Evolution and molecular basis of substrate specificity in a 3-ketoacyl-CoA synthase gene cluster from *Populus trichocarpa*

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Very long chain fatty acids (VLCFAs) are precursors to sphingolipids, glycerophospholipids, and plant cuticular waxes. In plants, members of a large 3-ketoacyl-CoA synthase (KCS) gene family catalyze the substrate-specific elongation of VLCFAs. Although it is well understood that KCSs have evolved to use diverse substrates, the underlying molecular determinants of their specificity are still unclear. In this study, we exploited the sequence similarity of a KCS gene cluster from *Populus trichocarpa* to examine the evolution and molecular determinants of KCS substrate specificity. Functional characterization of five members (PtKCS1, 2, 4, 8, 9) in yeast showed divergent product profiles based on VLCFA length, saturation, and position of the double bond. In addition, homology models, rationally designed chimeras, and site-directed mutants were used to identify two key regions (helix-4 and position 277) as being major determinants of substrate specificity. These results were corroborated with chimeras involving a more distantly related KCS, PtCER6 (the poplar ortholog of the *Arabidopsis* CER6), and used to show that helix-4 is necessary for the modulatory effect of PtCER2-like5 on KCS substrate specificity. The role of position 277 in limiting product length was further tested by substitution with smaller amino acids, which shifted specificity toward longer products. Finally, treatment with KCS inhibitors (K3 herbicides) showed varying inhibitor sensitivities between the duplicated paralogs despite their sequence similarity. Together, this work sheds light on the molecular mechanisms driving substrate diversification in the KCS family and lays the groundwork for tailoring the production of specific VLCFAs.

Very long chain fatty acids (VLCFAs) have a backbone of 20 or more carbons and are precursors for important cellular lipids such as sphingolipids, glycerophospholipids, and triacylglycerols (1–3). VLCFAs and their derivatives regulate a variety of physiological and developmental processes in eukaryotes (4–7). In higher plants, VLCFAs play an additional role in the biosynthesis of cuticular waxes, a hydrophobic film deposited on aerial surfaces that provides protection against environmental stressors such as desiccation, infection, and herbivory (8–10). Among most characterized plants, the predominant components of cuticular waxes are aliphatic, functionalized hydrocarbons derived from VLCFAs (11).

VLCFAs are biosynthesized by the iterative two-carbon elongation of C16-C18 long chain fatty acyl-CoAs (12). In plants, a key enzyme in VLCFA elongation is the β- or 3-ketoacyl-CoA synthase (KCS), an integral membrane protein that forms parts of the microsomal fatty acid elongation (FAE) complex (13, 14). The FAE complex consists of four core enzymes that each catalyze one of the four steps in VLCFA elongation (15). The first reaction, catalyzed by KCS, condenses a fatty acyl-CoA with malonyl-CoA to generate a 3-ketoacyl-CoA. Subsequent reduction of the carbonyl by ketoacyl-CoA reductase, dehydration of the hydroxyl group by hydroxyacyl-CoA dehydratase, and reduction of the double bond by enoyl-CoA reductase yield a fatty acyl-CoA extended by two carbons (16). This extended very long chain fatty acyl-CoA can be used for another round of elongation; however, this depends on the substrate specificity of the condensing enzyme.

Previous work has shown that KCS catalyzes the only substrate-specific step in the FAE complex, whereas the other three enzymes are generalists (17). Because of this, KCSs have expanded into a large, multigenic family with diverse substrates that have yet to be fully explored (18–20). To date, the substrates of characterized KCSs include fatty acids with lengths ranging from C16-C36, containing double bonds or hydroxyl groups (20–25). The spatiotemporal expression of KCSs with different substrate specificities modulates the VLCFA profiles of different tissues. For example, expression of KCS16 in *Arabidopsis* trichomes results in the production of exceptionally long VLCFAs (up to C38), whereas seed-specific expression of KCS18/FAE1 results in the accumulation of shorter unsaturated gondoic (C20:1) and erucic (C22:1) acid in storage triacylglycerols (17, 25). The epidermis expressed *Arabidopsis* KCS6/CER6/CUT1 produces VLCFAs longer than C26 that are important precursors to cuticular waxes, particularly in the stem (26, 27). The substrate specificity of CER6 has also been shown to be modulated by a binding partner, CER2, which modifies the elongation capability of CER6 to even longer VLCFAs (14, 28, 29). CER2s and their paralogs (termed CER2-likes) are part of the functionally diverse BAHD-acyltransferase family and have been characterized in Arabidopsis, rice, maize, sacred lotus, and poplar (28–33).
Substrate specificity of a KCS gene cluster

While studies on other fatty acid biosynthetic and modifying enzymes have explored the molecular basis of substrate specificity, such studies in KCSs are limited by the lack of structural resources and commercially available very long chain fatty acyl-CoA substrates, and insolubility of membrane proteins (34–36). A way around this is to heterologously express KCS enzymes in yeast (Saccharomyces cerevisiae) and compare their VLCFA profiles to the empty control strain (37). The differences in VLCFA product profiles are indicative of the distinct substrate specificities of the expressed KCS. The yeast elongase complex is compatible with plant KCSs and can complete the other three reactions of the FAE complex (i.e., reduction, dehydration, and reduction). Previous work has shown that substrate preference from in vitro assays with purified enzymes and assays comparing yeast profiles in vivo are comparable, indicating that heterologous expression in yeast can be used to assess activity toward different VLCFAs (12, 37, 38). With this approach, chimeric proteins expressed in yeast were used to compare the substrate specificity between the FAE1 orthologs from Arabidopsis thaliana and Brassica napus. The first 173 residues of the N-terminal region were shown to convert the product ratio of C20:1 to C22:1 from one KCS to another (37). It is presently unknown whether these results are generalizable to other KCSs, and regions that affect specificity toward unsaturated substrates or substrates longer than C22 have yet to be identified. Interestingly, distant members of the KCS family show differential inhibitor specificity toward K3 herbicides, although the molecular basis of this inhibition is also still unclear (38).

