Cloning of a Coconut Endosperm cDNA Encoding a 1-Acyl-sn-Glycerol-3-Phosphate Acyltransferase That Accepts Medium-Chain-Length Substrates

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Immature coconut (Cocos nucifera) endosperm contains a 1-acyl-sn-glycerol-3-phosphate acyltransferase (LPAAT) activity that shows a preference for medium-chain-length fatty acyl-coenzyme A sub- 
strates (H.M. Davies, D.J. Hawkins, J.S. Nelsen [1995] Phytochemistry 39: 989-996). Beginning with solubilized membrane preparations, we have used chromatographic separations to identify a polypeptide with an apparent molecular mass of 29 kD, whose presence in various column fractions correlates with the acyltransferase activity detected in those same fractions. Amino acid sequence data obtained from several peptides generated from this protein were used to isolate a full-length clone from a coconut endosperm cDNA library. Clone pCGN5503 contains a 1325-bp cDNA insert with an open reading frame encoding a 308-amino acid protein with a calculated molecular mass of 34.8 kD. Comparison of the deduced amino acid sequence of pCGN5503 to sequences in the data banks revealed significant homology to other putative LPAAT sequences. Expression of the coconut cDNA in Escherichia coli conferred upon those cells a novel LPAAT activity whose substrate activity profile matched that of the coconut enzyme.

The fatty acyl groups found in membrane phospholipids of higher plants are predominantly 16 or 18 carbons in length. However, the seed TAGs of many plants contain large amounts of fatty acyl groups different from those found in the phospholipids, suggesting that mechanisms exist for partitioning of specific fatty acids into the TAG fraction. Seed storage TAGs are synthesized in the ER from acyl-CoA and glycerol-3-P in a series of reactions termed the Kennedy pathway (reviewed by Stymne and Stobart, 1987; Frentzen, 1993). The first step of this pathway is the acylation of the sn-1 position of glycerol-3-P (catalyzed by glycerol-3-P acyltransferase) to form 1-acyl sn-glycerol-3-P, also termed LPA. The sn-2 position of LPA is subsequently acylated to yield PA in a reaction catalyzed by 1-acyl sn-glycerol-3-P acyltransferase (EC 2.3.1.51). This enzyme is more commonly known as LPAAT. Formation of TAG is completed by dephosphorylation of PA to produce diacylglycerol and the transfer of a third acyl group to the sn-3 position of the glycerol backbone by diacylglycerol acyltransferase. Synthesis of precursors of membrane phospholipids can also occur via a similar set of enzymatic steps with the exception of the final acylation (reviewed by Frentzen, 1993). Thus, sn-3 acylation is the only known enzymatic reaction unique to TAG production.

Although these basic pathways have been characterized, the underlying mechanisms regulating the specific partitioning of fatty acids into TAGs are not well understood. In addition, the distribution of individual fatty acids along the glycerol backbone of both TAGs and phospholipids is not random (Stymne and Stobart, 1987). The specificities of the acyltransferases, particularly LPAAT, are believed to play a key role in this nonrandom distribution (Ichihara et al., 1987; Oo and Huang, 1989; Frentzen, 1993). Several studies have demonstrated a correlation between the in vitro substrate specificities of LPAAT assayed in developing seed tissue and the fatty acid composition of the TAGs present in the mature seed (reviewed by Frentzen, 1993). These studies have also indicated that LPAAT is more selective with respect to substrates than the other two acyltransferases of the Kennedy pathway.

The selectivity of the LPAAT involved in TAG synthesis has clear implications for efforts to develop oils with very high levels of specific fatty acids. In particular, the preference of the Brassica LPAAT for 18:1-CoA and its lack of activity on 12:0- or erucyl-CoA (Oo and Huang, 1989; Cao et al., 1990) suggests a theoretical 67 mol% limit for accumulation of these fatty acids in TAGs. Expression of LPAAT from species that can utilize these substrates may permit the incorporation of these fatty acids into the sn-2 position of Brassica TAGs.

