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Review

A historical perspective on the regulation of cellulose biosynthesis



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ABSTRACT

Cellulose is a β -1,4 linked glucose polymer that is synthesized by higher plants, algae and even by some bacteria and animals, making it the most abundant polymer on earth. As the major load bearing structure of the plant cell wall, it is hugely important in terms of plant growth and development, and in recent years it has gained interest for its biotechnological applications. Naturally, there has been a large concerted research effort to uncover the regulatory mechanisms underpinning cellulose synthesis. During the last century, several major breakthroughs in our understanding of cellulose synthesis in algae, bacteria, and plants have been pivotal in advancing the field of cellulose research, improving the likelihood that cellulose synthesis could be feasibly adapted for sustainable purposes. In this review, we will summarize the major hypotheses and advancements made during the last century on the regulation of cellulose biosynthesis, focussing on *Arabidopsis thaliana*.

1. Introduction

For centuries, cellulose has been widely recognized in terms of its economic potential and biological influence. Cellulose is an essential multi-purpose resource that is heavily used in construction, paper manufacturing, textile production, and as a source of fuel. More recently, cellulose has been recognized as a potential feedstock for renewable biofuels and other sustainable products. All plant cells deposit cell walls that contain cellulose. As a result, cellulose is the most abundant organic polymer on earth, contributing between 150-170 billion tons of carbon to the biosphere per year through carbon sequestration (Engelhardt, 1995). During growth, plant cells develop a primary cell wall that consists of three main polymers: cellulose, hemicellulose (typically xyloglucan), and pectin, contained within an aqueous matrix (Cosgrove & Jarvis, 2012). Once cells cease expanding, specialized cell types can deposit a thicker, stronger secondary cell wall that is reinforced by the hydrophobic polymer lignin. Cellulose tends to be more abundant in secondary cell walls that are comprised of up to 50 % cellulose (Meents, Watanabe, & Samuels, 2018).

Despite its huge importance, cellulose research consisted of a relatively finite, insignificant field a hundred years ago. Since the 1950s, several major breakthroughs in our understanding of cellulose synthesis and regulation have turned this on its head and it is now a thriving field of research. Studies historically focussed on characterizing the structure of cellulose microfibrils and synthetic mechanisms in cellulose-rich unicellular organisms, including the green algae, *Valonia* and *Oocystis*

and the bacteria Acetobacter xylinum. In addition, fibers from Gossypium (Cotton) and Boehmeria (Ramie) were also used. Ground-breaking findings from these organisms were applied to higher plants on the basis that the intrinsic properties of cellulose were shared, igniting the study of cellulose synthesis in higher plants. The development of the herbaceous species, Arabidopsis thaliana, as a model plant for genetic research in the 1980s caused a noticeable shift from studying the biophysical aspects of cellulose to a genetic and cell biology led approach, especially regarding the dynamics of the cellulose synthase complex (CSC). In Arabidopsis, significant advancements have been somewhat restricted to the primary cell wall, since Arabidopsis undergoes limited secondary growth, though some notable contributions have been made (Strabala & Macmillan, 2013). Poplar became a genetic model for studying cellulose in secondary cell walls and is frequently used to validate assumptions made from Arabidopsis, and other species, regarding secondary cell wall formation. Poplar can also produce a gelatinous, 'G-layer' that is composed almost entirely of cellulose during tension wood formation, which has greatly supplemented studies of cellulose synthesis (Felten & Sundberg, 2013).

The increasing availability of biological tools combined with the development of highly sensitive techniques have been largely responsible for the significant progress made in the study of cellulose synthesis in higher plants. Together these have confirmed many of the original hypotheses made and answered, at least partially, many of the outstanding questions regarding cellulose synthesis. In this review, we will focus on how our understanding of the regulation of cellulose

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synthesis has developed during the last century, with particular focus on i) how cellulose is synthesized?; ii) when it is synthesized?; iii) CSC trafficking; and iv) how it is regulated? We will cover the main hypotheses regarding cellulose synthesis, and the significant advancements that have been made to support these in *Arabidopsis*, though contributions from other species will be included where relevant. We regret that due to space limitations we cannot cover every aspect of cellulose synthesis, but some excellent reviews are widely available (Brown & Saxena, 2000; Delmer, 1999; Guerriero, Fugelstad, & Bulone, 2010; Haigler & Roberts, 2018; Lampugnani et al., 2019; Li, Bashline, Lei, & Gu, 2014; Somerville, 2006; Wolf, Hematy, & Hofte, 2012).

2. How is cellulose synthesized?

The long-standing use of cellulose as a feedstock for the pulping and energy industries naturally called for a more thorough understanding of the physical structure, biochemistry, and synthesis of microfibrils. Early studies on the physical aspects of cell walls and cellulose crystallinity relied on a combination of polarizing microscopy, transmission electron microscopy (TEM) and X-ray diffraction analysis of algae and bacteria. Even with the limited resources available, many of these assumptions were held to be true in higher plants when they were later reinforced by genetics.

2.1. Structure of cellulose

While cellulose was first described as a polymer in the 1920s, the crystal structure of cellulose was not resolved until fifty years later. X-ray diffraction analysis of cellulose from ramie fibers and the algae, *Valonia ventricosa*, revealed that cellulose is a crystalline β -1,4 linked glucose polymer (Gardner & Blackwell, 1974). More specifically, cellulose is a two-fold helical structure of alternating cellobiose units, as β -1,4 glycosidic linkages require a 180° rotation of consecutive molecules (Hermans, de Booys, & Maan, 1943). The 120° rotation of β -1,4 bonds is thought to facilitate the inversion of glucose molecules during synthesis (Delmer, 1999). Multiple isoforms of cellulose exist (I—IV), although the most labile form, cellulose I, is produced almost exclusively in nature (Delmer, 1999). Physical and chemical deformations of cellulose I can produce cellulose II—IV isoforms that are inherently more stable.

The idea of the microfibril was first coined by Preston, Nicolai, and Millard (1948) who observed in electron micrographs and X-ray diffraction analyses of V. ventricosa that cellulose consists of multiple glucan chains bound together. Characterizing the structure of cellulose I was initially problematic and diverse X-ray diffraction patterns of cellulose I were reported amongst research groups (Preston, 1974). Assessing the structure more intricately with solid-state nuclear magnetic resonance (ssNMR) revealed that cellulose I exists as two distinct forms, Iα and Iβ, (Atalla & Vanderhart, 1984). Cellulose Iα exhibits a triclinic structure composed of one chain and cellulose Iβ contains two parallel chains within a monoclinic structure. Cellulose Iα microfibrils predominate in algae and bacteria, whereas in higher plants and tunicates, cellulose I_β- tends to be more abundant, though microfibrils are often comprised of both forms. Within the microfibril, parallel glucan chains are stabilized by intra- and inter-molecular hydrogen bonds. Hydrophobic van der Waals forces can also form between glucan sheets, particularly in aqueous environments (Cousins & Brown, 1997) and so are perhaps more relevant in primary cell walls that have a high water content. In secondary cell walls, cellulose is held together by a higher degree of intra-molecular hydrogen bonding, creating a rigid, crystalline polymer that invokes strong structural support to the cell. In tension wood, cellulose is almost purely crystalline, which is likely related to the production of wood under high tensile stress (Foston et al., 2011).

The properties of cellulose can be measured in terms of its crystallinity; width; degree of polymerization; and cross-sectional shape, to name a few variables. Unsurprisingly, considerable variation in cellulosic properties exists between species, cell types, and even within the microfibril itself. Variations in the width of cellulose microfibrils have been interpreted as differences in the number of glucan chains, the extent of bundling and interactions with non-cellulosic polysaccharides. Measuring microfibrils with a diversity of techniques, including atomic force microscopy (AFM), small-angle neutron scattering (SANS) and wide-angle X-ray scattering (WAXS), have demonstrated that individual microfibrils are consistently 3-4 nm wide, across different species and cell wall types (Fernandes et al., 2011; Song, Zhao, Shen, Collings, & Ding, 2020; Zhang, Zheng, & Cosgrove, 2016). Close microfibril spacing can cause neighbouring microfibrils to associate into larger bundles, that span up to 50 nm in width in secondary cell walls (Fernandes et al., 2011; Song et al., 2020; Thomas et al., 2013; Zhang, Zheng et al., 2016). Detailed examination of microfibrils with AFM has revealed the sheer extent of bundling, particularly in onion primary cell wells where up to ²/₃ of the microfibril length coalesces with other microfibrils (Zhang, Zheng et al., 2016). Wide cellulose microfibrils also tend to accompany a higher degree of polymerization (DP). In primary cell walls, cellulose DP can range from 500-8,000 and in cotton secondary cell walls, cellulose DP can exceed 15,000 (Brett, 2000). Far longer microfibrils of up to 23, 000 DP have been observed in algae that secrete cellulose outside of the cell (Brown, 1996), indicating that microfibril elongation may be partially restricted by the biophysical and spatial constraints of cell walls. An important caveat is that these estimates of cellulose DP have not yet been verified in the intact cell wall, and so may not be representative of true microfibril DPs. The biological significance of DP and what triggers the termination of chain elongation is unknown, but chain length is likely to be an important determinant of cell wall function and architecture (Somerville, 2006).

Uncovering the structure of cellulose microfibrils formed the building block for all future studies on cellulose, as it can be used as a tool to make logical inferences about the underlying synthetic mechanisms and architecture of cell walls. In particular, the width and cross-sectional shape of cellulose microfibrils has been used to predict the size and arrangement of synthetic complexes and the orientation of microfibrils has informed models of cell expansion. While it has been firmly established that these features of cellulose are highly influential, how many of these cellulosic properties are determined remains an open question.

2.2. Cellulose synthase complexes (CSCs) - Structure

Once the structure of cellulose microfibrils was largely characterized, the next main focus was to identify the protein complex responsible for cellulose synthesis. Uncovering the structure of the synthetic complex was a major breakthrough in cellulose research (Table 1). Twenty years after Roelofsen (1958) correctly predicted that microfibrils extend from the growing tip by large enzyme complexes, linear structures matching that description were identified in the plasma membrane of the algae, *Oocystis apiculate* by freeze-fracture TEM (Brown & Montezinos, 1976). As they were situated at the base of microfibril imprints they were referred to as 'terminal complexes'. This was arguably the first indication that cellulose synthesis was highly distinct from other polysaccharides that are synthesized in the Golgi, emphasizing that the production of cellulose in such close proximity to the cell wall has functional significance.