Recently, a large screen of cuticular waxes from a collection of wild genotypes of Populus trichocarpa led to the identification of a KCS with substrate preference toward monounsaturated VLCFAs (PotriKCS1/PtKCS1) (24). This study found an alkene homologous series (ranging in length from C\textsubscript{23} to C\textsubscript{34}) as major components of the abaxial wax of P. trichocarpa leaves. Moreover, striking variation in the content of alkenes across samples was found, with most genotypes showing an increase in alkene accumulation as leaves expanded and a few genotypes failing to accumulate this compound class in their wax. When grown in a common garden, alkene-accumulating genotypes exhibited larger weight and diameter and greater resistance to the leafspot, suggesting that this trait confers an important advantage to P. trichocarpa. Comparison of the transcriptomes of contrasting genotypes pointed at PtKCS1 as the candidate gene responsible for the differences. The expression of PtKCS1 was found to be associated with the accumulation of very long chain alkenes in leaf cuticular waxes. When expressed in yeast, PtKCS1 showed substrate preference toward monounsaturated VLCFAs, consistent with a proposed pathway where these VLCFAs are decarboxylated to yield alkenes. Interestingly, PtKCS1 is part of a tandem duplicated cluster of eight KCS-encoding genes on chromosome (Chr) 10, which share a high degree of homology (24). Unlike PtKCS1, expression of the closely related PtKCS2 in yeast (S. cerevisiae) showed higher relative preference toward the elongation of saturated VLCFAs. Notably, an evolutionary hallmark of specialized metabolism is the duplication of biosynthetic genes, after which subfunctionalization or neofunctionalization can result in diversified gene functions (39).

Although the activity of several KCSs has been previously characterized, little is known about how substrate specificity evolves immediately following gene duplication and which amino acid substitutions drive these changes. To address these questions, we examined the Chr 10 cluster of KCSs from P. trichocarpa. First, we explored the evolutionary history of this gene cluster, revealing ancient and recent duplication events. Then, elongation assays of five paralogs (PtKCS1, 2, 4, 8, 9) were performed to compare substrate preferences toward saturated or monounsaturated VLCFAs of varying lengths. Based on these results, chimeric proteins and site-directed mutagenesis were used to identify two key regions (helix-4 and residue 277) affecting substrate specificity in this clade. Furthermore, these regions were confirmed to alter substrate specificity in a more distantly related KCS, PtCER6, and affect the synergistic relationship between PtCER6 and a CER2 (PtCER2-like)5. We used these findings to rationally engineer longer VLCFA products by substituting smaller amino acids at position 277. Finally, we showed that the paralogs of the gene cluster exhibited differential inhibition by K3 herbicides despite their sequence similarities. Our results shed light on the molecular basis of KCS substrate specificity, highlighting the plasticity and epistatic nature of substrate binding in KCSs.

Results

Evolution of a KCS gene cluster in P. trichocarpa

In P. trichocarpa (v3.0), Chr 10 harbors a tandem duplicated gene cluster encoding eight KCS ORFs within a 215 kb region (Fig. 1A). To examine the origin of the gene cluster and relationship between the paralogous genes, we first conducted a phylogenetic analysis of all KCSs from P. trichocarpa. This analysis identified another closely related KCS on Chr 8 (Fig. 1, A and B). Since Chr 8 and 10 arose from a whole genome duplication (WGD) event approximately 60 million years ago, the most parsimonious explanation is that following WGD, the copy on Chr 10 expanded by tandem duplication (40). The phylogenetic tree indicates that the PtKCS1 clade can be further divided into two subclades based on sequence similarity: subclade A (PtKCS1, 3, 5, 7, 9, 11) and subclade B (PtKCS2, 4, and 8). Furthermore, subclade B genes are more similar to the copy on Chr 8 than to adjacent subclade A genes on Chr 10 in terms of coding sequence, promoter sequence, and expression profiles, indicating that the subclade divergence preceded the WGD event (Figs. 1B, S1, and S2). Therefore, phylogenetic evidence supports a model in which tandem duplication of an ancestral KCS, followed by WGD, gene loss on Chr 8, and additional independent tandem duplications on Chr 10 have led to the current architecture of the gene cluster in P. trichocarpa (Fig. 1C). In addition to multiple KCS copies, this region has a high density of repetitive elements (Fig. 1D). Notably, we found that the gene cluster is disproportionately rich in Helitrons (DNA-transposons) relative to adjacent regions on Chr 10 (Fig. S3). Additional evidence is
needed to distinguish whether the high density of Helitrons is due to active transposition, by-products of uneven crossovers during cell division, or a combination of both events. Nevertheless, these observations suggest that this locus is prone to retaining repetitive elements.

As a result of the independent duplication events, members of the PtKCS1 clade have varying degrees of relatedness, sharing 74% to 99% amino acid identity between the nine copies (Table S1). Previous work has demonstrated that PtKCS1 has a higher preference toward unsaturated VLCFAs relative to PtKCS2, evidence of functional divergence in this clade (24). Thus, we hypothesized that other PtKCS1 clade members could serve as a model to understand the evolution of KCS substrate specificity on different timescales. Additionally, their high sequence similarity could help narrow down the molecular determinants of KCS substrate specificity.

**PtKCS1 clade members have divergent VLCFA substrate specificities**

KCS enzymes specialize in the substrates they can use for the condensing reaction. Hence, the VLCFA product profile observed in yeast will be reflective of the differences in substrates of the heterologously expressed KCS. To investigate the extent of substrate specificity divergence in the PtKCS1 clade, we chose to compare the elongation activity of five members: PtKCS1, 2, 4, 8, 9 (Fig. S4). Elongation activity was assayed using heterologous expression in yeast followed by fatty acid methyl ester (FAME) analysis (37). Yeast accumulates endogenous VLCFAs (predominantly C26 and minor levels of C20 and C20:1) as a result of the native ELO2 and ELO3 enzymes that share no homology to KCSs (Fig. 2A) (12, 41). However, KCS activity can be detected by the increase in accumulated VLCFAs relative to the empty vector strain. In yeast, the five enzymes synthesized varying amounts of saturated and

**Figure 1. PtKCS1 gene cluster in P. trichocarpa.** A, diagram of the locus on Chr 8 and 10 containing PtKCS1 clade members (bolded). B, maximum-likelihood phylogenetic tree of all KCS protein sequences in P. trichocarpa, excluding sequences shorter than 200 amino acids. Bootstrap values are represented by line color. Scale bar for branch length represents 0.2 amino acid substitutions per site. The asterisk denotes the gene corresponding to PtCER6. C, proposed evolutionary model of the PtKCS1 gene cluster from an ancestral KCS. D, kernel density of annotated transposable elements across a 3 Mb region spanning Chr 10: position 9,000,000...12,000,000, on the P. trichocarpa v3.0 genome. A gene map is shown beneath the graph, with red bars representing KCS-encoding genes. Chr, chromosome.
monounsaturated VLCFAs ranging from C20-C30 in length (Fig. 2, A and B). Using authentic standards, we identified two series of monounsaturated products: one corresponding to the cis-ω9 double bond configuration and the other to the cis-ω7 configuration (Fig. S5). The substrates and products of the PtKCS1 clade members can be described in terms of three properties: carbon length, unsaturation, and position of the double bond. Subclade A members (PtKCS1 and 9) generally showed higher accumulation of shorter products relative to PtKCS2 and 8 from subclade B (Fig. 2B). In terms of unsaturation, PtKCS2 and 8 showed preference toward the production of saturated VLCFAs and cis-ω7-monounsaturated substrates. Conversely, subclade A members (PtKCS1 and 9) showed higher production of cis-ω9-monounsaturated fatty acids. The only exception was PtKCS4, which accumulated both shorter and cis-ω9-monounsaturated VLCFAs (like subclade A) but also high levels of saturated products (like subclade B). Remarkably, due to their high sequence similarity (99%) and homology, PtKCS2 and PtKCS4 only differ at five amino acids. Four of these residues account for higher activity toward monounsaturated VLCFAs, demonstrating that KCS substrate specificity can evolve rapidly following gene duplication.

