Functional Role of Long-Chain Acyl-CoA Synthetases in Plant Development and Stress Responses

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Fatty acids (FAs) play vital roles in plants as components of lipid membranes that demarcate cells and organelles, as sources of stored energy in the form of neutral lipids, and as signaling molecules that elicit plant responses to adverse conditions. The activation of FAs through the formation of acyl-CoA intermediates by acyl-CoA synthetase (ACS) family enzymes is required for their synthesis and degradation. Long-chain ACSs (LACSs) represent a small subgroup of ACS enzymes that specifically convert long-chain or very-long-chain FAs into corresponding thioesters for multiple lipid-associated processes. Alteration of LACS activity often results in pleiotropic phenotypes such as male sterility, organ fusion, aberrant cuticular structure, delayed seed germination, altered seed oil content, and plant capacity to respond to various environmental stresses. This review provides a comprehensive analysis of LACS family enzymes including substrate specificity, tissue-specific expression patterns, and distinct subcellular localization highlighting their specific roles in lipid synthesis and degradation, the effects of altered LACS activity on plant development, the relationship between LACS activity and stress resistance, and the regulation of LACS activity. Finally, we pose several major questions to be addressed, which would advance our current understanding of LACS function in plants.

Keywords: long-chain acyl-CoA synthetases, lipid, biosynthesis, regulation, stress, metabolism

INTRODUCTION

Fatty acids (FAs) are nearly ubiquitous in plant cells where they are incorporated into various types of lipids including phospholipids and other membrane lipids, triacylglycerols (TAGs), as well as epidermal cuticular lipids and suberin, and thus are important for maintaining membrane integrity, providing energy for various metabolic processes, and forming surface barriers against abiotic and biotic stresses (Fulda et al., 2004; Schnurr et al., 2004; Li-Beisson et al., 2013; Grevengoed et al., 2014; Jessen et al., 2015; Fich et al., 2016; Ingram and Nawrath, 2017). The shunting of FAs into different metabolic pathways initially requires activation to high-energy CoA thioesters by the activity of acyl-CoA synthetases (ACSs), also known as FA:CoA ligases. The formation of acyl-CoAs mediated by ACS is an ATP-dependent process that proceeds as a two-step reaction. In the first step, the α-phosphorous of ATP is attacked by the partially negatively charged oxygen on the carbonyl on an FA forming an acyl-AMP intermediate and releasing pyrophosphate. In the second step, the thiol group of CoA conducts a nucleophilic attack on the carbonyl of the acyl-AMP, forming a fatty acyl-CoA and releasing AMP (Grevengoed et al., 2014).
Long-chain ACSs (LACSs) as a subgroup of ACS family preferentially activate long-chain (LCFAs; C16–C18) or very-long-chain FAs (VLCFAs; ≥C20) (Shockey et al., 2002; Lü et al., 2009; Shockey and Browse, 2011). Here, LACS enzymes differ in terms of substrate specificity, tissue-specific expression pattern, and subcellular distribution, confining each member to specific metabolic pathways. Furthermore, since LACS activity is closely related with production traits such as oil quality and quantity, and stress resistance, in recent years, research on LACS has also been carried out in economically relevant crop species. Herein, we present a concise review of progress in understanding LACS function in plants and further pose major unanswered questions about LACS enzymes that need to be addressed in order to deepen our understanding of these critical enzymes.

SUBSTRATE PREFERENCE

Despite being a small subgroup of ACS enzymes in Arabidopsis, AtLACSs show a surprising diversity in substrate specificity albeit restricted to LCFAs or VLCFAs (Shockey et al., 2002). AtLACS isoform divergence in substrate specificity is easily related to their diversity in function and differential subcellular localizations (Table 1). AtLACS1, a major isoform involved in the synthesis of wax, activates VLCFAs (20:0–30:0) with the highest specificity for triacananoic acid (30:0) (Lü et al., 2009) (Table 1). AtLACS1 shows the next highest activity with palmitic acid (16:0). In agreement with this, mutants with loss of function of AtLACS1 have substantial reductions in 16:0-derived cutin monomers (Lü et al., 2009). AtLACS2 is mainly responsible for cutin synthesis, preferring hydroxylated over unsubstituted FAs, with enhanced activity toward hydroxylated 16:0 (Schnurr et al., 2004) (Table 1). AtLACS4, AtLACS8, and AtLACS9 appear to play redundant roles in wax and cutin synthesis, showing catalytic activities with LCFAs similar to those of AtLACS1 and AtLACS2 but displayed lower activities with hydroxylated versus unsubstituted FAs (Shockey et al., 2002; Fulda et al., 2004; Zhao et al., 2019). Additionally, simultaneous mutation of AtLACS4 and AtLACS9 led to a significant reduction of linoleic acid (18:2) at the sn2 position of MGDG, indicating that 18:2 might also be a preferred substrate of AtLACS4 or AtLACS9 (Jessen et al., 2015). Peroxisome-localized AtLACS6 and AtLACS7 mediate TAG degradation during seed germination, activating FAs stored in seed TAGs with enhanced activity for eicosenoic acid (20:1), which is predominantly found in Arabidopsis seed lipids (Table 1) (Fulda et al., 2002, 2004; Shockey et al., 2002).