Terminal complexes were subsequently identified in a whole host of different species, including bacteria, higher plants and tunicates (Kimura & Itoh, 1996). However, they were not uniform in their shape, abundance, or organization (Tsekos, 1999). In contrast to the linear complexes described in *O. apiculate*, freeze-fractured membranes of maize and pine seedlings revealed that terminal complexes in higher plants consist of rosette-shaped particles with six-fold symmetry (Haigler & Brown, 1986; Mueller & Brown, 1980). Re-examination of rosettes in the moss, *Physcomitrella patens*, has suggested that rosette lobes can be triangular and the six-fold symmetry can be lost due to unequal spacing between lobes (Nixon et al., 2016). Sin tobacco BY-2 cells the

Table 1A summary of the major hypotheses made regarding the regulation of cellulose synthesis.

| Original hypothesis | Studies | Current status | Studies |
|---|--|--|--|
| Cellulose is synthesized by a terminal complex in the plasma membrane | Observational (Roelofsen, 1958) | Widely accepted | Freeze-fracture TEM of <i>Oocystis apiculate</i> , maize and pine (Brown & Montezinos, 1976; Haigler & Brown, 1986; Mueller & Brown, 1980) |
| 2. Terminal complex arrangement facilitates the coalescence of glucan chains | TEM of green algae cell walls (Giddings et al., 1980; Herth, 1983) | Partially confirmed: Conflicting evidence | Live-cell imaging of <i>Arabidopsis</i> cell walls and <i>in vitro</i> studies of cellulose synthesis (Cho et al., 2017; Li et al., 2016; Purushotham et al., 2016; Watanabe et al., 2015) |
| 3. Cellulose is synthesized from terminal complexes | Sequencing analysis of the <i>Bcs</i> operon in <i>Acetobacter xylinum</i> and <i>CESA</i> genes in cotton (Pear et al., 1996; Saxena et al., 1990; Wong et al., 1990) | Widely accepted | Immunolabelling of CESA proteins in freeze fractured azuki bean complexes (Kimura et al., 1999) |
| 4. CSCs are composed of a 'hexamer of hexamers' that synthesize cellulose microfibrils containing 36 chains | Hypothesis based on TEM structure (Herth, 1983) | Modified: Microfibrils consist of 18–24 chains. More evidence suggests the CSC is a hexamer of trimers, synthesizing an 18-chain microfibril | Physical studies of microfibril widths in mung bean, freeze fracture of <i>Physcomitrella patens</i> , stoichiometry of <i>Arabidopsis</i> and <i>in vitro</i> trimer formation in poplar (Gonneau et al., 2014; Hill et al., 2014; Newman & Hemmingson, 1990; Nixon et al., 2016; Vandavasi et al., 2016) |
| 5. Each CESA protein synthesizes one glucan chain | CSC crystallography in <i>Rhodobacter</i> sphaeroides (Morgan et al., 2013) | Recent evidence:In <i>PttCesa8</i> homotrimers, each CESA particle associates with a single glucan molecule | Cryogenic-EM structure of <i>PttCesa8</i> homotrimers produced <i>in vitro</i> (Purushotham et al., 2020) |
| 6. Three distinct CESA isoforms are required for cellulose synthesis | CESA mutant analysis in <i>Arabidopsis</i> (Desprez et al., 2007; Taylor et al., 2003) | <u>Modified</u> : Some cross-over between isoforms | Functional complementation in <i>Arabidopsis</i> (Carroll et al., 2012; Li et al., 2013) |
| 7. CESA proteins have 8 transmembrane domains | Sequencing analysis of <i>A. xylinum</i> and cotton (Pear et al., 1996; Saxena et al., 1990; Wong et al., 1990) | Modified: CESA proteins have 7 transmembrane domains | Mutational analysis and functional complementation in <i>Arabidopsis</i> and <i>P. patens</i> and structural analysis of <i>PttCESA8</i> homotrimers (Slabaugh et al., 2016; Purushotham et al., 2020) |
| CESA transmembrane domains form a channel for glucan chain release | CSC crystallography in <i>R.sphaeroides</i> (Morgan et al., 2013) | Recent evidence:In <i>PttCesa8</i> homotrimers, the transmembrane domains of each CESA particle forms a channel | Cryogenic-EM structure of <i>PttCesa8</i> homotrimers produced <i>in vitro</i> (Purushotham et al., 2020) |
| Cellulose microfibrils are extended by the stepwise addition of glucose | CSC crystallography in R. RRR. sphaeroides (Morgan et al., 2013) | Not confirmed in plants | |
| 10. Microfibrils are simultaneously crystallized and polymerized | Calcofluor white interference in A. xylinum (Benziman et al., 1980) | Accepted with limited further study | |
| 11. The rosette structure promotes the crystallization of glucan chains | Hypothesis based on TEM structure (Herth, 1983) | Accepted on little empirical evidence | Mutational studies in <i>Arabidopsis</i> and poplar (Arioli et al., 1998; Harris et al., 2012; Purushotham et al., 2016) |
| 12. Polymerization drives CSC movement | Observational (Herth, 1983) | Widely accepted | Live-cell imaging and biophysical modelling in Arabidopsis (Diotallevi & Mulder, 2007; Paredez et al., 2006) |
| 13. Multi-net growth hypothesis | TEM of Nitella and Tradescantia (Roelofsen & Houwink, 1951) | Not universally accepted: Cannot explain anisotropy in all tissue types | AFM and FESEM in <i>Arabidopsis</i> (Marga, Grandbois, Cosgrove, & Baskin, 2005; Wiedemeier et al., 2002; Xin et al., 2020) |
| 14. Alignment hypothesis | TEM and polarizing microscopy (Green, 1962; Ledbetter & Porter, 1963) | <u>Partially confirmed:</u> Not representative of all tissue types | Live-cell imaging in <i>Arabidopsis</i> (Himmelspach et al., 2003; Paredez et al., 2006; Sugimoto et al., 2003) |
| 15. Direct-guidance model | TEM and live-cell imaging in <i>Arabidopsis</i> (Heath, 1974; Paredez et al., 2006) | <u>Updated</u> : CSI1/POM2 links CSCs with microtubules in primary cell walls | Y2H and csi1/pom2 mutant analysis in Arabidopsis (Bringmann et al., 2012; Gu et al., 2010) |
| CSCs are assembled in the Golgi Microtubules define CSC | TEM and freeze fracture of Zinnia elegans (Haigler & Brown, 1986) TEM of Coleus and Z. elegans (Haigler & | In question: ER assembly has been proposed but evidence is scarce Updated: Microtubules coincide with Golgi | Live-cell imaging in <i>Arabidopsis</i> (Gardiner et al., 2003; Paredez et al., 2006; Park et al., 2019) Live-cell imaging in <i>Arabidopsis</i> (Crowell et al., |
| delivery | Brown, 1986; Hepler & Newcomb, 1964) | pausing events and the insertion of SmaCC/ MASCs | 2009; Gutierrez et al., 2009) |
| 18. CSCs are recycled | Live-cell imaging and mutagenesis in <i>Arabidopsis</i> (Bashline et al., 2013, 2015) | Partially confirmed: Evidence of CME, but it is not known if they are recycled | |

transmembrane region spans 25 nm, similar to measurements made by Mueller and Brown (1980), and the cytosolic region is twice as wide, ranging between 45–50 nm (Bowling & Brown, 2008).

Many researchers have repeatedly suggested that differences in the morphology of terminal complexes may be responsible for the diversity in microfibril architecture. In particular, the arrangement of terminal complexes has been linked with the extent of crystallization and microfibril bundling in different types of cell wall (Tsekos, 1999) (Table 1). Octagonal arrays and linear rows of rosette complexes in the secondary cell walls of *Micrasteria denticulata* and *Spirogyra* respectively, produce microfibril bundles consisting of more glucan chains than a single rosette in primary cell walls (Giddings, Brower, & Staehelin, 1980Herth, 1983). The closer arrangement of multiple terminal complexes in secondary cell walls may be necessary to facilitate a higher

degree of inter-molecular hydrogen bonding between chains. Indeed, in *Arabidopsis*, dispersed complexes produce widely spaced cellulose microfibrils in primary cell walls, whereas in secondary cell walls, dense regions of complexes synthesize highly aggregated crystalline microfibrils (Li et al., 2016). Interestingly, *in vitro* studies of cellulose synthesis have suggested that adjacent microfibrils can spontaneously coalesce to form thicker microfibril bundles in the absence of a rosette complex. Although this indicates that microfibrils may self-assemble in the cell wall, whether these microfibrils resemble *in vivo* structures was not quantified (Cho et al., 2017; Purushotham et al., 2016) and so more rigorous assessment is required to draw this conclusion with great certainty.

The location of terminal complexes at the ends of microfibrils and the high density of rosettes in areas undergoing secondary cell wall

deposition (Herth, 1985) made terminal complexes primary candidates for cellulose biosynthesis, however, this evidence was purely circumstantial. Fifteen years after terminal complexes were first visualized in green algae, genes with cellulose synthetic ability were cloned from the bacteria, A. xylinum (Saxena, Lin, & Brown, 1990; Wong et al., 1990). The bacterial operon encodes four bacterial cellulose synthase (Bcs) genes, BcsA/B/C/D, that are members of the glycosyltransferase 2 (GT2) family. Homologs in higher plants were found by screening a cotton cDNA library for sequence similarities with A. xylinum (Pear, Kawagoe, Schreckengost, Delmer, & Stalker, 1996). Although the genes from cotton exhibited low sequence homology with A. xylinum, as these proteins could bind to UDP-glucose in vitro they were putatively named cellulose synthase (CESA) genes. Immuno-labelling of freeze-fractured terminal complexes in azuki beans with CESA-specific antibodies, made the vital connection between cellulose synthesis and terminal complexes and identified CESAs as a component of the terminal complexes (Kimura et al., 1999) (Table 1). As a result, terminal complexes are commonly referred to as cellulose synthase complexes (CSCs).

The exact number of CESA proteins that occur in CSCs has been widely debated and is still an outstanding question in the field (Table 1). Originally, it was speculated that each rosette subunit contains a hexamer of CESA proteins that each synthesize a single chain, producing a 36-chain microfibril (Herth, 1983). Each CESA protein is still proposed to synthesize a single glucan chain, based on strong homology between the catalytic domain of cotton CESA proteins and the Bcs complex of Rhodobacter sphaeroides, that produces a single chain (Morgan, Strumillo, & Zimmer, 2013; Sethaphong et al., 2013). Recent evidence corroborates this hypothesis, as single CESA isoforms purified from rice and poplar arecapable of synthesizing cellulose in vitro (Olek et al., 2014; Purushotham et al., 2016). However, the 36-chain model has been widely rejected as 3 nm wide microfibrils are simply too narrow to accommodate 36 chains (Fernandes et al., 2011; Newman, Hill, & Harris, 2013; Thomas et al., 2013), and 45-50 nm wide cytoplasmic domains of CSCs are predicted to contain a maximum of four CESA proteins per rosette subunit (Bowling & Brown, 2008).