To better understand KCS activity toward the different classes of VLCFAs, we generated a metabolite-metabolite correlation map, showing that the products of these KCSs segregated into three main clusters: (1) saturated and cis-ω7 monounsaturated, (2) cis-ω9 monounsaturated, and (3) saturated VLCFAs with a chain length > C26 (Fig. 2C). These results suggest that the substrate specificity of these KCSs is predominantly determined by the presence and position of the double bond as opposed to chain length, apart from VLCFAs longer than C26. Furthermore, the tight correlation of cis-ω7 monounsaturated products with saturated products likely results from the double bond being closer to the terminal methyl end of the VLCFA; this is opposed to cis-ω9 VLCFAs which contain a kink closer to the middle of the hydrocarbon tail and thus may be more sterically constrained.
Homology modeling and ligand docking of PtKCS1, 2, and 4 show differences in the putative substrate-binding site

To gain insight into the putative substrate-binding site of the KCS, we built a homology model of the cytosolic portion of PtKCS1 using Alphafold2 with templates from the closely related Type III polyketide synthases (PKSs; Fig. 3A, Tables S2 and S3) (42, 43). The predicted PtKCS1 structure shows the characteristic five layer αβαβα fold that is conserved between Type III PKSs, 3-ketoacyl-ACP synthases, and thiolases (44–46). To identify putative substrate-binding domains, we docked docosanoic acid (a C22:0 fatty acyl chain) and malonyl-CoA (the extender substrate) onto the homology model (Fig. 3A). Malonyl-CoA and docosanoic acid bind to distinct regions of the protein but converge on the catalytic triad as expected for the condensation reaction. Though a free fatty acid is not generated in the proposed KCS reaction mechanism, docosanoic acid represents the fatty acyl chain that is covalently attached to the active site cysteine (47). The fatty acid docks to a large hydrophobic tunnel that extends from the active site and is only accessible through the core of the protein. Comparison with models of PtKCS2 and 4 shows that the binding tunnel in PtKCS1 is kinked compared to PtKCS2 and 4, which could explain the relative preference of PtKCS1 for cis-mono-unsaturated compared to saturated substrates (Fig. 3B). The binding tunnels of PtKCS2 and 4 are similarly shaped but differ near the end of the tunnel around helix-4 (residues 131–141).

The amino acid sequences of PtKCS1, 2, and 4 show distinct regions with varying degrees of conservation (Fig. 3C). For

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**Figure 3.** Structural and sequence comparisons of PtKCS1, 2, and 4. **A,** homology model of the cytosolic portion of PtKCS1 docked with docosanoic acid (black) and malonyl-CoA (magenta), with the catalytic triad shown as yellow sticks. Alpha helix 4 and position 277 are highlighted by an orange dotted line. **B,** comparison of the binding tunnel in PtKCS1, 2, and 4 homology models, shown by surface representation and colored according to hydrophobicity. Docosanoic acid (black) is docked to each model with residues of the catalytic triad shown as yellow sticks. **C,** sequence alignment of PtKCS1, 2, and 4 including predicted secondary structures from PtKCS1. The five amino acid differences between PtKCS2 and 4 are highlighted in turquoise, and the three catalytic residues are highlighted in yellow. Predicted transmembrane helices are labeled (TM).
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example, the C-terminal half of the proteins is more conserved compared to the N terminus (Fig. 5). When plotted on the homology model of PtKCS1, most substitutions line the putative VLCFA binding tunnel by helices-4, 6, 7, and 11 (Fig. 5). Conversely, helix-5 and 12, which are positioned by the CoA moiety, contain few substitutions between paralogs. As recognition of CoA thioesters is a conserved feature across KCSs and even Type III PKSs, mutations are more likely to be deleterious in this region.

**Site-directed mutagenesis of two residues switches the substrate preference of PtKCS4 to 2**

PtKCS2 and 4 have divergent substrate preferences despite only having four amino acid differences in the cytosolic domain and one in the transmembrane domain (Fig. 3). These substitutions are sufficient to confer relative preferential elongation of shorter, monounsaturated VLCFAs in PtKCS4 and relative preferential elongation of longer saturated VLCFAs in PtKCS2, especially C28 (Fig. 2A). To determine the effect on substrate specificity, we used site-directed mutagenesis to convert each of the four different residues on PtKCS4 to the corresponding residue from PtKCS2 (Fig. 4A). Mutagenesis of R283C or T137K reduced the production of monounsaturated VLCFAs, while mutagenesis of T137K on helix-4 additionally increased production of saturated C28 VLCFA (Fig. 4, A and B). To test whether the combined effect of two substitutions could convert the overall product profile to that of PtKCS2, we generated a double mutant, PtKCS4T137K,R283C. By changing these two residues, the overall substrate preference of PtKCS4 was almost completely converted to that of PtKCS2 (Fig. 4 and Table S4). Notably, there was a synergistic increase in C28 in the double mutant compared to either mutant alone, showing these positions have epistatic effects on substrate specificity.

**Identifying the molecular determinants of substrate specificity between PtKCS1 and 2**

Compared to PtKCS2 and 4, PtKCS1 has a higher relative preference toward cis-monounsaturated compared to saturated VLCFAs (Fig. 2). To test whether the two sites we identified in PtKCS2 and 4 are also necessary for substrate specificity between PtKCS1 and 2, we generated site-directed mutants at these positions, corresponding to residues 131 and 277 in PtKCS1 (Fig. 5, Group I). The activity toward cis-monounsaturated fatty acids was reduced in the single and double mutants, showing that these sites also affect substrate specificity between PtKCS1 and 2 (Fig. 5 and Table S5). The combined effect of the double mutant on accumulation of cis-monounsaturated VLCFAs was greater than either single mutant alone, further indicating that these two positions have synergistic effects. Though the PtKCS1W277C mutant only had a minor effect on monounsaturated VLCFAs, there was a large increase in C28, characteristic of PtKCS2 activity. In contrast, mutating residue 275 (PtKCS1R275L) resulted in a product profile identical to PtKCS1, showing that not all divergent positions near the binding tunnel affect substrate specificity.

