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Introduction

n-3 Polyunsaturated fatty acids (PUFAs) such as eicosapentaenoic acid (EPA, C20:5*n*-3) and docosahexaenoic acid (DHA, C22:6*n*-3) are important for brain development, tissue formation and repair in mammals.^{1,2} They also have significant effects in the prevention of asthma, cancer, depression, obesity and immune cardiovascular disorder.^{3–5} However, the limited natural sources of *n*-3 PUFAs come mainly from microalgae oil and fish oil.⁶ As an alternative source, microorganisms have been used to produce EPA and DHA.⁷

Characterization and molecular docking of new $\Delta 17$ fatty acid desaturase genes from *Rhizophagus irregularis* and *Octopus bimaculoides*[†]

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Fatty acid desaturases are key enzymes in the biosynthesis of n-3 polyunsaturated fatty acids (PUFAs) via conversion of n-6 polyunsaturates to their n-3 counterparts. In this study, we reported the characterization and molecular docking of Δ 17 desaturases from *Rhizophagus irregularis* and *Octopus* bimaculoides. These two new desaturase genes were screened using the known $\Delta 17$ desaturase gene (oPaFADS17) from Pythium aphanidermatum as a template. Analysis of their genes revealed that the sequences of oRiFADS17 and oObFADS17 contained the typical His-rich motifs (one HXXXH and two HXXHH). They were then expressed in Saccharomyces cerevisiae INVSc1 to examine their activities and substrate preferences. Our results show that the two candidate n-3 desaturases possess a strong $\Delta 17$ desaturase activity, exhibiting remarkable increase in desaturation activity on C20 fatty acids compared to C18 fatty acids. To the best of our knowledge, oRiFADS17 desaturase has greater (3-4 fold) catalytic activity for C18 substrates than other reported Δ 17 desaturases and oObFADS17 is the first reported Δ 17 desaturase in sea mollusks. Characterization of these two new desaturases will be of greater value for genetic engineering in industrial production of eicosapentaenoic acid (EPA, C20:5n-3) and docosahexaenoic acid (DHA, C22:6n-3). Due to lack of crystal structure information about n-3 desaturases, for the first time, the view of their predicted structures, binding pockets and substrate tunnels was clearly observed based on molecular docking. This will contribute to strengthening our understanding of the structure-function relationships of n-3 fatty acid desaturases.

> The *n*-6/*n*-3 fatty acid metabolic flux in microbial cells is tightly regulated by its intrinsic synthesis pathways ($\Delta 6\text{Des}/\Delta 6\text{Elo}/n3\text{Des}$ or $\Delta 9\text{Elo}/\Delta 8\text{Des}/n3\text{Des}$), and the key regulator n3Des is located at the branch point of metabolic flux.^{8,9} It can convert *n*-6 PUFAs linoleic acid (LA, C18:2*n*-6), γ -linoleic acid (GLA, C18:3*n*-6), dihomo- γ -linoleic acid (DGLA, C20:3*n*-6) and arachidonic acid (AA, C20:4*n*-6) into *n*-3 PUFAs α -linoleic acid (ALA, C18:3*n*-3), stearidonic acid (STA, C18:4*n*-3), eicosatetraenoic acid (ETA, C20:4*n*-3) and eicosapentaenoic acid (EPA, C20:4*n*-3), respectively, and is crucial for the synthesis of *n*-3 long-chain PUFAs (LC-PUFAs).^{10,11}

> Some reports have involved studies of various types of n3Des from a wide range of species including microalgae, cyanobacteria, yeast, fungi and plants.^{6,7,12} n3Des from various species were shown to exhibit a broad *n*-6 fatty acid substrate specificity with different catalytic efficiencies and preferences in the presence of various fatty acid substrates.¹³ The first yeast n3Des gene cloned from *Saccharomyces kluyveri* showed a dramatic preference for LA over AA substrate,^{14,15} and the same preference was found in the bifunctional $\Delta 12/n3$ desaturases from *Fusarium moniliforme, Fusarium graminearum*, and *Magnaporthe grisea*.¹⁶ FADS15 and maw3 from *Mortierella alpina* also showed a strong preference for LA.^{17,18} In 2018, two new n3Des isolated