The substrate specificity of the LACS enzymes has been extensively studied in plant species besides Arabidopsis. They also preferentially utilize LCFAs or VLCFAs. For example, GhACS1 from cotton (Gossypium hirsutum), sharing high amino acid sequence identity with AtLACS4 and AtLACS5, is required for normal microsporogenesis (Wang and Li, 2009). In vitro enzyme activity analysis revealed that GhACS1 prefers LCFAs with the highest activity toward oleic acid (18:1) (Wang and Li, 2009). MdLACS2 from apple (Malus domestica), an ortholog of AtLACS2, was shown to catalyze the formation of 16:0 CoA (Zhang et al., 2020a). BnLACS2 from rapeseed (Brassica napus), involved in seed oil production, exhibited a substrate preference for 14:0, 16:0, 18:0, 18:1, and 22:1 (Ding et al., 2020). These studies demonstrate that LACS enzymes play conserved roles in higher plants. However, given that lipid metabolic processes vary among different plant species, LACS proteins will exhibit species-dependent substrate specificities. For example, α-linolenic acid (18:3) was enriched in flax (Linum usitatissimum) seed oil; recently, LuLACS8A from flax was identified to show high activity toward 18:3 (Xu et al., 2018). RcACS2 from Ricinus communis is found to preferentially activate ricinoleic acid (12-OH 18:1), which accumulates to very high levels in castor oil (He et al., 2007). TpLACS4 in Thalassiosira pseudonana preferentially catalyzes the formation of several molecular species of polyunsaturated VLCFA CoAs (Tonon et al., 2005) (Table 1).

EXPRESSION PATTERN

In Arabidopsis, most AtLACS genes show broad expression patterns (Figure 1); i.e., they are highly expressed in multiple organs including root, stem, leaf, flower, and germinating seedlings (Shockey et al., 2002; Zhao et al., 2010). Different from these AtLACS genes, AtLACS3 is only expressed in root, stem, leaf, and flower, whereas AtLACS5 is exclusively detected in flower (Shockey et al., 2002). Moreover, analysis of Arabidopsis eFP Browser data1 reveals that some AtLACS genes exhibit tissue-specific expression patterns. For example, AtLACS1, AtLACS2, AtLACS3, and AtLACS6 are specifically expressed in leaf epidermal cells where cuticular lipids are produced; and AtLACS2, AtLACS3, and AtLACS9 are highly expressed in root endodermal cells where suberin is synthesized (Figure 1), suggestive of their possible roles in apoplastic lipid synthesis in these tissues.

The expression pattern of LACS genes has also been extensively investigated in other plant species such as B. napus and M. domestica, Helianthus annuus, and Glycine max (Table 2). A total of 34 BnLACSs genes are found in the B. napus genome, 18 of which are expressed in developing seeds (Xiao et al., 2019), implicating major roles for these genes in lipid metabolism. These BnLACS genes show diversified expression patterns. Several genes display broad expression patterns including two BnLACS4s, two BnLACS8s, and one BnLACS9, quite similar to their closest Arabidopsis orthologs (Xiao et al., 2019), whereas some genes show tissue-specific expression patterns. For example, several BnLACS5 genes are specifically expressed in buds, anthers, and stamens (Xiao et al., 2019), resembling AtLACS5. One cotton ortholog, GhACS1, also displays an expression pattern similar to AtLACS5, indicating the conserved function of these LACS5s in flower development (Wang and Li, 2009). Another research group also comprehensively checked the expression of a number of MdLACS genes in apple and found that all tested genes are highly expressed in pericarp tissues where wax and cutin are actively produced (Zhang et al., 2018), suggesting their possible roles in cuticle synthesis.

