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Increasing hydroxy fatty acids in transgenic seeds

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The Phosphatidylcholine Diacylglycerol Cholinephosphotransferase Is Required for Efficient Hydroxy Fatty Acid Accumulation in Transgenic Arabidopsis¹

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Abstract

We previously identified an enzyme, phosphatidylcholine diacylglycerol cholinephosphotransferase (PDCT), that plays an important role in directing fatty acyl fluxes during triacylglycerol (TAG) biosynthesis. The PDCT mediates a symmetrical inter-conversion between phosphatidylcholine (PC) and diacylglycerol (DAG), thus enriching PC-modified fatty acids in the DAG pool prior to forming TAG. We show here that PDCT is required for efficient metabolism of engineered hydroxy fatty acids in *Arabidopsis* seeds. When a fatty acid hydroxylase (FAH12) from castor (*Ricinus communis*) was expressed in *Arabidopsis* seeds, the PDCT-deficient mutant accumulated only about half the amount of hydroxy fatty acids compared to that in the wild type seeds. We also isolated a PDCT from castor encoded by the *RcRODI* gene. Seed-specific co-expression of this enzyme significantly increased hydroxy fatty acid accumulation in wild type-FAH12 and in a previously produced transgenic *Arabidopsis* line co-expressing a castor diacylglycerol acyltransferase 2 (*RcDGAT2*). Analyzing the TAG molecular species and regiochemistry, along with analysis of fatty acid composition in TAG and PC during seed development, indicate that PDCT acts *in planta* to enhance the fluxes of fatty acids through PC and enrich the hydroxy fatty acids in DAG, and thus in TAG. In addition, PDCT partially restores the oil content that is decreased in FAH12 expressing seeds. Our results add a new gene in the genetic toolbox for efficiently engineering unusual fatty acids in transgenic oilseeds.

Vegetable oils consist principally of triacylglycerols (TAG) and have wide applications in human consumption and more recently in renewable biofuels and industrial materials (Gunstone, 1998; Durrett et al., 2008; Napier and Graham, 2010; Lu et al., 2011). The fatty acid (FA) composition in TAG determines the quality and thus the uses of plant oils. While edible oils should contain as little saturated FAs as possible and a significant proportion of polyunsaturated FAs (Riediger et al., 2009), the industrial oils require low polyunsaturated FAs (PUFAs) for desirable oxidative stability or high homogeneity of certain FAs (Dyer and Mullen, 2008; Dyer et al., 2008; Pinzi et al., 2009). For effective improvements of plant oils, we need to understand the fundamental aspects of how plant FAs are synthesized and accumulated in seed oils.

In plants, FA synthesis occurs exclusively in plastids and produces mostly oleic acid (18:1; number of carbon : number of double bonds) and a small amount of palmitic acid (16:0) and stearic acid (18:0) that are esterified to the acyl carrier protein (Ohlrogge and Browse, 1995). In oilseeds, these FAs are almost entirely (>95%) exported outside the plastid, and are converted into acyl-CoA to be used for glycerolipid synthesis in the endoplasmic reticulum (ER) (Browse and Somerville, 1991; Ohlrogge and Browse, 1995). In *Arabidopsis* seeds, 18:1-CoA may be elongated into 20:1-CoA to 22:1-CoA by a fatty acid elongase (FAE1) on the ER membrane (Kunst et al., 1992; Kunst and Samuels, 2003); or may be incorporated into the membrane lipid phosphatidylcholine (PC) to be desaturated to linoleic acid (18:2) and subsequently to linolenic acid (18:3) by the oleate desaturase FAD2 and the linoleate desaturase FAD3, also located on the ER (Browse et al., 1993; Sperling et al., 1993; Okuley et al., 1994). These reactions are the major causes of the wide variations of FA composition in TAG of many oilseeds.

The *de novo* PC synthesis pathway incorporates FAs through diacylglycerol (DAG) by the sequential *sn*-1 and *sn*-2 acylations of glycerol-3-phosphate followed by dephosphorylation of phosphatidic acid (Kennedy, 1961). DAG is converted into

PC by a reversible CDP-choline:diacylglycerol cholinephosphotransferase (CPT) (Slack et al., 1983; Goode and Dewey, 1999). Recent metabolic analyses indicate that the majority of newly synthesized FAs may enter PC through a process called acyl editing, which involves a very rapid cyclic deacylation and reacylation of PC (Williams et al., 2000; Bates et al., 2007; Bates et al., 2009). The genes involved in this process have not been identified, but the enzymes may include the acyl-CoA:lysophosphatidylcholine acyltransferases and phospholipase A2s. The PC-acyl editing cycle incorporates nascent 18:1 into PC and releases modified, e.g., polyunsaturated 18:2 and 18:3 from PC. This process generates an acyl-CoA pool that is enriched in polyunsaturated FAs to be utilized for glycerolipid synthesis in the ER.

The TAG synthesis is completed by the acylation of the *sn*-3 hydroxyl of DAG. This step may be catalyzed by an acyl-CoA:diacylglycerol acyltransferase (DGAT) using the acyl-CoA pool (Hobbs et al., 1999), or by a phospholipid:diacylglycerol acyltransferase (PDAT) that directly transfers the *sn*-2 acyl group from PC onto DAG to form TAG (Dahlqvist et al., 2000). The DAG precursor may be from *de novo* synthesis through a series of acylations of glycerol-3-P or may be derived from PC. Recent results demonstrate that the latter provides the predominant source (Bates and Browse, 2011). The high-flux exchange between the PC and DAG pools appears to be mainly catalyzed by the phosphatidylcholine diacylglycerol cholinephosphotransferase (PDCT), although the reverse reaction of CPT may also have some roles. The PDCT enzyme, discovered in Arabidopsis encoded by the *RODI* locus, catalyzes the inter-conversion between DAG and PC by phosphocholine headgroup exchange (Lu et al., 2009). Through reactions of PDCT, the acyl groups on DAG enter PC, and then return to DAG after they are desaturated or otherwise modified on PC.

This latest model of TAG synthesis described above highlights the central roles of PC as the source of greatly diversified FAs in membrane and storage lipids. This has also been reinforced by results in transgenic plants producing some modified fatty acids (mFAs) such as those containing hydroxy, epoxy, conjugated diene or

cyclopropane groups. These mFAs are synthesized by the FAD2-related enzymes acting on the *sn*-2 acyl groups on PC (Drexler et al., 2003; Jaworski and Cahoon, 2003; Lu et al., 2006; Napier and Graham, 2010). Despite high levels of mFAs occurring in many native species, attempts to produce high yields of mFAs in a crop species, or in *Arabidopsis* as a model, have largely been unsuccessful (Jaworski and Cahoon, 2003; Lu et al., 2006; Lu and Kang, 2008). It was shown that inefficient removal of mFAs from PC and incorporation into TAG was responsible for very low contents of conjugated FAs in TAG in transgenic *Arabidopsis* seeds (Cahoon et al., 2006). This notion was further supported by an experiment showing a castor PDAT that facilitated the transfer of hydroxy fatty acids (HFA) from PC into TAG in transgenic *Arabidopsis* (van Erp et al., 2011). The hydroxy FA, ricinoleic acid (12-hydroxyoctadec-cis-9-enoic acid; 18:1OH), constitutes approximately 90% of total FAs in castor (*Ricinus communis*) seeds. The 18:1OH is produced by the hydroxylation of 18:1 that is esterified to the *sn*-2 position of PC (Bafar et al., 1991) by the oleate Δ 12-hydroxylase (FAH12) (van de Loo et al., 1995). Seed-specific expression of FAH12 in *Arabidopsis* or the oilseed crop camelina (*Camelina sativa*) only resulted in up to ~17% total HFAs including 18:1OH and its derivatives (Broun and Somerville, 1997; Smith et al., 2003; Lu et al., 2006; Lu and Kang, 2008). Co-expressing a PDAT (*RcPDAT1A*) from castor with FAH12 significantly increased the HFA levels up to ~27% (van Erp et al., 2011). Analyzing the FA composition in different lipids during seed development indicated that increased HFA content in TAG was accompanied by a significant decrease of HFA on PC (van Erp et al., 2011). This result is in agreement with the role of PDAT transferring the HFAs from *sn*-2 PC onto DAG to form TAG, and supports the proposal that PDAT plays an important role in castor for accumulating a high level of HFAs (Bafar et al., 1991; Dahlqvist et al., 2000). In addition, co-expressing FAH12 with the *RcDGAT2* greatly increased HFA accumulation, and also resulted in lowered HFA on PC during seed development, supporting a possible relationship between efficient removal of mFA from PC and their incorporation into TAG (Burgal et al., 2008; van Erp et al., 2011).

