Identification of Amino Acids Conferring Chain Length Substrate Specificities on Fatty Alcohol-forming Reductases FAR5 and FAR8 from Arabidopsis thaliana*

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Background: Fatty alcohols produced by fatty acyl reductases (FARs) have important biological roles. Results: Expression of Arabidopsis FAR5 and FAR8 mutants in yeast revealed amino acids important for their stability or substrate specificity. Conclusion: Amino acids 355 and 377 are 16:0 versus 18:0 chain length specificity determinants of FAR5 and FAR8. Significance: Engineered FAR enzymes with desired substrate specificities will enable production of high value products.

Fatty alcohols play a variety of biological roles in all kingdoms of life. Fatty acyl reductase (FAR) enzymes catalyze the reduction of fatty acyl-coenzyme A (CoA) or fatty acyl-acyl carrier protein substrates to primary fatty alcohols. FAR enzymes have distinct substrate specificities with regard to chain length and degree of saturation. FAR5 (At3g44550) and FAR8 (At3g44560) from Arabidopsis thaliana are 85% identical at the amino acid level and are of equal length, but they possess distinct specificities for 18:0 or 16:0 acyl chain length, respectively. We used Saccharomyces cerevisiae as a heterologous expression system to assess FAR substrate specificity determinants. We identified individual amino acids that affect protein levels or 16:0-CoA versus 18:0-CoA specificity by expressing in yeast FAR5 and FAR8 domain-swap chimeras and site-specific mutants. We found that a threonine at position 347 and a serine at position 363 were important for high FAR5 and FAR8 protein accumulation in yeast and thus are likely important for protein folding and stability. Amino acids at positions 355 and 377 were important for dictating 16:0-CoA versus 18:0-CoA chain length specificity. Simultaneously converting alanine 355 and valine 377 of FAR5 to the corresponding FAR8 residues, leucine and methionine, respectively, almost fully converted FAR5 specificity from 18:0-CoA to 16:0-CoA. The reciprocal amino acid conversions, L355A and M377V, made in the active FAR8-S363P mutant background converted its specificity from 16:0-CoA to 18:0-CoA. This study is an important advancement in the engineering of highly active FAR proteins with desired specificities for the production of fatty alcohols with industrial value.

Primary fatty alcohols are aliphatic compounds that contain a hydroxyl group at the terminal position. In nature, long chain (C16 and C18) and very long chain (>C18) primary fatty alcohols are found either as free alcohols or in a combined state, such as wax esters, alkyld hydroxycinnamates, and ether lipids (1). Primary fatty alcohols and their derivatives have a wide range of biological roles in bacteria, fungi, insects, plants, and animals. Some important functions of free and esterified fatty alcohols in nonplant organisms are as follows: 1) as an energy storage reserve in microorganisms such as Euglena gracilis, Acinetobacter sp., and some marine microalgae; 2) as a constituent of sex pheromones in some insect and reptile species; 3) as a uroepigal gland secretion in birds to maintain feather condition; 4) in the heads of sperm whales for echolocation and possibly to regulate buoyancy; and 5) as ether lipids, which play a variety of roles and are highly abundant in animal nervous system tissue (2–10). In plants, fatty alcohols and their derivatives are found primarily as chemical constituents of three extracellular lipid-pheonomic barriers as follows: cuticle coating the aerial surfaces of plants, suberin present in the cell walls of various external and internal tissue layers, and sporopollenin found in the outermost layer of the pollen spore coat (1, 11). The functions of these polymeric surface lipid barriers are to protect plants from abiotic and biotic stresses. In a unique example, fatty alcohols are found in the seeds of the jojoba plant (Simmondsia chinensis) in the form of wax esters as a lipid energy reserve (12–14).

Fatty acyl reductases (FARs)² catalyze the four-electron reduction of fatty acyl-coenzyme A (CoA) or fatty acyl-acyl carrier protein to a primary fatty alcohol in a NADPH-dependent reaction. This is a two-step process involving the production of an unreleased aldehyde intermediate (15–17). FAR proteins share two distinct domains as follows: a Rossmann-fold NAD(P)H binding domain at the N terminus and a fatty acyl-

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2 The abbreviations used are: FAR, fatty acyl reductase; CoA, coenzyme A; GC, gas chromatography; 16:0-OH, 16:0 primary fatty alcohol; 18:0-OH, 18:0 primary fatty alcohol.
CoA reductase (FAR_C) domain at the C terminus (Fig. 1A) (1). The presence of the Rossmann-fold structure is typical of intermediate short-chain dehydrogenase/reductase proteins (18). Within the Rossmann-fold domain is a conserved GXXGXX(G/A) motif, which is believed to mediate NAD(P)H binding. The Rossmann-fold domain contains the active site motif, YXXXK, where the tyrosine (Y) and lysine (K) residues are predicted to play direct roles in catalysis based on kinetic studies with other reductases (19, 20).