Residues 131 and 277 affected substrate preference but were not sufficient to entirely convert the substrate specificity of

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**Figure 4. Substrate specificity of site-directed mutants between PtKCS4 and PtKCS2.** A, diagrams of the site-directed mutants and respective production of cis-ω9-monounsaturated, saturated, and C28 VLC-FAMEs expressed in nmol per gram of lyophilized yeast. Reference lines indicate levels of VLC-FAMEs in the empty vector (gray), PtKCS2 (salmon), and PtKCS4 (blue). Data represent means of three biological replicates ± SEM. B, close-up view of the region in PtKCS2 and 4 where docosanoic acid (black) is docked. Amino acid differences between PtKCS2 and 4 are labeled and shown in stick representation. The residues of the catalytic triad are shown as yellow sticks. Alpha helix 4 and position 283 are highlighted by an orange dotted line. VLC-FAME, very long chain fatty acid methyl ester.
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PtKCS1 to PtKCS2. This is expected since PtKCS2 and 4 share 99% identity, whereas PtKCS1 and 2 only share 74% identity. Therefore, we looked to identify the molecular basis of substrate specificity between PtKCS1 and 2 using a fusion protein approach guided by natural polymorphisms between paralogs, hydrophobicity plots, and homology models (Fig. 5A). Werationally designed chimeric proteins by fusing varying lengths of the PtKCS1 N terminus to the PtKCS2 C terminus (Fig. 3A; Group II). PtKCS21-129, which only contains the transmembrane domain from PtKCS1, showed identical activity to PtKCS2, indicating that this region does not affect substrate specificity, consistent with previous reports that substrates are delivered cytosolically and not through the membrane (37) (Fig. 5). The first 129 residues swapped between PtKCS1 and 2 were also not sufficient to modify substrate specificity (as shown by PtKCS21,129). Conversely, PtKCS21-277 showed almost identical activity to PtKCS1, demonstrating that the first 277 residues from PtKCS1 are sufficient for converting the substrate specificity of PtKCS2 to PtKCS1. We generated a reciprocal fusion with the N terminus of PtKCS2 fused to the C terminus of PtKCS1 (PtKCS11,283), but this abolished the elongation activity in yeast, which could be due to unfavorable interactions in the tertiary structure.

To further narrow down the specific regions required for PtKCS1 substrate specificity, we generated chimeras to swap smaller regions of PtKCS2 with the corresponding region from PtKCS1 (Fig. 5A; Group III). PtKCS21,283 was still sufficient to confer PtKCS1 substrate specificity though there was a minor loss of activity. Surprisingly, taking the fusion with reduced activity (PtKCS21,181) and replacing residues 274 to 277 by helix-11 (generating PtKCS21,181,280-283) was sufficient to convert the substrate specificity to PtKCS1. Narrowing the region down to only 25 amino acid differences between PtKCS1 and PtKCS2 in the PtKCS21,283, 280-283 caused a large reduction in activity, but this fusion was still sufficient to increase the relative preference toward monounsaturated substrates. This fusion contains substitutions that span helices-4, 5, 6, and 11. Though the transmembrane domain does not affect substrate specificity, it appears to be essential for KCS activity in yeast, which was further confirmed by a transmembrane domain deletion (PtKCS1del1-101) that showed no elongation activity.

Interestingly, only nine substitutions around helix-4 (PtKCS21,129-143) were sufficient to increase the production of monounsaturated fatty acids (predominantly C20:1 and C22:1) without affecting activity toward saturated VLCFAs. Therefore, helix-4 is important in conferring activity toward monounsaturated substrates, but additional regions by helices-5, 6, and 11 are important in discriminating against saturated products. These combined analyses identify the key regions (in the N terminus and at position 277) that contribute to substrate preference between PtKCS1 and 2.

The N-terminal region controls substrate specificity in a more distantly related KCS

The results aforementioned show that helix-4 and site 277 are key determinants of substrate specificity in the PtKCS1 clade. To test whether these results can be extrapolated to more distantly related KCSs, we cloned PtCER6 (Potri.010G125300), sharing 59% amino acid identity with PtKCS1, for subsequent fusion and site-directed mutagenesis studies (Fig. 1B). Based on sequence similarity, PtCER6 is the putative poplar ortholog of AtCER6, a KCS from Arabidopsis which produces mainly C28 VLCFAs in yeast (28). Functional characterization of PtCER6 showed similar accumulation of predominantly C28 VLCFAs, markedly different from the broad product profile of PtKCS1 (Fig. 6A and Table S6). Reciprocal swaps of the N-terminal region between PtKCS1 and PtCER6 (PtCER61,279 and PtKCS11,283) resulted in empty control levels of VLCFA accumulation, likely due to the disruption of epistatic interactions on the protein tertiary structure as observed in the PtKCS11,283 mutant between PtKCS1 and 2 (Figs. 6, A–C and 5). However, swapping helix-4

![Figure 5. Substrate specificity of fusion proteins and site-directed mutants between PtKCS1 and PtKCS2.](image)

Diagrams of the fusion proteins and site-directed mutants between PtKCS1 and PtKCS2. Reference lines indicate levels of VLC-FAMEs in the empty vector (gray), PtKCS1 (teal), and PtKCS2 (salmon). The asterisk on PtKCS21,181, 280-283 indicates that the actual position of the second swapped region is 274 to 277, but this corresponds to position 280 to 283 on the rest of the Group III fusions. Data represent means of three biological replicates ± SEM. VLC-FAME, very long chain fatty acid methyl ester.
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Figure 6. Effect of helix-4 and site 277 on the substrate specificity of PtCER6. A, diagrams of chimeras and site-directed mutants between PtKCS1 and PtCER6 and their respective production of cis-9-mono-unsaturated, saturated, and C28 VLC-FAMEs expressed in nmol per g of lyophilized yeast. Data represent means of three biological replicates ± SEM. B, representative total ion chromatograms of PtCER6, PtKCS1, and PtCER6(Y258W). C, chain length distribution of saturated VLC-FAMEs produced by chimeras expressed with or without PtCER2-like5. Data represent means of three biological replicates ± SEM. VLC-FAME, very long chain fatty acid methyl ester.