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from plants – ShFAD3 from *Salvia hispanica* and PfFAD3 from *Perilla frutescens* – were used for ALA production based on their C18 substrate preference.¹² In contrast, the yeast n3Des gene Ppfad3 isolated from *Pichia pastoris* GS115 showed no substrate preference with nearly the same conversion rates towards each added substrate.¹⁹ Nevertheless, a new type of n3Des – sdd17 derived from an EPA-rich fungus *Saprolegnia diclina* and Pi-D17 from *Phytophthora infestans* – were found in their ability to exclusively desaturate C20 fatty acids.^{20,21} Three new Δ 17 desaturase enzymes isolated from *Pythium aphanidermatum*, *Phytophthora sojae* and *Phytophthora ramorum* were also reported with preferences for the C20 substrates.^{22,23}

Although various n-3 fatty acid desaturases have been identified, few $\Delta 17$ Des have been isolated compared to reported Δ 15Des. In this study, we reported the two new Δ 17 desaturases oRiFADS17 from Rhizophagus irregularis and oObFADS17 from Octopus bimaculoides. They were identified from the National Center for Biotechnology Information database using the known Δ17 desaturase oPaFADS17 gene (codon-optimized desaturase PaD17; GenBank accession numbers KT372000) as a template. Two additional hypothetical desaturase genes oAcFADS17 from Aplysia californica and oBgFADS17 from Biomphalaria glabrata - were chosen and synthesized altogether. To examine their activities and substrate preferences, all of them were expressed in Saccharomyces cerevisiae INVSc1. Based on bioinformatics analysis including topology model and molecular docking, the views of their structures, binding pockets and substrate tunnels were also clearly observed. The characterization and docking models of new Δ17Des - oRi-FADS17 and oObFADS17 - would contribute to elucidating the structure-function relationships of n3Des, and it would also provide a powerful new choice for genetic engineering of oleaginous organisms to produce the desired long-chain fatty acids in a sustainable way.

Materials and methods

Strains and plasmids

Escherichia coli Top 10 was supplied by our laboratory. The plasmid pYES2-oPaFADS17 was stored in Top 10.²⁴ The oAc-FADS17, oBgFADS17, oObFADS17 and oRiFADS17 genes were synthesized by Nanjing Genscript Biotechnology Co Ltd; *S. cerevisiae* yeast strain *INVSc1* was obtained from Invitrogen (Shanghai, China); and plasmid pYES2/NT C (Invitrogen) was used for heterologous expression of these genes.

GenBank accession numbers. *R. irregularis* codon-optimized Δ 17 desaturase oRiFADS17:MH028784.

O. bimaculoides codon-optimized $\Delta 17$ desaturase oObFADS17:MH028785.

Media and cultivation conditions

The isolated clones that contained these genes were cultivated in lysogenic broth medium with 100 μ g mL⁻¹ of ampicillin at 37 °C and shaken at 200 rpm. The yeast transformants were incubated with SC-U medium, which was synthetic minimal defined medium for yeast and contained 0.67% yeast nitrogen base, 2% raffinose or glucose, 0.01% amino acids (leucine, lysine, threonine, tryptophan, adenine, arginine and cysteine), 0.005% amino acids (aspartic acid, proline, serine, tyrosine, valine, histidine, isoleucine, methionine and phenylalanine) and 2% agar. Extra 2% galactose and 1% raffinose were supplemented during the induction period.

Sequence comparison, phylogenetic analysis

We used the amino acid sequence of oPaFADS17 as a template to BLAST these n3Des from the databases of the National Center for Biotechnology Information servers (https:// blast.ncbi.nlm.nih.gov/Blast.cgi). Four hypothetical desaturase proteins were then selected and compared with 16 other reported desaturases and aligned using the Clustal Omega web server, ver. 6. A GenBank accession number was provided for each sequence in Table 2. They included a diverse superfamily of enzymes from various strains, including *M. alpina* 1S-4, *M. alpina* ATCC32222, *F. moniliforme*, *M. grisea*, *F. graminearum*, *S. kluyveri*, *P. pastoris*, *C. elegans*, *P. frutescens*, *S. hispanica*, *S. diclina*, *P. aphanidermatum*, *P. ramorum*, *P. sojae*, *P. infestans*, *P. parasitica*, *R. irregularis*, *O. bimaculoides*, *B. glabrata* and *A. californica*. The sequences underlined in red were three His-rich motifs.

Phylogenetic analysis showed the relationship among the four hypothetical proteins and 16 reported desaturases. The phylogenetic tree was developed using the substitution model, and gaps were treated using the method of pairwise deletion. This work was created with Microsoft Word, Clustal Omega, and Adobe Illustrator.