FAR enzymes have distinct substrate specificities with regard to acyl chain length and degree of acyl chain saturation (1). The specificity of the FAR is often critical to the physical properties of the final biosynthetic product. An example of this comes from the European corn borer moth, *Ostrinia nubilalis*, which is represented by two races differentiated by the ratios of cis- and trans-isomers in their sex pheromone chemical mixtures. It is the preference of FAR enzymes, expressed in the pheromone gland of the two moth races, for fatty acyl substrate containing either a cis or a trans double bond that has led to reproductive isolation of the races (21). In recent years, there has been increased interest in engineering FAR proteins that produce fatty alcohols with desired chemical qualities for industrial use; however, little is known about the amino acid residues responsible for conferring substrate specificity. A three-dimensional crystal structure of a FAR protein has not yet been elucidated, and no other investigations have yet been carried out to determine the specific residues influencing substrate preference.

The *Arabidopsis thaliana* genome encodes for eight FAR proteins (FAR1 to FAR8), each with a distinct substrate specificity for saturated fatty acyl precursors with chain lengths ranging from C16 to C30 (17, 22–25). FAR5 (At3g44550) and FAR8 (At3g44560) are located in tandem on chromosome 3 and encode for proteins that are 85% similar at the amino acid level. Identical amino acids are highlighted in black, and physiochemically similar amino acids are highlighted in gray. Arrows indicate domain swap sites, and asterisks indicate site-specific mutagenesis sites.

**FIGURE 1.** FAR structural domains and protein sequence alignment of *Arabidopsis* FARs and FAR8. A, schematic of the structural domains of FAR proteins. FAR enzymes, minus possible subcellular localization signals (e.g. plastid targeting), are ~500 amino acids in length and contain an NAD(P)H binding Rossmann-fold domain (light gray) and a FAR_C domain (dark gray) at the N and C termini, respectively. The GXXGXX(G/A) predicted NADPH-binding motif as well as the predicted active site motif YXXXK are indicated (where X represents any amino acid). B, protein sequence alignments of FAR5 and FAR8, which are 85% similar at the amino acid level. Identical amino acids are highlighted in black, and physiochemically similar amino acids are highlighted in gray. Arrows indicate domain swap sites, and asterisks indicate site-specific mutagenesis sites.
### Table 1

**List of oligonucleotide primers used in this study**

For means forward, and Rev means reverse.

| Primer name | Sequence (5’ to 3’) |
|-------------|---------------------|
| **FARS/FAR8 amplification** | |
| FARS_BamHI_For | GAGGATCCATGGAACCTTCATGTTGTTCTAAT |
| FARS_XhoI_Rev | GCGCTCGAGTTTCTCCTAAGACGTCAG |
| FAR8_BamHI_For | GAGGATCCATGGAACCTTCATGTTGTTCTCA |
| FAR8_XhoI_Rev | GCGCTCGAGTTTCTCCTAAGACGTCAG |

| **Domain swap constructs** | |
| FAR8-FAR8_388_For | CTGCGTTGTGTTCACGCTCTAACGAGGCTTTGTTACA |
| FAR8-FAR8_388_Rev | TATTTTGGTTCGTTACTTTACGAAA |
| FAR8-FAR8_353_For | GAGGAGAGTCCGCAAAATACGGGTTCTCC |
| FAR8-FAR8_353_Rev | TGAACGAGACCGCAATTTCAAGCCTGTCCTCC |
| FAR8-FAR8_344_For | TCAATCTCACCAAAACCCAGTCACATTTGGTAGATGA |
| FAR8_FAR5_353_Rev | GCCATTACGAAAAACTCTTTGGGTTCTG |
| FAR8_FAR5_344_Rev | TATTTTGGGTTCTCCTTACGAAA |
| FAR8_FAR5_388_Rev | TATTTTGGGTTCTCCTTACGAAA |
| FAR8_FAR5_388_For | TATTTTGGGTTCTCCTTACGAAA |