We now recognize that the flux of acyl groups through PC poses a bottleneck for the accumulation of mFAs in transgenic seeds (Bates and Browse, 2011; van Erp et al., 2011). The PDCT catalyzes the shuffling of acyl groups between PC and DAG, thus providing a potential mechanism for the removal of polyunsaturated or other mFAs on PC for them to be incorporated into TAG. We show here that PDCT is required for efficient HFA accumulation in transgenic *Arabidopsis*. Very low levels of HFAs were detected when FAH12 was expressed in seeds of the *Arabidopsis rod1* mutant, which contains a lesion in PDCT (Lu et al., 2009). A castor PDCT enzyme encoded by the *RcROD1* gene increased HFA levels when co-expressed with FAH12. Our results demonstrate that PDCT plays an important role in seed lipid metabolism, and provides a useful tool for engineering HFA and possibly other mFAs as well in transgenic plants.

RESULTS

PDCT Is Required for Efficient HFA Accumulation

The castor fatty acid hydroxylase (FAH12) acts on PC to convert 18:1 into 18:1OH, which will then be channeled into TAG through several possible pathways (van Erp et al., 2011). The PDCT may play an important role in this process by turning over PC into HFA-containing DAG. To test this possibility, we expressed the FAH12 in the *Arabidopsis rod1* mutant and the wild type (Col-0) under control of the seed-specific phaseolin promoter as described previously (Lu and Kang, 2008). The plasmid used for transformation contained a DsRed marker, and transgenic T1 seeds were selected by screening for red fluorescence. T1 plants were grown, and 10-15 transgenic lines from each background were selected based on a ratio of fluorescent to nonfluorescent seeds of 3:1 in the T2 seeds, which suggested single insertions for the transgene *FAH12*. The fatty acid composition of transgenic red seeds was determined by analyzing fatty acid methyl esters using gas chromatography. Four novel fatty

acids were detected compared to wild type seeds, which corresponded to the HFAs previously identified as ricinoleic acid (18:1OH), densipolic acid (18:2OH), lesquerolic acid (20:1OH) and a small amount of auricollic acid (20:2OH) (Broun and Somerville, 1997; Smith et al., 2003). The total HFA contents in each of the transgenic lines ranged from 4-5% in the *rod1* mutant, compared with ~10% in the wild type background (Fig. 1). Three independent transgenic lines each from the *rod1* and WT backgrounds were raised to reach homozygosity for the transgene. Analysis of fatty acid composition in the T4 seeds confirmed the results in the T2 seeds (Table 1). These results indicated that only about half of the hydroxy fatty acids were incorporated into TAG in the *rod1* mutant compared to the wild type, and that this inefficiency was presumably caused by the loss of the PDCT function in the *rod1* mutant.

A Castor ROD1 (*RcROD1*) with PDCT Activity

Results presented above indicated that PDCT is required for an efficient metabolism of hydroxy fatty acids in transgenic *Arabidopsis* seeds. This raised the possibility that overexpressing PDCT may increase HFA accumulation in TAG by facilitating the conversion of HFA-containing PC into DAG. Considering that a PDCT enzyme from castor may have been co-evolved with FAH12 to contribute to the high content of ricinoleic acid in castor oil (Lu et al., 2006; van Erp et al., 2011), we decided to use the castor PDCT in our transgenic experiments. The *Arabidopsis* PDCT is encoded by the *ROD1* (At3g15820; *AtROD1*) gene (Lu et al., 2009). We identified a sequence from an expressed sequence tag (EST) collection of castor seeds (E. Cahoon, person. comm.) that showed homology to *AtROD1*. Full length cDNA sequence of this putative castor *ROD1*, designated *RcROD1*, was compiled by comparing the castor cDNA (Lu et al., 2007) and genomic (<http://castorbean.jcvi.org/index.php>) sequences. The deduced *RcROD1* (EMBL-Bank: EQ973818.1; UniprotKB: B9RV74) contains 285 amino acids as shown in

supplemental Figure S1. Despite great variations between the two sequences of *RcROD1* and *AtROD1*, especially at the N- and C- termini, the central regions have over 70% identical residues. The putative transmembrane domains and the catalytic triad (histidine, histidine and aspartate) are also conserved in both *ROD1* sequences (Fig. S1).

To determine whether *RcROD1* possesses the PDCT activity, the *RcROD1* cDNA sequences were cloned into the yeast expression vector p424 GPD and transfected the *Saccharomyces cerevisiae* strain HJ091 (*cpt1::LEU2 ept1-*) to extract microsomal proteins for enzyme activity assay as described previously (Lu et al., 2009). When incubated with dioleoyl- ^{14}C glycerol and non-radioactive soybean PC, microsomal preparations from HJ091 cells expressing *RcROD1* and *AtROD1* both were able to synthesize radiolabelled PC. No activity was detected in the control microsomes that were transformed with the empty vector (Fig. 2A). These assays indicate that *RcROD1*, like *AtROD1*, has PDCT activity that synthesizes ^{14}C -PC by transferring the phosphocholine headgroup from PC to ^{14}C -DAG. In our assay conditions, *RcROD1* had even higher PDCT activity at ~5.3 nmol/mg protein/min compared to *AtROD1* at ~3.5 (Fig. 2B). To test whether the castor PDCT prefers HFA-containing substrates, we incubated yeast microsomal preparations of *RcROD1* and *AtROD1* with dioleoyl- ^{14}C glycerol and di18:1OH-PC prepared from castor bean (a gift from S. Stymne). As shown in Figure 2, PDCT activity was detected for both enzymes using these PC substrates, but lower activity was found in reactions incubated with di18:1OH-PC. The PDCT activity of *AtROD1* and *RcROD1* was detected at 1.4 and 2 nmol/mg protein/min, respectively. These results were not indicative of the 18:1OH-PC as the preferred substrates for either PDCTs from castor or Arabidopsis in our assay conditions.

The function of *RcROD1* as a PDCT was further confirmed by its ability to restore the seed fatty acid composition in the Arabidopsis *rod1* mutant to the wild type level when the *RcROD1* cDNA was introduced into the mutant (Table 1).

RcROD1 Enhances HFA accumulation in TAG in Arabidopsis

To determine whether overexpression of PDCT may increase HFA accumulation, the castor genes *RcROD1* and *FAH12* were co-expressed in the *rod1* mutant and wild type Col-0 plants under the seed specific promoters (Supplemental Fig. S2). As described above, transgenic seeds were selected based on DsRed expression, and the red T2 seeds were analyzed for the fatty acid composition. The levels of HFA content in the *rod1* transformants containing both *RcROD1* and *FAH12* were similar to those in the WT background expressing *FAH12* alone (Fig. 1; Table 1). This is consistent with the results that *RcROD1* is able to restore fatty acid composition of the *rod1* mutant to the wild type level. Interestingly, co-expression of *RcROD1* with *FAH12* in the WT Arabidopsis increased the HFA contents from ~10% to ~20%, at the expense of mostly 18:2 and 18:3 polyunsaturated FAs as well as 20:1. The data presented in Table 1 is from the T4 seeds of three homozygous lines as determined by the segregation ratios of the DsRed marker. These results demonstrated that *RcROD1* was able to significantly increase HFA accumulation when co-expressed with the castor *FAH12* in Arabidopsis seeds, and this increase was stably inherited over multiple generations.

