity was tested by western blot against microsomal protein fractions. For each antibody, a corresponding \( PpCESA \) overexpression line was used as a positive control, and the KO line was used as a negative control. Figure 2 (left panel) shows that anti-\( PpCESA3 \) recognizes a band that has the predicted size for a \( PpCESA \) (120 kD) in both \( Act1pro::3xHA-PpCESA3 \) and wild-type. The band was not detected in \( ppcesa3\text{KO} \). We were unable to design an antibody that distinguishes \( PpCESA6 \) and \( PpCESA7 \), which differ by three amino acids. Thus, we made an antibody that can recognize both. Anti-\( PpCESA6/7 \) detected a 120 kD band in \( Act1pro::3xHA-PpCESA7 \) and wild-type, but not in \( ppcesa6/7\text{KO} \) (Figure 2, middle panel).
The anti-PpCESA8 detected a 120 kD band in Act1pro::3xHA-PpCESA8 and Gd11. However, it also weakly detected band around 120KD in ppcesa8KO (Figure 2, right panel). When ppcesa3/8KO is used as a negative control, no band was detected. This indicates that anti-PpCESA8 has weak cross-reactivity with PpCESA3 in addition to detecting PpCESA8.

**Protein expression profiling of the PpCESAs**

We used western blotting to examine the protein expression patterns for PpCESA3, PpCESA8, and PpCESA6/7 at different developmental stages. Figure 3 shows that none of these proteins were detectable in the Day-6 wild-type cultures consisting of pure protonema. PpCESA3, PpCESA8, and PpCESA6/7 were detected in Day-10 cultures, which contain protonema, emerging gametophore buds, and young gametophores. Finally, much larger amounts of these three PpCESAs were detected in Day-21 cultures, which contain numerous fully developed leafy gametophores. This shows that PpCESA3, PpCESA8, and PpCESA6/7 exhibit similar expression profiles, with highest expression in the gametophores, consistent with roles in gametophore development.

**Interactions between the PpCESAs**

Based on the similarity of their mutant phenotypes, correlated gene expression, and protein expression profiles, we hypothesized that PpCESA3, PpCESA6, PpCESA7, and PpCESA8 physically interact with each other to form hetero-oligomeric complexes. To address this question, we carried out Co-IP experiments on detergent-solubilized protein extracts from the 15-day-old leafy gametophores of transgenic *P. patens lines* that expressed HA-tagged PpCESAs under the control of their native
promoters in their cognate mutant backgrounds. Complementation of the mutant phenotype was verified for *PpCESA8pro::HA-PpCESA8 (ppcesa6/7KO, Figure S2 and S3)*. However we could not verify complementation for *PpCESA3pro::HA-PpCESA3*, since we have not detected a phenotype for *ppcesa3KO*. For the *PpCESA3pro::HA-PpCESA3* line (*Figure 4A*), blotting with anti-PpCESA3 showed that the IP antibody (anti-HA) successfully precipitated HA-PpCESA3 from the lysate of *PpCESA3pro::HA-PpCESA3*. When blotted with anti-PpCESA6/7, the target proteins were found in the IP eluate indicating that PpCESA6 and/or PpCESA7 were co-precipitated with HA-PpCESA3. Likewise, PpCESA8 was also detected in the IP eluate by anti-PpCESA8, suggesting a co-precipitation with HA-PpCESA3. In the Co-IP assay for *PpCESA8pro::HA-PpCESA8* (*Figure 4B*), blotting with anti-PpCESA8 verified that anti-HA pulled down the HA-PpCESA8. PpCESA6 and PpCESA7 were also detected in the IP eluate indicating co-precipitation with the primary target. Interestingly, anti-PpCESA3 also detected a band in the eluate. Similar results were observed with the reciprocal experiment in which anti-HA was used to pull down HA-tagged PpCESA7 from the protein extracts of the *PpCESA7pro::HA-PpCESA7* transgenic line (*Figure 4C*). Again, HA-tagged PpCESA7 was precipitated, and PpCESA3 and PpCESA8 were co-precipitated as expected. For the control experiment, Co-IP was carried out for wild-type *P. patens* (Gd11), which does not produce HA-tagged proteins (*Figure 4D*). As a result, none of the PpCESAs were immuno-detected indicating the IP antibody is specific.
Discussion