| **Point mutation constructs** | |
| FAR8_S363P_For | TTACGAAAAACCCTGCGCGCTG |
| FAR8_S363P_Rev | ACTGCAGCCACGCGTTTGGGTTG |
| FAR8_P363S_For | TTACGAAAAACCCTGCGCGCTG |
| FAR8_P363S_Rev | ACTGCAGCCACGCGTTTGGGTTG |
| FAR8_K242I_For | GAGACCGGATCCGTTCTG |
| FAR8_K242I_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_Y238F_For | GAGACCGGATCCGTTCTG |
| FAR8_Y238F_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_V377M_For | TGTATTGCTTTGGTACGTTG |
| FAR8_V377M_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_A355L_For | GAGACCGGATCCGTTCTG |
| FAR8_A355L_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_L355A_For | GAGACCGGATCCGTTCTG |
| FAR8_L355A_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_I347T_For | GAGACCGGATCCGTTCTG |
| FAR8_I347T_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_T347I_For | GAGACCGGATCCGTTCTG |
| FAR8_T347I_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_M377V_For | GAGACCGGATCCGTTCTG |
| FAR8_M377V_Rev | TTACGAAAAACCCTGCGCGCTG |
| FAR8_BamHI_For | GAGGATCCATGGAACCTTCATGTTGTTCTCA |
| FAR8_BamHI_Rev | GCGCTCGAGTTTCTCCTAAGACGTCAG |
| FAR8_XhoI_For | GAGGATCCATGGAACCTTCATGTTGTTCTCA |
| FAR8_XhoI_Reverse | GCGCTCGAGTTTCTCCTAAGACGTCAG |

#### EXPERIMENTAL PROCEDURES

**Materials**—All chemical reagents were from Sigma unless otherwise stated. [1-14C]Palmitoyl-CoA and [1-14C]stearoyl-CoA were from PerkinElmer Life Sciences and Amersham Biosciences, respectively.

**Construction of Yeast Expression Plasmids Containing Arabidopsis FARS and FAR8 Variants**—The coding regions of FARS and FAR8 were cloned between the BamHI and XhoI restriction enzyme sites of a modified version of pYES2 (Invitrogen) containing enzyme sites of pYES2-His6/T7. The insert in each construct was verified by DNA sequencing using flanking sequencing primers.

**Expression of FARS and FAR8 Variants in Yeast**—The pYES2-His6/T7 plasmids containing wild-type and mutant FARS and FAR8 coding regions were transformed into S. cerevisiae yeast strain W303-1A (MATa his3Δ1 leu2 trp1-289 ura3-52) using the method described previously (28). Yeast transformants were selected on synthetic dropout (SD) media plates (2% d-glucose, 0.67% yeast nitrogen base, 0.01% adenine, 0.002% histidine, 0.002% tryptophan, 0.01% leucine, and 2% agar) lacking uracil (SD−Ura). Four yeast colonies for each construct, including empty vector, were separately inoculated into 3 ml of SD−Ura liquid media and grown for 24 h at 30 °C and 250 rpm. The Abs600 for each culture was measured, and a volume of culture required to inoculate 3 ml of induction SG−Ura liquid media and grown for 24 h at 30 °C and 250 rpm. The Abs600 of 0.4 was removed and centrifuged. The resulting yeast pellets were washed twice with 1.5 l of sterile water and then resuspended in 3 ml of SG−Ura media. The yeast strains were then grown for 2 days at 30 °C and 250 rpm prior to lipid extraction and analysis of protein levels by Western blotting.
Analysis of Protein Levels in Yeast by Protein Immunoblotting—A volume of each galactose-induced yeast culture equal to an A600 of 2.5 was harvested by centrifugation. The supernatants were removed and the pellets stored at −80 °C. Protein samples were prepared according to Ref. 29 using phosphate-containing loading buffer (62.5 mM sodium phosphate buffer, pH 7.0, 2% SDS, 10% glycerol, 5% β-mercaptoethanol, 0.0001% bromphenol blue) prior to separation on a 12% SDS-polyacrylamide gel at 150 V in 1 × running buffer (25 mM Tris, pH 8.3, 186 mM glycine, 0.1% SDS). The proteins were transferred to nitrocellulose membranes by wet-transfer electrophoresis. The nitrocellulose membranes were blocked overnight at 4 °C in blocking solution consisting of 5% fat-free skim milk in TBST (137 mM NaCl, 25 mM Tris base, pH 7.6, 0.1% Tween 20). The nitrocellulose membranes were then incubated with a 1:50,000 dilution of horseradish peroxidase-conjugated anti-mouse secondary antibody (EMD4-Biosciences) was added to detect the T7 epitope-tagged proteins; the membranes were washed four times for 5 min each with TBST. The membranes were then incubated with a 1:50,000 dilution of T7 Tag monoclonal mouse antibody (catalogue number 69522-3, EMD4Biosciences) was added to detect the T7 epitope-tagged proteins; the membranes were washed four times for 5 min each with TBST. The membranes were then incubated with a 1:1 mixture of Lumigen TMA-6 Solution A (contains Tris buffer in 3.2% v/v ethanol) and Lumigen TMA-6 Solution B (proprietary substrate in Tris buffer) (GE Healthcare) for 5 min in the dark and imaged with a FluorChemQ (Alpha Innotech). Western blots with yeast microsomes were done with 15 µg of protein and revealed with the ECL Western blotting detection kit (Amersham Biosciences).