RcROD1 Increases HFA-TAGs and Changes Their Molecular Species

To gain insight in biochemical mechanisms of the PDCT effects on HFA metabolism in transgenic seeds, we analyzed the HFA-containing TAG and its regiochemical composition in our transgenic lines. There are four molecular species of TAG in *FAH12* transgenic Arabidopsis seeds, namely 0-, 1-, 2- and 3-HFA TAG, according to the number of HFAs esterified onto the glycerol backbone. These TAGs are present in all the *FAH12* expressing lines in the *rod1* mutant or WT backgrounds. All classes of HFA-TAGs are significantly reduced in the *rod1*-*FAH12* seeds compared to *FAH12* expression in WT or *RcROD1*-complemented *rod1* seeds. The

RcROD1 overexpression in the Col-FAH12 seeds significantly lowers the 0-HFA TAG proportion and increases the amount of all HFA-TAG species, especially the 2- and 3-HFA TAGs (11.3% and 4.5%, respectively), which are more than doubled compared to Col-FAH12 (5.5% and 2.1%, respectively) (Fig. 3). These results suggest that *RcROD1* effectively metabolizes HFAs into TAG in FAH12 expressing Arabidopsis seeds.

We further analyzed regiochemical composition of HFA containing TAGs from the above transgenic lines. Figure 4 shows the HFAs as a percentage of FAs at the *sn*-2 versus the *sn*-1/3 position for individual 1- and 2-HFA TAG molecular species. In 1-HFA-TAG, PDCT did not significantly affect the percentage of HFAs in *sn*-2 and *sn*-1/3 positions. However, PDCT caused dramatic changes in the 2-HFA-TAGs. Compared to Col-FAH12 lines, the percentage of HFAs in the *rod1*-FAH12 lines decreased in the *sn*-2 position (from 31% to 24.5%) and increased in the *sn*-1/3 position (from 69% to 75.5%). Overexpression of *RcROD1* in the Col-FAH12 line increased HFA percentage in the *sn*-2 position from 31% to 42% and decreased in the *sn*-1/3 position from 69% to 58% (Fig. 4B). Together with data in Figure 3, these results indicate that PDCT activity of *RcROD1* effectively mobilize HFAs from PC into DAG and increase HFA contents in TAG of the transgenic Arabidopsis seeds. These results are the expected because the hydroxylase works on *sn*-2 PC, and thus we would expect PDCT to produce *sn*-2 HFA DAG and thus more *sn*-2 HFA TAG.

Changes of HFA in PC and TAG during Seed Development

The FAH12 hydroxylase acts on 18:1 at the *sn*-2 position of the membrane lipid PC (Bafar et al., 1991). Castor seeds can efficiently transfer HFAs into TAG while keeping a very low level of HFAs in membrane lipids. Since PDCT catalyzes the inter-conversion of PC and DAG, we reasoned that the increased HFA accumulation in TAG by expressing *RcROD1* could have resulted from efficient HFA migration from PC to DAG, which was then turned into TAG. Therefore, we measured the HFA

contents in PC and TAG in the above transgenic lines during TAG synthesis. Total lipids were extracted from seed samples of 7 to 18 days after flowering (DAF) and separated by thin-layer chromatography (TLC). The PC and neutral lipid (TAG) fractions were recovered from the TLC plates and analyzed by GC. HFA accumulation in TAG increased during seed development, and reached the maximum levels of 5% and 10% in 15-18 DAF in the *rod1* and WT backgrounds, respectively (Fig. 5A, C). As was observed previously in transgenic *Arabidopsis* expressing FAH12 (Thomaeus et al., 2001), we also found that HFAs transiently accumulated in PC during seed development, and reached a maximum of 5-6% in the mid-late stage of seed development (~15 DAF) (Fig. 5B, D). There was no significant difference between the *rod1* mutant and Col backgrounds.

We then analyzed the *RcROD1* expressing lines for fatty acid composition in PC and TAG during seed development. As shown in Figure 5, HFA contents in PC decreased significantly in both *rod1* and WT backgrounds, and were apparently associated with increased levels of HFA in TAG. At 15 to 16 DAF, PC in *rod1*-FAH12 seeds contained 4.9% HFA, compared with 3.6% in the seeds of *rod1*-FAH12-*RcROD1* lines (Fig. 5B). At the same time, HFA contents in TAG increased from 6.1% to 8.1% (Fig. 5A). Similarly, PC in Col-FAH12 seeds contained 6.2% HFAs, compared with 5.4% in Col-FAH12-*RcROD1* seeds (Fig. 5D). The HFAs in TAG increased from 8.3% to 16% during the same period of seed development (Fig. 5C).

Expression of *RcROD1* in the CL37 and CL7-*RcDGAT2* Backgrounds

It appears that overexpressing *RcROD1* may significantly increase the HFA accumulation in transgenic *Arabidopsis* seeds. This could have been resulted from the enhanced conversion of PC into HFA-containing DAG. It has been shown previously that the castor enzyme *RcDGAT2* prefers such DAG molecules and dramatically increases HFA accumulation in TAG when co-expressed with FAH12 in *Arabidopsis*

seeds (Burgal et al., 2008). To explore the possibility of synergistic effects of *RcROD1* and *RcDGAT2* on HFA levels, the above *RcROD1* construct was transformed into the CL7-*RcDGAT2* line, which was produced by co-transformation of *RcDGAT2* and *FAH12* on the Arabidopsis *fae1* background (Burgal et al., 2008). As a control, *RcROD1* was also transformed into the *FAH12*-containing CL37 line, which was produced on the same background (Lu et al., 2006). Because of the mutation at the *FAE1* locus in the *fae1* mutant (Kunst et al., 1992), the CL37 and CL7-*RcDGAT2* lines do not accumulate very long chain (>18) fatty acids including 20:1 and 20:1OH, and the seed oils contain 17% and 25% HFAs respectively in our analysis (Table 2).

The *RcROD1* transgenic plants in these backgrounds were grown, and 11 and 7 homozygous lines in CL37 and CL7-*RcDGAT2* were obtained for analysis by GC to determine their fatty acid composition. The results are shown in Table 2, which compare the HFA levels in the CL7-*RcDGAT2*-*RcROD1* triple transgenic and the CL37-*RcROD1* double transgenic with the CL37 control. Expression of *RcROD1* increased HFA levels in the CL37 background from 17% to 22%, and further increased in the CL7-*RcDGAT2* from 25% to 28.5%. These results clearly indicated that *RcROD1* may significantly increase HFA accumulation in transgenic Arabidopsis seeds.

We also conducted regiochemical analysis of HFA containing TAGs in CL37-*RcROD1*, CL7-*RcDGAT2*-*RcROD1* and compared with CL37 and CL7-*RcDGAT2*. Similar to the results shown in Figure 4, we found that the percentage of HFAs in the *sn*-2 position of the 2-HFA-TAGs increased in the *RcROD1* lines in both CL37 and CL7-*RcDGAT2* backgrounds (Fig. 6). These results strongly support the role of the PDCT in transferring HFAs from PC to DAG by phosphocholine headgroup exchange.

The castor PDCT is encoded by the *RcROD1* gene which shows close homology to the Arabidopsis *AtROD1*. Surprisingly however, overexpressing *AtROD1* using the same vector in the CL7-*RcDGAT2* lines did not result in significant increases of HFA

contents (data not shown).

