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pfaB products determine the molecular species produced in bacterial polyunsaturated fatty acid biosynthesis

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Abstract

When pDHA4, a vector carrying all five pfaA-pfaE genes responsible for docosahexaenoic acid (DHA;

22:6) biosynthesis in Moritella marina MP-1, was coexpressed in Escherichia coli with the individual

pfaA-pfaD genes for eicosapentaenoic acid (EPA; 20:5) biosynthesis from Shewanella pneumatophori

SCRC-2738, both polyunsaturated fatty acids were synthesized only in the recombinant carrying pfaB

for EPA synthesis. Escherichia coli coexpressing a deleted construct comprising pfaA, pfaC, pfaD, and

pfaE for EPA and pfaB for DHA produced EPA and DHA. Both EPA and DHA were detected in

bacteria that inherently contained pfa genes for DHA. These results suggest that PfaB is the key enzyme

determining the final product in EPA or DHA biosynthesis.

Running Head: Bacterial polyunsaturated fatty acid molecular species

Keywords

Docosahexaenoic acid; Eicosapentaenoic acid; Moritella marina strain MP-1; Polyunsaturated fatty

acid; pfaB gene.

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Introduction

Bacteria synthesize polyunsaturated fatty acids (PUFAs), such as eicosapentaenoic acid (EPA; 20:5) and docosahexaenoic acid (DHA; 22:6), in a mode of polyketide biosynthesis that differs from the combined fatty acid elongation and aerobic desaturation system commonly observed in eukaryotes (Metz *et al.*, 2001). Because there are some benefits, including the need for fewer reducing equivalents such as NADPH and simple fatty acids consisting solely of EPA or DHA, in using this bacterial system to produce PUFA rather than using the aerobic system (Okuyama *et al.*, 2007), the factor(s) that determine the final molecular species in the system must be identified for the practical production of these PUFAs.

The bacterial polyketide synthase-type multienzyme complex responsible for the biosynthesis of EPA and DHA is composed of proteins encoded by five *pfa* genes: *pfaA*, *pfaB*, *pfaC*, *pfaD*, and *pfaE* (Okuyama *et al.*, 2007). Four of the *pfa* genes of the DHA-producing bacterium *Moritella marina* MP-1, *pfaA*, *pfaB*, *pfaC*, and *pfaD*, are clustered on the chromosome (Tanaka *et al.*, 1999), whereas *pfaE* is located separately (Orikasa *et al.*, 2006a). However, in the EPA-producing bacterium *Shewanella pneumatophori* SCRC-2738 and some other EPA-producing bacteria, all five genes are clustered in close proximity (see Fig. 1). The number of acyl carrier protein (ACP) repeats in the *pfaA* gene product is five in *M. marina* MP-1 and six in *S. pneumatophori* SCRC-2738. The 3-ketoacyl synthase (KS) domain is present in the *pfaB* product of *M. marina* MP-1, but not in that of *S. pneumatophori* SCRC-2738 (Orikasa *et al.*, 2004).

Although the structures of the *pfa* genes and their encoded proteins from EPA-producing and DHA-producing bacteria are very similar, the bacterial syntheses of EPA and DHA have been generally considered mutually exclusive; that is, bacteria that produce EPA do not produce DHA and vice versa. Some deep-sea or Antarctic bacteria produce only EPA (Bowman *et al.*, 1998; Fang *et al.*, 2003; Nichols *et al.*, 1997), as do EPA-producing *S. pneumatophori* SCRC-2738 (Hirota *et al.*, 2005) and *S.*

marinintestina IK-1 (Satomi et al., 2003). Conversely, some deep-sea bacteria produce only DHA (DeLong & Yayanos, 1986; Hamamoto et al., 1995). However, this criterion is not absolute, because some bacteria from the intestinal contents of marine fish produce both EPA and DHA (Watanabe et al., 1997; Yano et al., 1998; Yano et al., 1994). In most of these cases, DHA is the primary product (approximately 5% or more of total fatty acids) and EPA is a minor component (less than 1% of the total fatty acids). So far, only EPA or DHA has been produced in recombinant E. coli cells carrying the pfa genes responsible for EPA or DHA biosynthesis, respectively (Orikasa et al., 2004; Orikasa et al., 2006a). These findings suggest that the molecular species involved in the bacterial biosynthesis of EPA or DHA must be determined by one or more of the pfa gene product(s). However, as demonstrated previously, the pfaE product has been excluded as a candidate determinant because the pfaE genes of S. pneumatophori SCRC-2738 and M. marina MP-1 are compatible with the other pfa genes of both bacterial species (Orikasa et al., 2006a; 2006b). The number of ACP domain repeats in the pfaA product is not involved in determining the molecular species of PUFA (Okuyama et al., 2007). Nonetheless, the details of how these highly similar structures preferentially synthesize either EPA or DHA have not yet been clarified.