Updated models now predict that CSCs that are composed of a hexamer of trimers or tetramers, producing 18- or 24-chain microfibrils, respectively (Table 1). SANS, WAXS and ssNMR examination of secondary cell walls in spruce and primary cell walls in celery collenchyma are consistent with a 24-chain model (Fernandes et al., 2011; Thomas et al., 2013) whereas an 18-chain model is favored in mung bean primary cell walls (Newman et al., 2013). Assuming all CESA proteins within a CSC are active, evidence from studies of *Arabidopsis* leans towards an 18-chain model, as equimolar concentrations of CESA proteins (Gonneau, Desprez, Guillot, Vernhettes, & Hofte, 2014; Hill, Hammudi, & Tien, 2014) and the formation of *CESA1* homotrimers in solution are

both incompatible with a 24-chain model (Vandavasi et al., 2016). In *Arabidopsis*, models predict that CSCs composed of 18 CESA proteins contain either hetero or homo-trimers. Each lobe contains either three distinct or identical CESA isoforms, based on the 1:1:1 stoichiometry of CESA proteins, the formation of homotrimers *in vitro* and the requirement of three CESA isoforms for a functioning CSC *in vivo* (Fig. 1A, B).

Due to the range of techniques, species and cell wall types adopted by different studies, it is hardly surprising that there is no consensus amongst research groups. It is also plausible that both the 18- and 24chain model are correct under different circumstances, since microfibril diameters can vary (Martinez-Sanz, Pettolino, Flanagan, Gidley, & Gilbert, 2017). For example, in poplar tension wood, individual microfibrils are twice as wide in the G-layer compared to the adjacent secondary cell wall layers (Müller, Burghammer, & Sugiyama, 2006) and in fruit tissues, the microfibril diameter can be as low as 1 nm (Niimura, Yokoyama, Kimura, Matsumoto, & Kuga, 2009). Measuring the width of microfibrils and estimating the number of CESA proteins as a proxy for the number of glucan chains is not ideal, as microfibrils frequently interact with other matrix components and CESA proteins are not necessarily all active within a rosette. However, deciphering accurate CSC compositions in different cell walls and species may not be possible until CSCs and CESAs can be examined at higher resolution.

2.3. CSCs - architecture

Discovering CESA proteins hugely expanded our capabilities for studying cellulose synthesis, as it became possible to identify CESA homologs by sequencing analysis in species where the complex had not been visualized. During this time, *Arabidopsis* had gained popularity as a molecular model and so became the preferred study system for cellulose synthesis. CESA homologs were successfully identified in *Arabidopsis* by screening mutant populations for cellulose deficiencies. CESA proteins were first described in the primary cell walls of the *radial swelling mutant*, *rsw1* (Arioli et al., 1998), which exhibited stunted growth and reduced cellulose content at 31 °C and in three *irregular xylem mutants*, *irx1/3/5*, exhibiting deformed secondary cell walls in vessels (Turner & Somerville, 1997). After the *Arabidopsis* genome was sequenced, a total of 10 CESA genes (*CESA1-10*) were identified (Richmond, 2000).

With the identification of multiple CESAs, it was soon realized that CSCs were not made up of a homogenous population of CESA proteins. CESA proteins can be divided into two distinct families depending on the type of cell wall. In *Arabidopsis*, *CESA1*, *CESA3* and *CESA6/CESA6*-like proteins (*CESA2*, *CESA5*, *CESA9*) are required in the primary cell wall (Persson et al., 2007) and in secondary cell walls, *CESA4*, *CESA7* and *CESA8* are indispensable for plant growth (Taylor, Howells, Huttly, Vickers, & Turner, 2003). Three distinct isoforms are required to form a

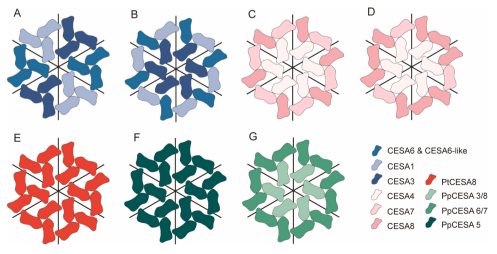


Fig. 1. Predicted arrangement of CESA proteins within the CSC. In Arabidopsis primary cell walls, CESA proteins exist as either (A) homotrimers or (B) heterotrimers within the CSC (Hill et al., 2014), (C) Models of CSCs in the secondary cell walls of Arabidopsis and spruce predict that CESA proteins have a 1:1:1 stoichiometry (Zhang, Dominguez et al., 2018). In the secondary cell walls of poplar, CSCs have a (D) 3:2:1 stoichiometry of CESA8:4:7 in normal wood, and a (E) 8:3:1 stoichiometry in tension wood (Zhang, Dominguez et al., 2018). Models of CSCs in Physcomitrella patens predict a h (F) homo-oligomer composition of PpCESA5 in primary cell walls (Goss, Brockmann, Bushoven, & Roberts, 2012) and a (G) hetero-trimer composition in secondary cell walls (Norris et al., 2017; Scavuzzo-Duggan

functioning complex as individually mutating each of these CESA isoforms causes severe defects in cell wall synthesis (Desprez et al., 2007; Taylor et al., 2003). CESA1 and CESA3 are essential for primary cell wall synthesis because cesa1 and cesa3 mutants are gamete lethal, whereas cesa6 mutants can still function due to partial redundancy with CESA6-like proteins, though they still exhibit a severe lack of cellulose and growth defects (Persson et al., 2007). Recent genetic work has shed uncertainty on the rigid distinction between primary and secondary cell wall CESAs, since primary cell wall CESAs can form functional complexes with secondary cell wall CESAs in both poplar and Arabidopsis (Carroll et al., 2012; Li, Lei, & Gu, 2013; Song, Shen, & Li, 2010). Furthermore, primary cell wall CESAs can interact with secondary cell wall CESAs both in vitro and in vivo, and pCESA7::CESA1 can partially rescue cesa8 knock-outs (Carroll et al., 2012). CESA6-like proteins are also important in synthesizing specialized secondary cell walls, as cellulose defects are apparent in the seed coat of cesa2, cesa5, cesa9 mutants and in the mucilage of cesa5 mutants (Mendu et al., 2011). As cellulose synthesis is vital for plant growth, some promiscuity between CESA binding may exist to ensure cellulose production is maintained. Mixed complexes may represent 'intermediates' that facilitate the rapid changeover between primary and secondary cell wall synthesis. Whether functional compatibility between CESA isoforms is merely due to the high conservation between CESA catalytic domains is uncertain, as currently there is no evidence these mixed complexes are formed in

It is not known why the composition of CESAs in the CSC differs between primary and secondary cell walls, but it must be essential to warrant such a significant energy investment in the changeover of CESA isoforms between cell wall types. The ability of single CESA isoforms to synthesize cellulose causes further confusion as to why multiple isoforms are needed (Purushotham et al., 2016). Only 25 % of the microfibrils produced in vitro are crystalline, so perhaps microfibrils synthesized in the absence of other CESAs are structurally defective. Differences between the composition of the CSC in primary and secondary cell walls is ultimately driven by evolution, since the common ancestor of moss and seed plants exhibited a rosette-CSC comprised of a single CESA isoform (Roberts & Bushoven, 2007). Both moss and seed plants evolved two classes of CESA proteins independently, stressing that a variety of isoforms evolved to fulfil separate functions in different cellular environments and under different regulatory pathways. In P. patens, PpCESA5 is required for primary cell wall formation in leaf gametophores (Fig. 1F), whereas PpCESA3/8 and PpCESA6/7 are required for secondary cell wall deposition in stereids that resemble tracheary elements (Fig. 1G). Convergent evolution of hetero-oligomeric CSCs suggests that the specialization of CESA was a fundamental requirement for synthesizing cellulose under different in vivo conditions (Li et al., 2019)

Attempts to tease apart the different functions of CESA by systematic mutagenesis have demonstrated that while their precise functions are not fully understood, CESAs clearly impart unequal roles in cellulose synthesis. Mutating catalytic motifs in different CESA proteins differentially impacts cellulose synthesis indicating CESA proteins vary in their catalytic ability. For instance, a cesa8 mutant exhibits severe reductions in cellulose content, whereas only mild decreases are reported for cesa4 (Kumar, Atanassov, & Turner, 2017). A popular hypothesis is that CESA isoforms may determine crystallinity, because cellulose contains a higher proportion of crystalline cellulose than primary cell walls. In particular, CESA8 may be fundamental for mediating crystallinity, as not only does is it appear more catalytically active than other isoforms in the CSC (Kumar et al., 2017), but it is more abundant than CESA4 and CESA7 in poplar secondary cell walls that have a high degree of crystallinity (Zhang, Dominguez et al., 2018). In Arabidopsis and Norway spruce, CESA proteins are expressed in equimolar concentrations with a stoichiometry of 1:1:1 (Fig. 1C). However, in poplar the stoichiometry of CESA8:4:7 is 3:2:1 (Fig. 1D) (Zhang, Dominguez et al., 2018). In poplar tension wood, a significant increase in PtCESA8b mRNA transcripts

produces a more exaggerated shift in stoichiometry of 8:3:1 that coincides with cellulose that is almost completely crystalline (Fig. 1E) (Zhang, Dominguez et al., 2018). Interestingly, *PtCESA7* transcripts do not change in abundance and *PtCESA4* and *PtCESA8a* decreases, indicating different CESA isoforms confer different roles in tension wood.

In higher plants, CESA proteins have 8 transmembrane domains separated by a large catalytic cytosolic loop between the 2nd and 3rd domain. The 8 transmembrane domain model has recently been challenged by Slabaugh et al. (2016) who proposed that the 5th domain is an interfacial helix, making CESA a 7 transmembrane domain protein (Table 1). The absence of this transmembrane domain relocates a loop with a conserved FxVTxK motif to the cytoplasm. Here, it might assist in substrate binding as it does in its bacterial counterpart. Based on the crystal structure of the BcsA-BcsB complex, the transmembrane domains of CESA proteins are predicted to form a channel through which newly synthesized glucan chains are released (Morgan et al., 2013). Recent structural analysis of PttCESA8 homotrimers indicates that the transmembrane domains of each CESA forms a continuous channel across the membrane, similar to the bacterial complex (Purushotham, Ho, & Zimmer, 2020). Furthermore, these channels appear to converge in the center of the trimer to facilitate the secretion and coalescence of nascent glucan chains, suggesting higher plant CESAs and the BcsA-BcsB complex share a common mechanism for cellulose synthesis.