***Rc*ROD1 Partially Restores Oil Accumulation in FAH12 Seeds**

It has been shown that expressing fatty acid hydroxylases in transgenic *Arabidopsis* caused a significant reduction of oil accumulation in seeds (Dauk et al., 2007; Bursal et al., 2008; Bates and Browse, 2011). It has been demonstrated that the flux of DAG through PC represents a major bottleneck for the accumulation of unusual FAs in TAG of transgenic *Arabidopsis* seeds (Bates and Browse, 2011). Given its important role in converting PC into DAG, we reasoned that PDCT may help relieve this metabolic bottleneck and increase oil content in HFA accumulating seeds. We measured seed oil content in *Rc*ROD1 transgenic lines and compared that with the FAH12 lines. The CL37 contained significantly reduced amount of oil, and we measured it at 4.4 µg/seed. The oil content in CL37-*Rc*ROD1 seeds is increased to 5.4 µg/seed (Fig. 7B). Although the level is not as high as in the CL7-*Rc*DGAT2 seeds (Bursal et al., 2008; van Erp et al., 2011), this substantial increase indicate that *Rc*ROD1 can partially relieve the bottleneck of HFA-TAG accumulation. This is further supported by the measurement of oil content in *rod1*-FAH12, *rod1*-FAH12/*Rc*ROD1 lines (Fig. 7A). The *rod1*-FAH12 seeds contain 4.6 µg/seed, complementing with *Rc*ROD1 in this line increases oil content to 6.5 mg/seed, nearly as high as the Col-FAH12 line at 7.4 µg/seed. These results also support our conclusion that PDCT is required for efficient metabolism of HFAs in transgenic *Arabidopsis* seeds.

Discussion

Triacylglycerols stored in seeds display an immense diversity of fatty acid (FA) composition among different plant species. Efficient utilization of such a rich resource in food and industrial applications has been hindered by a lack of understanding of the mechanisms of FA modification during TAG synthesis. The traditional biochemical

and genetic studies continue to yield fruitful insights in this area. One of the recent breakthroughs is the elucidation of the importance of the dynamic metabolism of phosphatidylcholine (PC) in developing oilseeds. In a metabolic labeling study using developing *Arabidopsis* seeds, Bates et al. demonstrated that the majority of *de novo* DAG was used for PC synthesis instead of being converted into TAG, suggesting that the major flux of TAG synthesis in *Arabidopsis* utilizes a *de novo* DAG to PC followed by a PC-derived DAG to TAG pathway (Bates and Browse, 2011). Therefore, the PC molecules are not only a major component of cellular membranes, but also provide a predominant source of DAG for synthesizing TAG. Since PC is the enzymatic substrate of FA desaturation and other types of modification such as hydroxylation, the incorporation of FAs into PC and the subsequent removal of modified fatty acids from PC will greatly affect the final composition of TAG. In support of this, it has been demonstrated that in the castor hydroxylase (FAH12) expressing *Arabidopsis*, hydroxy fatty acids containing *de novo* DAG was not efficiently converted to PC, thus representing a major bottleneck for the accumulation of HFAs in TAG of transgenic *Arabidopsis* seeds (Bates and Browse, 2011). In another study, it was shown that co-expressing a castor PDAT (*RcPDAT1A*) with the FAH12 significantly increased the HFA accumulation to 27%, compared to 17% when expressing the FAH12 alone in transgenic *Arabidopsis* seeds (van Erp et al., 2011). The PDAT directly transfers the HFAs from the *sn*-2 position of PC to the *sn*-3 position of DAG, producing TAG. These studies suggest that efficient mechanisms are required for the acyl fluxes through PC in genetic engineering of fatty acid modifications in oilseeds.

In our previous study using forward genetics (Lu et al., 2009), we discovered a PDCT in *Arabidopsis* that controls roughly 40% of 18:1-CoA entering PC for desaturation. The interconversion of PC and DAG mediated by PDCT also transfers the desaturated 18:2 and 18:3 from PC into DAG, which will then form TAG. Our results and others indicate that the PDCT reaction is responsible for most of the interconversion of DAG to PC in developing seeds (Bates et al., 2009; Lu et al.,

2009). As a result, the PDCT deficient *rod1* mutant in *Arabidopsis* accumulated increased 18:1 and concomitantly decreased polyunsaturated 18:2 and 18:3 in seed TAG. Here we show that PDCT is also required for efficient metabolism of HFAs in transgenic *Arabidopsis* seeds. Expressing the FAH12 in the *rod1* mutant only accumulated 5.5% HFAs in TAG, about half of the amount in the WT background, which is ~10%. Co-expressing the FAH12 with the castor PDCT encoded by *RcROD1* in the wild type *Arabidopsis* seeds significantly increased the accumulation of HFAs, at a doubled 20% compared to expressing the FAH12 alone (Table 1).

These results are in agreement with the function of PDCT that catalyzes the inter-conversion between PC and DAG. Overexpressing the castor PDCT (*RcROD1*) enhances the acyl groups entering PC, where they are desaturated or hydroxylated when expressing FAH12, and then returning to DAG. This symmetrical reaction (one DAG molecule is generated for each DAG consumed) produces 18:1OH-DAG and 18:2OH-DAG, along with the regular 18:2- and 18:3-DAG, resulting in enriched HFA-DAGs in the DAG pool and subsequently increase the accumulation of these modified fatty acids in TAG. This hypothesis is supported by several lines of evidence obtained in this study.

First, we show that the increased HFA content in TAG of the *RcROD1* expressing lines is associated with lowered HFA amounts in PC (Fig. 5), a similar effect as seen in the *RcPDAT1A* expressing lines (van Erp et al., 2011). This indicates that PDCT is required for efficient removal of HFAs from PC after they are produced.

Second, *RcROD1* significantly increases the amount of HFA-containing TAGs, especially the 2- and 3-HFA-TAGs, in the Col-FAH12 lines shown by the TAG molecular species analysis (Fig. 3). Regiochemical analyses indicate that *RcROD1* increases HFAs on the *sn*-2 position while decreases on the *sn*-1/3 positions in the 2-HFA-TAGs (Fig 4). Similar changes are also detected in the CL37 and CL7-*RcDGAT2* lines (Fig. 6). This suggests that the stereochemistry of 1-HFA-DAG pool contains more *sn*-2 HFA in the *RcROD1* lines. The HFAs are synthesized from FAH12 acting on acyl groups esterified at the *sn*-2 position of PC. Overexpression of

RcROD1 may enhance the conversion of the *sn*-2 HFA-PC into DAG and therefore result in more *sn*-2 HFA-DAGs in the DAG pool. The acyltransferases DGAT or PDAT then incorporate a second HFA onto the *sn*-3 in DAG and form 2-HFA-TAGs. An alternative scenario is that *RcROD1* enhances the flux of *sn*-1-HFA-DAG into PC (by passing the bottleneck) and allowing production of *sn*-1,2-HFA-PC, which is then acylated with a normal FA by DGAT or PDAT.

Third, we noticed that the increased HFAs in the Col-FAH-*RcROD1* lines are at the expense of C18 PUFAs (18:2 + 18:3) and 20:1, the products of the desaturation and the elongation of 18:1, respectively (Table 1). The decrease of PUFAs is expected because the competition of FAH12 with FAD2 for the 18:1 substrate. It is interesting that the 20:1 level is also decreased. The very long chain FA (20:1) is synthesized from 18:1-CoA outside PC, probably most of which are products directly exported from plastids. Therefore the elongation and the incorporation into PC are two competing pathways for the 18:1 substrates. We have shown that the PDCT is responsible for about 40% of the 18:1 that enters PC (Lu et al., 2009). Overexpressing the castor PDCT may direct more 18:1 to enter PC for desaturation and hydroxylation, and reduce the flow towards the elongation pathway. The elongation pathway is blocked in the CL37 and CL7-*RcDGAT2* lines that were generated in the *fae1* background. The CL37 seeds already contain 17% HFAs, which is substantially higher than the levels in the Col-FAH12 lines in this study and the lines reported previously (Smith et al., 2003). However, *RcROD1* still significantly increases HFA levels in the CL37 and CL7-*RcDGAT2* lines (Table 2).