The obstacle in this task has been the lack of cloned genes for LPAAT with the desired specificities. Direct biochemical purification of an LPAAT involved in TAG biosynthesis has been difficult owing to its association with membranes. Although attempts at solubilization have been made (Hares and Frentzen, 1991; Taylor et al., 1992), they have not previously led to identification of proteins that could be associated with the enzyme activity. The only LPAAT sequences so far reported have been obtained by a genetic complementation approach. An Escherichia coli mutant defective in LPAAT activity has been isolated (Coleman, 1990), and the corresponding LPAAT gene (plsc) has been

Abbreviations: CHAPS, 3-[3-cholamidopropyl]-dimethylammonio]-1-propane-sulfonate; HA, hydroxylapatite; LPA, lysophosphatic acid; LPAAT, lysophosphatic acid acyltransferase; PA, phosphatic acid; TAG, triacylglycerol; 12:0, lauroyl; 18:1, oleoyl.

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Materials

Immature coconut (Cocos nucifera) seeds of desired developmental stage were obtained from local retail stores as detailed by Davies et al. (1995). Proteolytic enzymes and restriction endonucleases were from Boehringer Mannheim. Unless otherwise indicated, chemicals and chromatography media were purchased from Sigma. Oligonucleotide primers were synthesized on an Applied Biosystems model 394 DNA and RNA synthesizer.

Enzyme Assays

Radioactive substrates were prepared, and the solubilized LPAAT activity was assayed as described by Davies et al. (1995). Unless otherwise stated, coconut LPAAT activity was measured using [1-14C]12:0-CoA and 12:0-LPA.

Synthesis of 12:0-CoA Sepharose

12:0-CoA was covalently coupled to Sepharose 4B with 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide HCl and 6-amino hexanoic acid-Sepharose 4B according to the instructions provided with the product (Sigma). The amount of covalently coupled 12:0-CoA was 4 mg/mL of support (determined by GLC analysis after derivatization to produce fatty acid methyl esters).

Chromatography and LPAAT Protein Identification

Preparation of a coconut endosperm membrane fraction and solubilization of LPAAT using CHAPS were carried out as described by Davies et al. (1995). The 250,000g supernatant fraction, which contained LPAAT activity, was diluted 2-fold with a buffer to yield a solution with the following components: 50 mM Hepes, 20% (w/v) glycerol, 0.5 M NaCl, 1.125% CHAPS, 5 mM β-mercaptoethanol, 100 μM Pefabloc (Boehringer Mannheim), 1 μM leupeptin, and 1 μM pepstatin, at pH 7.5. The buffering solution used throughout chromatographic separations was 50 mM Hepes, 20% (w/v) glycerol, 1% (w/v) CHAPS, and 5 mM β-mercaptoethanol (buffer A). Where indicated, buffer A was supplemented by the inclusion of NaCl and/or potassium phosphate. In all cases, the final pH of the buffer was adjusted to 7.5 with NaOH.

The diluted supernatant fraction was applied to a 2.5-× 3-cm column of Red 120 agarose equilibrated with 0.5 M NaCl in buffer A. The column was washed with equilibration buffer and LPAAT activity was eluted with 2.5 M NaCl in buffer A. Fractions from the Red 120 agarose column that contained LPAAT activity were pooled and applied to a 1.5-× 6.6-cm column of Bio-Rad HA HT equilibrated with 1 M NaCl in buffer A. The HA column was washed with 2.5 bed volumes of equilibration buffer. LPAAT activity was eluted during application of a 10-bed-volume linear gradient of 0 to 100 mM potassium phosphate in buffer A that also contained 1 M NaCl. Fractions from the HA column that had LPAAT activity were pooled, diluted 2.5-fold with buffer A, and applied to a 1.5-× 1.1-cm column of 12:0-CoA Sepharose 4B equilibrated in 0.4 M NaCl in buffer A. After the column was washed with 10 bed volumes of equilibration buffer, LPAAT activity was eluted with buffer A containing 2.5 M NaCl. Fractions having LPAAT activity were pooled, diluted 2.5-fold in buffer A, and concentrated 65-fold in an Amicon (Beverly, MA) stirred cell fitted with a YM30 membrane. A portion of the concentrate was applied to a Superose 12 (Pharma
cia) size-exclusion column equilibrated with 1 M NaCl in buffer A and LPAAT activity was eluted with the same buffer. Proteins in the eluted fractions were concentrated by ultrafiltration (Centricon 30, Amicon). Samples were adjusted to 30 mM DTT and 2% SDS prior to electrophoresis on Tris-Gly 10 to 15% acrylamide gradient gels (Novex, San Diego, CA). Proteins were visualized using a silver-staining protocol (Blum et al., 1987).