1http://efp.ucr.edu/cgi-bin/relative.cgi
| Species                  | Protein | Substrate specificity | Subcellular localization | Function                                | References                          |
|--------------------------|---------|-----------------------|--------------------------|-----------------------------------------|-------------------------------------|
| Arabidopsis thaliana     | AtLACS1 | 16:0, 16:1, 18:1, 18:2, 20:0 (Shockey et al., 2002); 16:0, 18:0, 20:0, 22:0, 24:0, 26:0, 28:0, 30:0↑ (Lü et al., 2009) | ER                        | Wax, cutin, TAG, and trypbine           | Lü et al., 2009; Zhao et al., 2010, 2019; Yang et al., 2017 |
|                          | AtLACS2 | 16:0, 16:1, 18:1, 18:2 (Shockey et al., 2002); 16:0, 16-OH16:0↑ (Schnurr et al., 2004); 16:0, 18:0, 20:0, 22:0, 24:0, 26:0, 28:0, 30:0↑ (Lü et al., 2009) | ER                        | Cutin, Wax, TAG, related with hypoxia response | Schnurr et al., 2004; Suh et al., 2005; Lü et al., 2009; Yang et al., 2017; Zhao et al., 2019 |
|                          | AtLACS3 | 16:0, 16:1↑, 18:1, 18:2 | Unidentified             | Unidentified                           | Shockey et al., 2002                |
|                          | AtLACS4 | 16:0, 16:1, 18:1, 18:2↑ (Shockey et al., 2002) | ER                        | Wax, TAG, trypbine, and glycerolipid    | Zhao et al., 2010, 2019; Jessen et al., 2011 |
|                          | AtLACS5 | 16:0, 16:1↑, 18:1, 18:2 (Shockey et al., 2002) | Unidentified             | Unidentified                           | Shockey et al., 2002                |
|                          | AtLACS6 | 16:0, 16:1, 18:1, 18:2, 20:1↑ (Shockey et al., 2002); 14:0, 16:0, 18:1, 18:2, 18:3 | Peroxisome                | β-oxidation                           | Fulda et al., 2002                  |
|                          | AtLACS7 | 16:0, 16:1, 18:1, 18:2, 20:1↑ (Shockey et al., 2002); 14:0, 16:0, 18:1↑, 18:2, 18:3, 20:1↑ (Fulda et al., 2002) | Peroxisome                | β-oxidation                           | Fulda et al., 2002                  |
|                          | AtLACS8 | 16:0, 16:1, 18:1, 18:2 | Unidentified             | Unidentified                           | Shockey et al., 2002; Zhao et al., 2010, 2019 |
|                          | AtLACS9 | 16:0, 16:1, 18:1, 18:2 | Unidentified             | Wax, TAG, and glycerolipid             | Shockey et al., 2002; Zhao et al., 2010, 2019 |
| Brassica napus           | BnLACS2 | 14:0, 16:0, 18:0, 18:1, 22:1 | ER                        | TAG                                     | Ding et al., 2020                   |
| Glycine max              | GmACSL2 | 14:0, 16:0, 18:0, 18:1, 22:1 | Peroxisome                | β-oxidation                           | Yu et al., 2014                     |
| Gossypium hirsutum       | GhACS1  | 14:0, 16:0, 18:0, 18:1, 20:0 | ER, PM                    | Microporogenesis                       | Wang and Li, 2009                   |
| Helianthus annusus       | HaLACS1 | 16:1, 18:1, 18:2, 18:3 | Unidentified             | Plastid envelop                        | Aznar-Moreno et al., 2014           |
| Malus domestica          | MdLACS2 | 16:0, 16:1, 18:1, 18:2 | Unidentified             | Unidentified                           | Aznar-Moreno et al., 2014           |
| Oryza sativa             | OsLACS9 | Unidentified           | Unidentified             | Wax                                     | Zhang et al., 2020a                 |
| Thalassiosira pseudonana | TpLACSA | 16:0, 18:3, 18:4, 20:4↑, 20:5↑, 22:6↑ | Unidentified             | Wax                                     | Zheng et al., 2019                  |
| Zea mays                 | ZmCER8  | 18:1, 18:2, 18:3↑ | Unidentified             | TAG                                     | Xu et al., 2018                     |
| Linum usitatissimum      | LuLACS8A| 12:0, 16:0, 18:1, 18:2, 18:3, 12-OH18:1↑, 11-OH12:0, 12-OH12:0 | Unidentified             | TAG                                     | He et al., 2007                     |
| Ricinus communis         | RcACS2  | 12:0, 16:0, 18:1, 18:2, 18:3, 12-OH18:1↑, 11-OH12:0, 12-OH12:0 | Unidentified             | TAG                                     | He et al., 2007                     |
|                          | RcACS4  | Unidentified           | Unidentified             | Unidentified                           |                                     |

Arrows indicate the enhanced activity. LACSs, long-chain acyl-CoA synthetases; ER, endoplasmic reticulum; TAG, triacylglycerol.
SUBCELLULAR LOCALIZATION

To minimize futile cycles, LACS orthologs in mammals are distributed across many different cellular compartments such as plasma membrane (PM), lipid droplet, mitochondria, peroxisome, and endoplasmic reticulum (ER) (Soupene and Kuypers, 2008). In plants, the LACS proteins are confined to fewer subcellular compartments including ER, plastid, peroxisome, and PM (Figure 2). Plastids are the places where de novo synthesis of FAs occurs. Several LACS orthologs are known to localize to the plastid envelope (Figure 2), including AtLACS9, HaLACS1 from sunflower (H. annuus), and OsLACS9 from rice (Oryza sativa) (Schnurr et al., 2002; Aznar-Moreno et al., 2014; Jessen et al., 2015; Kitajima-Koga et al., 2020). They are implicated to function in the activation of de novo synthesized LCFAs in plastids. Moreover, AtLACS9 is also identified to be involved in lipid trafficking between plastid and ER (Jessen et al., 2015). ER is the compartment where membrane lipids, cuticular lipids, and TAG are produced. Most identified AtLACSs localize to this organelle including AtLACS1, AtLACS2, AtLACS4, and AtLACS8 (Figure 2). Similarly, one AtLACS2 ortholog from B. napus known for its roles in rapeseed oil synthesis, BnLACS2, is also found in the ER (Ding et al., 2020). Surprisingly, GhACS1 from cotton (G. hirsutum), despite sharing high amino acid sequence identity with AtLACS4 and AtLACS5, displays different localization patterns being found not only in ER but also in PM, implicating dual roles in two compartments (Wang and Li, 2009). Distinct from the LACS proteins involved in lipid synthesis, LACS proteins mediating FA degradation usually localize to peroxisomes. To date, three LACS proteins are found to reside in peroxisome, including AtLACS6, AtLACS7, and one soybean protein GmACSL2 (Fulda et al., 2002, 2004; Yu et al., 2014). Their localization often requires peroxisomal targeting signal (PTS) sequences. For instance, AtLACS6 localization is targeted by a type 2 PTS (PTS2), whereas AtLACS7 localization is driven by both PTS1 and PTS2 (Fulda et al., 2002). Moreover, AtLACS6, AtLACS7, and GmACSL2 have been identified to be involved in lipid degradation, providing energy for seedling development (Fulda et al., 2004; Yu et al., 2014). Taken together, it can be surmised that LACSs residing in different subcellular compartments play distinct roles in lipid synthesis and degradation. We found it intriguing that despite sharing high sequence identity with each other, LACS proteins sometimes display different subcellular expression patterns. For example, the dual localization of GhACS1 to ER and PM is different from that of its Arabidopsis ortholog AtLACS4, suggesting overlapping but also distinct functions for these LACS orthologs in different species. Further study of LACS in diverse plant species is required to comprehensively clarify the function of LACS genes during lipid metabolism.
## TABLE 2 | Expression pattern of LACS genes from different plants.