Our results of *RcROD1* in different transgenic lines, along with previous reports (Smith et al., 2003; Lu et al., 2006; Burgal et al., 2008; van Erp et al., 2011), point out the complexity of the interactions among different mechanisms that channel HFAs into TAG. PDCT enhances the acyl fluxes through PC and thus reduces the pathway towards elongation outside of PC. The rapid PC-DAG turn around may also have an impact on the acyl editing by reducing the HFA-CoA substrates available for DGAT2. In this regard, it is noteworthy that overexpression of *RcROD1* results in reduced

HFAs in *sn*-1/3 positions and increased content in the *sn*-2 position (Fig. 6). It is interesting to note the different stereochemistry observed in the 1-HFA-TAG for the HFA expressing Col (and *rod1*) and the CL37 (and CL7-*RcDGAT2*) lines (Fig. 4, 6). The distribution of HFAs on *sn*-2 and *sn*-1/3 position is opposite between these two groups of lines. It is possible that the *fae1* in the CL37 and CL7-*RcDGAT2* lines at least partly contributes to a shift of more HFAs into the *sn*-2 position in the 1-HFA-TAG. More studies, such as metabolic labeling experiments, are needed to understand the interactions between these competing pathways.

Finally, to support our hypothesis, we show that the *RcROD1* partially compensates for the decreased oil accumulation in *FAH12* expressing seeds. It has been shown that the *FAH12* transgenic seeds contain reduced oil content (Dauk et al., 2007; Burgal et al., 2008; Bates and Browse, 2011). This may have resulted from the buildup of inefficiently used HFAs that cause feedback inhibition of FA synthesis or activate the FA β -oxidation (Moire et al., 2004; Bates and Browse, 2011). The *rod1*-*FAH12* lines accumulate higher amount of HFAs in PC during seed development (Fig. 5), and the seeds contain much reduced HFA and total FAs (Fig. 7). The FA content in CL37 is also significantly reduced (van Erp et al., 2011). Expressing *RcROD1* in *rod1*-*FAH12* and CL37 significantly increases seed oil content (Fig. 7), although not to the control levels in WT or *RcDGAT2* expressing lines. These results suggest that the castor PDCT efficiently converts PC, especially HFA-PC, into DAG and subsequently into TAG.

The castor and Arabidopsis *ROD1* homologs both possess the PDCT activity. In this study, we have used a castor PDCT in our experiments to co-express with the *FAH12*, and resulted in increased HFA accumulation in all lines tested including the WT (Col), CL37 and CL7-*RcDGAT2* (Table 1, 2). However, the Arabidopsis PDCT failed to produce such effects. This result was surprising since we expected that over-expression of either PDCT would enhance the acyl fluxes through PC. We have previously proposed that for efficient metabolism of HFAs in transgenic seeds, one needs to co-express other enzymes from the same native species (e.g., castor) that

may have coevolved with FAH12 (Lu et al., 2006). This hypothesis has been supported by co-expressing the *RcDGAT2* and *RcPDAT1A* with FAH12 in transgenic *Arabidopsis* seeds (Burgal et al., 2008; van Erp et al., 2011). These results are in agreement with the observations that suggest a preference for HFA-containing substrates of the castor enzymes (Dahlqvist et al., 2000; Burgal et al., 2008). Our results of *RcROD1* seemingly also support the co-evolution theory, but we cannot demonstrate the substrate preference for HFA-containing PC of the castor PDCT (Fig. 3) due to limited availability of substrates. However, the symmetrical reaction of DAG-PC conversion may not necessarily require substrate selectivity of the PDCT enzyme (i.e. 18:2/3-, 18:1OH-PC over 18:1-PC; 18:1-DAG over 18:2/3-, 18:1OH-DAG), since the desaturases and the hydroxylase will enrich the PC pool with the polyunsaturated and hydroxylated FAs. On the other hand, although the *RcROD1* protein has been detected in castor endosperm (Brown et al., 2012), previous results suggested that the castor PDCT unlikely preferred HFA-containing substrates since microsomal preparations from castor endosperm rapidly turned the glycerol labelled di-ricinoleoyl-DAG to TAG whereas dioleoyl-DAG was preferentially used in PC synthesis (Bafar et al., 1991). Moreover, Dahlqvist et al. showed that castor bean microsomes preferentially metabolised ricinoleoyl-PC to TAG (probably via PDAT) whereas oleoyl-PC was preferentially metabolised to DAG (probably via PDCT) (Dahlqvist et al., 2000).

Nevertheless we show in this report that PDCT is required for efficient metabolism of hydroxy fatty acids in the FAH12 transgenic *Arabidopsis* seeds. Co-expressing a castor PDCT with FAH12 significantly increases the HFA accumulation. The strong PDCT activity conferred by the *RcROD1* gene may enhance the fluxes of fatty acids through PC, thus partly overcome the bottleneck of modified fatty acid accumulation in transgenic seeds proposed by previous research (Bates and Browse, 2011).

Materials and methods

Construction of Plasmid Vectors

For co-expressing the FAH12 and *RcROD1* in Arabidopsis seeds, the castor open reading frame sequences *RcROD1* and *RcFAH12* were cloned into a binary vector pBinGlyRed2, kindly provided by Edgar Cahoon (University of Nebraska – Lincoln), which contains a DsRed selection marker (Clontech, Mountain View, CA, USA) behind the constitutive cassava vein mosaic virus (CVMV) promoter (Verdaguer et al., 1996). *RcROD1* was placed behind the seed specific glycinin promoter, and the *RcFAH12* was driven by a seed-specific napin promoter (Supplemental Fig. S2). To express *RcROD1* in the CL7-*RcDGAT2* line, the *RcROD1* sequences were cloned into the pGATE-PHAS-DsRed vector under the seed-specific phaseolin promoter as described previously (Lu et al., 2006).

Plant Material and Transformation

Arabidopsis (*Arabidopsis thaliana*) ecotype Columbia (Col-0), mutant *rod1* (Lu et al., 2009), transgenic line CL37 expressing FAH12 in the *fae1* mutant background (Lu et al., 2006), and the CL7-*RcDGAT2* co-expressing FAH12 and *RcDGAT2* (Burgal et al., 2008) are used in this study. All Arabidopsis plants were grown in controlled-environment chambers at 22°C under a 16-h photoperiod of 150 $\mu\text{mol quanta m}^{-2} \text{sec}^{-1}$ photosynthetically active radiation. Arabidopsis were transformed by the floral dipping procedure (Clough and Bent, 1998) following electroporation of individual cDNA clones into *Agrobacterium tumefaciens* strain GV3101 (pMP90). Transgenic seeds were identified by examination under green illumination using a red filter (Stuitje et al., 2003; Lu et al., 2006).

Lipid Extraction

Total lipids were extracted from mature seeds or developing seeds on ice as described before (Blight and Dyer, 1959) by suspending in 3 ml of a chloroform/methanol (1/2; v/v) mixture and then seeds were ground in mortar. Ground seeds and solvent were transferred to a 10 ml glass tube and the mixture made up to 5.8 ml by the addition of 1ml chloroform and 1.8ml of 0.9% NaCl, so the final relative volume of chloroform/methanol/0.9% NaCl is 1:1:0.9. The chloroform phase was transferred to a clean glass tube and evaporated to dryness under a stream of N₂. Lipids were resuspended in a small volume of chloroform and stored at -20 °C.

PC Analysis

Phosphatidylcholine (PC) was resolved by TLC of the total lipids with a solvent system consisting of CHCl₃: MeOH: H₂O:30% ammonium hydroxide (65:35:3:2.5 v/v/v/v) (Cahoon et al., 2006). Lipid bands were visualized under UV light after staining with 0.005% primulin in 80% acetone, and TAG and PC were collected for GC analysis as below.