...with the homologous region from PtKCS1 (PtCER6106-118) was sufficient to increase production of monounsaturated fatty acids by 3.4-fold and levels of C20-C24 VLCFAs at the expense of C28 VLCFAs, resembling the product profile observed in the PtKCS1 gene cluster (Fig. 6B). A single mutation in PtCER6 (PtCER6(Y258W)) also caused a shift toward shorter VLCFAs characteristic of PtKCS1 but was not sufficient to increase specificity toward unsaturated products. Our results show that helix-4 appears to be a major determinant for specificity across KCSs. Furthermore, this region could be changed without deleterious effects on activity, which may be explained by the placement of helix-4 near the surface of the homology model and relative isolation from other secondary structures (Fig. 3A).

The elongation activity of certain KCSs is modulated by a binding partner, CER2, through an unknown biochemical mechanism (29). In Arabidopsis, the current model proposes that AtCER2/CER2-like proteins can extend the chain length of AtCER6 products beyond C28. To test whether PtCER2s could extend the chain length specificity of different poplar KCSs and identify which regions on the KCS are important for this synergistic interaction, we coexpressed a previously characterized PtCER2-like5 with different KCSs and chimeras (Fig. 6A) (33). When PtCER2-like5 was coexpressed with PtCER6, the product specificity was shifted toward C30 VLCFAs (Fig. 6C and Table S6). Coexpression with PtKCS1 did not shift the products toward longer VLCFAs. Remarkably, the product specificity of PtKCS11-260 and PtCER6(Y258W) was partly modified by PtCER2-like5 toward longer VLCFAs, indicating that the first N-terminal 1 to 260 residues of PtCER6 are sufficient for the modulatory effects of PtCER2-like5 on KCS substrate specificity. Furthermore, constructs containing helix-4 from PtKCS1 (PtCER6106-118 and PtCER6106-118, Y258W) abolished the effect of PtCER2-like5 on extending the length of PtCER6 products, indicating this structure is necessary for the modulatory effect of PtCER2-like5 on PtCER6. Interestingly, coexpression of PtCER2-like5 increased the total production of VLCFAs in PtKCS1, PtCER6106-118, and the empty vector control, an effect which was reversed in PtCER6 chimeras containing the Y258W mutation. Although the increase could be due to interactions of PtCER2-like5 with the endogenous ELO enzymes, these results suggest an additional role of CER2s in regulating VLCFA biosynthesis that is independent of the role in extending elongation beyond C28 (Fig. 6C).

Engineering longer VLCFAs by substituting smaller amino acids at position 277

Since position 277 (position 283 in PtKCS2 and 4) appears to have an important effect on the length of KCS products, we tested whether products could be rationally engineered toward
longer VLCFAs by substituting smaller amino acids at this position. We mutated the amino acids at this position in PtKCS1 (PtKCS1_W277V) and PtKCS4 (PtKCS4_R283G). While both mutants had a similar substrate specificity to the WT, the levels of C28 increased at the expense of shorter VLCFAs, suggesting that the residue at 277 can act as a molecular ruler to regulate product length (Figs. 7A and S9).

Next, we tested whether these mutants had activity toward VLCFAs with additional functional groups. The fatty acids in yeast that are available to KCSs as substrates are saturated and monounsaturated fatty acids (predominantly C18:0 and C18:1), but KCSs could have activity toward other substrates, such as polyunsaturated or hydroxylated fatty acids. To test this, yeast expressing KCSs were exogenously fed with linoleic, linolenic, and ricinoleic acid, corresponding to C18:2, C18:3, and ω-3, and hydroxylated fatty acids, respectively (Fig. 7B). Most KCSs, including the PtKCS1_W277V mutant, had little to no activity toward these substrates beyond the background activity of the empty control (Fig. 7C). Remarkably, the PtKCS4_R283G mutant could uniquely elongate diunsaturated fatty acids up to C26:2 and triunsaturated fatty acids up to C24:3, corresponding to four and three cycles of elongation, respectively. Together, these results show that KCSs can be engineered to tailor VLCFA biosynthesis and introduce novel substrate specificities.

### Members of the PtKCS1 gene cluster are differentially inhibited by K3 herbicides

Acetamides, chloroacetamides, oxacylamides, and tetrazolinones are part of the widely used class K3 herbicides that target KCSs to inhibit VLCFA biosynthesis in plants (48, 49). Since the duplicated KCSs exhibited divergent substrate preferences, we wanted to test whether they could also be differentially inhibited by K3 herbicides, which has been shown for distantly related KCSs from Arabidopsis (38). Since endogenous yeast elongases are not affected by K3 herbicides, we tested the effect of four inhibitors (alachlor, anilofos, fenazamide, and flufenacet) on the elongase activity of PtKCS1, 2, and 4 using the yeast system (38). The substrate preferences of PtKCS1, 2, and 4 cover the major differences in activity observed in our studies, which is why they were chosen for the herbicide assay. The effect of inhibition varied between the three paralogs (Fig. 8A). Flufenacet and fentrazamide had the greatest effect on reducing VLCFA elongation in all three KCSs, whereas alachlor and anilofos slightly reduced activity. PtKCS1 was the most sensitive to inhibition, although there were also minor differences between PtKCS2 and 4. Since PtKCS2 and 4 only differ by five amino acids, these results imply that at least one of these residues is involved in conferring sensitivity to alachlor and anilofos.

To assess the inhibitor sensitivity of the different KCSs, we assayed PtKCS1, 2, and 4 in the presence of increasing concentrations of fentrazamide. PtKCS2 and 4 showed similar dose-response curves, whereas PtKCS1 was more sensitive to fentrazamide at lower concentrations (Fig. 8B). Since PtKCS1 and PtKCS2 have a different sensitivity to fentrazamide, we wanted to test whether KCS inhibition could be uncoupled from substrate specificity. Therefore, we repeated the experiment with PtKCS2_1-277, a chimera between PtKCS1 and PtKCS2. The dose-response curve of PtKCS2_1-277 was more like PtKCS2, despite having nearly identical substrate specificity to PtKCS1 (Figs. 8C and 5). This shows that it is possible to uncouple inhibitor sensitivity from substrate specificity. Comparing PtKCS1, PtCER6, and PtCER6_106-118 also showed differential inhibitor sensitivity of each enzyme to fentrazamide, although PtCER6_106-118 was more similar to PtCER6 (Fig. S10).

### Discussion

Gene duplication plays an important role in the evolution of new gene functions, particularly in the diversification of specialized metabolites (39). In P. trichocarpa, PtKCS1 is part of a tandem duplicated gene cluster on Chr 10 that was recently identified as a key enzyme in the biosynthesis of alkenes for cuticular waxes (24). In this study, we used the PtKCS1 gene cluster as a model to study the molecular mechanisms driving evolution of substrate specificity. We first examined the origin of the gene cluster to uncover a history of tandem duplication events, WGD, and gene loss (Fig. 1C).