| Species                  | Gene name               | Gene ID                      | Expression pattern                                                                 | References                    |
|--------------------------|-------------------------|------------------------------|-------------------------------------------------------------------------------------|-------------------------------|
| *Brassica napus*          | BnLACS1-2               | BnaAnng39590D                | Highly expressed in leaves and buds                                                 | Xiao et al., 2019             |
|                          | BnLACS1-3               | BnaC04g51420D                | Expressed in all organs except for roots, highly expressed in leaves and buds, siliques, and pericarps |                               |
|                          | BnLACS1-9               | BnaCnng60230D                | Highly expressed in leaves, moderately expressed in flowers and buds                |                               |
|                          | BnLACS1-10              | BnaA05g00640D                | Highly expressed in buds, moderately expressed in flowers                           | Xiao et al., 2019; Ding et al., 2020 |
|                          | BnLACS2-1/BnLACS2       | BnaA05g16170D                | Highly expressed in buds and developing seeds, moderately expressed in leaves and buds, lowly expressed in roots and stems |                               |
|                          | BnLACS2-2               | BnaC05g51350D                | Highly expressed in buds                                                            | Xiao et al., 2019             |
|                          | BnLACS4-1               | BnaA01g13470D                | Expressed in all organs, highly expressed in leaves and flowers                     |                               |
|                          | BnLACS4-2               | BnaC01g15670D                | Expressed in all organs, highly expressed in leaves and flowers                     |                               |
|                          | BnLACS5-1               | BnaC09g26090D                | Highly expressed in buds, anthers, and stamens                                      |                               |
|                          | BnLACS5-2               | BnaA09g22150D                | Highly expressed in buds, anthers, and stamens                                      |                               |
|                          | BnLACS5-3               | BnaC02g28920D                | Highly expressed in buds, anthers, and stamens                                      |                               |
|                          | BnLACS5-4               | BnaA02g21860D                | Highly expressed in buds, anthers, and stamens                                      |                               |
|                          | BnLACS5-9               | BnaC03g34500D                | Highly expressed in leaves and flowers                                              |                               |
|                          | BnLACS6-2               | BnaA03g29320D                | Expressed in all organs, highly expressed in leaves and flowers                     |                               |
|                          | BnLACS6-8               | BnaC03g44430D                | Expressed in all organs, highly expressed in leaves                                 |                               |
|                          | BnLACS8-2               | BnaA03g57930D                | Expressed in all organs, highly expressed in leaves and flowers                     |                               |
|                          | BnLACS9-3               | BnaC06g20910D                | Highly expressed in buds, moderately expressed in flowers                           |                               |
|                          | BnLACS9-4               | BnaA07g20920D                | Highly expressed in buds                                                            |                               |
| *Malus domestica*         | MdLACS1                 | MD09121286100                | Highly expressed in pericarps, moderately expressed in stems                        | Zhang et al., 2018            |
|                          | MdLACS2.1               | MD05G1070880D                | Highly expressed in pericarps                                                       |                               |
|                          | MdLACS2.2/MdLACS        | MD10G1086400D                | Expressed in all organs, highly expressed in pericarps and young leaves, moderately expressed in roots, stems, flowers, fruits, seeds, sarcocarps, and climax leaves | Zhang et al., 2018, 2020a     |
|                          | MdLACS4.1               | MD06G1106100D                | Highly expressed in pericarps                                                       | Zhang et al., 2018            |
|                          | MdLACS4.2/MdLACS4       | MD14G1128200D                | Expressed in all tissues, highly expressed in pericarps, lowly expressed in roots, stems, young leaves, mature leaves, flowers, fruits, seeds, and sarcocarps | Zhang et al., 2018, 2020b     |
|                          | MdLACS6.1               | MD13G1188800D                | Expressed in all tissues, highly expressed in pericarps, climax leaves, and stems    | Zhang et al., 2018            |
|                          | MdLACS6.2               | MD16G1189300D                | Expressed in all tissues, highly expressed in pericarps                             |                               |
|                          | MdLACS8.1               | MD09G1129700D                | Highly expressed in pericarps                                                       |                               |
|                          | MdLACS8.2               | MD17G1118500D                | Highly expressed in pericarps                                                       |                               |
|                          | MdLACS9.1               | MD08G1163100D                | Highly expressed in pericarps                                                       |                               |
|                          | MdLACS9.2               | MD15G1349800D                | Expressed in all tissues, highly expressed in pericarps                             |                               |
| *Gossypium hirsutum*      | GhACS1                  | DQ174259                     | Highly expressed in anther                                                          | Wang and Li, 2009             |
|                          | GhACS2                  | DQ174260                     | Highly expressed in germinating seedlings and young leaves, moderately expressed in developing seeds and flowers, lowly expressed in roots, stems, and senescent leaves | Yu et al., 2014              |
| *Glycine max*             | GmACSL2                 | Glyma_05G151200              | Highly expressed in germinating seedlings and young leaves, moderately expressed in developing seeds and flowers, lowly expressed in roots, stems, and senescent leaves |                               |
| *Oryza sativa*            | OsLACS9                 | Os12g0102350D                | Highly expressed in shoots, moderately expressed in leaves                           | Ding et al., 2020             |
| *Helianthus annuus*       | HaLACS1                 | HM490305                     | Highly expressed in seeds                                                           | Aznar-Moreno et al., 2014    |
|                          | HaLACS2                 | HM490306                     | Highly expressed in cotyledons, stems, and leaves                                    |                               |
| *Ricinus communis*        | RaACS2                  | DQ300358                     | Highly expressed in germinating seeds                                               | He et al., 2007               |
|                          | RaACS4                  | DQ300359                     | Ubiquitously expressed in all organs                                                |                               |
LONG-CHAIN ACS ASSOCIATED WITH CUTICULAR LIPID BIOSYNTHESIS