Our results are consistent with the hypothesis that the *P. patens* CSCs that synthesize the midrib secondary cell wall are obligate hetero-oligomeric, with members from both clade A and clade B. This hypothesis was suggested by the similar mutant phenotypes of *ppcesa3/8KO* and *ppcesa6/7KO* showing decreased cellulose deposition in the midribs of the gametophore leaves (Norris et al., 2017). Here, it is further supported by 1) co-regulation of *PpCESA3*, *PpCESA8*, and *PpCESA7* gene expression (Figure 1), 2) coordinated accumulation of PpCESA3, PpCESA8, and PpCESA6/7 proteins in leafy gametophores (Figure 3), and 3) detection of physical interactions between PpCESA3, PpCESA8, and PpCESA6/7 by Co-IP (Figure 4).

*Arabidopsis* has 10 *CESA* genes that are specialized for primary and secondary cell wall synthesis (Taylor et al., 1999; Fagard et al., 2000; Scheible et al., 2001; McFarlane et al., 2014). Phylogenetic studies show that the *CESA* families in *P. patens* and *Arabidopsis* are similar in size (Yin et al., 2010; Carroll & Specht, 2011; Harholt et al., 2012) Thus, it is imaginable that the *P. patens* CSCs are also hetero-oligomeric (Roberts et al., 2012). However, because *P. patens* CESAs diversified and specialized independently from seed plant CESAs (Roberts & Bushoven, 2007), this hypothesis must be tested independently.

In *Arabidopsis*, the genes that encode CESA isoforms that function within the same CSCs are co-expressed (Persson et al., 2005; Brown et al., 2005; Manfield et al., 2006). Here we show that, similar to the *Arabidopsis* CESAs, the expressions of the *PpCESA3*, *PpCESA8*, and *PpCESA6/7* genes are also co-regulated (both *PpCESA3* and *PpCESA8* have down-regulated expression when *PpCESA6* and *PpCESA7* are
knocked out, and vice versa). One exception is up-regulation of \textit{PpCESA8} to compensate for the loss of \textit{PpCESA3} (Norris et al., 2017). Taking results of phenotype analysis into consideration, it was suggested that \textit{PpCESA3} and \textit{PpCESA8} have interchangeable functions and may compete for the same positions in the CSCs (Norris et al., 2017). In addition, our Co-IP results show that \textit{PpCESA3} is co-immunoprecipitated with \textit{PpCESA8} and vice versa (Figure 4A). \textit{PpCESA8} appears to be dominant over \textit{PpCESA3} (in the amount of protein) in wild-type \textit{P. patens}. This is based on the observation that \textit{ppcesa8KO} has an obvious reduction in the leaf midrib cellulose deposition, which is not shown in \textit{ppcesa3KO} (Norris et al., 2017).

In \textit{Arabidopsis}, loss of a single CESA usually is enough to cause either obvious morphological defects or even lethal developmental defects (Taylor et al., 2003; Persson et al., 2007), which means \textit{Arabidopsis} CESAs are functionally distinct. In contrast, the PpCESAs show less functional differentiation. With the exception of \textit{PpCESA5} (Goss et al., 2012), we have to knock out at least two \textit{PpCESAs} from the same clade to observe a strong phenotype. In \textit{P. patens}, CESAs within the same sub-clade (A-clade or B-clade) appear to be functionally interchangeable. The major functional differences might only exist between PpCESAs from different sub-clades. For instance, the mutant phenotype of \textit{ppcesa3/8KO}s can be rescued by expressing \textit{PpCESA8}, \textit{PpCESA3} or \textit{PpCESA5} under control of the \textit{PpCESA8} promoter, but cannot be rescued by expressing any of the clade-B \textit{PpCESAs} using the same \textit{PpCESA8} promoter (Norris et al., 2017). The "no gametophore" phenotype of \textit{ppcesa5KO}s can be rescued by the expression of \textit{PpCESA3} or \textit{PpCESA8} driven by the \textit{PpCESA5} promoter, but again, it cannot be rescued by expression of any of the clade-
B PpCESAs driven by the PpCESA5 promoter (Scavuzzo-Duggan et al., in revision). In contrast, the AtCESAs have very limited interchangeability, with partial rescue only of atcesa3 by AtCESA3pro::AtCESA7 and atcesa8 with AtCESA8pro::AtCESA1 (Carroll et al., 2012).