Using this gene cluster as a model for KCS evolution, we compared the substrate preference of five paralogs that share varying degrees of relatedness. The paralogs were promiscuous...
Substrate specificity of a KCS gene cluster

Figure 8. Differential inhibition of PtKCS1 clade members with K3 herbicides. A, proportion of total VLC-FAMEs in the respective herbicide treatments (2 μM) relative to the uninhibited and empty vector controls. Proportion represents the mean of 3 biological replicates ± SE. The structures of the four herbicides, alachlor, anilofos, flufenacet, and fentrazamide are shown above. B, log-inhibitor response curves of PtKCS1, 2, and 4 in the presence of increasing fentrazamide concentration. C, log-inhibitor response curves of PtKCS1, 2, PtKCS21-277 in the presence of increasing fentrazamide concentration. VLC-FAME, very long chain fatty acid methyl ester.

ptKCS2. On the other hand, PtKCS2 has specialized toward again in subclade B enzyme PtKCS4 after diverging from This specialization toward unsaturated fatty acids has occurred where subclade A enzymes have increased their relative preference of these genes, corroborating the promiscuity of 51). These studies provide insight into the ancestral substrate preference based on length, unsaturation, and position of the double bond. In general, subclade A members had a higher relative preference for monounsaturated substrates compared to subclade B, apart from PtKCS4. Previous work on the closest orthologs to the PtKCS1 clade in A. thaliana (AtKCS2/20) and Helianthus annuus (HaKCS1) found a broad product specificity characteristic of the poplar clade, accumulating predominantly saturated C20-C26 VLCFAs (38, 50, 51). These studies provide insight into the ancestral substrate preference of these genes, corroborating the promiscuity of this clade toward shorter VLCFAs. In poplar, the gene cluster provides an example of specialization after gene duplication, where subclade A enzymes have increased their relative preference toward unsaturated VLCFAs compared to subclade B. This specialization toward unsaturated fatty acids has occurred again in subclade B enzyme PtKCS4 after diverging from PtKCS2. On the other hand, PtKCS2 has specialized toward longer products, up to C30 VLCFAs (Fig. 2B). Members of the gene cluster appear to have undergone different variations of subfunctionalization and neofunctionalization, specializing in the production of shorter versus longer, saturated versus monounsaturated VLCFAs.

Despite the extensive literature on characterizing different KCSs using plant mutants and heterologous expression, little is known about the molecular basis of KCS substrate specificity. Studies have been partly hindered by the recalcitrance of the in vitro assay, owing to both the limited availability of commercially available very long chain acyl-CoA substrates and insolubility of the membrane-bound KCSs. Furthermore, we found that the addition of a C-terminal His-tag to KCSs caused a significant reduction in activity (Fig. S11). As such, our results focus on the comparison of product profiles as opposed to enzyme kinetics.

Generating fusion proteins to identify the molecular determinants of substrate specificity revealed that the first 277 residues of the N-terminal domain were sufficient to switch the substrate specificity from PtKCS2 to PtKCS1 (Fig. 5). A prior genome survey of KCSs from 28 plant species showed higher diversity on the N-terminal region, suggesting that this region is under more relaxed purifying selection (19). The previous study on KCS substrate determinants compared orthologs of FAE1 from A. thaliana and B. napus, also finding that the N-terminal region (up to helix-4) was key (37). Using different KCSs (PtKCS1, 2, 4, and PtCER6), we narrowed down the substrate determining regions in the N-terminal domain and identified an additional region at position 277 affecting substrate specificity. Site-directed mutants and fusion constructs showed that substitutions by helix-4 and at site 277 had major effects on activity toward VLCFAs based on unsaturation and length. Using this knowledge, the rational substitutions of smaller amino acids at site 277 were able to shift the products toward longer VLCFAs in PtKCS1 and 4, as well as introduce novel activities toward elongation of the linolenic acid, a triunsaturated ω-3 fatty acid (Fig. 7C).

It is worth noting that during the process of constructing the chimeras between PtKCS1 and PtKCS2, a few constructs failed to modify the VLCFA profile of yeast (Fig. 5). Notably, PtKCS11-283 lost activity on saturated and unsaturated fatty acids, whereas the reciprocal PtKCS21-277 was active. Similarly, PtKCS21-181, PtKCS2124-207, 280-283 and PtKCS2124-170, 280-283 were inactive, suggesting that there are key interactions between the region around 124 to 170 amino acids with the C-terminal half that are disrupted in these constructs. When this region was narrowed down to helix-4 (130–143 amino acids), constructs were active. This suggests that there are incompatible combinations between the two halves of PtKCS1 and PtKCS2 and that helix-4 is unusual in its modular property, that is, it can be replaced without destabilizing these interactions. Another interesting observation was that though helix-4 and residue 277/283 had clear effects on substrate specificity, combination mutants of these two regions for PtKCS1/PtKCS2 and PtKCS1/PtCER6 did not necessarily have an additive or synergistic effect (Figs. 5 and 6). These results again show that other regions on the protein are seemingly
epistatic and contribute to the structure of the substrate-binding tunnel. This makes sense considering the size of the binding site spans almost half of the entire protein (Fig. 3).

By modifying the more distantly related PtCER6 (59% amino acid identity to PtKCS1) with residues from PtKCS1, we further supported the role of helix-4 and site 277 in other KCSs. Furthermore, the coexpression of PtCER2-like5 with chimeras of PtCER6 and PtKCS1 showed that PtCER2-like5 requires the N-terminal region of PtCER6 for its modulatory effect on VLCFA elongation. Though these data likely suggest that PtCER6 interacts with PtCER2-like5 through the N-terminal domain (particularly helix-4), additional structural and interaction studies are required to test this. The interaction between CER2-like5 with condensing enzymes is not fully understood; yet at this point, it seems that the ability of CER2-like5 to shift the chain length profile is limited to AtCER6 and phylogenetically related KCSs (AtCER60, WLS4 in rice, and PtCER6 in poplar) (29, 38). Structural models for CER2-like show that the N-terminal 100 residues that differentiate the modulatory activity of Arabidopsis CER2-like5 on AtCER6 are positioned on the surface of the protein (2). However, it remains unknown if this region physically interacts with the FAE complex, binds any of the substrates of KCS, or whether CER2-like5 physiologically interacts with other KCS whose elongation it does not modulate. Future studies should consider physical interaction assays to determine if the increase in total VLCFA binding to substrate speciﬁcity, KCS inhibition also seemed to be easily modified by few substitutions. Using a fusion between PtKCS1 and 2 (PtKCS21-283), we showed that substrate speciﬁcity could be uncoupled from inhibitor sensitivity to fentrazamide (Fig. 8C). Considering that the most closely related enzymes to KCSs (chalcone synthases) are covalently inhibited by K3 herbicides, it is likely that KCSs are also inactivated in this manner, and the properties of the active site pocket could change without affecting the terminal end of the large binding tunnel (52). Understanding the molecular basis of this inhibition could be useful for engineering transgenic crops that are resistant to certain herbicides or developing herbicides that target speciﬁc KCSs.