Cuticle mainly consists of cutin and wax. Cutin is a polyester mainly composed of glycerol and C16/C18 FA derivatives (Fich et al., 2016). Wax is composed of VLCFAs and corresponding derivatives in Arabidopsis (Lee and Sub, 2013). The synthesis of cutin and wax requires LACS activity. Cutin biosynthesis starts in plastid where de novo FA synthesis occurs, and then the generated LCFAs are transported from plastid to ER for final modification. During this process, LACSs facilitate the formation of LCFA-CoAs prior to the further modification of LCFA for integration into the cutin polymer (Fich et al., 2016). AtLACS2 is identified to be essential for this process. Its mutation resulted in drastic reduction of dicarboxylate cutin monomers, predominant components of Arabidopsis cutin (Bessire et al., 2007), and a significant reduction of total stem cutin loads, suggesting a predominant role for AtLACS2 in cutin synthesis under normal growth conditions. In addition, a recent study reported that the AtLACS2 activity also affects plant sensitivity to submergence by changing cuticle permeability (Xie et al., 2020). AtLACS1, sharing high amino acid sequence similarity with AtLACS2, is also involved in cutin synthesis, but its role is subordinate (Lü et al., 2009; Weng et al., 2010). Simultaneous suppression of AtLACS1 and AtLACS2 activity resulted in more severe cuticle defects than observed in either parental mutant, signifying their overlapping roles for the generation of LCFA-CoA esters (Lü et al., 2009; Weng et al., 2010). Interestingly, our recent study has shown that AtLACS1, rather than AtLACS2, specifically coordinates with CER17/ADS4 for the synthesis of cutin in upper, younger portions of inflorescence stems, demonstrating an organ and developmental-specific function for AtLACS1 in cutin biosynthesis (Yang et al., 2017). Apart from AtLACS1 and AtLACS2, AtLACS4 is also involved in cutin biosynthesis; the total cutin amount of Arabidopsis lacs1 lacs2 lacs4 triple mutant is much lower than that of either single or double mutants (Zhao et al., 2019), indicative of functional redundancy among these isoforms.

Just like cutin, wax synthesis also proceeds through two processes occurring in two different compartments, i.e., the synthesis of LCFAs in plastid, and the elongation and modification of VLCFAs in ER. LACS catalyzes the binding of CoA with LCFA or VLCFA prior to the modification of VLCFA. Several AtLACSs are identified to play roles in this process, including AtLACS1, AtLACS2, AtLACS4, AtLACS8, and AtLACS9 (Lü et al., 2009; Weng et al., 2010; Zhao et al., 2019), which function redundantly in wax synthesis given that higher-order mutants often possess severer defects in wax accumulation than their parental lines (Lü et al., 2009; Weng et al., 2010; Zhao et al., 2019). Among these proteins, AtLACS1 (also known as CER8) is the main isoform responsible for wax biosynthesis, since decreasing AtLACS1 activity alone leads to a drastic reduction of most wax components including alkanes, aldehydes, 2-alcohols, and ketones, finally resulting in the dramatic reduction of total wax amounts (Lü et al., 2009; Weng et al., 2010).

LONG-CHAIN ACSs RELATED WITH TAG SYNTHESIS

Triacylglycerol accumulates to very high levels in developing seeds serving as an energy reserve for seed germination and seedling growth. TAG biosynthesis starts with de novo FA synthesis in plastids. Plastid-derived LCFAs are transported into ER and esterified into a glycerol backbone. The activation of LCFA to CoA thioesters by LACSs provides acyl-CoAs for TAG synthesis. Several AtLACSs are known for their involvement in TAG synthesis. AtLACS1, AtLACS2, AtLACS4, AtLACS8, and AtLACS9 are highly expressed in developing seeds, suggesting their potential role in seed oil production (Zhao et al., 2010). AtLACS1 and AtLACS9 were the first two genes identified to be involved in TAG synthesis (Zhao et al., 2010), although AtLACS1 and AtLACS9 are localized in two different compartments (Zhao et al., 2010). Among nine Arabidopsis LACS proteins, AtLACS9 is the only isoform residing in the outer membrane of plastids. Its loss of function alone has little effect on TAG synthesis, whereas the introduction of AtLACS1 defects in an atlacs9 background causes a moderate reduction of FA content, indicative of overlapping roles in TAG synthesis (Zhao et al., 2010). Besides AtLACS1 and AtLACS9, other AtLACS genes highly expressed in seeds include AtLACS2, AtLACS4, and AtLACS8, which are also reported to be involved in TAG synthesis. Decreased oil content was detected in seeds of double mutants including atlacs1 atlacs2, atlacs1 atlacs4, atlacs4 atlacs8, and atlacs4 atlacs9 (Zhao et al., 2010, 2019). Higher-order mutants show a greater reduction in oil; the TAG content of atlacs1atlacs2atlacs4 triple mutant seeds is far lower than that of either parental double mutant (Zhao et al., 2019).