PCR amplification and sequence analysis for oAcFADS17, oBgFADS17, oObFADS17 and oRiFADS17

The oAcFADS17, oBgFADS17, oObFADS17 and oRiFADS17 genes were amplified by polymerase chain reaction (PCR) with forward primers and reverse primers including restrictive endonuclease sites *Eco*RI and *Xho*I, respectively (Table 1). The amplified PCR products were then subcloned into the vector pYES2/NT C, and the positive clones were selected and confirmed by DNA sequencing in both directions (Sunny Biotechnology Co Ltd, Shanghai, China).

Yeast transformation, heterologous expression in *S. cerevisiae* and determination of substrate preference

The constructed plasmids pYES2-oAcFADS17, pYES2oBgFADS17, pYES2-oObFADS17 and pYES2-oRiFADS17 were transformed into *INVSc1 via* the lithium acetate transformation method. On SC-U selective plates, the recombinant yeast clones that exhibited uracil prototrophy were selected and identified by colony PCR. The yeast transformant with a pYES2-oPaFADS17 vector was used as a control strain.

To acquire insight into the substrate specificities of these recombinant proteins, the transformants were cultured for 48 h at 20 °C in SC-U media. Various fatty acid substrates were supplemented into induction buffer with fatty acids (0.1 mM *cis*-LA, 0.1 mM *cis*-DGLA or 0.1 mM *cis*-AA). Cells were collected *via* filtration by centrifugation for 30 s at 12 000

Table 1 Primers used in this study

Primer name	Restriction enzyme	Oligo nucleotide sequence $(5'-3')^a$	Function
Rid-F	<i>Eco</i> RI	CCG <u>GAATTC</u> ATGTCGCCCTTGGAGC	oRiFADS17 amplification for expression in S. cerevisiae
Rid-R	XhoI	CCG <u>CTCGAG</u> TTACTGGTTCTTCTCCTTCTGG	
Obd-F	EcoRI	CCG <u>GAATTC</u> ATGGTCACCGCTATCC	oObFADS17 amplification for expression in S. cerevisia
Obd-R	XhoI	CCG <u>CTCGAG</u> TTACTTGTACCAGTGGATGAGG	
Oac-F	EcoRI	CCG <u>GAATTC</u> ATGGAGCCCACCCCCCCGTCTCG	oAcFADS17 amplification for expression in S. cerevisiae
Oac-R	XhoI	CCG <u>CTCGAG</u> TTACTTGTAGAAGTGGATG	
Obg-F	EcoRI	CCG <u>GAATTC</u> ATGGAGACCGCTCCTGC	oBgFADS17 amplification for expression in S. cerevisia
Obg-R	XhoI	CCGCTCGAG TTACTTGTAGAGGTGG	

^a Underlined sequences indicate the additional restriction sites.

Table 2 Literature summary of catalytic efficiency of n-3 desaturases from various species with LA/GLA/DGLA and AA substrates

Strain name	The conversion rate of n3Des on LA (%)	The conversion rate of n3Des on GLA (%)	The conversion rate of n3Des on DGLA (%)	The conversion rate of n3Des on AA (%)	Locus
Saprolegnia diclina	_	_	4.98	25.9	AY373823
Phytophthora infestans	_	_	_	30.94	CAJ30870.1
Pythium aphanidermatum	_	5.97	28.85	56.46	FW362186.1
Phytophthora ramorum	_	4.70	31.02	37.12	FW362214.1
Phytophthora sojae	_	6.18	35.45	48.79	FW362213.1
Phytophthora parasitica	7.11	5.63	25.34	49.70	KT372001
Caenorhabditis elegans	11	_	0	≈56	CELE_Y67H2A.8
Pichia pastoris	36.5	33.8	35.1	35.3	EF116884
Mortierella alpina 1S-4	11.50	8.90	3.60	6.70	AB182163
Mortierella alpina ATCC32222	62.37	63.99	41.82	23.67	AGZ84120.1
Fusarium moniliforme	49.30	15.70	17.50	9.80	DQ272516
Fusarium graminearum	17.40	3.10	5.80	2.80	EAA75859
Magnaporthe grisea	18.6	3.40	4.6	4.7	XP 362963
Saccharomyces kluyveri	22.0	10.0	3.9	1.60	AB118663
Salvia hispanica	16.9	_	_	_	KX610653
Perilla frutescens	10.6	_		_	KX880389

g min⁻¹, washed twice with distilled water, and freeze-dried. The dried cells were used for fatty acid analysis and determination of substrate preference and the remainder was stored at -80 °C.