Taken together, including this study, the current evidence indicates that secondary cell wall in the moss P. patens are synthesized by an obligate hetero-oligomeric CSC assembled from PpCESAs from both clade A and clade B. Our findings combined with phylogenetic analysis (Roberts & Bushoven, 2006) suggest that hetero-oligomeric CSCs arose in both mosses and seed plants through independent evolution. Norris et al. (2017) showed that diversification of the CESAs happened independently through both subfunctionalization and neofunctionalization in mosses and vascular plants, and these events are associated with convergent evolution of secondary wall structure. This indicates that selection pressure favored cellulose-rich secondary cell walls for better mechanical support to colonize land in both lineages. Here, our results indicate that selection for secondary cell walls with specialized cellulose microfibril textures might have favored emergence of hetero-oligomeric CSCs through convergent evolution in different land plants. Alternatively, the independent evolution of hetero-oligomeric CSCs in the seed plant and moss lineages could be explained by constructive neutral evolution.

Characterization of the PpCESAs shows some consistency with the theory of constructive neutral evolution which can be used to explain the evolution of the hetero-oligomeric CSCs. According to the hypothesis, after ancestral gene duplication, simple and high-probability mutations are considered to be a sufficient cause leading
to the increased complexity of a multi-protein complex (Doolittle, 2012; Finnigan et al., 2012). Most of these mutations are thought to be insufficient to cause changes in protein biochemical output. However, mutations occurring at the protein-protein binding interface can cause the mutant proteins to lose the ability to interact with the others members in the complex. In that case, a hetero-oligomeric complex eventually might evolve by complementary loss of asymmetric interactions of certain protein subunits in the original homo-oligomeric complex (Doolittle, 2012; Finnigan et al., 2012). We have shown that ppces6/7KOs phenocopy ppcesa3/8KOs; both show defects in secondary cell walls (Norris et al., 2017). This suggests that clade-A PpCESAs and clade-B PpCESAs carry out non-overlapping functions after neo-functionalization of the common ancestral PpCESA. The distinct functions of these PpCESAs might be caused by mutations at the binding sites, according to the theory above. This assumption is further supported by the Co-IP results here together with results of yeast two hybrid assay showing PpCESA8 cannot interact with itself (unpublished data). In addition, the promoter-swap assays mentioned above suggest there is no major functional difference among the paralogues from the same PpCESA clade, which is also consistent with constructive neutral evolution. To continue to test this theory, more precise characterization of PpCESAs need to be carried out to identify the binding sites of these PpCESAs.

Our study also indicate a possibility that PpCESA5 can form homo-oligomeric CSCs (Li et al., unpublished). We propose this hypothesis based on the distinct isoform function (Goss et al., 2012), unique gene expression pattern as well as the fact that mosses and seed plants derived from the common ancestor which had a single CESA
and consequently homo-oligomeric CSCs (Roberts & Bushoven, 2007). If this hypothesis is proved to be true, it will be another evidence supporting the constructive neutral evolution. But, no matter what the answers will be, implications provided by these studies will be helpful for understanding the roles of the different CESAs composing seed plant CSCs.