GC Analysis

The fatty acids of lipid fractions and the whole Arabidopsis seeds were derivatized to fatty acid methyl esters (FAMES) in 1 ml of 2.5% sulfuric acid in methanol at 80°C for 90 min (Browse et al., 1986). FAMES were injected into a Shimadzu 2010 GC fitted with a narrowbore column (HP-Innowax; 30 m × 0.25 mm i.d. × 0.25 µm; Agilent Technologies). The oven temperature was programmed at 190°C initially followed by an increase of 20°C/min to 250°C and maintained for 9 min.

Characterization of TAG Species and Regiochemical Analysis of 1- and 2-HFA-

TAG

Neutral lipids were separated by TLC (silica gel G60 plates; EM Separations Technology) with hexane/diethyl ether/acetic acid (140:60:2, v/v/v) (Smith et al., 2003) as the developing solvent system. Castor oil was used to produce standards of TAG species with hydroxy fatty acids as described previously (Smith et al., 2003; van Erp et al., 2011); Lipid bands were visualized under UV light after staining with 0.01% primulin in 80% acetone. The fractions of TAG with zero, one, two, or three hydroxy fatty acid residues, namely 0-HFA-TAG, 1-HFA-TAG, 2-HFA-TAG and 3-HFA-TAG respectively, were scraped from the TLC plate into glass tubes and FAMES were prepared for GC analysis as above. Calculation of the percentages of HFAs at the sn-2 position and at the sn-1/3 position was done as described (van Erp et al., 2011).

Seed Oil Content Measurements

Seed oil content was determined by GC analysis of FAMES following the method of Li et al. (Li et al., 2006) using 30 seeds and 68 µg of triheptadecanoin as a TAG standard.

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Figure legends

Figure 1. Hydroxy fatty acid levels in Arabidopsis seeds transformed with the castor fatty acid hydroxylase FAH12. The lines are *rod1* mutant, Col (WT), and *rod1*

transformed with *RcROD1*. The data represent averages \pm SE from 10-15 independent T2 lines.

Figure 2. Castor *ROD1* (*RcROD1*) encodes a PDCT enzyme. A, Radio-TLC image of PDCT assays. Microsomes from HJ091 *S. cerevisiae* cells transfected with p424-*AtROD1* (At) or p424-*RcROD1* (Rc) were incubated with [¹⁴C-glycerol]di18:1-DAG and Soy or 18:1OH-PC (1mM). V, control with empty p424 vector. B, The PDCT activity measured as the amount of radioactive PC formed. Data represent means of three independent reactions \pm SE.

Figure 3. Molecular species composition of HFA containing TAGs of *rod1* FAH12, *rod1* FHA12/*RcROD1*, Col FAH12, and Col FAH12/*RcROD1* seeds. 0-, 1-, 2-, and 3-HFA represent TAG molecular species with zero, one, two, or three HFAs, respectively (no stereochemistry implied). The HFAs represent the sum of ricinoleate (18:1-OH) and densipolate (18:2-OH) and a small amount of lesquerolic acid (20:1-OH). The data represent averages of three replicates \pm SE.

Figure 4. Regiochemical analysis of 1- and 2-HFA-TAG species in *rod1* FAH12, Col FAH12, and Col FAH12/*RcROD1* seeds: mol % HFA at the *sn*-2 position compared with the *sn*-1/3 position in 1- and 2-HFA-TAG. A, 1-HFA-TAG. B, 2-HFA-TAG. White bars, *rod1* FAH12; slashed bars, Col FAH12; black bars, Col FAH12/*RcROD1*. The data represent averages of three replicates \pm SE.

Figure 5. Changes in the percentage of hydroxy fatty acids in PC and TAG in transgenic *Arabidopsis* lines during seed development. A and B, *rod1* mutant transformed with FAH12 alone or together with *RcROD1*; C and D, Wild type (Col) transformed with FAH12 or together with *RcROD1*. The data represent averages of three replicates \pm SE.

Figure 6. Regiochemical analysis of 1- and 2-HFA-TAG species in CL37, CL37 *RcROD1*, CL7 *RcDGAT2*, and CL7 *RcDGAT2/RcROD1* seeds: mol % HFA at the *sn*-2 position compared with the *sn*-1/3 position in 1- and 2-HFA-TAG. A and C, 1-HFA-TAG. B and D, 2-HFA-TAG. White bars, CL37 (A, B) or CL7 *RcDGAT2* (C, D); black bars, CL37 *RcROD1* (A, B) or CL7 *RcDGAT2/RcROD1* (C, D). The data represent averages of three replicates \pm SE.

Figure 7. Seed oil content in Col, *rod1* and FAH12 transformed lines. FAs (total μ g per seed) were determined for each line listed in Table 1 and 2. The data (averages \pm SE) represent three replicates for each line.

Table 1. Fatty acid composition of transgenic *Arabidopsis* seeds of T4 lines expressing castor *RcFAH12* and *RcROD1*

The transgenic lines of these genes were obtained from independent transformation events. Data represent measurements of three homozygous lines (mean \pm SE)

Line	fatty acid composition (wt % of total)									Total
	16:0	18:0	18:1	18:2	18:3	20:1	18:1OH	18:2OH	20:1OH	HFA*
Col-0	8.0 \pm 0.1	2.7 \pm 0.1	15.8 \pm 0.2	30.6 \pm 0.2	18.7 \pm 0.1	21.3 \pm 0.1				
<i>rod1</i>	7.9 \pm 0.1	2.5 \pm 0.1	33.5 \pm 0.2	15.5 \pm 0.1	13.1 \pm 0.1	25.3 \pm 0.1				
<i>rod1</i> RcROD1	7.2 \pm 0.5	3.0 \pm 0.5	16.2 \pm 1.7	29.7 \pm 1.2	17.6 \pm 0.4	20.7 \pm 0.9				
<i>rod1</i> FAH12	12.4 \pm 0.6	5.2 \pm 0.2	38.3 \pm 0.8	14.5 \pm 0.3	5.9 \pm 0.1	15.4 \pm 0.5	2.8 \pm 0.1	1.2 \pm 0.1	1.5 \pm 0.1	5.5 \pm 0.1
<i>rod1</i> FAH12/RcROD1	10.1 \pm 0.1	4.8 \pm 0.2	25.5 \pm 0.3	24.3 \pm 0.2	11.6 \pm 0.5	11.1 \pm 1.5	6.2 \pm 0.2	3.0 \pm 0.4	0.6 \pm 0.1	9.7 \pm 0.6
Col FAH12	10.1 \pm 0.4	4.7 \pm 0.3	19.6 \pm 0.4	22.2 \pm 0.2	10.7 \pm 0.9	17.9 \pm 0.2	6.3 \pm 0.1	2.1 \pm 0.1	1.6 \pm 0.1	9.9 \pm 0.3
Col FAH12/RcROD1	10.0 \pm 0.5	5.1 \pm 0.3	24.2 \pm 0.7	17.5 \pm 1.2	5.2 \pm 0.1	14.9 \pm 0.6	12.7 \pm 0.2	4.6 \pm 0.5	2.6 \pm 0.1	19.9 \pm 0.5

* Excluding small amounts of 20:2OH.

Table 2. Fatty acid composition of transgenic *Arabidopsis* seeds expressing *FAH12* and *RcROD1*

CL7 and CL37 are the *FHA12* transformed lines in the *Arabidopsis fae1* mutant background. The transgenic lines of these genes were obtained from independent transformation events. Data represent measurements of three homozygous lines (mean \pm SE).