Western blotting

To analyze the translation levels of these recombinant desaturases, 20 μ L of the collected cells were mixed with 5× SDS lysate buffer and heated in boiling water for 10 min for western blotting. The quantitative analysis method of bovine serum albumin was used to determine the volume of equivalent samples for SDS-PAGE gel loading. The gel was transferred to a polyvinylidene fluoride membrane (60 mA, 5 h), which was then incubated with mouse THETM His tag antibody (1 : 5000) and special HRP-conjugated goat anti-mouse IgG (1 : 2000). Finally, the membrane was developed and fixed with the chemiluminescence reagent for 1 min, and photographs were taken with FluorChem FC3.

Lipid extraction and fatty acid analysis

Lipid extraction and methyl esterification of free fatty acids was conducted as described previously.²⁵ Fatty acid methyl esters

(FAMEs) profiles were then analyzed by GC-2010 (Shimadzu Co. Japan) with a 30 m \times 0.32 mm DB-wax column (Agilent) and an FID detector. The samples were measured with the following temperature program: 40 °C for 5 min, increase to 120 °C at 20 °C min⁻¹, hold for 3 min, ramp to 190 °C at 5 °C min⁻¹, ramp to 220 °C at 4 °C min⁻¹ and hold 220 °C for 20 min.

Topology prediction

Prediction of membrane-spanning helices and topology models for the four hypothetical proteins and oPaFADS17 were executed with the PolyPhobius program (https://omictools.com/ polyphobius-tool) HMMTOP (http://www.enzim.hu/hmmtop/ html/submit.html), TMHMM (http://www.cbs.dtu.dk/services/ TMHMM/) online websites and I-TASSER algorithms (https:// zhanglab.ccmb.med.umich.edu/I-TASSER/) (Fig. S1 and S2⁺).²⁶

Homology modeling of three-dimensional structures and molecular docking of oObFADS17 and oRiFADS17

A template of crystal structure of human stearoyl-CoA hSCD1 desaturase was used to construct homology models of the oObFADS17 and oRiFADS17. The Schrödinger Suite shareware

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was used for molecular docking and the docking models were performed using PYMOL software (Fig. 7). Their models were docked with the representative substrates LA-COA and AA-COA. Two metal zinc-ions bound to the substrate-binding pockets were also depicted as grey or yellow spheres. The view of their predicted structures, binding pockets and substrate tunnels were clearly observed.

Results

Biosynthetic process and literature summary of catalytic efficiency of n3Des

Desaturases are large complex biological protein families that catalyze a wide range of fatty acids in the biosynthesis of PUFAs. Although they achieve these transformations *via* various biosynthetic processes ($\Delta 6 \text{Des}/\Delta 6 \text{Elo}/n3 \text{Des}$ or $\Delta 9 \text{Elo}/\Delta 8 \text{Des}/n3 \text{Des}$), n3Des which converts *n*-6 polyunsaturates to their *n*-3 counterparts, is the key enzyme for the synthesis of long-chain fatty acids such as EPA and DHA (Fig. 1).^{9,16}

Some genes that encode n3Des were recently cloned and sequenced from various organisms. They consist mainly of Δ 15Des and Δ 17Des based on their substrate preferences and exhibit broad *n*-6 fatty acid substrate preferences and specificities. A review of the catalytic efficiencies of these reported n3Des is shown in Table 2, which illustrates their obvious preferences for C18 or C20 *n*-6 fatty acid substrates. FADS15 in *M. alpina* (Δ 15 desaturase) had the highest catalytic activity towards LA, whereas oPaFADS17 (Δ 17 desaturase) had the highest catalytic activity towards AA.

Sequence comparison, phylogenetic analysis

The protein sequences of the four candidate $\Delta 17$ desaturase proteins from *A. californica*, *B. glabrata*, *O. bimaculoides* and *R.*

irregularis had 416, 402, 419 and 397 amino acids, respectively. As illustrated in Fig. 2, sequence comparison analysis suggested that these ORFs shared greater sequence similarities and homologies to Δ 17 desaturases than to Δ 15 desaturases. The results also show that oRiFADS17 is 44% identical to Pi-D17, oObFADS17 is 46% identical to Ps-D17, oBgFADS17 is 45% identical to Pr-D17 and oAcFADS17 is 43% identical to oPa-FADS17. The lighter shaded areas were conserved residues, and the blue shaded areas were consensus residues. These new genes were also predicted to have three typical His-rich desaturase motifs (underlined with red thick lines), suggesting they owned the characteristic of Δ 17 desaturases.