**Materials and methods**

**Culture conditions**

Wild-type and all transgenic *P. patens* lines used in this study were maintained on BCDAT plates and propagated by subculturing weekly as described (Roberts et al., 2011). To produce growing leafy gametophores, explants of 7-day-old protonemal tissue was transferred onto BCD plates and cultured for 15 days before being harvested for experiments.

**Vector construction and transformation**

All the PpCESA KO lines used in this study were created previously. Construction of knockout vectors and transformations for making those lines were described in Norris et al. (2017). Transformation for making quadruple ppcesa4/6/7/10 KOs was described in an unpublished manuscript (Li et al., unpublished).

PpCESA overexpression lines used as positive controls for testing antibody specificity were selected from transformations of ppcesa5KO-2 with vectors driving expression of 3X-HA-tagged PpCESA3, PpCESA7 or PpCESA8 under control of the rice Actin1 promoter (Scavuzzo-Duggan et al., in revision).
Expression vectors for HA-tagged *PpCESAs* under control of their native promoters were created using Multi Site Gateway Pro (Invitrogen, Grand Island, NY, USA). The *HA-PpCESA5*, *HA-PpCESA7*, and *HA-PpCESA8* coding sequences were amplified from cDNA clones pdp24095, pdp38142, and pdp39044 (RIKEN BRC), respectively, using forward primers containing a single HA tag coding sequence flanked by an attB5 site and a reverse primer flanked by an attB2 site (Supplemental Table 2). *HA-PpCESA3* was amplified from a cDNA clone describe previously (Scavuzzo-Duggan et al, in revision) using appropriate primers (Supplemental Table 2). PCR amplification was catalyzed by Phusion High-Fidelity PCR Master Mix (New England Biolab) in 50 L reactions with an initial denaturation (98°C, 30 sec); 35 cycles of denaturation (98°C, 7 sec), annealing (68°C, 15 sec), and extension (72°C, 40 sec); and a final extension (72°C, 5 min) and the products were cloned into pDONR p5-p2 vector (Invitrogen) to create entry clones. To create the *PpCESA*promoter::*HA-PpCESA* vectors, the HA-CESA entry clones along with an entry clone containing the corresponding native promoter (Tran & Roberts, 2016) were inserted into the Si3-pTH-GW destination vector (Tran & Roberts, 2016) using LR Clonase II Plus (Invitrogen). *PpCESA3pro::HA-PpCESA3* and *PpCESA5pro::HA-PpCESA5* were linearized with SwaI. *PpCESA8pro::HA-PpCESA8* was linearized with PciI.

RNA extraction and Real-time quantitative PCR (RT-qPCR)

Total RNA extraction from gametophore leaves followed by cDNA conversion was carried out as described (Tran & Roberts, 2016). RT-qPCR analysis was performed using the ΔΔCt method (Livak & Schmittgen, 2001) of relative quantification with a Roche Lightcycler 480, using SYBR Green I Master Mix (Roche) to monitor doubled
strand DNA synthesis. Primers for *PpCESA* detection were as used in Tran & Roberts (2016), and primers for reference genes, *actin* and *v-Type h+ translocating pyrophosphatase*, were as used in Bail et al. (2013).

**Generation of monoclonal anti-PpCESAs**

Peptide antigens, designed to regions of each PpCESA for the purpose of raising antibodies specific for each isoform (Table S2), were synthesized chemically and injected into New Zealand white rabbits (Covance Inc., Princeton NJ USA). For PpCESA6 and PpCESA7, it was not possible to generate two unique peptides in order to raise antibodies to differentiate these isoforms. The peptides were conjugated, via the cysteine residue, to Sulfolink Immobilization resin (Thermo Fisher Scientific) according to the manufacturer’s instructions. The purification of PpCESA antibodies from total serum was carried out by affinity chromatography. Briefly, 10 mL of serum, buffered with WB (20 mM NaHPO₄, pH7.2, 50 mM NaCl) was incubated with the resin-linked peptides for 18h at 4°C. The resin was loaded into a column and the flow through was passed over the resin twice. The resin was washed with 20 mL of WB followed by 10 mL of WB containing an additional 250 mM NaCl. Antibodies were eluted from the resin using 5 mL of EB (100 mM glycine, pH 2.5). Fractions of 250 μL containing NB (50 μL 1 M Tris-Cl, pH 8.0) were collected and mixed immediately to neutralize pH. Fractions containing PpCESA antibodies were identified by absorbance at 280 nm and combined. Glycerol was added to 30% and CESA antibodies were stored at -80°C. The specificity of each antibody was tested by western blotting against *P. patens* protein extracts.
Protein expression profiling of the PpCESAs