Protein Sequencing

Large-scale preparations of protein for sequencing were made from 200 g of coconut endosperm. The proteins were solubilized and chromatographed over Red 120 agarose and HA as described above with no increase in the column bed sizes. Appropriate fractions from the HA column were pooled, and the proteins were prepared as described above and electrophoretically separated on 12% acrylamide gels (Novex, San Diego, CA). Immediately following electrophoresis, protein was transferred (Towbin et al., 1979) either to a nitrocellulose membrane for enzymatic digestion or to ProBlott (Applied Biosystems) for N-terminal sequence analysis. Protein blotted to nitrocellulose was digested with either trypsin or a combination of asparagin
yl-N protease and chymotrypsin (Aebersold, 1989). Peptides obtained from enzymatic digestions were sepa
rated via reverse-phase HPLC on a C18 column (Vydac, Hesperia, CA) with a gradient of 7 to 52.5% acetonitrile in the
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0.1 mM sodium phosphate, pH 2.2 (Rosner and Robbins, 1982). Sequencing was performed on an Applied Biosystems 373A protein sequencer with an on-line 120A phenylthiohydantoin analyzer.

Library Construction and Screening

Total RNA was isolated from immature coconut endosperm using a modification of a hexadecyltrimethylammonium bromide DNA isolation procedure as described by Jones et al. (1995). A coconut endosperm cDNA library was prepared using the UniZap system (Stratagene) with the following modifications to the synthesis of first-strand cDNA. Forty micrograms of total RNA from coconut endosperm were heated to 65°C for 20 min and chilled on ice prior to the addition of reaction components. The first-strand synthesis was carried out in a 50-μL reaction volume as recommended by Stratagene with the substitution of 0.1 mM sodium phosphate, pH 2.2 (Rosner and Robbins, 1982). Sequencing was performed on an Applied Biosystems 373A protein sequencer. Sequences were analyzed using the Intelligenetics Suite, version 5.3 (Intelligenetics, Inc., Mountain View, CA), and MacVector, version 4.5 (Kodak). Homology searches were performed at the National Center for Biotechnology Information using the BLAST Network Service. DNA sequence alignment was carried out using Megalign software (DNASTAR, Inc., Madison, WI). Protein structure prediction was performed using TopPred II (Claros and von Heijne, 1994).

Expression of Coconut LPAAT in Escherichia coli

The NdeI, Nhel, and BamHI sites of pET3a (Rosenberg et al., 1987) were replaced by SalI, BamHII, and PstI to create pCGN7645 (M. Lassner, personal communication). The LPAAT cDNA in pCGN5503 was modified by PCR to insert a SalI restriction site immediately upstream of the ATG start codon and a BamHI site immediately downstream of the TAA stop codon. The LPAAT-coding region was inserted as a SalI/BamHI fragment into pCGN7505 to create pCGN5505 for expression in E. coli under control of a T7 promoter.