Phylogenetic analysis also demonstrated the evolutionary relationship amongst these selected n3Des from various organisms using the neighbour-joining method (Fig. 3). According to the clear separation of the branches of these proteins identified by the phylogenetic tree, they were proposed to be roughly distinguished into different lineages: $\Delta 15$ desaturases, $\Delta 17$ desaturases and hypothetical $\Delta 17$ desaturases. As illustrated in Fig. 3, the last group comprised the four new genes. They were found from mycorrhizal fungi and sea mollusks, and were thus quite different from other n3Des. According to the bootstrap values at this branch point, the genome evolution of the phylogenetic relationships revealed significant differences in their species origins from the other reported $\Delta 17$ desaturases. In conclusion, it was deduced that they belonged to $\Delta 17$ desaturases.

Cloning and expression of four candidate $\Delta 17$ desaturase genes in S. cerevisiae

Owing to the bioinformatics analysis in the public databases, the oAcFADS17, oBgFADS17, oObFADS17 and oRiFADS17 genes were chosen and amplified by PCR with forward primers and



Fig. 1 Schematic diagram of the fatty acid pathways for EPA and DHA synthesis.

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reverse primers including the restrictive endonuclease sites *Eco*RI and *Xho*I, respectively (Table 1); these fragments were then subcloned into the vector pYES2/NT C (Fig. 4a). After lithium acetate transformation, their positive yeast clones heterologously expressed in *INVSc1* were obtained. The result from the map of DNA agarose electrophoresis indicated that they were successfully expressed (Fig. 4b).

Western blot analysis of recombinant proteins was conducted to reveal successful transcription of the four candidate $\Delta 17$ desaturases and of the control, which was also a complementary proof for evaluation of transforming activity (Fig. 5a). The target 46 kD bands on the membrane were marked with an arrow. The gray scanning analysis result showed no obvious differences among the translation levels of the recombinant proteins (Fig. 5b).



Fig. 3 Phylogenetic analysis showing the relationship among 16 selected desaturases and 4 hypothetical proteins.



Fig. 4 Construction and verification of *INVSc1* plasmids. (a) Construction of *INVSc1* plasmids pYES2-oAcFADS17, pYES2-oBgFADS17, pYES2-oDbFADS17, pYES2-oRiFADS17. (b) Verification of *INVSc1* transformants. M: maker; lane 1: control; lanes 2-5: pYES2-oAcFADS17, pYES2-oBgFADS17, pYES2-oObFADS17, pYES2-oRiFADS17, pYES2-oRiFADS17.

Characterization of activities and substrate specificities of four candidate $\Delta 17$ desaturases

The transformants were constructed to confirm that the four candidate proteins have $\Delta 17\text{Des}$ activity. Their lipids were extracted as previously described,²⁵ and the fatty acids compositions of four candidate proteins were analyzed as shown in Table 3 and Fig. 6a. The fatty acid experiments indicated that the two $\Delta 17$ desaturases oRiFADS17 and oObFADS17 were capable of catalyzing LA, GLA, DGLA and AA conversion to ALA, SDA, ETA and EPA. Furthermore, they both exhibited stronger preference for C20 PUFA substrates DGLA and AA. The conversion rates of oRiFADS17 desaturase were $58.5\% \pm 3.2\%$ for AA and $35.1\% \pm 1.6\%$ for LA and those of oObFADS17 were 22.9% $\pm 2.6\%$ and $3.8\% \pm 0.2\%$, respectively. According to the conversion rate in Fig. 1, oRiFADS17 desaturase in this study had higher conversion rate (3–4 fold) of the LA substrate than

the control oPaFADS17 desaturase and other reported Δ 17 desaturases, while containing a high conversion rate for AA (nearly 60%) at the same time (Fig. 6b). In addition, oObFADS17 was found to be the first reported Δ 17 desaturase from sea mollusks, although it converted little AA compared with oRi-FADS17. However, another set of predicted Δ 17 desaturase oAcFADS17 from *A. californica* and oBgFADS17 from *B. glabrata* did not have any detectable desaturase activity. Their fatty acid compositions of the oAcFADS17 and oBgFADS17 strain.