Line	fatty acid composition (wt % of total)							Total
	16:0	18:0	18:1	18:2	18:3	18:1OH	18:2OH	HFA
CL37	12.2 \pm 0.3	5.6 \pm 0.5	38.3 \pm 1.2	19.8 \pm 0.9	5.3 \pm 0.4	13.1 \pm 0.4	4.1 \pm 0.2	17.2 \pm 0.2
CL37 <i>RcROD1</i>	11.6 \pm 1.1	5.7 \pm 0.5	33.7 \pm 1.3	19.6 \pm 0.9	6.1 \pm 0.3	18.3 \pm 0.6	4.9 \pm 0.3	23.3 \pm 0.6
CL7 <i>RcDGAT2</i>	10.6 \pm 0.3	5.4 \pm 0.4	30.1 \pm 1.5	20.7 \pm 1.0	6.2 \pm 0.4	20.1 \pm 0.6	4.6 \pm 0.3	24.7 \pm 0.6
CL7 <i>RcDGAT2/RcROD1</i>	13.1 \pm 0.3	4.8 \pm 0.2	23.1 \pm 0.5	21.4 \pm 0.5	7.6 \pm 0.3	23.1 \pm 0.4	5.4 \pm 0.3	28.5 \pm 1.0

Supplemental Data:

Figure S1. Comparison of deduced amino acid sequences of *At*ROD1 and *Rc*ROD1. Identical residues are shaded. Putative transmembrane domains are underlined. The putative catalytic triads are shown in boxes.

Figure S2. The open reading frame nucleotide sequence of *Rc*ROD1 and the plasmid construct for co-expressing *Rc*ROD1 and *FAH12*.

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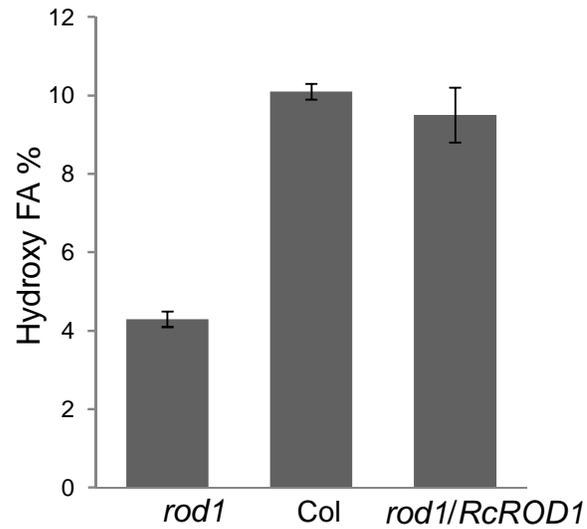


Figure 1. Hydroxy fatty acid levels in Arabidopsis seeds transformed with the castor fatty acid hydroxylase FAH12. The lines are *rod1* mutant, Col (WT), and *rod1* transformed with *RcROD1*. The data represent averages \pm SE for 10-15 individual T2 plants for each line.

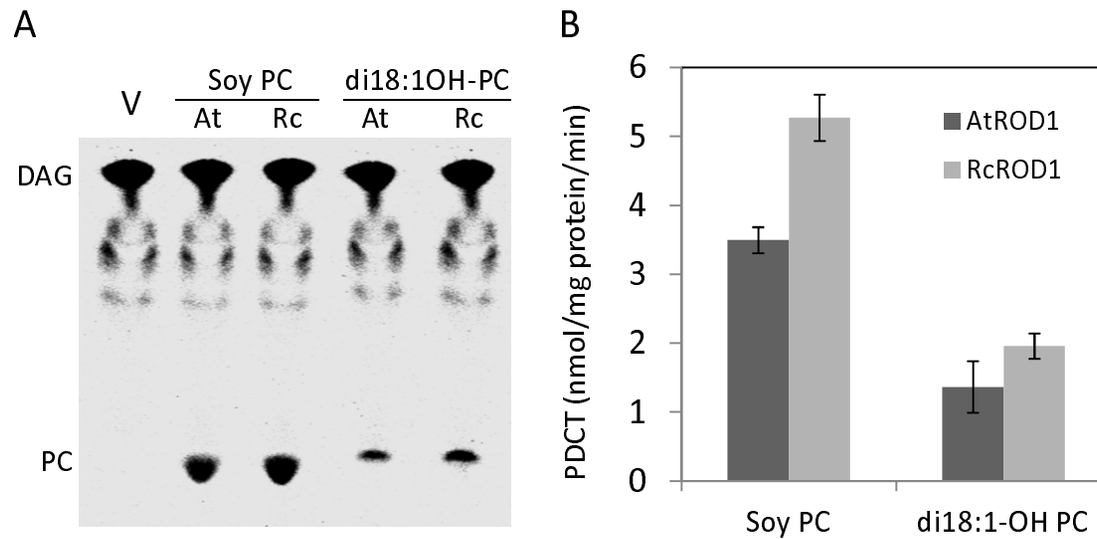


Figure 2. Castor *ROD1* (*RcROD1*) encodes a PDCT enzyme. A, Radio-TLC image of PDCT assays. Microsomes from HJ091 *S. cerevisiae* cells transfected with p424-*AtROD1* (At) or p424-*RcROD1* (Rc) were incubated with [¹⁴C-glycerol]di18:1-DAG and Soy or di18:1OH-PC (1mM). V, control with empty p424 vector. B, The PDCT activity measured as the amount of radioactive PC formed. Data represent means of three independent reactions \pm SE.

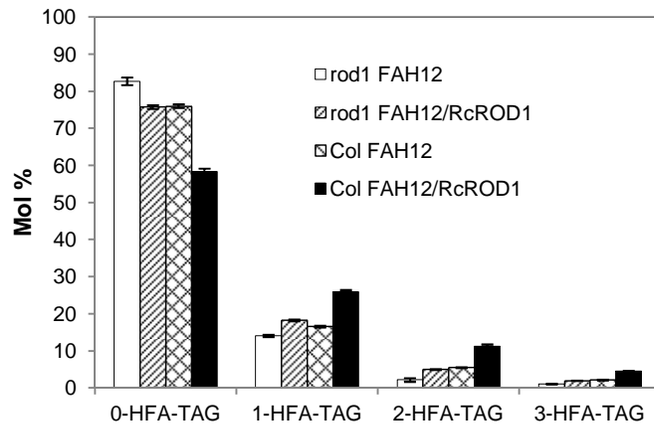


Figure 3. Molecular species composition of HFA containing TAGs of *rod1* FAH12, *rod1* FAH12/RcROD1, Col FAH12, and Col FAH12/RcROD1 seeds. 0-, 1-, 2-, and 3-HFA represent TAG molecular species with zero, one, two, or three HFAs, respectively (no stereochemistry implied). The HFAs represent the sum of ricinoleate (18:1-OH) and densipolate (18:2-OH) and a small amount of lesquerolic acid (20:1-OH). The data represent averages of three replicates \pm SE.