To produce samples for enzyme assay, E. coli cultures were grown at 37°C to an A600 of 0.5 and induced with isopropyl β-D-thiogalactopyranoside to 0.4 mM for 2 h. Pelleted cells were suspended in 50 mM Hepes, 1 mM NaCl, 10 mM EDTA, 100 μM Pefabloc, 1 μM leupeptin, 1 μM pepstatin A, 5 mM β-mercaptoethanol, pH 7.5, and were broken by sonication. The samples were centrifuged at 5,000g for 15 min. The supernatant fractions were centrifuged for 2 h at 134,000g, and the pellet membranes were suspended in 50 mM Hepes, 200 mM NaCl, 20% (w/v) glycerol, 5 mM β-mercaptoethanol, pH 7.5. Membrane fractions were assayed for acyl-CoA substrate specificities with 12-O-LPA and various acyl-CoA species as described by Davies et al. (1995).

RESULTS

Identification of a Protein Associated with LPAAT Activity

Starting with the solubilized 12:0-CoA LPAAT activity from coconut endosperm (Davies et al., 1995), several chromatographic methods were screened for utility in enriching LPAAT activity. Inclusion of a high concentration of CHAPS (1%) in the column buffers was necessary to prevent aggregation of the enzyme. Also, at least 400 mM NaCl was required to maintain LPAAT activity. We were unsuccessful in attempts to use either ion-exchange or hydrophobic interaction methods, perhaps because of these detergent and salt requirements. From among several dye resins capable of adsorbing the LPAAT activity under conditions of high ionic strength, Red 120 agarose was chosen for the initial chromatographic step. Approximately 80% of the
CHAPS-solubilized protein was excluded from the Red 120 agarose at 0.5 M NaCl, whereas the LPAAT activity was completely adsorbed. Subsequent elution with 2.5 M NaCl resulted in recovery of 80 to 100% of the applied LPAAT activity (Fig. 1A). All of the LPAAT activity bound to the HA and was eluted with a potassium phosphate gradient (Fig. 1B). Three protein peaks were always obtained during the phosphate gradient elution, but the relative proportions of these peaks varied among preparations, depending on the maturity of the coconut tissue. The LPAAT activity was always associated with the first of the three peaks. The combination of chromatography over Red 120 agarose and HA separated the LPAAT activity from approximately 98% of the protein. Recovery of activity from the membrane fraction through the HA column was approximately 45%. The activity of the HA-purified fraction had the same specificity for medium-chain acyl-CoA substrates as did the original solubilized membrane fraction.

From SDS-PAGE analysis of the HA-purified fraction, it was not possible to unambiguously associate one of the remaining proteins with LPAAT activity. Chromatography over 12:0-CoA Sepharose (Fig. 1C) allowed recovery of LPAAT activity while removing several contaminating proteins. Active fractions from the 12:0-CoA column were pooled, concentrated, and applied to a Superose 12 size-exclusion column. The LPAAT activity profile and SDS-PAGE protein-banding patterns of the fractions from this column are shown in Figure 2. Band patterns were compared from fraction to fraction to identify bands whose intensities increased and decreased in concert with LPAAT activity. LPAAT activity was most closely correlated with a 29-kD protein.
protein band having an electrophoretic mobility of 29 kD. The correlation between LPAAT activity and the presence of the 29-kD protein in specific fractions was also observed upon chromatography on Blue A agarose, thiophilic affinity agarose, and rechromatography on HA (data not shown).

**Protein Sequencing**

Although purification over the 12:0-CoA and Superose 12 columns was necessary for initial identification of the 29-kD band, only the Red 120 and HA chromatographic steps, in combination with SDS-PAGE, were used for preparation of the protein for sequencing. These two chromatographic steps removed most of the contaminating proteins that migrated near the 29-kD candidate on a 12% polyacrylamide SDS gel. It was not possible to obtain an N-terminal sequence from the intact protein, which suggested that the N terminus was blocked. Amino acid sequence was obtained from a total of 11 peptides (Fig. 3).