Analysis of Fatty Alcohol Content of Transgenic Yeast by Gas Chromatography (GC)—Yeast cells from 2 ml of each galactose-induced culture were harvested by centrifugation, and the supernatant was poured into a separate glass tube. 10 µg of 1-pentadecanol (15:0-OH) was added as an internal standard to both the pellet and supernatant. The yeast supernatant was then extracted twice with 1 ml of 2:1 chloroform/methanol and once with 1 ml of chloroform, and the organic phases were combined and washed with 2.5 ml of 0.9% NaCl (w/v) before being evaporated under nitrogen gas at 37 °C. Both the yeast pellet and supernatant samples were resuspended in 3 ml of 1 M methanolic-HCl, and lipids were transmethylated at 80 °C for 90 min. 0.9% NaCl was added to each sample, and the lipids were extracted twice into 500 µl of hexane. The pooled hexane extracts were dried under nitrogen gas at 37 °C. The lipids were then resuspended in 100 µl of N,O-bis(trimethylsilyl) trifluoroacetamide plus 100 µl of pyridine and incubated at 110 °C for 15 min for silylation of free hydroxyl groups. The samples were then re-dried down as before and resuspended in a 1:1 (v/v) mixture of hexane/toluene. Lipids were quantified with a Varian GC450 equipped with a split-splitless injector, HP-1 column (30 m length, 0.25 mm inner diameter, 0.10 µm film thickness), and a flame ionization detector. One µl of each sample was injected using a 10:1 split ratio. The carrier gas was helium with a constant flow rate of 2 ml/min. The column oven was held initially at 150 °C for 5 min, then ramped at 10 °C/min to 300 °C, and held for 8 min, for a total run time of 28 min.

RESULTS

Amino Acids of FAR5 and FAR8 Important for Protein Stability and Enzymatic Activity—We examined the activities and substrate specificities of the FAR5 and FAR8 enzymes using S. cerevisiae as a heterologous system. The open reading frames for both genes were cloned into the yeast expression vector pYES2-His6/T7 downstream of the GAL1 promoter to allow high level protein induction by galactose. The expressed FAR proteins contained a T7 epitope tag at the N terminus to enable detection using Western blots. The empty pYES2-His6/T7 vector was used as a negative control. The endogenous acyl-CoA pool of S. cerevisiae has high and nearly equal amounts of 16:0-CoA and 18:0-CoA (31), thus providing relevant substrates for heterologously expressed FAR5 and FAR8 variants.

Analysis of the internal lipid content of 2-day-old galactose-induced yeast cultures by GC demonstrated that FAR5 produced high levels of 18:0 primary fatty alcohol (2.07 µg per unit of absorbance) and that the protein was highly expressed in yeast as determined by Western blot analysis (Figs. 2 and 3). In contrast, FAR8 did not produce any fatty alcohols under the same conditions, and the protein was barely detected in the Western blot (Figs. 2 and 3). However, when transgenic yeasts expressing FAR8 were grown for 4 days prior to lipid analysis, low levels of 16:0 fatty alcohol were detected (data not shown). Similarly, very small amounts of 16:0-OH were sometimes detected in yeast expressing FAR5 for 2 days (e.g. Fig. 5A), and these amounts increased after 4 days of FAR5 expression albeit to levels less than 4% of the accumulated 18:0-OH (data not shown).