In CESA proteins, the catalytic domain contains the highly conserved motifs (D, D, DxxD and QxxRW) common to all GT2 enzymes (Sethaphong et al., 2013). Point mutations in these motifs in Arabidopsisand cotton have verified that they perform distinct roles in catalysis and substrate binding. The first two aspartates (D, D) are involved in the binding of UDP-glucose substrates, DxxD acts as a base for glucan chain extension and the QxxRW motif as a binding site for the final glucan residues in the chain (Saxena et al., 2001). Within the catalytic loop there is a plant-conserved region (P-CR) between D and DxxD and a hypervariable region (HVR) at the other end of the domain (Pear et al., 1996). The HVR contains a class-specific region (CSR) specific to each isoform, that is widely homologous between different species, yet highly diverse amongst different isoforms (Vergara & Carpita, 2001). Intuitively, the CSR is thought to determine CESA isoform specificity. However, chimeric studies in Arabidopsis and moss have firmly established that the CSR is largely interchangeable between different CESA isoforms. Swapping domains between different CESA proteins does not compromise catalytic ability and chimeric CESA proteins can successfully rescue the corresponding mutants, indicating that class specificity is neither restricted, nor defined by these regions (Hill, Hill, Roberts, Haigler, & Tien, 2018; Kumar et al., 2017; Scavuzzo-Duggan et al., 2018; Sethaphong et al., 2016; Wang, Howles, Cork, Birch, & Williamson, 2006).

Alternatively, the CSR and P-CR are speculated to mediate CESA interactions, CESA positioning in the CSC and rosette formation, since both these regions are absent from bacteria that do not form rosettes. Indeed, when CSR and P-CR regions are included in structural models of the CESA catalytic domain, they diverge significantly from the structure of Bcs (Olek et al., 2014; Purushotham et al., 2016). Studies combining mathematical modelling with low-resolution SANS and SAXS analysis provide conflicting evidence for the roles of P-CR and CSR in CSC formation. In rice, the CSR region is predicted to assist in OsCESA8 dimerization and the P-CR region is implicated in dimer-dimer interactions (Olek et al., 2014), whereas in Arabidopsis, the P-CR region of AtCESA1 is predicted to recruit non-CESA proteins and the CSR is implicated in trimer-trimer assembly (Vandavasi et al., 2016). Resolving the crystal structure of the P-CR region in OsCESA8 revealed that it consists of two α-helices linked by a large extended loop (Rushton et al., 2017). Incorporating the crystal structure into previous SAXS-based models predicts that the P-CR is located in the catalytic core close to the active site (Rushton et al., 2017), which is easily reconciled with the dimerization of CESA proteins (Olek et al., 2014). Discrepancy between dimer- and trimer-models is likely caused by a low homology of CSR

between different isoforms and the purification of CESA isoforms under different experimental conditions. As these studies report the formation of homodimers and homotrimers from single CESA isoforms *in vitro*, it cannot be discounted that hetero-dimers and -trimers may be formed in the presence of other CESA proteins *in vivo* due to the high conservation of catalytic domains. While the precise role of P-CR and CSR in CSC assembly is ambiguous, it can be concluded that these regions mediate CESA interactions in different capacities.

The N-terminal contains a zinc-finger domain that can dimerize with the same or different CESA proteins under redox conditions in cotton (Kurek, Kawagoe, Jacob-Wilk, Doblin, & Delmer, 2002)., A recent study of the *Arabidopsis* acylome revealed that the zinc-finger is likely to be inactive in *CESA4* and *CESA8* since the acylation of key cysteine residues compromise its ability for metal ion binding (Kumar, Carr, & Turner, 2020). On the contrary, mutating key cysteines in the zinc-finger of *CESA7* greatly impairs its function, suggesting *CESA7* may be essential for maintaining the integrity of the CSC (Kumar et al., 2020). With more intricate examination of CESA structures, the exact functions of CESA domains and individual CESA isoforms in the CSC are starting to unravel.

Continuing advancements in sequencing technology have facilitated the identification of CESA homologs in a huge diversity of eukaryotes and prokaryotes. Assigning function to CESA homologs represents the rate-limiting step as functional genetic analysis can take years in some species, particularly trees. Expression analysis has been used as an indirect indicator of CESA function in various tissues and points of development. However functional genomics is needed to definitively assign function to these orthologs. So far this has been achieved in several commercially important species such as rice, maize, poplar and eucalyptus. Further quantification is needed, particularly in woody species that have multiple copies of CESA isoforms that presumably have distinct roles in wood formation (Zhang, Dominguez et al., 2018).

2.4. Crystallization and polymerization

Due to the lability of cellulose I, it was reasoned that crystallization and polymerization must be co-ordinated for cellulose I to acquire stability in the cell wall (Saxena & Brown, 2005). Inhibiting crystallization with Calcofluor white in *A. xylinum* increases the rate of polymerization by four-fold, suggesting that not only are these processes tightly coupled, but that crystallization limits polymerization (Benziman, Haigler, Brown, White, & Cooper, 1980). In R. sphaeroides, newly synthesized glucan chains are elongated by the stepwise addition of glucose units (Morgan et al., 2016) - a mechanism thought to be shared with plants (Table 1). In higher plants, the close proximity of rosette subunits likely facilitates simultaneous crystallization with the coalescence of glucan chains (Table 1), since the loss of CSC organization is often concurrent with an increase in amorphous cellulose. For example, mutating the catalytic subunit or transmembrane domains of AtCESAs decreases crystalline cellulose (Arioli et al., 1998; Harris et al., 2012) and removing the zinc-finger domain in pttcesa8 mutants produces amorphous cellulose exclusively, reinforcing that the structure of the rosette is mandatory for crystallization (Purushotham et al., 2016).

Continuous chain elongation was predicted to drive the movement of CSCs through the plasma membrane (Herth, 1983) (Table 1). A later study also suggested that the continuous synthesis of cellulose from CSCs generates the driving force to propel its movement, based on the migration of *YFP::CESA6* in the plasma membrane observed with spinning disc confocal microscopy (Paredez, Somerville, & Ehrhardt, 2006). Biophysical modelling of CSC movement based on crystallization and polymerization alone, predicted that the CSC could move in the plasma membrane at a speed of 10^{-9} – 10^{-8} m s⁻¹ (Diotallevi & Mulder, 2007), similar to reported values of 5– 8×10^{-9} m s⁻¹ (Paredez et al., 2006).

2.5. Non-catalytic genes involved in cellulose synthesis

Identifying non-catalytic genes essential for cellulose synthesis was relatively straightforward in bacterial genomes, where functionally related genes often cluster together. For plants that have more complex genomes, candidates were initially identified using forward genetic screens with cellulose biosynthesis inhibitors. One of the first non-CESA genes to be identified was the putative membrane-spanning endo-1,4 β-D-glucanase, KORRIGAN (KOR1) (Nicol et al., 1998). Determining the precise role of KOR1 has not been possible, because mutating KOR1 causes a range of phenotypes including reduced crystalline cellulose (Maloney & Mansfield, 2010); altered CSC velocity (Vain et al., 2014); and perturbed microfibril orientation (Lei et al., 2014). Since the evolution of KOR1 pre-dates the appearance of CESA in green algae, KOR1 may have been fully responsible for synthesizing cellulose in primitive life forms (Lampugnani et al., 2019). As KOR1 is still functional in higher plants it must have had a selective advantage, possibly by assisting with cellulose synthesis in conjunction with CESA proteins. With the exception of CESA7, KOR1 can bind to all cell wall CESA proteins in yeast two hybrid (Y2H) assays (Mansoori et al., 2014) and fluorescent tagging of KOR1 revealed that it associates with CSCs in the Golgi, TGN, secretory vesicles and the plasma membrane (Lei et al., 2014; Vain et al., 2014). Together, this strongly indicates that KOR1 is a permanent resident of the CSC that modulates CSC function throughout its lifespan. Due to its tight association with the CSC, defects observed in kor1 mutants may be an indirect consequence of gene perturbation, further complicating the assignment of KOR1 function. Another early non-CESA gene identified was the glycosyl-phosphatidyl inositol-anchored protein, COBRA (COB) (Benfey et al., 1993; Roudier et al., 2005). COB has been described as a 'scaffold' for maintaining microfibril orientation and binding in Arabidopsis (Roudier et al., 2005). COB evolved alongside CESA, coinciding with the shift in linear arrays to rosette-shaped CSCs (Lampugnani et al., 2019), and therefore may be important in synthesizing glucan chains in close proximity to one another.

Significant advancements in genetic techniques have now made it possible to identify genetic candidates based on their physical interactions with CSC machinery. Many CESA-interacting proteins have been discovered from Y2H assays, GFP-TRAP, co-immunoprecipitation combined with mass-spectrometry, in vitro pull downs and biomolecular fluorescence complementation (BiFC). In particular, key proteins integral for maintaining the relationship between CSC and underlying microtubules have been described including, CELLULOSE SYNTHASE MICROTUBULE UNCOUPLING PROTEIN (CMU) that prevents the lateral displacement of microtubules from the hypothesized pressure generated by CSC migration (Liu et al., 2016); CELLULOSE-SYNTHASE INTERAC-TIVE PROTEIN 1 (CSI1/POM2) that links CSCs with microtubules and marks regions for CSC exocytosis (Bringmann et al., 2012; Gu et al., 2010; Zhu, Li, Pan, Xin, & Gu, 2018); and COMPANION OF CELLULOSE SYNTHASE (CC) that promotes microtubule dynamics for CSC localization under specific stress conditions (Endler et al., 2015). CMU, CSI1 and CC were some of the most recent cellulose-related genes to evolve, appearing in a group of Charophyceae algae known as Zygnematophyrae (Lampugnani et al., 2019). The evolution of a specialized microtubule band involved in cytokinesis in Zygnematophyrae strongly suggests that CMU, CSI1 and CC evolved for the succinct co-ordination of microfibrils and microtubules - a feature that was retained by higher plants (Lampugnani et al., 2019).

In general, the roles of non-catalytic proteins in cellulose synthesis have been described in the context of primary cell walls. However, there is increasing evidence that many of these genes have reciprocal or divergent functions in secondary cell walls. For instance, *kor1* mutants exhibit defects in vessel secondary cell wall formation in *Arabidopsis* (Szyjanowicz et al., 2004) and *KOR1* can physically interact with secondary cell wall CESAs (Mansoori et al., 2014; Vain et al., 2014). The role of *CSI1* in secondary cell walls is disputed (Zhu, Xin, & Gu, 2019). No cellulose defects are apparent in *csi1* mutants (Gu et al., 2010), yet it

is abundant in induced *Arabidopsis* tracheary elements (Derbyshire et al., 2015) and in *pom2-4* mutants, xylem vessels have irregular wall patterns and *CESA7* is mis-aligned with microtubules (Schneider et al., 2017). An alternative isoform of *COB, COBL4*, may be specifically involved in producing highly crystalline cellulose in secondary cell walls. *COBL4* is upregulated in secondary cell walls (Brown, Zeef, Ellis, Goodacre, & Turner, 2005) and tension wood (Andersson-Gunnerås et al., 2006) and the *COBL4* homolog in rice, *BRITTLE CULM1* (*bcl*), can bind to crystalline microfibrils (Liu et al., 2013). As many non-catalytic genes clearly participate in various aspects of both primary and secondary cell wall formation, perhaps assigning precise functions is not possible or biologically accurate.