The wild type *P. patens* were used in this assay. The 6-day-old protonema growing on BCD plates were collected for protein extraction of the first time point. The "spot plates" were set up by transferring additional protonema at the day 7 onto BCD media to growing for gametophores. The tissues on the "spot plates" were collected at the day 10 for protein extraction of the second time point. Later, the "spot plate" tissues were collected again at the day 21 and used for protein extraction of the last time point. Microsomal protein isolation and western blot analysis were proceeded as described in the Scavuzzo-Duggan et al. (2015).

Co-immunoprecipitation

Co-immunoprecipitation (Co-IP) experiments were performed according to the method from previous studies with some modifications (Desprez et al., 2007; Song, Shen, & Li, 2010b). Squeeze-dried 15-day-old moss tissue (0.5 g) containing mostly leafy gametophores was ground in liquid nitrogen, and the powder was transferred to a 2 mL centrifuge tube with 1 mL of ice-cold IP buffer [20mM Tris.HCl, pH7.5; 150mM NaCl; 5mM MgCl₂; 10% sucrose; 1% glycerol; 1mM EDTA; 1.5% CHAPS (Sigma, St. Louis, MO, USA); 1% protease inhibitor cocktail (Sigma, P9599); 1% phosphatase inhibitor mixture 2 (Sigma, P5726), and 3 (Sigma, P0044); and 1% polyvinylpyrrolidone]. The tube was incubated on ice for 30 min and centrifuged at 20,000 x g for 30 min to pellet insoluble debris. The supernatant was transferred to a new 1.5 mL centrifuge tube with 25 μL of Pierce Anti-HA Magnetic Beads (Thermo Fisher Scientific, Waltham MA USA) and rotated (8 RPM) for 50 min on an end-over-end rotator (Thermo Fisher Scientific). Beads were then collected with a magnetic
stand (Thermo Fisher Scientific), and the unbound sample was removed. 400 µL of TBS-T buffer (Scavuzzo-Duggan et al., 2015) was added to the tube and gently mixed. Beads were collected again by magnetic stand, and the supernatant was discarded. This step was repeated twice. For the last wash, 400 µL of ultrapure water was added to the tube and gently mixed. Beads were collected on a magnetic stand, and the supernatant was removed. For elution, 50 µL of 2X SDS-PAGE sample buffer (Scavuzzo-Duggan et al., 2015) and 50 µL of ultrapure water were added to the tube, and gently mixed. The tube was incubated at 95°C-100°C for 10 min. Finally, beads were magnetically separated, and initial input (total protein), unbound fraction, wash, and IP eluate were stored at -20°C for up to three months and used for western blot analysis. Gel electrophoresis and western blot using anti-PpCESA3, anti-PpCESA6/7, and anti-PpCESA8 antibodies were carried out as described in Scavuzzo-Duggan et al. (2015).

Statistical analysis

One-way Analysis of Variance (ANOVA) followed by post-hoc Tukey Honest Significant Difference (HSD) test was performed using "R" programming (Vienna, Austria; http://www.R-project.org/) to identify the potential significant difference in each assay.
Supplemental Materials

Table. S1. Primers for amplification of HA-tagged *PpCESAs*.

Table. S2. Peptide antigens, designed to regions of each PpCESA, used to raise specific antibodies for each PpCESA isoform.