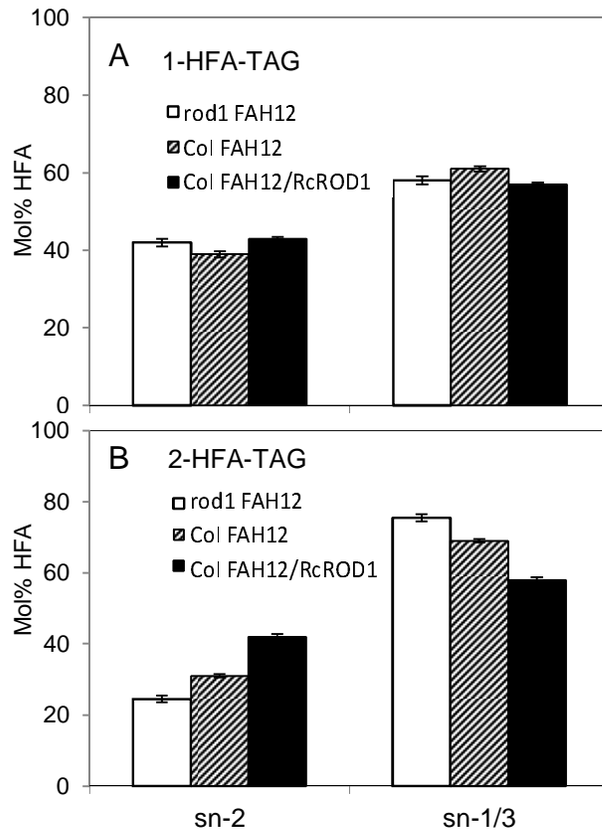


Figure 4. Regiochemical analysis of 1- and 2-HFA-TAG species in *rod1* FAH12, Col FAH12, and Col FAH12/*RcROD1* seeds: mol % HFA at the *sn-2* position compared with the *sn-1/3* position in 1- and 2-HFA-TAG. A, 1-HFA-TAG. B, 2-HFA-TAG. White bars, *rod1* FAH12; slashed bars, Col FAH12; black bars, Col FAH12/*RcROD1*. The data represent averages of three replicates \pm SE.

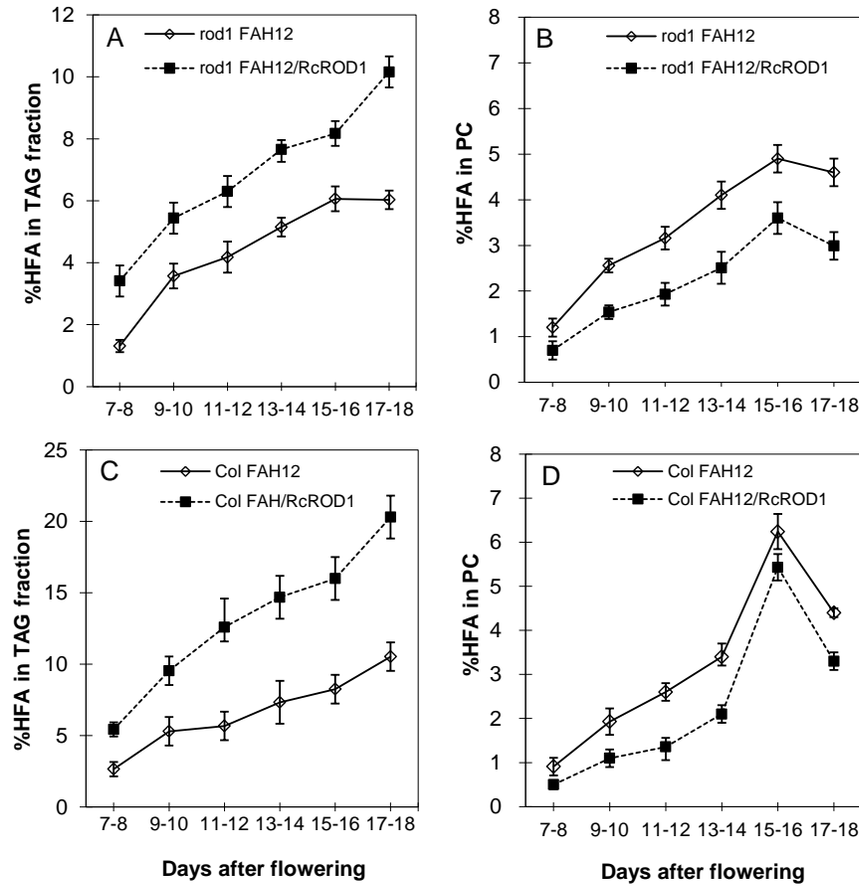


Figure 5. Changes in the percentage of hydroxy fatty acids in PC and TAG in transgenic Arabidopsis lines during seed development. A and B, *rod1* mutant transformed with FAH12 alone or together with *RcROD1*. C and D, Wild type (Col) transformed with FAH12 or together with *RcROD1*.

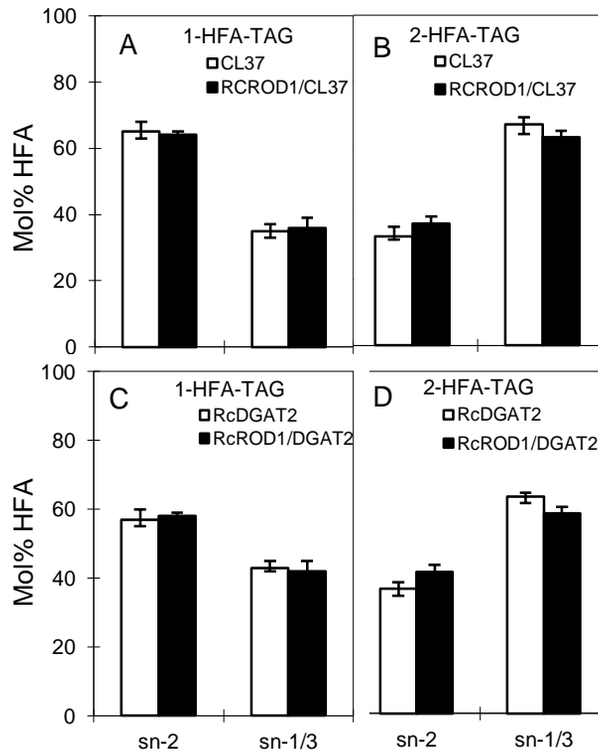


Figure 6. Regiochemical analysis of 1- and 2-HFA-TAG species in CL37, CL37 *RcROD1*, CL7 *RcDGAT2*, and CL7 *RcDGAT2/RcROD1* seeds: mol % HFA at the *sn-2* position compared with the *sn-1/3* position in 1- and 2-HFA-TAG. A and C, 1-HFA-TAG. B and D, 2-HFA-TAG. White bars, CL37 (A, B) or CL7 *RcDGAT2* (C, D); black bars, CL37 *RcROD1* (A, B) or CL7 *RcDGAT2/RcROD1* (C, D). The data represent averages of three replicates \pm SE.

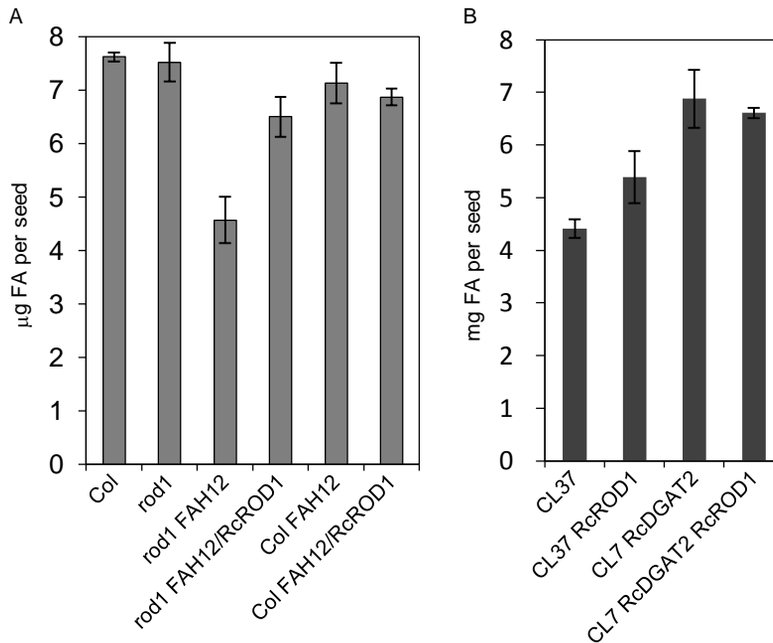


Figure 7. Oil content in Col, *rod1* and FAH12 transformed seeds. FAs (total µg per seed) were determined for each line listed in Table 1 and 2. The data (averages \pm SE) represent three replicates for each line.