Long-chain ACS involvement TAG synthesis has been characterized not only in Arabidopsis but also in economically relevant plant species such as B. napus. These genes play...
conserved roles across plant species (Xiao et al., 2019; Ding et al., 2020). Several LACS orthologs from *B. napus* are found to be highly expressed in high oil content cultivars (Xiao et al., 2019). Furthermore, overexpression or knockout of one *AtLACS2* ortholog in *B. napus* impacted seed oil content (Ding et al., 2020). These results demonstrate that LACSs display conserved roles in TAG synthesis in the identified plant species, but we cannot ignore that TAG FA compositions vary among species. For example, oleic acids are the predominant FA constituent of TAG in rapeseed oil, whereas linoleic acids dominate *Arabidopsis* seed oil (Guo et al., 2019; Ding et al., 2020). As such, to precisely interpret the function of each LACS in TAG biosynthesis, multiple factors need to be considered, including plant species, substrate preference, tissue expression pattern, subcellular localization, and protein-level interactions with other enzymes involved in TAG synthesis.

**LONG-CHAIN ACS MUTATIONS CAUSE PLEOTROPIC PHENOTYPES DURING PLANT DEVELOPMENT**

Owing to the functional redundancy of LACS isoforms in *Arabidopsis*, most single *atlacs* mutants display no visible phenotypes, but *atlacs2* is one exception. Given LACS2’s primary function in cutin synthesis and ubiquitous pattern of expression, its deficiency leads to multiple defects in plant growth including reduced leaf size, overall plant size and seed sets, and lower rates of germination and seedling establishment (Schnurr et al., 2004). To further comprehend the roles of *Arabidopsis* LACS genes in plant growth and development, several research groups have generated higher-order mutants defective in the expression of multiple *AtLACS* genes. These mutants display much more severe phenotypes than their corresponding parental lines (Fulda et al., 2004; Weng et al., 2010; Zhao et al., 2019). For example, *atlacs4*, *atlacs8*, and *atlacs9* single mutants do not exhibit visible phenotypes; however, *atlacs4 atlacs8* and *atlacs4 atlacs9* double mutants display phenotypes reminiscent of the *atlacs2* single mutant (Jessen et al., 2015; Zhao et al., 2019). Furthermore, the *atlacs4 atlacs8 atlacs9* triple mutant exhibits lethality (Zhao et al., 2019), suggesting that LACS activity is vital for plant growth. Additionally, we noticed that *atlacs1atlacs2* double mutant displays organ fusion (Weng et al., 2010), which resembles the phenotype of several cuticle-deficient mutants such as *lacerata*, *bodyguard*, and *atp-binding cassette proteins g11 and g13* (Ingram and Nawrath, 2017). These studies indicate that LACS genes participate in the formation of organ boundaries.

Besides the aforementioned phenotypes, male fertility is also closely related with LACS activity. AtLACS1 and AtLACS4 are reported to be associated with the synthesis of both wax and pollen lipids (triphine). Triphine is a mixture of very-long-chain lipids including alkanes, alkenes, primary alcohols, secondary alcohols, aldehydes, and free FAs. Simultaneous suppression of AtLACS1 and AtLACS4 activity led to significant reduction in wax as well as triphine lipids resulting in conditional male sterility, albeit could be rescued by high humidity (Jessen et al., 2011). However, it seems that AtLACS1 and AtLACS4 have opposing roles in the triphine synthesis given that triphyne levels are apparently decreased in the *atlacs1* mutant but are dramatically increased in the *atlacs4* mutant (Jessen et al., 2011). It is unclear why high levels of triphyne accumulate in the *atlacs4* mutant. The authors proposed that the accumulation of triphyne in *atlacs4* mutant might be due to the defective lipid transport from tapetal cells to developing pollen grains (Jessen et al., 2011). The roles of LACS in male fertility seem to be conserved, since GhACS1 in cotton, an ortholog of AtLACS4 and AtLACS5 is preferentially expressed in developing anther, and cotton plants with reduced GhACS1 activity produce defective microspores (Wang and Li, 2009).

Different from the roles of those AtLACS isoforms involved in lipid synthesis, AtLACS6 and AtLACS7 are peroxisome-located and involved in lipid degradation, providing energy for seed germination and other energy-requiring physiological processes (Shockey et al., 2002). Thus, simultaneous deficiency of both genes delays post-germination seedling growth due to the lack of sufficient energy (Fulda et al., 2004).