With the exception of Arabidopsis FAR8, all plant FAR proteins reported have a proline at relative position 363 of FAR5 and FAR8. FAR8 instead has a serine at this position (Fig. 1A). Unlike native FAR8 protein, the FAR8-S363P variant revealed that FAR8-S363P accumulated to higher levels...
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FIGURE 2. Gas chromatograms of internal lipids of yeast expressing Arabidopsis FAR5, FAR8, or FAR8-S363P. The empty vector pYES2-His6/T7 tag acted as a negative control. Transformants were cultured in galactose media to induce protein expression. Fatty acids were transmethylated to their corresponding methyl esters, and fatty alcohol hydroxyl groups were derivatized to trimethylsilyl ethers before separation by GC and detection by flame ionization. The peaks corresponding to pentadecanol (15:0-OH) internal standard (IS), 16:0-OH and 18:0-OH are indicated, as well as saturated and mono-unsaturated C16 and C18 fatty acids (FA).

than its native counterpart (Fig. 3B). For much of our subsequent mutational analyses, this active FAR8-S363P mutant was used as the model of 16:0 fatty alcohol production. We subsequently refer to this active FAR8 variant as FAR8R, where R stands for “resurrected.” The reciprocal amino acid conversion in FAR5, proline to serine conversion at position 363, resulted in a 30% decrease in 18:0-OH production (Fig. 3A), although the FAR5-P363S and native FAR5 protein levels in yeast were comparable (Fig. 3B). It was possible that FAR8 or FAR8R had activity for a shorter acyl chain length than 16:0, with the most likely possibility being 14:0. Therefore, we also tested whether FAR5, FAR8, and/or FAR8R could generate 14:0-OH by feeding yeast 14:0 fatty acid using a previously described protocol (32). No 14:0-OH was produced by FAR5 or FAR8 and only a very small amount by FAR8R in our feeding experiments (data not shown). We cannot rule out other potential substrates for FAR8, but the most likely scenario is that FAR8 is indeed a low activity FAR due to the serine rather than proline at position 363 and that the specificity of FAR8R is nearly strict for 16:0-CoA.

The 18:0 and 16:0 fatty alcohols produced by the FAR5 and FAR8R variants, respectively, were found not only within the yeast cells but also secreted into the media (Fig. 4). Of the total fatty alcohols produced by FAR5 and FAR8R in yeast, 47 and 48%, respectively, were found secreted into the media (Fig. 4). For ease of screening purposes (below), we only quantified the fatty alcohols found in the yeast pellet. We verified that no bias was introduced by excluding the secreted fatty alcohols in our screens by measuring the internal and secreted fatty alcohols of FAR5 and FAR8R, as well as some key FAR5 and FAR8 variants that alter substrate specificity as described below (Fig. 4). The chain length distributions of fatty alcohols produced by these FARs expressed in yeast were nearly identical whether examining just internal fatty alcohols or examining total fatty alcohols (internal plus secreted).

We next investigated the proposed roles of tyrosine 238 and lysine 242 in the predicted YXXXK active site motif of the Rossmann-fold in FAR proteins (Fig. 1) (1, 22). Two separate site-specific substitutions were made in the FAR5 coding sequence to convert tyrosine 238 to a phenylalanine (FAR5-Y238F) and separately lysine 242 to an isoleucine (FAR5-K242I) in the encoded protein. These same amino acid substitutions were used to investigate the active site of alcohol dehydrogenase from Drosophila melanogaster (19), revealing the importance of the hydroxy group of tyrosine and the amino group of lysine in catalysis. Phenylalanine differs from tyrosine only by the presence of a hydroxyl group on the aromatic ring. Lysine and isoleucine are similarly sized amino acids, but isoleucine lacks the positive charge conferred by the amino group on lysine. Lipid analysis of both the FAR5-Y238F and FAR5-K242I variants expressed in yeast revealed no fatty alcohol production (Fig. 3A), although Western blot analysis of the mutant proteins
indicated that they were expressed to the same levels as the wild-type FAR5 (Fig. 3B).

Altogether, these data showed that FAR5 and FAR8 nearly exclusively produce 18:0-OH and 16:0-OH, respectively, that proline 363 is important for protein stability, and that tyrosine 238 and lysine 242 are necessary for catalytic activity.

**Domain Swaps between FAR5 and FAR8**—A series of domain swap chimeras involving FAR5 and FAR8 were then made to investigate the protein region responsible for conferring substrate specificity. An initial reciprocal pair of domain swaps were made between FAR5 and FAR8 at positions corresponding to amino acid residues 283/284 (Figs. 1B and 5A). The FAR5(1–283)FAR8(284–496) [1] chimera produced predominantly 16:0-OH. Conversely, the FAR8(1–283)FAR5(284–496) chimera [2] produced substantial amounts of 18:0-OH and very little 16:0-OH (Fig. 5A). This indicated that C16/C18 chain length specificity is dictated by amino acids between position 284 and the C terminus of the FAR5/FAR8 proteins. To further investigate this, domain swaps between FAR5 and FAR8 were made at amino acid residues 388/389 (Figs. 1B and 5A). We found that the FAR5(1–388)FAR8(389–496) chimera [3] produced nearly exclusively 18:0-OH, whereas the FAR8(1–388)FAR5(389–496) chimera [4] had low activity, but it produced more 16:0-OH than 18:0-OH (Fig. 5A). Collectively, the data from these four domain swaps indicate that C16/C18 chain length specificity is dictated by amino acids between positions 284 and 388 of the FAR5 and FAR8 proteins.