2.6. Biochemistry of cellulose synthesis

Studying the biochemical aspects of cellulose synthesis has been notoriously problematic over the last 30 years. A persistent problem has been that β -1,3 linked callose was preferentially synthesized over β -1,4 linked cellulose from plant membrane extracts, hampering efficient cellulose production (Amor et al., 1995). Moderate improvements were achieved from *in vitro* cultures of hybrid aspen that produced almost 50 % cellulose (Ohlsson et al., 2006) and microsome preparations of blackberries that yielded up to 1 mg cellulose (Lai-Kee-Him et al., 2002), but poor yields and callose contamination were still major concerns. Significant advances have recently been achieved from the heterologous expression of CESA isoforms from poplar and P. patens in yeast (Cho et al., 2017; Purushotham et al., 2016). Reconstituting PttCESA8 and PpCESA5 in proteoliposomes that mimic the lipid bi-layer environment proved essential for successful synthesis, as disrupting the bilayer with detergent eliminated catalytic activity. Radio-active tracing of UDP-[³H]-Glc, determined that catalysis was maintained for 90–150 min, a considerable improvement from previous in vitro reactions that terminated after 10 min (Amor et al., 1995). Whether these cellulose microfibrils are representative of microfibrils invivo presents the next major challenge.

Despite significant advancements in the synthesis of cellulose in vitro, the purification and reconstitution of the entire CSC has so far not been possible and remains a major research priority. Biochemical inferences of CESAs have been made from low resolution SAXS analysis that does not depend on protein crystallization and comparisons with the crystal structure of the BcsA-BcsB complex. Recent structural analysis of a PttCESA8homotrimer with cryogenic-EM suggested that plants and bacteria share a common mechanism for synthesizing cellulose (Purushotham et al., 2020). However, this mechanism may not be entirely indicative of CSC function in higher plants since it has not been established if these homotrimers exist in vivo. Furthermore, the cellulose microfibrils produced by recombinant PttCESA8homotrimers expressed in insect cells, do not resemblemicrofibrils produced by previous in vitroassays or microfibrils synthesized in vivo. Microfibrils were 4 times narrower (10-15 Å) than the 4.3 and 4.8 nm wide microfibrils produced by re-constituted PttCESA8 and PpCESA5 proteoliposomes (Cho et al., 2017; Purushotham et al., 2016), and microfibrils were amorphous, contrary to higher plants that contain a high proportion of crystalline cellulose. Whilst heterologous expression of CESAs in different systems may be the cause of this discrepancy, inconsistencies in the in vitrocellulose production of PttCESA8casts some doubt on the proposed mechanism of PttCESA8homotrimers. Nonetheless, the ability to study the structure of purified CESAs with cryogenic-EM, represents a major breakthrough in the study of CSC structure (Table 1) that will facilitate a more complete understanding of cellulose synthesis in the future.

3. When is cellulose synthesized?

Plant growth and shape is achieved predominately by cell expansion as opposed to cell division (McFarlane, Doring, & Persson, 2014). Cell expansion is permitted by internal stresses generated by turgor pressure

and the slow yielding of the primary cell wall (Cosgrove, 2016). Unsurprisingly, the organization of cellulose microfibrils and the cell wall architecture is tightly linked with cell expansion. Early hypotheses regarding the role of cellulose microfibrils in cell expansion were developed solely from TEM-based observations. While TEM is a useful tool for visualizing the cell wall architecture in its entirety, sample preparation can disrupt native cell wall structures. Developing techniques that preserve the cell wall architecture with higher fidelity, such as field emission scanning electron microscopy (FESEM) and AFM, meant that these predictions could be more rigorously scrutinized, but only in the innermost cell wall layer. Cell expansion studies have been fairly limited to cell-types with thin cell walls, which lend themselves to high resolution imaging, particularly the epidermal tissues from onion (Suslov, Verbelen, & Vissenberg, 2009) and the dark-grown hypocotyl and root elongation zone from *Arabidopsis*.

3.1. Cell elongation and expansion

Directional growth in plant cells is achieved by anisotropic expansion, whereby cells stretch longitudinally and undergo minimal lateral expansion. As the load bearing structure, cellulose microfibrils are important in generating differential resistance to turgor pressure and determining the direction of growth. Consequently, anisotropic expansion is highly reliant on efficient cellulose biosynthesis. In fact, many cellulosic biosynthetic genes were initially identified from mutants exhibiting abnormal cell elongation, such as cesa1^{rsw1} (Arioli et al., 1998); cesa6^{prc1-1} (Fagard et al., 2000); cob (Benfey et al., 1993); kor1 (Nicol et al., 1998); and pom1/2 (Hauser, Morikami, & Benfey, 1995). For some mutants, including cesa6^{prc1-1} and pom1, the microfibril deposition is not altered (Baskin, 2005; Pagant et al., 2002; Refregier, Pelletier, Jaillard, & Hofte, 2004) meaning these defects in anisotropic expansion may be a knock-on effect of cellulose perturbation caused by hormonal changes or compensational responses of other cell wall components.

Cellulose microfibrils was first connected with anisotropic growth from TEM-based observations of the primary cell walls of *Tradescantia* stamen hairs (Roelofsen & Houwink, 1951). In the newly formed central lamellae, cellulose microfibrils were deposited perpendicular to the direction of cell growth, whereas new lamellae deposited towards the outside of the cell tended to have a longitudinal orientation, parallel to the growth axis. Changes in microfibril orientation led to the concept of 'multi-net growth' (Table 1). Under the multi-net growth hypothesis, the progressive re-alignment of microfibrils towards the outer cell layers causes the cell to elongate (Roelofsen & Houwink, 1953). Identical observations were subsequently reported in algae (Tsekos, 1999) and *Arabidopsis* (Anderson et al., 2010). The transverse orientation of microfibrils was predicted to generate differential resistance to turgor pressure by physically restricting lateral expansion and promoting rapid longitudinal elongation (Green 1962, Green, 1960).

The multi-net growth hypothesis is one of the longest standing hypotheses in cellulose biosynthesis, but it has lost considerable backing as many of the conditions required by the multi-net growth hypothesis are no longer satisfied when complex tissues of higher plants are considered (Table 1). This is particularly true of cross polylamellate walls in the epidermis of hypocotyls, stems and coleoptiles that exhibit parallel microfibrils that alternate by 30-90° between successive lamellae (Chan et al., 2010; Zhang, Zheng et al., 2016). Furthermore, transverse microfibril orientation does not consistently induce anisotropy (Wiedemeier et al., 2002; Xin et al., 2020) and expansion can be achieved without the passive reorientation of microfibrils (Bashline, Lei, Li, & Gu, 2014; Marga, Grandbois, Cosgrove, & Baskin, 2005). In the stem and hypocotyl epidermis, cell elongation is achieved despite having longitudinally or randomly orientated microfibrils. To explain this discrepancy, it has been suggested that the inner cell layers control the direction of expansion by imparting the necessary axial force to the outer epidermis (Baskin, 2005), or by generating sufficient anisotropic

expansion than negates the isotropic expansion of the epidermis (Fujita et al., 2011). Indeed, examining the innermost cell wall layer of etiolated *Arabidopsis* hypocotyls with FESEM demonstrated that the transverse microfibril orientation of the inner regions could induce growth anisotropy of the outer epidermal layers (Chan et al., 2011; Crowell et al., 2011). Additionally, Baskin (2005) discovered that anisotropic expansion was reduced when microfibril alignment was not uniform within tissues, suggesting that the net alignment of microfibrils between cells is more crucial than within cells for determining the degree of anisotropic expansion. It is important to note that hypotheses regarding cell expansion are predominately tested in model systems and are unlikely to be representative of other cell types and developmental stages that differ in their extent of expansion, due to differences in turgor pressure and microfibril orientation.

3.2. Relationship with microtubules

In early studies of cellulose synthesis, one of the most frequent observations made was the relationship between cellulose microfibril orientation and cortical microtubules patterns (Hepler & Newcomb, 1964), 'Cortical cytoplasmic elements', later realized to be microtubules, were proposed to guide the positioning of nascent cellulose microfibrils (Green, 1962). Based on the parallel alignments of cortical microtubules with cellulose microfibrils, the 'alignment hypothesis' was developed (Ledbetter & Porter, 1963) (Table 1). The complementary association between microtubules and microfibrils was initially confirmed in TEM studies of green algae (Tsekos, 1999), and later by confocal microscopy, where YFP::CESA6 and RFP::TUA6 signals co-localized in the plasma membrane of Arabidopsis primary cell walls (Li et al., 2016; Paredez et al., 2006). Early studies showed that disrupting microtubule dynamics prevented cell elongation, providing a direct link between microfibril orientation and microtubules (Morejohn, 1991). Live-cell imaging provided further confirmation that CSCs are directly guided by underlying cortical microtubules, by demonstrating that the trajectories of CSCs and newly synthesized microfibrils were re-orientated to align with new patterns of microtubules, following microtubule disruption (Paredez et al., 2006).

Whilst many studies largely support the alignment model, there are some notable inconsistencies. It was soon realized that the relationship is not as simple as when first proposed since microtubules are not ubiquitously required for the alignment of CSCs and microfibrils (Chan & Coen, 2020; Mizuta & Okuda, 1987). In the innermost layers of the root and hypocotyl epidermis, the parallel trajectories of CSCs and the transverse orientations of cellulose microfibrils are maintained in the absence or disorder of microtubules (Himmelspach, Williamson, & Wasteneys, 2003; Sugimoto, Himmelspach, Williamson, & Wasteneys, 2003; Xin et al., 2020), suggesting that microfibril assembly is not reliant on microtubules. Under these circumstances, CSCs that are not linked with underlying microtubules may maintain their alignments by tracking previous microtubule trajectories, as observed with light microscopy (Chan & Coen, 2020). Alternatively, microtubules may influence cell expansion and cellulose synthesis by determining the extent of cellulose crystallinity. When cell expansion is stimulated at 29 °C, the proportion of crystalline cellulose simultaneously decreases, however when the abundance of microtubules is reduced in the temperature-sensitive mor1-1 mutant, cells can no longer expand and cellulose crystallinity content does not change at 29 °C (Fujita et al., 2011). Microtubules may modulate crystallinity by controlling the fluidity of the plasma membrane or the interaction with non-cellulosic components (Fujita, Lechner, Barton, Overall, & Wasteneys, 2012). It is also true that cellulose microfibrils may determine the distribution of cortical microtubules. Tobacco BY2 cells and Arabidopsis roots treated with cellulose biosynthesis inhibitors exhibit a dispersed, unordered microtubule array (Fisher & Cyr, 1998; Himmelspach et al., 2003) and in cesa2 and cesa6 mutants' cortical microtubules have a distorted alignment (Chu et al., 2007; Paredez, Persson, Ehrhardt, & Somerville, 2008).