To the best of our knowledge, pDHA3 is the only vector encoding the bacterial genes responsible for the biosynthesis of DHA, pfaA, pfaB, pfaC, and pfaD of M. marina MP-1 (Okuyama et al., 2007; Orikasa et al., 2006b). However, only DHA was produced in recombinant E. coli cells coexpressing the genes encoded by pDHA3 and pfaE from M. marina MP-1 (Orikasa et al., 2006b) or pfaE from other bacteria (Orikasa et al., 2006a; Sugihara et al., 2008). Various constructs are available that contain the pfa genes of S. pneumatophori SCRC-2738. In addition to cosmid and plasmid vectors that carry all five pfa genes, deletion constructs lacking one of the five pfa genes and clones containing individual pfa genes have been prepared (Orikasa et al., 2004; Orikasa, 2007; Yazawa et al., 1998). These clones have been used to increase the heterologous production of EPA (Orikasa et al., 2004;

Orikasa *et al.*, 2007) and to determine the physiological functions of EPA in *E. coli* recombinant systems (Nishida *et al.*, 2006a; 2006b; Okuyama *et al.*, 2008).

In this study, a vector that carries all five pfa genes derived from M. marina MP-1 was constructed (pDHA4; Fig. 1). To identify the factor(s) determining the final product in the polyketide mode of PUFA biosynthesis, the pfa genes on pDHA4 were coexpressed with individual pfa genes derived from S. pneumatophori SCRC-2738. The pfaB gene was also cloned from pDHA4 and expressed with a construct encoding all five pfa genes (pEPA Δ 1; see Table 1) and a mutant pfa gene construct encoding pfaA, pfaC, pfaD, and pfaE (pEPA Δ 6) for EPA biosynthesis.

Materials and methods

Bacterial cells and cultivation

Escherichia coli DH5α was the host bacterium for the recombinant *pfa* genes. It was grown in LB medium supplemented with the indicated antibiotics at 37°C for 16 h. Part (40 μL) of the culture was transferred to 2 mL of fresh LB medium and was cultivated with shaking at 180 rpm at 15°C for 96 h to stimulate DHA and/or EPA production. Bacterial strains and vectors used in this study are summarized in Table 1.

EPA-producing *S. pneumatophori* SCRC-2738 from the intestines of Pacific mackerel (Hirota *et al.*, 2005) and *Shewanella marinintestina* IK-1 from squid body (Satomi *et al.*, 2003) were cultivated in 3 mL of LB medium containing 3.0% NaCl at 20°C for 48 h. DHA-producing *M. marina* MP-1 from deep-sea waters (DeLong & Yayanos, 1986) was cultivated in the same medium at 15°C for 72 h. *Moritella marina* MP-1 and *E. coli* recombinant carrying pEPAΔ6 and pSTV28::*pfaB*(DHA) were cultivated in the presence and in the absence of cerulenin at indicated concentrations.

DNA procedures

To construct a vector carrying all five *pfa* genes responsible for the biosynthesis of DHA, pDHA3 harboring *pfaA*, *pfaB*, *pfaC*, and *pfaD* from *M. marina* MP-1 and a fragment of pET21a::*pfaE* harboring *pfaE* from the same bacterium were combined (Table 1). Polymerase chain reaction (PCR) amplified a DNA fragment that included *pfaE* and a T7 RNA polymerase binding site using pET21a::*pfaE* as a template, with the forward primer (5'-TCAAGGGCATCGGTCGACATC-3') and reverse primer (5'-CCGGATATAGGTCGACCTTTC-3'), in which *Sal*I sites were introduced by base modification and are underlined. The PCR-amplified 2.8 kbp DNA fragment containing *pfaE* was digested with *Sal*I and then purified by agarose gel electrophoresis. It was ligated into pDHA3, which had been digested with *Sal*I, and the recombinant DNA was introduced into *E. coli* DH5α cells by heat shock. Colony direct PCR was used to confirm the presence of the insert by using primers designed against the inner sequence of the *pfaE* gene (Orikasa *et al.*, 2006a): 5'-TGTTGTTTCTACCGCACCTT-3' for the forward primer and 5'-GTTGGCCGATTCATTAATGC-3' for the reverse primer. A positive clone was cultured to prepare the plasmid. Plasmid digested with *Sal*I gave two bands of 25 kbp and 2.8 kbp on agarose gels. The plasmid was designated pDHA4.