Current models suggest that the binding site of the KCS is flexible to accommodate for different substrates, which generates the broad product proﬁle observed for certain KCSs (37). We propose that this “flexible model” can be reﬁned to include a recognition domain for certain motifs on functionalized VLCFAs. For example, the carbon series of monounsaturated VLCFAs observed for PtKCS1 likely results from preferential recognition of the cis-ω9 motif (Fig. 2). Furthermore, fusions of PtKCS2 and PtCER6 containing helix-4 from PtKCS1 shows increased production of cis-ω9 products, suggesting that helix-4 confers partial recognition of this motif. Feeding 18:2 and 18:3 fatty acids to yeast expressing PtKCS4R283G also generated a carbon series of diunsaturated or triunsaturated VLCFAs, respectively. Therefore, a ﬂexible region between the active site and the recognition domain could explain how functionalized VLCFAs are extended to varying lengths (Fig. 5I2). Another explanation we propose here is the “ﬁxed model” whereby the ﬂexible substrates of varying lengths conform to ﬁt inside a ﬁxed binding tunnel containing the recognition domain. Supporting this model is the shape of the cavity in the PtKCS homology models, showing a large cavity at the active site that could potentially accommodate folding of the hydrocarbon chain (Fig. 3B).

Solved structures with bound substrates would help distinguish between these two models and further elucidate the mechanisms of KCS substrate speciﬁcity.

In general, KCS substrate preference was easily altered by perturbations to the substrate-binding domain. This is likely because substrate speciﬁcity of hydrocarbon chains is largely determined by the size and hydrophobicity of the binding tunnel unlike polar ligands, which are stabilized by speciﬁc ionic interactions (53). Substrate promiscuity is a key trait of enzymes involved in specialized metabolism (39). Using different mutants, we show that even a single substitution can drastically change KCSs substrate speciﬁcity. These minor changes, coupled with their promiscuity, demonstrate how KCSs might have rapidly evolved to accept new substrates. KCSs are closely related to Type III PKSs such as chalcone synthases, which also use diverse substrates. Unlike PKSs, however, the KCSs has been largely overlooked in terms of their molecular mechanisms and biocatalytic potential. There is recent evidence that certain KCSs can condense elongation cycle intermediates in a PKS-like manner, suggesting they are more versatile than previously thought (23). Using directed evolution or rational approaches to alter the substrate determining regions of KCSs has the potential to engineer novel, noncanonical substrate speciﬁcities.

Here, we show that the evolution of KCSs can happen rapidly following gene duplication, resulting in sub-functionalization and neofunctionalization. Using the paralogs of a duplicated KCS gene cluster as the basis for mutational analyses, we identiﬁed major determinants of speciﬁcity toward unsaturation and chain length. The PtKCS1 gene cluster is part of the larger KCS gene family that has unique and diverse substrate speciﬁcities. This work lays the foundation toward understanding the molecular mechanisms behind their incredible diversity.

**Experimental procedures**

**Materials**

The INVSc1 yeast strain, PYEDEST52 vector, and PYE3/CT: CER2-like5 construct were generously shared by Shawn D. Mansﬁeld at the University of British Columbia. The herbicides (alachlor, anilofos, fentrazamide, and flufenacet) were purchased from Sigma–Aldrich as PESTANAL analytical standards. External standards methyl heptadecanoate, methyl oleate, SUPELCO 37 Component FAME Mix, 11(E)-eicosenoic acid, 13(Z)-eicosenoic acid, and 14(Z)-eicosenoic acid.
were purchased from Sigma–Aldrich, Cayman Chemical, Acros Organics, and Tokyo Chemical Industry at ≥95% purity. The internal standard (nonadecanoic acid) was purchased from Agilent. Fatty acids used for feeding were purchased from Cayman Chemical and Sigma–Aldrich at ≥98% purity.

**Yeast expression assays**

were performed using the OneScript Plus cDNA Synthesis Kit (Applied Biological Materials). The following genes were used for synthesis: PtKCS8, Potri.010G080400; PtKCS4, Potri.010G079300; PtKCS9, Potri.010G080400; PtKCS5; Potri.008G160000; PtCER6, Potri.010G125300; PtCER2-like5, and Potri.005G052200. The KCSs were cloned from cDNA (apart from Potri.010G080400 which was gene synthesized by Bio Basic) for directional Gateway cloning into pENTR/D-TOPO (Invitrogen). All primers used in this study are listed in Table S7. The plasmids containing 2% raf EST52 vectors were inoculated into SC-Ura liquid media containing 2% glucose. The cultures containing PYESD-EST52 vectors were grown shaking at 30 °C for 24 h. In this study: (Applied Biological Materials). The following genes were used for synthesis: PtKCS8, Potri.010G080400; PtKCS4, Potri.010G079300; PtKCS9, Potri.010G080400; PtKCS5; Potri.008G160000; PtCER6, Potri.010G125300; PtCER2-like5, and Potri.005G052200. The KCSs were cloned from cDNA (apart from Potri.010G080400 which was gene synthesized by Bio Basic) for directional Gateway cloning into pENTR/D-TOPO (Invitrogen). All primers used in this study are listed in Table S7. The plasmids were transformed into *Escherichia coli* TOP10 competent cells, screened, and confirmed by Sanger sequencing. Constructs had the same amino acid sequence as the reference *P. trichocarpa* (v3.0) genome, except PtKCS1, which has a serine to proline mutation at position 125. The sequence-confirmed genes were transferred to the PYEDEST52 destination vector via the Gateway LR reaction and transformed into *E. coli* TOP10 cells. Vectors obtained from PCR-positive clones were transformed into INVSc1 yeast (Invitrogen) using the SC Easy Comp Kit (Invitrogen) and plated on Synthetic Complete minus Uracil (SC-Ura) media containing 2% (w/v) glucose. Individual transformants were selected and screened by PCR. Yeast containing the PYE3S/CT:PtCER2-like5 construct were made competent and transformed with PYEDEST52 vectors as described previously. Colonies were selected on Synthetic Complete minus uracil and tryptophan (SC-Ura-Trp) and screened by PCR. To each vial, 2 ml of a 1.5 M H2SO4 solution containing 50 ng/μl nonadecanoic acid (internal standard) was added, then heated to 90 °C for 1.5 h with periodic mixing. After cooling, 1.5 ml of pentane and 2 ml 0.9% NaCl was added to extract methyl esters. The organic phase was concentrated under N2 gas, resuspended in pentane, and analyzed by GC-MS.