Conversion rates in response to treatment at various temperatures

The effects of temperatures on fatty acid substrate preferences and the transforming activities of some n3Des have been investigated.^{13,23,24,27} To reveal the conversion rates in response to treatment at various temperatures, the yeast transformants were cultured 28 °C, 20 °C and 15 °C. As shown in Table 4, the conversion rates of the oPaFADS17, oObFADS17 and oRiFADS17 desaturase increased slightly as the incubation temperature decreased, but they did not change their substrate preferences after low-temperature treatment. Their responses were consistent with other n3Des such as maw3, FADS15 and CiFAD3, but not with oPPD17.^{13,23,24,27}

Topology prediction

To further facilitate the gathering of biological information of these desaturases and explain the differences in the membrane structures of these proteins, topology analysis of the sequences was carried out with structural prediction software on Clustal Omega web servers. Their structural arrangements were very similar to that of the control protein oPaFADS17. As a result of the membrane topology in Fig. S1 and S2,† their primary structures contained 2 large hydrophobic domains and 3 separate hydrophilic clusters but no confirmed important core cytb5 motif HPGG, unlike most $\Delta 6$ and $\Delta 9$ desaturases. Due to the high similarities in their amino acid sequences and membrane structures, they were predicted to show similar substrate chainlength and regioselectivity preferences. In contrast, extra AHs were predicted with a lower probability in oPaFADS17 (Cterminal) and oAcFADS17 (N-terminal). In addition, a signal peptide was found in the oBgFADS17 protein sequence (Nterminal). Therefore, it was deduced that the structural diversities may be the reason why these two genes were inactive.

Three-dimensional structures and molecular docking of oObFADS17 and oRiFADS17

To further validate the molecular structures of substrates of oObFADS17 and oRiFADS17, their homology models of 3dimensional structures were also predicted. On the basis of a recent crystal structure of the human stearoyl-CoA desaturase hSCD1, the molecular docking of oObFADS17 and oRiFADS17 with LA-CoA or AA-CoA substrate was successfully performed (Fig. 7a, b, d and e). The carrier portion of the substrate and binding pocket were clearly observed. Two zinc-ions bound to the pocket center were close to the *n*-6 carbon atom of each



Fig. 5 Verification of translation levels of yeast transformants. (a) Western blotting of recombinant proteins expressed in S. cerevisiae. M: marker; C: control protein oPaFADS17; arrows in the western blotting indicated the location of the target protein. (b) Western-blot gray scale result showed that the average translation levels of 4 hypothetical Δ 17 desaturases were almost the same amount.

substrate. From the kink and the narrow aperture of the substrate tunnel, it was assumed to present that how the substrate entered in or released out of the tunnel (Fig. 7c and f). Based on knowledge of these structures, the molecular mechanism of substrate specificity and preference would be further deeply investigated and this also enabled us to strengthen our understanding of the structure-function relationships of novel fatty acid desaturases.

Discussion

Several previous studies have reported substrate preferences and specificities in membrane-bound fatty acid desaturases such as Δ 4des Δ 6des, Δ 8des, Δ 9des and Δ 12des.²⁸⁻³³ The biological information including sequence alignment, phylogenetic analysis and topology prediction was applied to help define the structure-function relationship of these desaturases.

Table 3 Fatty acid compositions (% w/w) of the total lipid contents of yeast transformants expressing the control and pYES2-oAcFADS17, pYES2oBgFADS17, pYES2-oObFADS17, pYES2-oRiFADS17