To examine this further, we made two FAR5/FAR8 internal domain swap mutants spanning residues 284–388, inclusively (Fig. 5A). The FAR5(1–283)FAR8R(284–388)FAR5(389–496) [5] and FAR8(1–283)FAR5(284–388)FAR8R(389–496) [6] internal domain swap chimeras produced nearly exclusively 16:0-OH and 18:0-OH, respectively (Fig. 5A). This verified that the amino acid residues responsible for 16:0 fatty acyl-CoA versus 18:0 fatty acyl-CoA substrate specificity are between residues 284 and 388. To further narrow down the protein region responsible for conferring chain length specificity, two more reciprocal internal domain swap constructs were made spanning residues 354–388 (Fig. 5A). The FAR5(1–353)FAR8R(354–388)FAR5(389–496) chimera [7] produced nearly exclusively 16:0-OH. The FAR8(1–353)FAR5(354–388)FAR8R(389–496) chimera [8] appeared to have very low activity for either fatty acyl-CoA substrate, but it produced more 18:0-OH than 16:0-OH. This suggests that the amino acid residues responsible for dictating 16:0 versus 18:0 acyl chain length specificity in FAR5 and FAR8 are found within the 35 amino acids interval spanning residues 354–388.

The chimeras that contained FAR5 sequence at the N-terminal end [1 + 3] accumulated to higher levels in yeast than the chimeras that contained FAR8 at the N-terminal end [2 + 4] (Fig. 5B). The internal domain swap chimeras were generally expressed to similar levels as the FAR8R protein, with the exception of FAR8(1–353)FAR5(354–388)FAR8R(389–496) [8], which gave very little fatty alcohol production and had near undetectable protein accumulation, and FAR5(1–353)FAR8R(354–388)FAR5(389–496) [7], which generated high amounts of fatty alcohols and was expressed at high levels similar to that of FAR5 (Fig. 5B). Additionally, this latter chimera [7] accumulated to higher levels than the FAR5(1–283)-

*FIGURE 4. Amounts of total fatty alcohols produced by FAR5 and FAR8 variants expressed in yeast. A, internal fatty alcohol content of yeast cells (nonsecreted fatty alcohols). B, fatty alcohols found in supernatant of yeast cultures (secreted fatty alcohols). C, total fatty alcohols produced by yeast expressing a FAR variant (combined nonsecreted and secreted fatty alcohol content). Values are expressed in μg/A600 unit ± S.D. (n = 4). Yeast cultures were grown for 48 h in galactose media for protein induction before lipid extraction and analysis.*
FAR8(284–388)FAR5(389–496) chimera [5], which had FAR8 sequence between residues 284 and 353. We therefore speculated that some residues immediately N-terminal to position 354 are critical for protein stability. To test this, another FAR8-FAR5-FAR8 domain swap chimera was expressed that contained residues 345–388 of FAR5. This FAR8(1–344)-FAR5(345–388)FAR8(389–496) chimera [9] accumulated to relatively high levels (Fig. 5B), indicating that one or more amino acids between positions 345 and 353 are important for protein stability. Of the four amino acids that are different than FAR5 and FAR8 in this region, only position 347 is a nonconservative amino acid difference (Fig. 1B). Residue 347 is a threonine in FAR5, and it is an isoleucine in FAR8. We found that converting the isoleucine present at position 347 in FAR8 to the reciprocal threonine of FAR5 causes the resultant FAR8-R347T protein to accumulate to higher levels in yeast than the parent FAR8R (Fig. 3B). The amount of 16:0-OH produced by FAR8R-I347T was 1.9 times that of FAR8R (Fig. 3A). This was paralleled by decreased FAR5-T347I protein levels compared with wild-type FAR5 (Fig. 3B), again indicating that threonine 347 plays an important role in protein stability.