**FAME analysis**

Briefly, 40 mg of lyophilized yeast powder was weighed and transferred to a Teflon capped vial. To each vial, 2 ml of a 1.5 M H2SO4 solution containing 50 ng/μl nonadecanoic acid (internal standard) was added, then heated to 90 °C for 1.5 h with periodic mixing. After cooling, 1.5 ml of pentane and 2 ml 0.9% NaCl was added to extract methyl esters. The organic phase was concentrated under N2 gas, resuspended in pentane, and analyzed by GC-MS.

**GC-MS**

Samples were analyzed using the Agilent 5977A series GC/MSD system fitted with a 30 m × 320 μm × 1 μm HP-5 column. The initial injection temperature of 150 °C was held for 1 min, ramped to 280 °C at a rate of 4 °C/min, then held at 280 °C for 7 min with a helium gas flow rate of 1.2 ml/min. Compound peaks from the resulting chromatograms were identified using mass spectra and retention time comparison with authentic standards (Fig. S5). Peak areas were normalized to the weight of lyophilized yeast, internal and external standards (methyl heptadecanoate for saturated VLCFAs and methyl octadecenoate for unsaturated VLCFAs) to obtain compound abundance.

**Homology modeling and ligand docking**

A version of AlphaFold2 using MMseq2 homology searching was used to predict the structure of KCSs from amino acid sequences (42, 43). Models were generated using multiple Type III PKS templates (Protein Data Bank IDs: 6dx8, 3elh, 3xyx, 3d8d, 3wzx, 1xes, 1xet, 3tsy, 4b0n, 5wxx, 2h84, 1u0m, 3vs8, 5hwo, and 5hwc) (Tables S2 and S3). Most sites were modeled at greater than 90% confidence according to the local distance difference test (Fig. S13) (56). The transmembrane domain (residues 1–102 in PtKCS1) anchors the enzyme to the membrane of the endoplasmic reticulum but was not modeled here due to low confidence (47). PyMol (version 2.5.2) was used for visualization (57).

**Substrate specificity of a KCS gene cluster**

Total RNA from *P. trichocarpa* (accessions HALS30-6 and KLINE20-1) leaf tissue was extracted using TRIzol (Invitrogen), followed by reverse transcription to generate cDNA. The PCR products were then used as a template for complementary DNA (cDNA) synthesis using the OneScript Plus cDNA Synthesis Kit (Applied Biological Materials). The following genes were used for synthesis: PtKCS8, Potri.010G080400; PtKCS4, Potri.010G079300; PtKCS9, Potri.010G080400; PtKCS5; Potri.008G160000; PtCER6, Potri.010G125300; PtCER2-like5, and Potri.005G052200. The KCSs were cloned from cDNA (apart from Potri.010G080400 which was gene synthesized by Bio Basic) for directional Gateway cloning into pENTR/D-TOPO (Invitrogen). All primers used in this study are listed in Table S7. The plasmids were transformed into *Escherichia coli* TOP10 competent cells, screened, and confirmed by Sanger sequencing. Constructs had the same amino acid sequence as the reference *P. trichocarpa* (v3.0) genome, except PtKCS1, which has a serine to proline mutation at position 125. The sequence-confirmed genes were transferred to the PYEDEST52 destination vector via the Gateway LR reaction and transformed into *E. coli* TOP10 cells. Vectors obtained from PCR-positive clones were transformed into INVSc1 yeast (Invitrogen) using the SC Easy Comp Kit (Invitrogen) and plated on Synthetic Complete minus Uracil (SC-Ura) media containing 2% (w/v) glucose. Individual transformants were selected and screened by PCR. Yeast containing the PYE3S/CT:PtCER2-like5 construct were made competent and transformed with PYEDEST52 vectors as described previously. Colonies were selected on Synthetic Complete minus uracil and tryptophan (SC-Ura-Trp) and screened by PCR. To each vial, 2 ml of a 1.5 M H2SO4 solution containing 50 ng/μl nonadecanoic acid (internal standard) was added, then heated to 90 °C for 1.5 h with periodic mixing. After cooling, 1.5 ml of pentane and 2 ml 0.9% NaCl was added to extract methyl esters. The organic phase was concentrated under N2 gas, resuspended in pentane, and analyzed by GC-MS.

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Malonyl-CoA and docosanoic acid were docked to the model using AutodockVina (58). Template structures cocrystallized with their substrates or substrate analogs were used to guide the docking of these ligands, which can be more reliable compared to docking without a reference (59). The dimensions of the grid box were $22 \times 10 \times 9$ Å for docking docosanoic acid and $14 \times 13 \times 17$ Å for docking malonyl-CoA. The scoring for accepted ligand docking conformations was between $-5$ to $-6$ kcal/mol, with a docking RMSD of less than 4 Å. The overall structure of the KCS homology model was corroborated using the Phyre2 server (60).

**Phylogenetic and correlation analyses**

Amino acid sequences from the *P. trichocarpa* (v3.0) genome were obtained from Phytozome (61). MEGAX was used to align sequences (MUSCLE algorithm) and build the maximum-likelihood phylogenetic tree (62). To calculate the density of transposable elements, annotated repeat sequences on Chr 10 were obtained from Phytozome and kernel density was calculated along the sequence with a bin width of 15 kb. The metabolite-metabolite correlation analysis was conducted using R and METAGENassist (63). Levels of metabolites were normalized by the autoscaling method and then a correlation matrix was computed with nonparametric Spearman rank correlation.

**Data availability**

All data are contained within the article and supporting information.

**Supporting information**—This article contains supporting information (61, 62).

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**Author contributions**—J. Y. C. and E. G. V. conceptualization; J. Y. C. methodology; A. M. validation; J. Y. C. writing—original draft; J. Y. C. and E. G. V. writing—review & editing; E. G. V. supervision; E. G. V. funding acquisition.

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**Conflict of interest**—The authors declare that they have no conflicts of interest with the contents of this article.

**Abbreviations**—The abbreviations used are: cDNA, complementary DNA; Chr, chromosome; FAE, fatty acid elongation; FAME, fatty acid methyl ester; VLCFA, very long chain fatty acid; WGD, whole genome duplication.

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