	Transformants ^b				
Fatty acid	oPaFADS17	oAcFADS17	oBgFADS17	oObFADS17	oRiFADS17
PA, C16:0	25.9 ± 2.4	23.2 ± 2.7	26.7 ± 1.4	23.4 ± 1.9	25.5 ± 0.7
PA, C16:1	30.6 ± 0.8	29.5 ± 0.9	30.4 ± 2.9	32.3 ± 2.4	30.9 ± 2.3
SA, C18:0	4.6 ± 1.0	4.0 ± 0.2	4.5 ± 0.7	5.0 ± 0.4	5.2 ± 0.8
OA, C18:1	24.2 ± 3.5	23.7 ± 1.1	23.6 ± 2.7	24.5 ± 1.2	24.0 ± 2.7
LA, C18:2 <i>n</i> -6	5.9 ± 0.3	8.7 ± 1.1	7.4 ± 0.7	8.6 ± 0.5	5.1 ± 0.4
ALA, C18:3 <i>n</i> -3	0.6 ± 0.1	ND	ND	0.4 ± 0.1	2.9 ± 0.3
AA, C20:4 <i>n</i> -6	2.1 ± 0.5	5.1 ± 0.4	5.8 ± 0.4	4.7 ± 0.8	2.1 ± 0.3
EPA, C20:5 <i>n</i> -3	3.2 ± 0.2	ND	ND	1.3 ± 0.2	3.0 ± 0.2
LA conversion rate ^{<i>a</i>}	9.2 ± 1.6	_	_	4.4 ± 0.5	36.3 ± 4.0
AA conversion rate ^{<i>a</i>}	60.4 ± 0.8	—	—	21.7 ± 2.5	59.1 ± 4.7

^{*a*} Conversion rate = $100 \times [(\text{product})/(\text{product} + \text{substrate})]$. ND: not detected. ^{*b*} Substrate concentration was 0.1 mM LA or AA.



Fig. 6 Conversion rates of yeast transformants on different fatty acids. (a) Rate of substrate conversion of oPaFADS17, oAcFADS17, oBgFADS17, oObFADS17 and oRiFADS17 in *S. cerevisiae*. (b) Rate of substrate conversion of sdd17, Pi-D17, Pr-D17, Ps-D17, fat-1, oPa-FADS17, oRiFADS17 and oObFADS17.

Based on the application of multiple sequence alignment, four candidate ORFs for $\Delta 17$ desaturase proteins from *A. californica*, *B. glabrata*, *O. bimaculoides* and *R. irregularis* were found. They were then compared and aligned with 16 reported n3Des with various substrate preferences. Pair-wise comparison showed that they all contained iron-binding sites with an active center and three typical His-rich motifs (one HX₃H and two HX₂HH). Fig. 2 showed many lighter or darker shaded amino acid residues in their protein sequences, and a switch of even a single amino acid of these residues would result in a dramatic change in substrate preference and/or substrate characteristics.^{29,30,33} These conserved residues between the $\Delta 15$ and $\Delta 17$ desaturases would be alternative possibilities for further examination of their activities and substrate preferences.^{24,29,31} Collectively, the residues provide significant insight into the relationship between the structure and function of n3Des.

Until now, the three-dimensional and crystal structure of n3Des have not been determined. A membrane topology model was used for determination of the functional domains using comparative analysis.³⁴ On the basis of the high similarities in their amino acid sequences and the proposed topology models (Fig. S1 and S2†), proteins encoded by these genes were predicted to have Δ 17 desaturase activity. However, this method is not entirely correct, such as with another two candidate Δ 17 desaturase genes oAcFADS17 and oBgFADS17. It was assumed that the extra N-terminal AHs in oAcFADS17 might cause them to lose their catalytic activities. Considering the high sequence and species similarities with oObFADS17, they deserve further investigation.

The typical types of n3Des have been reported from a wide range of species including microalgae, cyanobacteria, yeast, fungi and plants.^{6,22,34} The phylogenetic tree in Fig. 3 shows that the top group of n3Des was derived from M. alpina, yeast and plants. The second cluster consisted of $\Delta 17$ desaturases from Pythium and Phytophthora and from nematodes. Interestingly, the $\Delta 17$ desaturases gene fat-1 from *Caenorhabditis elegans* was actually closer to the first category.35 This may be related to the source of species and genetic differences. The $\Delta 17$ desaturase oObFADS17 of the bottom group originated from sea mollusks together with oAcFADS17 and oBgFADS17, thus differed greatly from other n3Des. Although oAcFADS17 and oBgFADS17 could not catalyze each added n-6 fatty acid substrates, we believe that sea mollusks organisms are a new source of Δ 17Des. Characterization of oObFADS17 would not only deepen our knowledge of a natural source of n3Des, but also pave the way for elucidating of the molecular basis and the enigmatic enzyme mechanism.