A bi-directional interaction between microtubules and microfibrils is not necessarily incompatible with the alignment hypothesis, but it indicates that revisions need to be made. A more suitable model may be the 'cellulose-constraint' model proposed by Giddings and Staehelin (1991) whereby cortical microtubules constrain paths for CSC movement and cellulose microfibrils exert biophysical forces on cortical microtubules as part of a self-reinforcing feedback loop.

The next step in understanding the relationship between microtubules and microfibrils was to establish the basis of their association. Whilst live-cell imaging confirms that their trajectories are correlated, it does not indicate whether CSC and microtubules are in direct contact, or if other factors are involved. Heath (1974) proposed the popular 'direct guidance model', whereby CSCs directly interact with microtubules (Table 1). Genetic evidence now supports that CSCs indirectly interact with microtubules through a linker protein known as CSI1 (Gu et al., 2010; Li, Lei, Yingling, & Gu, 2015). CSI1 interacts with both microtubules and the catalytic domain of CESA proteins in vitro and RFP::CSI1 co-localizes with YFP::CESA6 in vivo (Bringmann et al., 2012; Gu et al., 2010; Li, Lei, Somerville, & Gu, 2012). In csi1-1 mutants, CSC trajectories are uncoupled from microtubules and CSC velocity is slower (Gu et al., 2010; Li et al., 2012), although when microtubules are removed, CSCs can maintain ordered trajectories. This may be explained if CSI1 is essential for the initial alignment of microtubules with CSCs, after which the trajectory does not depend on microtubule presence (Schneider et al., 2017). CC1 also directly interacts with microtubules and the CSC. Mutating two tyrosine residues essential for microtubule-binding in the CC1gene, disrupts the parallel alignment between CSCs and microtubules, suggesting CC1has an important role in maintaining the relationship between CSCs and microtubules (Kesten, 2019). A wealth of studies has convincingly demonstrated that CSCs and microtubules are co-dependent and both are important for cell anisotropy. Ultimately our ideas match those originally proposed by Green (1962), but the relationship is clearly more nuanced than first proposed and so these early hypotheses have been more rigorously scrutinized.

4. CSC trafficking

A huge breakthrough that facilitated the study of CSC trafficking was the development of live-cell imaging that allowed CSC dynamics to be visualized within the cell. Functional complementation of the non-lethal $cesa6^{prc1-1}$ mutant, with fluorescently tagged CESA6 proteins, enabled CSC movements in primary cell walls to be traced with confocal microscopy (Paredez et al., 2006). Studies are preferentially performed on dark-grown Arabidopsis hypocotyls, due to the high abundance of CSCs coupled with thin primary cell walls that enhance imaging resolution. On the other hand, high quality imaging of CSC movements in secondary cell walls, which can be deeply embedded within tissues, has been a much greater challenge. With the development of inducible lines, it is now possible to visualize tracheary elements with greater resolution (Yamaguchi et al., 2010). Together with live-cell imaging, proteomic analysis has been ground-breaking in identifying key proteins that interact with CSCs during trafficking.

4.1. CSC assembly

Terminal complex assembly was hypothesized to occur in either the ER or the Golgi, before being transported to the plasma membrane (Table 1). Evidence for Golgi assembly was first indicated from TEM-based observations of fully formed terminal complexes embedded in the Golgi, TGN and post-Golgi vesicles in algae (Brown, Franke, Kleinig, Falk, & Sitte, 1970; Giddings et al., 1980). For algal species that produce large linear complexes, such as *Erthyrocladia* and *Vaucheria*, assembly is partially completed at the membrane as vesicles containing single particles, multi-subunits and precursor-complexes all fuse with the membrane (Mizuta & Brown, 1992; Tsekos, 1999). In multicellular organisms, rosettes were first observed in the TGN and post-Golgi

vesicles in differentiating tracheary elements of *Zinnia elegans* mesophyll cells (Haigler & Brown, 1986).

Little progress has been made in uncovering how the CSC assembles, due to the difficulties in visualizing pre-Golgi processes, particularly in the ER. During live-cell imaging, YFP::CESA fluorescence is either very weak or undetectable in the ER (Crowell et al., 2009; Gutierrez, Lindeboom, Paredez, Emons, & Ehrhardt, 2009; Paredez et al., 2006), presumably due to the quenching of fluorescence deeper in the cell. In the cesa6D395N mutant, diffuse signals of YFP::CESA6 below the Golgi was interpreted as the retention of malformed CSCs in the ER (Park, Song, Shen, & Ding, 2019). However, as no ER marker was used, and the distribution of CESA1 and CESA3 was not examined in conjunction with CESA6, it is not possible to differentiate whether the entire CSC or single CESA6 proteins are retained in the ER (Park et al., 2019). In secondary cell walls, the co-localization of GFP::CESA4 and GFP::CESA8 with the ER binding protein, BiP, in cesa7^{irx3} mutants further supports the idea that incomplete CSCs cannot be transported from the ER (Gardiner, Taylor, & Turner, 2003). Despite the limited evidence, it is generally

accepted that CSCs are assembled in the ER where they would undergo quality control (Strasser, 2018). Dissecting specific molecular partners in CSC assembly in the ER is problematic, as ER-secreted proteins rely on a set of shared molecular chaperons for folding, so mutating these genes will likely exert pleiotropic effects unrelated to cellulose synthesis.

Assembled CSCs are assumed to be transported *via* direct streaming or in COPII vesicles to the Golgi where they are then modified before export (Neumann, Brandizzi, & Hawes, 2003). One study on *Arabidopsis* has indicated that CSCs may assemble in the Golgi with the assistance of Golgi-localized *STELLO(STL1/2)* proteins that have a glycosyltransferase (GT) domain. In *stl1 stl2* double mutants, primary and secondary CSCs were less abundant, CSC delivery rates were reduced and *CESA3* distribution was altered in the Golgi (Zhang, Nikolovski et al., 2016), which are all phenotypes consistent with defective Golgi assembly. Split-ubiquitin and BiFC assays confirmed that *STL1* and *STL2* could bind to all primary and secondary cell wall CESAs, but whether the precise interactions involve the *STELLO* GT domain was not tested (Zhang, Nikolovski et al., 2016). CSC assembly may also be facilitated by

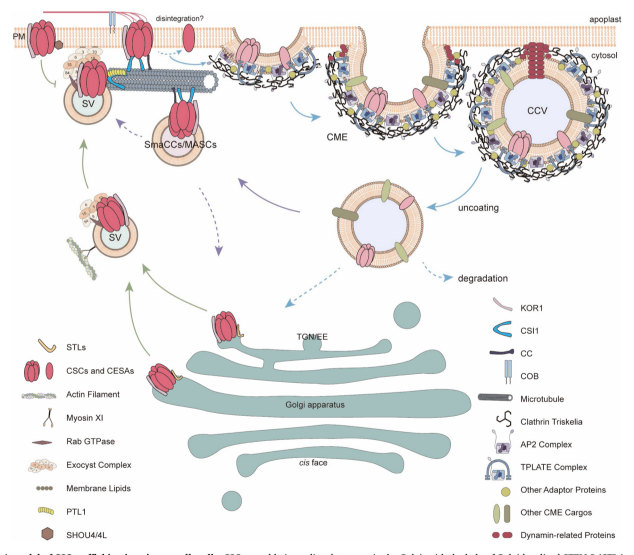


Fig. 2. A model of CSC trafficking in primary cell walls. CSC assembly is predicted to occur in the Golgi, with the help of Golgi-localized STELLO (STLs) proteins. CSCs are then transported from the TGN/EE and the Golgi via secretory vesicles (SV) and are delivered to specific sites on the plasma membrane that are marked by CSI1 proteins linked to microtubules. Myosin XI may also help deliver SV containing CSCs to the plasma membrane along actin filaments. Physical interactions between PATROL1 (PTL1) and the exocyst complex with SVs are required for the insertion of CSCs into the plasma membrane. SHOU4/4 L negatively regulates CSC delivery. Various non-CESA proteins are required for optimal cellulose biosynthesis, including COB and CC that associate with CSCs at the plasma membrane and KOR1 that additionally associates with the CSC during trafficking. Intact or degraded CSCs can be internalized into clathrin-coated vesicles (CCV) and undergo clathrin-mediated endocytosis (CME). Various CME components, such as the adaptor protein 2 complex (AP2), TPLATE complex (TPC) are essential for CME. Interna Internalized CSCs can be recycled back to the plasma membrane via SmaCCs/MASCs.

KOR1 and COB that co-localize with CESA proteins in the Golgi (Lei et al., 2014; Roudier et al., 2005; Vain et al., 2014), however, this has not been functionally assessed. Identifying how CESA proteins interact with accessory proteins in the Golgi or ER will fill in some of the crucial gaps in our understanding of CSC assembly.

4.2. CSC delivery

Although progress on CSC assembly has been slow, considerable knowledge has been gained in the trafficking of CSCs to the plasma membrane (Fig. 2). Early observations of intact CSCs in the Golgi and TGN/EE (Giddings et al., 1980; Haigler & Brown, 1986) were later reinforced by live-cell imaging of fluorescent CESA particles in the Golgi and TGN/EE (Crowell et al., 2009; Paredez et al., 2006). Therefore, trafficking of the CSC to the plasma membrane may occur from the Golgi via the TGN, or independently from the Golgi. The main route for CSC delivery is thought to occur via the Golgi, as various TGN markers, VHA-a1 and SYP61, do not co-localize with CSC membrane insertion events (Crowell et al., 2009). However, confirming this is problematic since the TGN/EE acts as both a secretory and recycling organelle, harbouring populations of both newly synthesized and recycled CSCs (Viotti et al., 2010). Two types of CSC trafficking vesicles have been identified depending on their microtubule associations: small cellulose synthase compartments (SmaCCs) (Gutierrez et al., 2009) and microtubule associated SmaCCs (MASCs) (Crowell et al., 2009). Partial co-localization of SmaCCs/MASCs with TGN/EE molecular markers implicates SmaCCs/MASCs in both the secretion of de novo synthesized CSCs, and CSC internalization. In general, SmaCCs/MASCs are regarded as endocytotic vesicles, though it has become evident that they confer multiple roles in CSC trafficking, complicating their study.