**Generation of an LPAAT DNA Probe**

Degenerate oligonucleotides to be used as PCR primers were designed to regions of sequenced peptides obtained from the 29-kD protein. The relative orientation and spacing of the individual peptides within the protein were not known; therefore, many combinations of primer pairs were used under several reaction conditions. PCR amplification of coconut first-strand cDNA with primers 4865 and 4988 (Fig. 3) resulted in a 271-bp product. Subcloning and sequencing of this PCR product revealed a sequence encoding the known peptides at each end, as well as three additional peptide sequences obtained from the 29-kD protein.

**Isolation of pCGN5503**

The 271-bp PCR product was radiolabeled and used to probe a coconut endosperm cDNA library. Approximately 240,000 plaques were screened, and 29 hybridizing plaques were purified. The cDNAs in each were recovered as excised plasmids. The single-strand DNA sequence of...
the 3' end of each cDNA insert indicated that, although various sites of polyadenylation were used, all clones represented transcripts from a single gene.

The complete sequence of one of the longest clones, pCGN5503, is shown in Figure 3. Beginning with the ATG codon at position 259, the 1325-bp cDNA contains an open reading frame encoding 308 amino acids. This ATG is believed to be the translational start site because of the presence of stop codons in all three frames upstream and the similarity to plant consensus start sites (Lütcke et al., 1987). There is a consensus poly(A) addition signal (Proudfoot and Brownlee, 1976) 18 bp 5' to the site of polyadenylation in this cDNA. The long open reading frame contains the PCR product of primers 4865 and 4988 as well as the other peptide sequences indicated in Figure 3. The calculated molecular weight of the protein encoded by pCGN5503 is 34,806 and the estimated pI is 9.79.

Comparison of the deduced 308-amino acid sequence from pCGN5503 to protein sequence databases revealed significant homology to the E. coli LPAAT encoded by the plsc gene (Coleman, 1992) and the closely related sequence from Salmonella typhimurium (Luttinger et al., 1991) and also to a putative LPAAT from yeast, SLC7 (Nagiec et al., 1993). Figure 4 displays some of these regions of homology. In addition, smaller localized homologies were seen to other acyltransferases from E. coli, such as 2-acylglycerophosphoethanolamine acyltransferase (Jackowski et al., 1994) and glycerol-3-P acyltransferase (Lightner et al., 1983). The only putative higher plant LPAAT, pMAT1, recently cloned from maize (Brown et al., 1994), was not in the top 20 matches to the coconut sequence.

The hydrophobicity profile of the protein encoded by pCGN5503 is shown in Figure 5; two transmembrane stretches are predicted, consisting of amino acids 64 to 84 and 131 to 151.

Expression of pCGN5503 in E. coli

To verify the identification of the cDNA insert of pCGN5503 as the coconut medium-chain-specific LPAAT, pCGN5505 was constructed to express the open reading frame in E. coli using a T7 RNA polymerase-based system. Membrane fractions of cells expressing the coconut cDNA had higher activity with medium-chain substrates, especially 12:0-CoA, than did those of control E. coli, which preferred 18:1-CoA (Fig. 6A). When the E. coli background activity was subtracted from pCGN5505 cultures, the resulting activity profile with medium-chain acyl-CoA substrates was similar to that obtained from membrane fractions of immature coconut endosperm supplied with these substrates (Fig. 6B). In addition, the LPAAT activity of the E. coli-expressed enzyme was found to be specific for acyl-CoA versus acyl-ACP substrates (data not shown).

DISCUSSION

In this paper we present the direct biochemical purification and molecular cloning of a higher plant LPAAT involved in TAG biosynthesis. Chromatographic purification of hydrophobic enzymes involved in plant lipid synthesis has traditionally been unsuccessful. However, we were able to establish conditions that maintained the coconut LPAAT in a soluble and active state (Davies et al., 1995). Although we did not purify the LPAAT to homogeneity, we did achieve sufficient enrichment to permit correlation of enzyme activity with the staining intensity of a particular polypeptide band on SDS gels. Cloning of pCGN5503 depended on the amino acid sequence data obtained from proteolytic fragments generated from this candidate polypeptide. Verification that pCGN5503 encodes a coconut medium-chain length preferring LPAAT was demonstrated by detection of that activity in membrane fractions obtained from E. coli cultures expressing the enzyme.