Reciprocal Amino Acid Substitutions in FAR5 and FAR8—To determine individual amino acid residues of FAR5 and FAR8 that are responsible for dictating chain length substrate specificity, site-specific substitutions were made within the 44-amino acid region (residues 345–388) of FAR5 and FAR8 identified by internal domain swaps to “flip” substrate specificity and be important for protein stability (Fig. 6A). Within this region, there are 13 amino acids that differ between FAR5 and FAR8, and we targeted eight of these residues for site-specific mutagenesis. Of these eight amino acids, positions 347 and 363 had already been examined (see above) and found to be important for protein stability/activity but not influencing chain length specificity. In all cases, we converted a given amino acid found at a certain position in one FAR to the reciprocal amino acid found at that same position in the other FAR.

Within the 345–388 region, there are two triplet blocks of amino acids, residues 355–357 (triplet I) and 377–379 (triplet II), that are not conserved between FAR5 and FAR8 (Fig. 6A). We individually mutated each of the six amino acids in triplets I and II of FAR5 to the corresponding FAR8 amino acid, and we
found that amino acid conversions at positions 355 and 377 of triplet I and II, respectively, resulted in the most 16:0-OH production (Fig. 6B), with the other four conversions causing little or no 16:0-OH production (data not shown). The FAR5-A355L single mutant produced 16:0-OH to levels similar to that of FAR8 and produced 18:0-OH to about 66% of that observed with wild-type FAR5 (Fig. 6B). The FAR5-V377M single mutant also produced both 16:0-OH and 18:0-OH, but the amount of 16:0-OH produced was about 25% of FAR8, whereas the amount of 18:0-OH produced was similar to wild-type FAR5 (Fig. 6B). By combining the mutations, we were able to generate a FAR5 variant, FAR5-A355L/V377M, which produced mostly 16:0-OH (Fig. 6B). The FAR5-A355L/V377M mutant had 37% greater 16:0-OH yield than FAR8, although it produced a small amount of 18:0-OH, ~6% of the amount produced by wild-type FAR5. These single and double FAR5 variants were all expressed to similar levels as wild-type FAR5 in yeast (Fig. 6B).

We then examined the effects of reciprocal amino acid substitutions at positions 355 and 377 of FAR8 (Fig. 6C). The FAR8-R355A and FAR8-R-M377V mutants had greatly reduced levels of 16:0 fatty alcohol production, ~8 and 18% of FAR8 levels, respectively. These mutants also produced 18:0-OH, albeit at very low levels compared with FAR5. Because the I347T mutation had a major effect on FAR8 protein stability/activity (see above), we combined this change with the M377V change. The resultant FAR8-R-I347T/M377V double mutant produced about twice the amount of 16:0-OH than the parent FAR8, although slightly less than FAR8-R-I347T (Figs. 3A and 6C). FAR8-R-I347T/M377V additionally produced high levels of 18:0-OH (0.63 g per unit of absorbance). As predicted, FAR8-R-I347T/M377V protein accumulated at much higher levels in yeast than FAR8-R-M377V, likely accounting for the increased fatty alcohol production. In an attempt to further convert FAR8 substrate preference from 16:0-CoA to 18:0-CoA, we added a third L355A mutation to create FAR8-R-I347T/L355A/M377V. This triple mutation nearly completely converted FAR8 specificity to 18:0-CoA (Fig. 6C). The FAR8-R-I347T/L355A/M377V mutant produced 18:0-OH to levels about 60% of wild-type FAR5. The FAR8-R-I347T/L355A/M377V mutant was expressed to similar levels as wild-type FAR5 (Fig. 6B).
M377V mutant produced a small amount of 16:0-OH, but it was only about 30% of that found with FAR8. In summary, we altered FAR8 chain length specificity from 16:0-CoA nearly completely to 18:0-CoA by changing amino acids at two positions, 355 and 377, to that of the corresponding FAR5 amino acids, but amino acids changes were also required at positions 347 and 363 to allow for high protein accumulation.