CSC delivery was observed in early TEM observations whereby vesicles containing terminal complexes frequently coincided with cortical microtubules underlying cell wall thickenings (Haigler & Brown, 1986; Hepler & Newcomb, 1964). Live-cell imaging of CESA3 and CESA6 with tubulin further corroborated the tight overlap between cortical microtubules and CSC delivery events in primary cell walls (Crowell et al., 2009; Gutierrez et al., 2009). Microtubules mark specific sites for CSC delivery as disrupting intact microtubules networks causes the random insertion of SmaCCs/MASCs into the membrane (Gutierrez et al., 2009; Paredez et al., 2006), and newly delivered CSCs track microtubule arrays after photo bleaching (Crowell et al., 2009). CSC insertion into the plasma membrane also coincides with Golgi pausing events immediately beneath sites of cortical microtubules (Crowell et al., 2009), implicating the Golgi in mediating CSC delivery. Although microtubule distribution also coincides with sites of secondary cell wall deposition in developing vessels, removing microtubules does not influence Golgi pausing events in the delivery of CESA7 (Wightman & Turner, 2008). Instead, transverse actin defines CSC delivery sites and actin cables indirectly transport CSCs to the plasma membrane, since actin depolymerization prevents CSC delivery and halts Golgi movement of CSCs (Wightman & Turner, 2008). While actin is not required for CSC insertion in primary cell walls (Sampathkumar et al., 2013), it may play a role in delivering CSCs to the membrane, as disrupting actin polymerization causes CESA3 and CESA6 Golgi bodies to aggregate beneath the membrane (Crowell et al., 2009; Gutierrez et al., 2009). Microtubules and actin involvement in CSC delivery may be spatially separated because subcortical SmaCCs have reduced velocity when actin cables were disrupted (Gutierrez et al., 2009). In the subcortical regions, actin may be responsible for the movement of the Golgi to sites of microtubules in the cell cortex that define domains for secretion. Recently, the actin-based motor protein, myosin XI, was implicated in the trafficking of CSCs in primary cell walls (Fig. 2). In triple xi1 xi2 xi3 mutants and wild-type plants treated with myosin inhibitors, CSC delivery is reduced and vesicles containing CSC vesicles accumulate below the membrane (Zhang, Cai, & Staiger, 2019). Failed delivery was attributed to defective vesicle tethering and fusion due to the overlap of XI with CESA6 fluorescent signals near the

membrane. Whether this is an indirect effect of defective Golgi pausing is unclear

Trafficking of any protein complex relies on four key phases - vesicle budding, cytosolic transport, tethering and ultimately fusion with the destination membrane. From studies on bacteria, yeast and animals it is obvious that delivery is co-ordinated by a complex interplay of proteins. Key players that have been identified include, Rab GTPases that target vesicles towards the destination membrane and mediate the fusion of the two membranes; tethering factors such as soluble N-ethylmaleimidesensitive factor attachment protein receptor (SNAREs) that assist with fusion; and the exocyst complex. Identifying candidates specific for CSC trafficking has been challenging due to the vast genetic redundancies of these families in Arabidopsis, combined with the fact that Rab GTPases and SNAREs associate with multiple cargoes (Uemura et al., 2012; Vernoud, Horton, Yang, & Nielsen, 2003). Some candidates have been identified, including the Golgi-localized Rab-H1B, whereby CESA6 has reduced motility and impaired exocytosis in loss of function mutants (He et al., 2018). However, their direct involvement in CSC trafficking has not been clear. Co-purification of CESA6 with the syntaxin SYP61, a sub-family of SNAREs, implicated SYP61 vesicles in the tethering of CESA6 to the membrane (Drakakaki et al., 2012). However, as SYP61 is a major component of post-Golgi vesicles it is unlikely to be specific for CSC trafficking. Ideally the purification of CESA specific compartments such as SmaCCs/MASCs would provide more insight into CSC delivery. However, due to their small size and low fluorescence signal this may prove difficult unless aggregated populations are extracted. Furthermore, extracting SmaCCs/MASCs during different points of CSC trafficking may be informative of the different genetic components involved in recycling and delivery, allowing the identification of specific markers of these processes.

Unsurprisingly, the precise temporal and spatial insertion of CSCs into the membrane is under tight genetic control. Coimmunoprecipitation of CSI1 identified several genetic components that assist with CSC delivery in primary cell walls (Fig. 2), including PATROL 1 (PTL1) and the exocyst subunits, SEC5B and SEC10 (Zhu et al., 2018). Mutagenesis combined with in vitro pull downs demonstrated that CSC delivery relies on intricate physical interactions between CESA6, CSI1, PTL1 and exocyst subunits. A model was developed for CSC delivery by carefully examining the temporal and spatial localization of these proteins during a live-cell imaging time-course (Zhu et al., 2018). CSII defines the domain in the plasma membrane for delivery and possibly acts as a direct tether of SmaCCs to the membrane, since SmaCC formation is reliant on the interaction between CSI1 and microtubules (Lei et al., 2015). After CSI1 interacts with the vesicle, PTL1 primes the vesicle for fusion by subunits of the exocyst complex, SEC5B and SEC10, that complete fusion. The association of PTL1 is fleeting but essential, as delivery rates are slower in ptl1, and ptl1 csi1 double mutants have an additive phenotype. Whether this mechanism is shared by secondary cell walls has not yet been established, although the accumulation of exocyst subunits and CSI1 during secondary cell wall deposition indicates that this could be a strong possibility (Derbyshire et al., 2015). Recently, SHOU4 proteins were identified as negatively regulators of CSC exocytosis (Polko et al., 2018). In shou4 shou4l double mutants enhanced CSC delivery is evident from an elevated density of CESA6 at the plasma membrane and an increase in amorphous cellulose content (Polko et al., 2018). Direct binding of the cytoplasmic domain of SHOU4 proteins with the catalytic domains of primary cell wall CESAs may cause the retention of CSCs in the cytoplasm, though this is speculative at present.

4.3. CSC endocytosis and recycling

The population of CSCs at the plasma membrane at any given time is determined by a balance between exocytosis and endocytosis and is often used as a proxy for the rate of cellulose synthesis. How this interplay is regulated is unknown, but it is possible that the plasma

membrane can monitor the density of CSC and subsequently mediate CSC delivery and recycling as part of a self-regulating feedback system. Supporting evidence has been provided from rabh1b CSC trafficking mutants that are defective in both exocytosis and endocytosis, suggesting that the two processes are inter-dependent (He et al., 2018). CSC recycling is inherently difficult to quantify, but it is widely believed to be a dominant process for several reasons. Firstly, CSCs have an average longevity of 30 min (Jacob-Wilk, Kurek, Hogan, & Delmer, 2006), yet typically, CSC membrane lifespan is only 7-8 min (Sampathkumar et al., 2013) suggesting CSC may be recycled several times before they are degraded. Secondly, as CSCs are large protein complexes, repeatedly constructing CSCs every 7 min would exert huge energetic costs on the cell. Finally, SmaCC/MASC populations tend to accumulate in cells not actively synthesizing cellulose, or cells under osmotic stress (Crowell et al., 2009; Gutierrez et al., 2009). In adverse conditions, SmaCCs/-MASCs may accumulate underneath the membrane as a 'temporary store' of CSCs that are rapidly recycled back once stress is alleviated. Furthermore, when protein and cellulose synthesis is inhibited, CESA3 accumulates in MASCs within 7 min suggesting internalization is more likely than de novo secretion (Gutierrez et al., 2009).

Clathrin-mediated endocytosis (CME) is a dominant process in Eukarvotes but it is only in the last two decades that its importance has been appreciated in plants (Holstein, 2002; Reynolds, Wang, Pan, & Bednarek, 2018). Clathrin coated vesicles were first observed in the TGN in the 1980s (Coleman, Evans, & Hawes, 1988) and 30 years later they were shown to be integral for CSC internalization (Fig. 2). Two components of CME machinery, AP2M and TWD40-2, have been implicated in CSC endocytosis (Bashline, Li, Anderson, Lei, & Gu, 2013; Bashline, Li, Zhu, & Gu, 2015). AP2M is homologous with a medium subunit of the adaptor protein complex 2 (AP2) that assists with docking and recruiting CME machinery (Bashline et al., 2013) and TWD40-2 is a potential member of a TPLATE complex (TPC) that is unique to plants (Gadeyne et al., 2014). In ap2m and twd40-2 mutants, reductions in endocytosis were inversely correlated with a higher density of CSC at the membrane (Bashline et al., 2013, 2015). Co-operation of TWD40-2 with AP2M is required for CME, as not only do they directly interact, but reductions in endocytosis and cellulose content are exacerbated in ap2m twd40-2double mutants (Bashline et al., 2015). However, they may confer distinct roles in CME, since hypocotyls exhibit reduced elongation in twd40-2 but have increased elongation in ap2m. TWD40-2 also has a much longer lifespan than AP2M during CME so it may be involved in scission from the membrane or quality control. Another TPC subunit, TML, can also interact with CESA6 catalytic units. tml knock-down lines exhibit similar decreases in cellulose content and an increased population of CSC at the membrane that is not attributable to increased delivery rates (Sanchez-Rodriguez et al., 2018). CME represents one route of CSC endocytosis, as SmaCCs/MASCs abundance is reduced, but not completely abolished in ap2m mutants (Lei et al., 2015). The TPLATE may have evolved to specifically tailor CME in plants or mediate endocytosis independently, so continued study of this complex will be revealing about CSC endocytosis.

CESAs have been described as *AP2M* cargo because *AP2M* can bind to *CESA6* and *CESA3* in split-ubiquitin assays and *in vitro* pull downs, and *mCHERRY::CESA6* patterns overlap with *YFP::AP2M* (Bashline et al., 2013). But discrepancies in their physical dimensions questions whether CSC can be internalized by CME, as the catalytic core of CSCs is 50 % wider than the lumen of typical CME particles (Bashline, Li, & Gu, 2014; Li et al., 2014). De-constructed complexes may be internalized as an increased number of CSC particles at the membrane coupled with decreased cellulose content in *twd40-2* mutants indicates CSC breakdown may start prior to internalization (Bashline et al., 2015). CSCs readily form monomers, dimers and trimers in solution under a range of conditions suggesting that CSCs may be easily broken down *in vivo* by local alterations in the membrane environment (Atanassov, Pittman, & Turner, 2009; Olek et al., 2014; Vandavasi et al., 2016). Alternatively, CSCs may appear larger if it is bound to other components that may be

released prior to endocytosis.

Whether endocytosed CSCs destined for proteolysis are trafficked to the vacuole directly or go via the TGN/EE is unclear, as both seem plausible (Crowell et al., 2009). Likewise, it is not known whether SmaCCs go to the TGN/EE before re-inserting CSCs into the membrane or bypass the TGN/EE altogether. At least some recycled CSCs pass through the TGN/EE as poor acidification of the TGN/EE in *det3* mutants causes defects in both secretion and recycling (Luo et al., 2015). As the TGN/EE is a sorting hub it would be convenient if all internalized CSCs travelled to the TGN/EE and were then exported for either recycling or degradation. Differentiating between populations of newly synthesized and recycled proteins that cross-over at the TGN/EE is a taxing question and has impeded research in this area. Determining the protein composition of vesicles involved at each stage of CSC trafficking may reveal markers that signify the destination of the vesicle, alleviating this problem. During the rapid changeover between primary and secondary cell wall synthesis, CSC exocytosis and endocytosis are temporally separated briefly, so could be probed to answer some of these outstanding questions. At the onset of secondary cell wall deposition in inducible VND7 tracheary elements, the tdTomato::CESA6 signal decreases in the membrane and increases at the Golgi, representing recently endocytosed primary cell wall CSCs. Once YFP::CESA7 starts appearing at the Golgi, the tdTomato::CESA6 signal disappears from the Golgi and a diffuse signal re-appears in the vacuole, indicating that the recently endocytosed tdTomato::CESA6 are transported to the vacuole during secondary cell wall deposition (Watanabe et al., 2018).