**LONG-CHAIN ACSs ARE ASSOCIATED WITH STRESS RESISTANCES INCLUDING DROUGHT, HYPOXIA, AND BIOTIC STRESSES**

Cuticle forms surface barriers against biotic and abiotic stresses. As described above, several LACSs have been identified to be involved in the synthesis cuticular lipids and thus are vital for plant stress resistance. The main function of cuticle is to prevent non-stomatal water loss. To date, several *Arabidopsis* LACS single or double mutants including *atlacs2, atlacs1 atlacs2, atlacs4 atlacs8*, and *atlacs4 atlacs9* are reported to show increased cuticle permeability, increased water loss, and the hypersensitivity to drought (Bessire et al., 2007; Lü et al., 2009; Weng et al., 2010; Zhao et al., 2019). Moreover, owing to functional redundancy, higher-order *atlacs* mutants often show increased sensitivity to drought. The *atlacs1 atlacs2 atlacs4* triple mutant is a prime example. The rosette water loss rate of the *atlacs1 atlacs2 atlacs4* triple mutant exceeds that of single or double mutants (Jessen et al., 2015; Zhao et al., 2019). Furthermore, the role of LACSs in drought tolerance has been described in species other than *Arabidopsis*. Zhang et al. (2020a,b) found that ectopic expression of two LACSs from apple (*MdLACS2* and *MdLACS4*) in *Arabidopsis* decreased epidermal permeability, reduced water loss, and enhanced the resistance of transgenic plants to drought and salt stress. In another study, Zhang et al. (2018) demonstrated that callus transformed with *MdLACS1* showed enhanced tolerance to PEG, NaCl, and ABA treatments. These results indicate that the roles of LACS activity in cuticle synthesis are indispensable for plant resistance to drought.

Long-chain ACS activity is also closely related with hypoxia response. RAP2.12, a member of subgroup VII ETHYLENE-RESPONSE FACTOR (ERFVII) transcription factors, is a key
AtLACS4 mutation in of RAP2.12 (Schmidt et al., 2018; Xie et al., 2020). Simultaneous AtLACSs have been identified to be related with the translocation synthesis requires the participation of LACSs enzymes. Several stimulus (Schmidt et al., 2018; Xie et al., 2020). Acyl-CoA triggering the dissociation of RAP2.12 from ACBPs upon hypoxia 2019; Xie et al., 2020). Acyl-CoAs are reported to act as signals to induce the expression of hypoxia responsive genes (Licausi et al., 2011; Schmidt et al., 2018; Schmidt and van Dongen, 2019; Xie et al., 2020). Mutations in the AtLACS2 gene are also reported to induce hypoxia responses via increased levels of polyunsaturated linolenoyl-CoA, which triggers the translocation of RAP2.12 in nucleus, thus inducing high expression of hypoxia response genes (Schmidt et al., 2018). Mutations in the AtLACS2 gene is required for regulating ERF-VII-mediated hypoxia signaling.

Long-chain ACS activity is not only required for plant resistance to abiotic stresses but also important for plant responses to biotic stress. For example, atlacs2 mutants display strong resistance to infection by a necrotrophic fungus, Botrytis cinerea (Bessire et al., 2007). The author proposed that loss of AtLACS2 function increased cuticle permeability and thus might facilitate the diffusion of signals and effector molecules across the cuticle, which triggers a protection mechanism against the attack of B. cinerea (Bessire et al., 2007). But different from the response to B. cinerea, AtLACS2 loss of function increases susceptibility to avirulent Pseudomonas syringae (Tang et al., 2007). Possibly, the atlacs2 mutant has reduced stomatal ledges like other cuticle mutants, allowing more Pseudomonas swim through the stomata and enter the leaf (Lü et al., 2009). Taken together, the alteration of LACS activity changes the cuticle structure that ultimately affects plant physiological responses.

**REGULATION OF LONG-CHAIN ACS ACTIVITY**

Long-chain ACSs provide various acyl-CoA pools for formation of the cuticle layer, the synthesis of membrane lipids and storage TAG, and β-oxidation. LACSs are known to be regulated at multiple levels. Until recently, more focus has been given to the transcriptional regulation of LACS genes. During cuticle biosynthesis, AtLACSs together with other cuticle-related genes are regulated by two types of transcription factors that have antagonistic functions. Transcription factors such as WAX INDUCER1/SHINE (WIN1), MYB16, MYB30, and MYB106 are identified to act as positive regulators of AtLACS2 (Kannangara et al., 2007; Raffaele et al., 2008; Oshima et al., 2013). **AtLACS2 expression is also negatively regulated by DECREASE WAX BIOSYNTHESIS1 (DEWAX1) and DEWAX2, two members of AP2/ethylene response element-binding factor (ERF)-type transcription factor family, which directly bind to the promoter region of AtLACS2 (Go et al., 2014; Kim et al., 2018)**. Just like DEWAXs, AtMYB41, one member of R2R3-MYB transcription factor, is also identified to be a negative regulator of AtLACS2, though it is unknown if their interaction is direct (Cominelli et al., 2008). Besides these transcription factors, two E3 ligases, HISTONE MONOUBIQUITINATION 1 and 2 (HUB1 and HUB2), are also reported to transcriptionally activate the expression of several cutin-related genes including AtLACS2 through remodeling chromatin structure (Menard et al., 2014). Taken together, the LACS genes are transcriptionally coregulated by multiple factors. These mechanisms of regulation appear to be conserved in other plant species. For example, PeSHN1, one poplar homolog of SHN1, is also identified to positively regulate the expression of LACS2 in poplar (Meng et al., 2019), and MdMYB30 is found to activate the expression of MdLACS2 in apple (Zhang et al., 2019).

Long-chain ACS activity can also be post-transcriptionally regulated. With the development of deep-sequencing technology, a large number of microRNAs have been identified in diverse plant species. Many of these microRNAs play important regulatory roles during plant development. Genes related with FAs were found to be regulated by microRNA in oil palm, including orthologs of AtLACS4 and AtLACS9, which were identified to be targeted by eg-miR444b and eg-miR397, respectively (Zheng et al., 2019). The regulation of LACS genes by microRNAs may also be a mechanism utilized by other plant species. The plethora of whole-genome sequences available from diverse plant species will facilitate studies of this nature.