The calculated mass of the protein encoded by pCGN5503 is 34.8 kD, whereas the corresponding partially purified protein migrates on SDS gels with an apparent mass of 29 kD. This discrepancy could be due to anomalous migration of the protein during SDS-PAGE; however, preliminary experiments using polyclonal antibodies raised
against a recombinant portion of the LPAAT protein suggest that the 29-kD protein has undergone a limited proteolysis during the purification process (data not shown). Further experiments using these LPAAT-specific antibodies to compare the electrophoretic migration of the protein in coconut endosperm extracts with the partially purified samples may resolve this question. The N-terminal sequence encoded by pCGN5503 does not show the characteristics of either a transit peptide of nuclear-encoded chloroplast proteins (de Boer and Weisbeek, 1991; von Heijne and Nishikawa, 1991) or those of a signal peptide (von Heijne, 1990), suggesting that the size discrepancy is not due to in vivo processing.

The protein encoded by pCGN5503 contains two hydrophobic regions predicted to be membrane-spanning domains using the TopPredII program, consistent with an integral membrane protein. Also, as has been observed for several proteins associated with lipid metabolism, the coconut LPAAT has a large net positive charge, which is thought to be important in interactions with both the acidic phospholipid membrane and the LPA substrate (Coleman, 1992). Several plant microsomal enzymes associated with lipid synthesis contain a motif of two Lys residues positioned three and five residues from the C terminus that is thought to be sufficient for retention of transmembrane ER proteins (Jackson et al., 1990). However, since not all ER proteins contain this motif, its significance is not clear (Yadav et al., 1993). The coconut LPAAT described in this paper represents another example of a microsomal protein that lacks this motif.

The preference of the LPAAT activity encoded by pCGN5503 for medium-chain-length acyl-CoA versus acyl-ACP substrates is consistent with that observed in developing coconut endosperm (Davies et al., 1995), a tissue that preferentially incorporates laurate (from 12:0-CoA) into the sn-2 position of TAG (Litchfield, 1970). We take this as evidence that we have cloned a cDNA encoding an LPAAT involved in TAG synthesis in coconut. This cDNA would apparently not account for the small amount of LPAAT activity with 18:1-CoA observed in coconut endosperm extract (Fig. 6B). Although LPAAT enzymes involved in cytoplasmic and plastidial membrane lipid synthesis, presumably having a preference for longer-chain fatty acids and/or acyl-ACP substrates, should also be present in developing coconut endosperm, we did not follow these activities in the chromatographic separations of medium-chain LPAAT activity. All of the cDNA clones obtained in this work appear to be derived from a single gene; under the conditions used in our screening, we did not detect homologous clones that could be associated with these other LPAAT activities.

The deduced amino acid sequence of the protein encoded by pCGN5503 contains significant homology to known or suspected LPAATs from bacteria and yeast. Over the region shown in Figure 4, the coconut sequence is 41% identical with both the E. coli and yeast sequences. In contrast, the homology of the protein encoded by the maize clone, pMAT1, to these same bacterial and yeast sequences is much more limited (Brown et al., 1994). In addition to the lack of sequence similarity between the proteins encoded by pCGN5503 and pMAT1, the hydrophobicity profiles of the proteins are also very different. Therefore, we believe that we have cloned a cDNA encoding a different enzyme than that encoded by pMAT1.

Transgenic rapeseed plants have been produced in which up to 50 mol% of the fatty acids present in the seed TAG are of medium-chain length (C12 and C14) (Kridl et al., 1993). As anticipated by analysis of rapeseed LPAAT specificities, it appears that the medium chains have been essentially excluded from the sn-2 position of the TAG in these lines (data not shown). With the availability of a cDNA encoding the medium-chain-specific LPAAT reported here, it should now be possible to test its effect on the sn-2 composition of transgenic rapeseed oil.

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