Fig. S1: RT-qPCR analysis of *PpCESA* expression in the KO mutants.

Fig. S2. Quantitative analysis of S4B fluorescence intensity in leaf midribs of *P. patens* wild-type (Gd11), *ppcesa8*KO-lox16 (*cesa8*KO), and *PpCESA8pro::HA-PpCESA8* (HA-CESA8).

Fig. S3. Quantitative analysis of S4B fluorescence intensity in leaf midribs of *P. patens* wild-type (Gd11), *ppcesa6/7*KO-lox23 (6/7KO), and *PpCESA7pro::HA-PpCES7* (HA-CESA7).

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### Figures

|                | CESA3 | CESA5 | CESA8 | CESA4 | CESA7 | CESA10 |
|----------------|-------|-------|-------|-------|-------|--------|
| 3KO            | NA    | NA    | **Red** | NA    | NA    | NA     |
| 8KO            | NA    | NA    | NA    | NA    | NA    | NA     |
| 3/8KO          | NA    | NA    | NA    | NA    | **Green** | **Green** | NA     |
| 4/10KO         | NA    | NA    | NA    | NA    | NA    | NA     |
| 6/7KO          | **Green** | **Green** | **Green** | NA    | NA    | NA     |
| 4/6/7/10KO     | NA    | NA    | NA    | NA    | NA    | NA     |

**Figure 1: Heat map of *PpCESA* expression in *PpCESA* knockout (KO) mutants.** Summary of results from RT-qPCR analysis of RNA isolated from 21-day old cultures of *PpCESA* KO lines normalized to *PpACT* and *PpVHP* reference genes. Three independent lines of each genotype were tested with two technical replicates. Green=significant down-regulation, *p*<0.05; gray=non-significant differences, *p*>0.05; red=significant up-regulation, *p*<0.05; NA: no expression). Graphs are shown in supplementary figure 1.
Figure 2: Antibody specificity test. Western blots of microsomal protein extracts from HA-tagged PpCESA overexpression lines (positive control), PpCESA knock out (KO) lines (negative control), and wild-type probed with anti-PpCESA3, anti-PpCESA6/7, and anti-PpCESA8 respectively. Molecular mass markers are given at left in kilodaltons. Black arrows indicate expected position of target bands (~120KD) detected by antibodies. Faint band in 8KO lane, but not 3/8KO line, indicates weak cross reactivity of anti-PpCESA8 with PpCESA3.
Figure 3: PpCESA protein expression in wild-type *P. patens*. Western blots of microsomal proteins isolated from wild-type *P. patens* cultures and probed with anti-PpCESA3, anti-PpCESA8, and anti-PpCESA6/7. Explants from protonema cultured on solid medium overlaid with cellophane for 6 days were cultured on solid medium without cellophane and harvested after 6 days (protonema only), 10 days (protonema and young gametophores) and 21 days (gametophores). Equal loading of protein (XXX g per lane) was verified by Ponceau S Staining.
Figure 4: Co-immunoprecipitation (Co-IP) of PpCESAs. Western blots of total protein lysates from lines expressing HA-PpCESAs under control of their native promoters (A-C) and wild-type (D) with unbound, wash and eluate from immunoprecipitation with anti-HA. Blots were probed with antibodies listed on the right of each graph.
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## Supplemental Materials

Table S1. Primers for amplification of HA-tagged *PpCESAs*.