Long-chain ACS activity is also regulated at the level of translation. Xu et al. (2019) reported that D238 in AtLACS9 is predicted to have been positively selected during evolution, which is naturally replaced by glutamic acid in other plant species. This substitution is identified to enhance enzyme activity possibly by creating a more favorable enzyme conformation, suggesting that some key amino acids are vital for enzyme activity. Site-directed mutagenesis or domain swapping or deletion will help to advance our understanding of LACS catalytic mechanism. In our work, we found that one E3 ligase (known as CER9) genetically interacts with AtLACSs since its deficiency shows additive effects with AtLACS1 or AtLACS2 on wax or cutin synthesis (Lü et al., 2012). We hypothesize that CER9 physically interacts with AtLACSs proteins, trapping them in the ER to maintain their normal activity or that CER9 targets a negative regulator of AtLACS for degradation. However, we cannot also eliminate the possibility that CER9 might target a thioesterase that would hydrolyze VLCFA-CoAs to VLCFAs. Clearly, experimental validation of these hypotheses is required. In addition to these intrinsic regulators, cellular energy status is also closely related with LACS activity. As such, energy-limited conditions will inevitably affect LACS activity. For example, LACS activity is apparently affected by hypoxia, a condition that limits energy that would be produced by aerobic respiration (Schmidt et al., 2018).
CONCLUSION AND PERSPECTIVES

Long-chain ACSs target LCFAs or VLCFAs to providing acyl-CoA substrates for lipid synthesis and degradation processes vital for plant normal growth. Herein, we summarize their tissue-specific expression patterns and subcellular localizations and provide a summary of their specific roles in different metabolic pathways. Lastly, we describe mechanisms by which the overexpression or knockdown of cutin-related genes often monomers, suggesting that some enzymes may be used for both glycerol (Philippe et al., 2020), with chemical analogy to cutin FAs, dioic acids, fatty alcohols, hydroxycinnamic acids, and glycerol (Philippe et al., 2020), with chemical analogy to cutin remain to be answered.

As a case in point, a function for LACS in regulating suberin biosynthesis remains unproven. Suberin comprises hydroxylated FAs, dioic acids, fatty alcohols, hydroxycinnamic acids, and glycerol (Philippe et al., 2020), with chemical analogy to cutin monomers, suggesting that some enzymes may be used for both cutin and suberin biosynthesis. Moreover, it is reported that the overexpression or knockdown of cutin-related genes often changes suberin profiles (Philippe et al., 2020). To date, the role of AtLACSs in cutin synthesis is well characterized, but their role in suberin synthesis is less known. Based on in vitro enzyme assays, AtLACS1, AtLACS2, and AtLACS4 are likely to participate in suberin biosynthesis (Philippe et al., 2020), but no direct evidence of this function has been obtained. Additionally, Geneinvestigator data show that AtLACS2, AtLACS3, and AtLACS9 are specifically expressed in the root endodermis where suberin distributes, implying a function in suberin synthesis. To verify these speculations, it will be necessary to determine the suberin composition and content of these LACS mutants by biochemical methods. This will help to provide a more complete understanding of the roles of LACS members in lipid metabolism.

Long-chain ACS proteins are known to have dual functions, acting as enzymes that activate LCFAs and VLCFAs, but also as lipid transporters. Most studies have focused on the role of LACS in activating FAs; however, a function for LACS in lipid transport in planta is less understood. Pulsifer et al. (2012) reported that the ectopic expression of AtLACS1, AtLACS2, and AtLACS3 in yeast complements the phenotype of a yeast fat1D mutant deficient in both very-long-chain ACS activity and exogenous FA uptake. It has also been proposed that AtLACS4 and AtLACS9 mediate the import of FAs generated in the ER into plastids according to the analysis of metabolic products in lacs mutants (Jessen et al., 2015). These studies cannot fully elaborate the role of LACS in lipid transport. Thus, comprehending LACS function in lipid transport in planta may better define subcellular lipid channeling routes and comprehensively elaborate the roles of LACS in plant lipid metabolism.

Though the function of LACS proteins is well identified in Arabidopsis, more studies are needed to understand the functional divergence of LACS during evolution and breeding. As shown in Table 1, LACSs usually display variable substrate preferences since FA constituents often vary among different species. For example, TpLACSA in T. pseudonana, which shares high sequence identity with AtLACS7 and AtLACS6, preferentially catalyzes the formation of acyl-CoAs from polyunsaturated FAs (PUFAs) including arachidonic acid (20:4), eicosapentaenoic acid (20:5), and docosahexaenoic acid (22:6) (Table 1). However, these PUFAs are not present in Arabidopsis and thus are not potential preferred substrates of ATLACS6 and ATLACS7. Additionally, LACS copy number varies greatly among the genomes of different species (Shockey et al., 2002; Zhang et al., 2018; Ding et al., 2020). Thirty-four LACS genes are found in B. napus, 11 LACS orthologs are present in M. domestica, and only nine LACS genes are found in Arabidopsis. These LACSs display different patterns of expression indicative of functional divergence during evolution. Thus, extensive research in species with divergent lipid compositions will expand our knowledge of LACS function in plants and provide a more comprehensive understanding of lipid metabolism across the plant kingdom.

AUTHOR CONTRIBUTIONS

HZ wrote the manuscript. DK and SL made critical revision. All authors contributed to the article and approved the submitted version.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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