This study used individual *pfa* genes for the biosynthesis of EPA or DHA. *pfaB*(DHA) gene of 2.5 kbp was excised from pDHA4 using restriction enzymes *Bam*HI and *Sma*I. Treatment of pDHA4 with *Bam*HI and *Sma*I provided four major DNA bands of 3.2 kbp, 5.6 kbp, 8.3 kbp, and 10.9 kbp on agarose gels. The band of 3.2 kbp, which was expected to contain the *pfaB* gene of 2.5 kbp, was extracted and purified, and then ligated into the pSTV28 plasmid, which had been digested with *Bam*HI and *Sma*I. The plasmid was designated pSTV28::*pfaB*(DHA).

pUC18::*pfaB*(EPA) and pUC18::*pfaD*(EPA) were prepared from the pSTV28::*pfaB*(EPA) and pSTV28::*pfaD*(EPA) vectors, respectively. Plasmids pSTV28::*pfaB*(EPA) and pSTV28::*pfaD*(EPA)

(Orikasa, 2007; Yazawa *et al.*, 1998) were digested with *Sfb*I and *Eco*RI and with *Bam*HI and *Eco*RI, respectively. The required DNA fragments were separated on agarose gels, purified and ligated into the pUC18 plasmid vector (Takara Bio), which had been digested with *Sfb*I and *Eco*RI to produce pUC18::*pfaB*(EPA) and with *Bam*HI and *Eco*RI to produce pUC18::*pfaD*(EPA). pUC18::*pfaA*(EPA) and pUC18::*pfaC*(EPA) have been previously described (Orikasa, 2007; Yazawa *et al.*, 1998).

Analysis of fatty acids

Fatty acids were converted to methyl esters and then analyzed as described (Orikasa *et al.*, 2006a). Identification of DHA and EPA was performed by comparing their retention time on gas–liquid chromatography with standards and by gas chromatography–mass spectrometry as described previously (Orikasa *et al.*, 2006a; Orikasa, 2007).

Results and discussion

Construction of a vector carrying the five *pfa* genes for DHA biosynthesis (pDHA4) and its combined expression with individual *pfa* genes for EPA

The five *pfa* genes—*pfaA*, *pfaB*, *pfaC*, *pfaD*, and *pfaE*—derived from DHA-producing *M. marina* MP-1 were designated *pfaA*(DHA), *pfaB*(DHA), *pfaC*(DHA), *pfaD*(DHA), and *pfaE*(DHA), respectively, to differentiate them from the *pfa* genes from EPA-producing *S. pneumatophori* SCRC-2738: *pfaA*(EPA), *pfaB*(EPA), *pfaC*(EPA), *pfaC*(EPA), and *pfaE*(EPA) (Table 1).

The pDHA4 vector was constructed by linking pDHA3 (pSTV29 carrying *pfaA*, *pfaB*, *pfaC*, and *pfaD* for DHA biosynthesis and five other irrelevant open reading frames) with *pfaE* derived from *M*.

**marina MP-1 (Fig. 1). The successful construction of pDHA4 was confirmed by the production of DHA

in recombinant *E. coli* DH5α (Table 2). No EPA was detected. The level of DHA was approximately 4% of the total fatty acid content when the cells were grown at 15 °C for 96 h. Table 2 shows the results of the combined expression of pDHA4 and either *pfaA*(EPA), *pfaB*(EPA), *pfaC*(EPA), or *pfaD*(EPA) in *E. coli* DH5α. In any combination, DHA was produced. However, EPA was detected in the cells coexpressing pDHA4 and *pfaB*(EPA). These results suggest that EPA was synthesized by an enzyme complex encoded by *pfaA*(DHA), *pfaC*(DHA), *pfaD*(DHA), and *pfaE*(DHA) and *pfaB*(EPA), and that the final EPA production was determined by the *pfaB*(EPA) product in this enzyme complex. The reason for the relatively low level of DHA (less than 1% of total fatty acids) in only the pDHA4 and *pfaB*(EPA) combination (Table 2) is unknown. It is considered that, in addition to the Pfa enzyme complex containing PfaB(EPA), these recombinant cell should have the complex of PfaA, PfaB, PfaC, PfaD, and PfaE for DHA synthesis derived only from pDHA4. When the two types of *pfaB* gene products in the enzyme complexes are compared, PfaB(EPA) may be more efficacious in producing EPA, even in the Pfa enzyme complex comprising PfaA, PfaC, PfaD, and PfaE for DHA and PfaB(EPA), than PfaB(DHA) is in producing DHA in the Pfa enzyme complex for DHA. Another possibility is that there is competition for the precursors of EPA and DHA biosynthesis between the two Pfa enzyme complexes.