According to the conversion rates (Table 3), the calculated conversion efficiencies of the three $\Delta 17$ desaturase enzymes PrD17, PsD17 and PaD17 were almost 6% for the LA substrate.⁶ In this study, the oRiFADS17 desaturase had nearly 34%

Table 4 Fatty acid compositions (% w/w) of yeast transformants expressing pYES2-oPaFADS17, pYES2-oObFADS17, pYES2-oRiFADS17 plasmids at 15 °C, 20 °C or 28 °C

	Conversion rate ^{a} (%)						
	LA, C18:2 <i>n</i> -6			AA, C20:4 <i>n</i> -6			
Transformants ^b	15 °C	20 °C	28 °C	15 °C	20 °C	28 °C	
oPaFADS17	13.4 ± 2.3	9.4 ± 1.4	7.0 ± 1.2	66.2 ± 2.4	60.7 ± 0.9	55.6 ± 4.1	
oObFADS17 oRiFADS17	$6.5 \pm 0.8 \ 37.6 \pm 1.5$	$4.4\pm0.5\ 34.2\pm3.1$	$rac{ m ND}{ m 26.2\pm1.0}$	$26.2\pm1.8\64.1\pm1.6$	$22.6 \pm 2.5 \\ 58.5 \pm 3.2$	$18.4 \pm 0.9 \\ 51.8 \pm 2.2$	

^{*a*} Conversion rate = $100 \times [(\text{product})/(\text{product} + \text{substrate})]$. ND: not detected. ^{*b*} Substrate concentration was 0.1 mM LA or AA.



conversion rate of the LA substrate (nearly 5 fold) and a higher conversion rate for AA (nearly 60%) at the same time (Fig. 6b). The highest conversion rate for AA can reach 66% when 0.1 mM AA was added at a lower temperature 15 °C. Generally speaking, although the substrate preference of oRiFADS17 was weaker compared with other Δ 17 desaturases, its comprehensive catalytic ability for *n*-6 PUFAs substrates increased significantly. Because of the strong preference for C18 substrates of endogenous n3Des or their limited catalytic efficiency *in vivo* in some oleaginous microorganisms, the newly identified Δ 17 desaturase could be an excellent choice for genetic engineering of organisms and have great potential to be used to produce EPA and DHA.^{18,22,23}

In contrast to the reported soluble acyl-ACP desaturases, little is known about the structure-guiding principle involved in substrate specificity and regioselectivity of n3Des, due to lack of the three-dimensional structure information. For the first time, the representative substrates LA-COA and AA-COA were chosen as the ligands, and the proposed structures of n3Des were clearly visualized based on molecular docking. Captures of the tunnel-like pockets were also shown in the hydrophobic interior of the n3Des. It was interpreted that how the substrate entered into or released from the tunnel, and this will help understand the crucial roles of active amino acids in regulating substrate specificity and enzyme activity.

Traditional source of n-3 PUFAs is diminishing due to overfishing and marine pollution, and a low dietary intake of n-3 PUFAs, especially EPA and DHA, is observed worldwide. Oleaginous microorganisms are increasingly attractive for their

promising potential as an alternative source. In their fatty acid pathways for EPA and DHA synthesis, $\Delta 17$ Des is more important than $\Delta 15$ Des because its ability to directly desaturate C20 fatty acids substrates. The newly identified $\Delta 17$ desaturases could therefore contribute to producing PUFAs for dietary supplementation or the prevention and treatment of numerous diseases.

Conclusion

In this study, two genes - oRiFADS17 from R. irregularis and oObFADS17 from O. bimaculoides – were confirmed as new $\Delta 17$ desaturases and share high protein sequence similarities with known $\Delta 17$ desaturases. They show a preference for C20 PUFA substrates DGLA and AA, and the conversion rate of oRiFADS17 from AA to EPA was up to 60% when expressed in S. cerevisiae. Furthermore, oRiFADS17 desaturase has greater catalytic activity for LA substrates than other reported $\Delta 17$ desaturases. oObFADS17 is also the first reported Δ 17 desaturase in a sea mollusk. The predicted 3-dimensional structures and molecular docking of oObFADS17 and oRiFADS17 with LA-CoA or AA-CoA substrate further supported our findings. The characterization of these two desaturases and the knowledge of their structures will contribute to strengthening our understanding of the relationship between the structure and function of novel fatty acid desaturases, and be of great value in PUFAs production for dietary supplementation and the prevention and treatment of numerous diseases.

Author's contributions

RSC Advances

CR designed the study, conducted the experiments and drafted the manuscript. HC and YQC helped propose and design the study, and revised the manuscript. HC, XT, ZG, JZ, HZ and WC assisted with revising the manuscript. All authors read and approved the final manuscript.

Conflicts of interest

The authors have no conflict of interest.

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