| Name                  | Sequence                                                                 | Tm (°C) | Amplicon Size (bp) | Description  |
|-----------------------|---------------------------------------------------------------------------|---------|--------------------|--------------|
| *HAPpCESA8*attB5      | GGGGACAACCTTTGTATACAAAAAGTGGCATGGGATACCCATACGATGTTCCAGATTACGCTATGGAGGCTAATGCGG | 76°C    | 3373               | HA-*PpCESA3* |
| *PpCESA3*attB2        | GGGGACCACCTTTGTACAAGAAAGCTGGTGAGAGCTTTATTATAGGTCG                          | 79°C    | 3378               | HA-*PpCESA7* |
| *HAPpCESA7*attB5      | GGGGACAACCTTTGTATACAAAAAGTGGCATGGGATACCCATACGATGTTCCAGATTACGCTATGGAGGCGAATGCGG | 79°C    | 3378               | HA-*PpCESA7* |
| *PpCESA7*attB2        | GGGGACAACCTTTGTACAAGAAAGCTGGTGAGAGCTTTATCCGACATTGCGA                      | 77°C    | 3370               | HA-*PpCESA8* |
| *HAPpCESA8*attB5      | See above                                                                |         |                    |              |
| *PpCESA8*attB2        | GGGGACAACCTTTGTACAAGAAAGCTGGTGAGAGCTTTATCCGGGCGACATTGCGA                 |         |                    |              |
Table S2. Peptide antigens, designed to regions of each PpCESA, used to raise specific antibodies for each PpCESA isoform.

| Peptide Antigen | Sequence                      |
|-----------------|-------------------------------|
| PpCESA3         | CPDHDQEKS$\uptheta$ILSTKDIEKR |
| PpCESA8         | CLDHYEKSSPIMSTKDIEKR          |
| PpCESA6/7       | CVIRQESDGP$\uptheta$LSN     |
Figure S1: RT-qPCR analysis of \textit{PpCESA} expression in the KO mutants. (A) \textit{PpCESA} expression levels relative to two reference genes (PpACT and PpVHP) in gametophore leaves of three clade-A \textit{PpCESA} KO mutants, \textit{ppcesa3KO}, \textit{ppcesa8KO}, and \textit{ppcesa3/8KO}. (B) The relative expression of \textit{PpCESAs} in the 21-day-old gametophore leaves from the three clade-B \textit{PpCESA} KO mutants, \textit{ppcesa4/10KO}, \textit{ppcesa6/7KO}, and quadruple \textit{ppcesa4/6/7/10KO} (QKO). Numbers on the y-axis represent relative transcript level. \textit{P. patens} lines used as a source of RNA are labeled along the x-axis. Error bars indicate standard error of the mean (\(n=3\)). “*”: p<0.05. Gd11 (wild type): Yellow bar; Clade-A \textit{PpCESA} KO: Blue bar; Clade-B \textit{PpCESA} KO: Gray bar.
Figure S2: Quantitative analysis of S4B fluorescence intensity in leaf midribs of *P. patens* wild-type (Gd11), *ppcesa8*KO-lox16 (8KO), and *PpCESA8pro::HA-PpCESA8* (HA-CESA8). Fluorescence intensity is significantly lower in *ppcesa8*KO-lox16 compared to wild-type and *PpCESA8pro::HA-PpCESA8*. The *PpCESA8pro::HA-PpCESA8* line was created by transforming the *ppcesa8*KO-lox16 line with the vector driving expression of HA-tagged PpCESA8 using *PpCESA8* native promoter. *PpCESA8pro::HA-PpCESA8* is not significantly different from wild-type indicating that the HA-tagged PpCESA8 successfully restored the function of the native PpCESA8.
Figure S3: Quantitative analysis of S4B fluorescence intensity in leaf midribs of *P. patens* wild-type (Gd11), *ppcesa6/*7KO-lox23 (6/7KO), and *PpCESA7pro::HA-PpCES7* (HA-CESA7). Fluorescence intensity of wild-type is the highest among the three. *Ppcesa7KO-lox23* has the lowest fluorescence intensity. *PpCESA7pro::HA-PpCES7* has an intermediate fluorescence intensity but significantly higher than the intensity of *ppcesa6/*7KO, indicating at least partial rescue by introducing the HA-tagged PpCESA7. The *PpCESA7pro::HA-PpCES7* line was created by transforming the *ppcesa7KO-lox23* line with the vector driving expression of HA-tagged PpCESA7 using *PpCESA7* native promoter.