**In Vitro FAR Assays Using Yeast Microsomes**—To provide additional support of our findings, we prepared microsomes from different transgenic yeast strains and performed **in vitro** assays using either $[1-^{14}C]$palmitoyl-CoA (16:0-CoA) or $[1-^{14}C]$stearoyl-CoA (18:0-CoA) as substrate. After 5 min of incubation in the presence of 5 μg of microsomal proteins, lipids were extracted and analyzed by thin layer chromatography. Microsomes containing FAR5 produced labeled fatty alcohols in the presence of $[1-^{14}C]18$-0-CoA, whereas no radioactivity associated with fatty alcohols was observed using microsomes from yeast transformed with empty vector, regardless of the labeled fatty acyl-CoA used (Fig. 7A). In the presence of $[1-^{14}C]16$-0-CoA, FAR5 produced very little fatty alcohol. Conversely, FAR5-A355L/V377M produced significant amounts of fatty alcohols with $[1-^{14}C]16$-0-CoA as substrate, in agreement with the results obtained **in vivo**. In contrast to the microsomes containing FAR5 variants, no fatty alcohols were detected when using microsomes from yeast transformed with FAR8R, FAR8R-I347T/M377V, or FAR8R-I347T/L355A/V377M, whatever the labeled fatty acyl-CoA used (Fig. 7A). The reason that these microsomes were inactive **in vitro** was most probably due to protein stability, as none of the FAR8 variants could be detected in the microsomes by Coomassie Blue staining or by Western blots in contrast to the FAR5 variants (Fig. 7B). The FAR8 variants may also be deficient in their ability to associate with membranes. These **in vitro** assays nevertheless allowed for quantification of the specific activities of FAR5, FAR5-A355L, and FAR5-A355L/V377M for 16:0-CoA and 18:0-CoA (Table 2). FAR5 was 20 times more active **in vitro** with 18:0-CoA than with 16:0-CoA. The FAR5-A355L single mutant was able to convert both 16:0-CoA and 18:0-CoA to the corresponding fatty alcohols, with its specific activity for 16:0-CoA being about half of that for 18:0-CoA. The FAR5-A355L/V377M double mutant had a clear preference for 16:0-CoA, with its specific activity for 16:0-CoA being 4.7 times that for 18:0-CoA (Table 2). Altogether, the **in vitro** assays using FAR5, FAR5-A355L, and FAR5-A355L/V377M were in good agreement with the results obtained **in vivo**.

**DISCUSSION**

FAR5 and FAR8 are located in tandem on the *Arabidopsis* genome, and the encoded proteins are 85% identical in sequence. It is likely that FAR5 and FAR8 are the result of a recent genome duplication event. We therefore speculate that the low native activity of FAR8 is a result of recently acquired mutations through its evolutionary history and that it may be on the way to being a pseudogene. The serine at position 363 of FAR8 rather than the conserved proline at this position may be the result of such a mutation. Although FAR5 and FAR8 are highly similar, they have distinct and nonoverlapping substrate specificities. Expression of *Arabidopsis* FAR5 in yeast produced relatively large quantities of 18:0-OH and very small amounts of 16:0-OH, although expression of the native *Arabidopsis* FAR8 produced exclusively 16:0-OH in very small quantities. To date, 16:0 fatty alcohols have not yet been detected, either in free or combined form, in *Arabidopsis* (25), although they are expected to be part of pollen exine based on the high activity of MS2/FAR2 for 16:0-acyl carrier protein (24). MS2/FAR2 is essential for pollen exine development (24, 33). FAR8 may be contribut-
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ing to 16:0-OH in pollen exine as low levels of FAR8 transcript were detected in developing and mature pollen using DNA microarray analysis (34). Regardless of the endogenous role of FAR8, it provided an ideal partnership with FAR5 to investigate amino acids important for dictating their distinct chain length substrate specificities.

As with other NAD(P)H-dependent oxidoreductases, the FAR enzymes possess a predicted YXXXK active site motif (35). The tyrosine and lysine residues present at this position, along with a serine residue that comes from another part of the protein, have been shown to play an important role in catalysis (19, 20). Deletion of the YXXXK motif in MS2/FAR2 results in a protein that cannot complement the pollen exine defects of ms2/far2 mutants (24). However, the individual roles of the tyrosine and lysine predicted to be directly involved in catalysis had not been previously investigated. We made two site-specific FAR5 mutants in which we converted tyrosine 238 to phenylalanine and lysine 242 to isoleucine (FAR5-Y238F and FAR5-K242I). The same substitutions were used to investigate the catalytic residues of FAR8 that would need to be altered to increase protein levels. However, as in FAR5, substitution of the serine at position 363 to isoleucine dramatically decreased its protein levels. However, without the benefit of an x-ray or NMR structure of the FAR substrate-binding site, it is difficult to determine precisely how amino acids 355 and 377 in FAR5 and FAR8 control binding of 16:0-CoA versus 18:0-CoA substrates. We attempted to use comparative homology modeling of FAR5 and FAR8 to gain further insights into the roles of these residues, but this approach was not feasible due to the lack of a similar enough modeling template. Because fatty alcohols are used in a variety of human applications, such as in cleaning detergents, cosmetics, and pharmaceutical formulations, food products, textiles and coatings, or high performance industrial lubricants (1), our study is a step forward in the engineering of FAR enzymes to produce fatty alcohols with desired specificities and to allow the production of renewable fatty alcohol-containing lipid products of high commercial value.

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