5. How is cellulose synthesis regulated?

Probing the molecular regulation of cellulose synthesis has only been possible in the last 20 years, due to significant advances in the generation of genetic mutants, genetic constructs, and next generation sequencing technologies. High-throughput sequencing has been used to explore the regulation of cellulose synthesis at multiple aspects including, genomic (DNA), transcriptional (mRNA), translational (proteins), and post translational processes (metabolites and small RNA), causing a marked shift in research focus from structural to molecular studies.

5.1. Transcriptional regulation

Since all cells have a primary cell wall and cells are continuously made throughout development, genes involved in primary cell wall synthesis are ubiquitously expressed (Hamann et al., 2004). As such, transcriptional regulators are likely to be housekeeping genes that are not specific for cellulose synthesis. Potential candidates have been identified in the ETHYLENE-RESPONSE-FACTOR (ERF) IIId and IIIe transcription factor family. Overexpressing ERF35produces thick cell walls with a primary cell wall composition in nst1 nst3 mutants that lack secondary cell walls Sakamoto, Somssich, & Mitsuda, 2018. Since many ERF transcription factors are co-expressed with CESA1, CESA3 and CESA6, and ERF34-ERF43 can physically bind to the promoters of primary cell wall CESA genes, the ERF transcription factor family may have a central role in regulating cellulose deposition in primary cell walls (Saelim et al., 2019; Sakamoto et al., 2018). Additionally, a brassinosteroid responsive transcription factor, BES1, can increase CESA expression by binding to the E-box (CANNTG) element in the promoters of CESA1, CESA3 and CESA6 (Xie, Yang, & Wang, 2011). However, BES1 is unlikely to be a specific activator of primary CESAs, as BES1 can simultaneously induce CESA4 and CESA8 expression.

In contrast to primary cell walls, the transcriptional network responsible for regulating cellulose synthesis during secondary cell wall formation has been extensively characterized in *Arabidopsis* (Yamaguchi et al., 2010; Zhong, Lee, Zhou, McCarthy, & Ye, 2008) and it is functionally conserved in woody species (Zhang, Xie, Tuskan, Muchero, & Chen, 2018) and grasses (Rao & Dixon, 2018). Two main transcription

factor families containing either NAC- or MYB domains, co-ordinate the expression of CESA biosynthetic genes. The NAC transcription factors, NAC SECONDARY WALL THICKENING PROMOTING FACTORS (NST1/2) and SECONDARY WALL-ASSOCIATED NAC DOMAIN PRO-TEIN (SND1) can activate cellulose synthesis in fibers, with snd1 nst1 double mutants exhibiting reduced cellulose content and impaired secondary cell wall formation (Zhong, Richardson, & Ye, 2007). NAC-domain transcription factors activate two downstream, transcription factors, MYB46 and MYB83, which are functionally redundant and MYB103 (Zhong et al., 2008). MYB103 can activate the expression of GUS reporter genes driven by the CESA8 promoter, implicating MYB103 as a specific regulator of cellulose deposition (Zhong et al., 2008). In cellulose-rich cotton fibers, MYB103 is one of the first transcription factors that is expressed during the changeover between primary and secondary cell wall deposition, providing further support that it is an important regulator of cellulose synthesis (MacMillan et al., 2017). Overexpressing MYB46 and MYB83 causes an increase in CESA expression that is accompanied by excessive cellulose deposition in ectopic cell walls, implicating MYB46/83 as direct activators of cellulose synthesis (Ko, Kim, & Han, 2009; McCarthy, Zhong, & Ye, 2009). MYB46 can specifically regulate CESA expression by binding to 8-bp MYB46-responsive cis regulatory elements (M46RE) in CESA promoters (Kim, Ko et al., 2013). Introducing CESA genes with point mutations in the M46RE into cesa mutants could not restore cellulose synthesis, demonstrating that MYB46 binding is crucial for regulating cellulose synthesis in Arabidopsis (Kim, Kim, Ko, Kim, & Han, 2013). Other direct targets of MYB46 include the CCCH zinc finger genes, C3H14 and C3H15 (Ko et al., 2009), which cause ectopic deposition of cellulose and upregulate CESA genes when overexpressed (Chai et al., 2015). More recently other transcription factor families have been implicated in cellulose synthesis regulation, such as WRKY and ERF. Cellulose deposition is stimulated in wrky12 mutants (Wang et al., 2010) or when the ERF transcription factor, PdSHINE2, is overexpressed in tobacco (Liu et al., 2017).

5.2. Post-translational regulation

Constitutive expression of CESA genes in primary cell walls implies that post-transcriptional regulation may be more important for regulating cellulose synthesis (Hamann et al., 2004). Arguably, the best studied form of post-translational regulation is phosphorylation (Speicher, Li, & Wallace, 2018). Phosphoproteomic analysis of primary CESA proteins demonstrated that many sites in the N-terminus and HVR of the central loop contain conserved serine (S) and threonine (T) residues that have the potential to be phosphorylated (Durek, Schudoma, Weckwerth, Selbig, & Walther, 2009; Nuhse, Stensballe, Jensen, & Peck, 2004). The effects of phosphorylation were first examined by mutating S and T sites to alanine (A) that eliminates phosphorylation, or glutamine (E) that mimics phosphorylation. Inhibiting phosphorylation at T¹⁶⁶A, S⁶⁸⁶A and S⁶⁸⁸A residues in the HVR of cesa1^{rsw1} mutants produced a variety of cellulose defective phenotypes, including reduced cellulose content, poor anisotropic cell expansion, reduced CSC velocity and the loss of bi-directional movement, which were all rescued when phosphorylation was restored (Chen, Ehrhardt, & Somerville, 2010). In contrast, permitting phosphorylation at S¹⁶²E, T¹⁶⁵E and S¹⁶⁷E in cesa1^{rsw1} mutants caused cellulose defective phenotypes, indicating that a balance between de-phosphorylation and phosphorylation finely tunes the regulation of CESA1 (Chen et al., 2010). Removing microtubules with oryzalin rescued the velocity and bi-directional movement of CESA1 at the membrane, supporting the idea that the phosphorylation of CESA proteins may modulate microfibril synthesis and anisotropic growth by its interaction with microtubules (Chen et al., 2010). Similar studies on CESA3 and CESA5 have reinforced that phosphorylation is critical for cellulose synthesis regulation in primary cell walls. For example, phosphorylation of S²¹¹A and de-phosphorylation of T²¹²E of CESA3 is crucial for maintaining anisotropy, deposition, bundling and bi-directional microtubule-based motility at the membrane (Chen et al.,

2016) and phosphorylating CESA5 alters the migration of CSC in a phytochrome dependent manner (Bischoff et al., 2011). Phosphorylation may also be important for mediating CESA endocytosis in the secondary cell wall since in vitro phosphorylation of CESA7 causes its degradation via the proteosome (Taylor and Turner, 2007). Identifying the corresponding protein kinases that activate phosphorylation has proved troublesome, with conflicting evidence in the literature and large genetic redundancies in kinase families (McFarlane et al., 2014). In Arabidopsis, the protein kinase BRASSINOSTEROID INSENSITIVE 2 (BIN2) can phosphorylate a CESA1 peptide, CESA1T157, in vitro (Sanchez-Rodriguez et al., 2017). In cesa1 bin2-1 double mutants, the CSC moves significantly faster at the membrane, implicating BIN2 as a negative regulator of cellulose synthesis in the primary cell wall (Sanchez-Rodriguez et al., 2017). BIN2 phosphorylation impacts the activity of the entire CSC, even though it cannot phosphorylate CESA3 or CESA6 peptides, demonstrating the importance of phosphorylation as a regulatory mechanism.

More recently it was revealed that secondary cell wall CESAs are heavily modified by the attachment of the fatty acid palmitate at conserved cysteine residues, also known as S-acylation. Mutating four cysteines in the VR2 and two cysteines in the C-terminal domain of CESA7 prevented the trafficking of CESA7 to the plasma membrane from the Golgi (Kumar et al., 2016). The role of S-acylation may be broadened to include other aspects of cellulose biosynthesis, since many important non-CESA proteins such as KOR1; CMU; CC; SHOU; PTL1; and CME components are also acylated (Kumar et al., 2020). Furthermore, heavy S-acylation of CESA3 and CSI1 suggests S-acylation may function in primary cell walls (Kumar et al., 2016, 2020). Although many of these assumptions have not been yet been functionally tested, it is probable that S-acylation is a dominant regulator of post-translational processes that we have only just begun to understand.

6. Significant achievements and future directions

Remarkably, many of the original hypotheses that were based on simple TEM observations and X-ray diffraction patterns in bacteria and algae, have stood the test of time and have been verified in higher plants by using a range of more accurate techniques (Table 1). Although revisiting other long-standing hypotheses with more sensitive techniques has revealed that some concepts are too simplistic to account for the diversity in cell wall architecture. Most notably the multi-net growth hypothesis is insufficient to explain anisotropy in all conditions and the relationship between CSCs and microtubules is not universally coupled. Significant progress in our capacities to study cellulose synthesis in vivo with live-cell imaging, AFM, FESEM and molecular genetics has resulted in some drastic changes in our understanding of some key aspects of cellulose synthesis, and in some cases has divided research groups. In the last 10 years, the 36-glucan chain model has been disregarded in favor of an 18-24 chain model, new models of cell elongation have been proposed and even the classic 8 transmembrane CESA-model has been brought into question (Table 1). While Arabidopsis has proved an invaluable model for enhancing our understanding of cellulose synthesis, these results need to be approached with caution as this system may not be representative of higher plants in general. Broadening the sample types may help settle variable findings between research groups and will strengthen the validity of hypotheses across higher plants. With many unanswered or modified hypotheses still requiring verification (Table 1), we can expect many great discoveries and changes in the field during this century. Adopting multidisciplinary strategies that link together the biophysical and biochemical properties of cellulose with underlying genetics and cell wall architecture, will be fundamental for this venture. Successful purification of CSCs, imaging the entire CSC in situ and assigning functions to microfibril properties are arguably the next major breakthroughs on the agenda in order to advance the study of cellulose synthesis, as such fundamental knowledge will be critical to eventually manipulate cellulose synthesis for desired use.

CRediT authorship contribution statement

Holly Allen: Conceptualization, Writing - original draft, Writing - review & editing. Donghui Wei: Conceptualization, Visualization, Writing - review & editing. Ying Gu: Conceptualization, Writing - review & editing, Funding acquisition. Shundai Li: Conceptualization, Project administration, Writing - review & editing.

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