Cloning of pfaB(DHA) and its combined expression with clustered pfa genes for EPA

The pfaB(DHA) gene was excised from pDHA4 and integrated into pSTV28 (designated pSTV28::pfaB(DHA)); Table 1). The E.~coli recombinant cells carrying pSTV28::pfaB(DHA) and pEPA Δ 6 produced EPA (9.2 \pm 0.8%) and DHA (0.2 \pm 0.1%) (Table 2), indicating that pfaB(DHA) has the potential to synthesize both DHA and EPA in combination with pfaA, pfaC, pfaD, and pfaE from S. pneumatophori SCRC-2738. These EPA and DHA contents increased to 13.8 \pm 0.9% and 0.6 \pm 0.1%, respectively, in cells grown with cerulenin at 5 μ g mL⁻¹ (Table 2), which inhibits the de novo synthesis of fatty acids (Omura, 1976), but not that of EPA (Allen et~al., 1999) or DHA (Morita et~al., 2005). No

DHA was detected with the coexpression of pEPA Δ 1 and pSTV28::pfaB(DHA) and this vector combination produced EPA at approximately 13% of total fatty acids (Table 2). This content of EPA is almost the same as that of *E. coli* DH5 α recombinant carrying a single pEPA Δ 1 vector (Orikasa *et al.*, 2004), suggesting that the expression of pSTV28::pfaB(DHA) has no effect of the formation of the Pfa enzyme complex for EPA in the *E. coli* DH5 α recombinant carrying both pEPA Δ 1 and pSTV28::pfaB(DHA).

Detailed analysis of fatty acids in EPA-producing and DHA-producing bacteria

Shewanella pneumatophori SCRC-2738 (Hirota et al., 2005) and S. marinintestina IK-1 (Satomi et al., 2003) have been reported to synthesize EPA (10%–20% of total fatty acids) but not DHA. In this study, too, only EPA was detected in these two species. Conversely, it has been reported that M. marina MP-1 produces DHA but no EPA (DeLong & Yayanos, 1986; Morita et al., 2005). However, reanalysis of its fatty acids showed that this bacterium has, in addition to DHA (8.1 \pm 0.2% of total), very low EPA (0.5 \pm 0.0%) at 15 °C, and the contents of both EPA and DHA increased to 1.0 \pm 0.2% and 19.6 \pm 0.8%, respectively, in cells grown with 0.5 μ g mL⁻¹ cerulenin. These results suggest that pfaB(DHA) functions in the production of both DHA and EPA in bacteria that have inherent pfa genes for DHA synthesis. EPA was not detected in recombinant E. coli carrying either pDHA4 (Table 2) or pDHA3 and pfaE (Orikasa et al., 2006b), which would be caused by the formation of undetectable levels of EPA in these systems.

In the polyketide biosynthesis mode, a double bond is anaerobically introduced into the elongating fatty acid (Metz *et al.*, 2001). The process requires three steps: condensation of the elongating fatty acid and the C2 donor (malonyl–ACP), introduction of a double bond by dehydration, and the subsequent geometric or positional isomerization of the double bond, although no direct experimental evidence of this has ever been reported. Ootaki *et al.* (2003) speculated that

3.6.9.12.15-octadecapentaenoic acid (18:5) is one of the intermediate fatty acids, at which point the pathway branches into EPA or DHA biosynthesis (Fig. 2). One cycle of C2 elongation of 18:5 without the introduction of further double bonds would allow the production of EPA, whereas two cycles of C2 elongation of 18:5 with the introduction of a single double bond would produce DHA. The KS domain of PfaB(DHA) may be specifically involved in the condensation step from C18 to C20 or C20 to C22 in DHA biosynthesis. The subsequent introduction of double bonds and the isomerization steps could be catalyzed by both PfaC(DHA) and PfaC(EPA) (Fig. 2). As indicated previously (Okuyama et al., 2007), a slight difference in the domain structures of PfaC(DHA) and PfaC(EPA) may alter their association with the KS domain of PfaB(DHA). In PfaC(EPA), all three dehydratase domains (HDs) are homologous to FabA, a 3-hydroxydecanoyl-ACP dehydratase, whereas PfaC(DHA) has two HDs similar to FabA and one domain similar to FabZ/FabA. FabZ is a 3-hydroxyacyl-ACP dehydratase (White et al., 2005). If the association between the KS domain of PfaB(DHA) and the FabZ/FabA domain of PfaC(DHA) predominates over its association with the corresponding FabA domain of PfaC(EPA), DHA would be preferentially produced as in the combination of pDHA4 and pUC18::pfaC(EPA). The KS domain structure of PfaB is conserved in Colwellia psychrerythraea 34H, which is expected to produce DHA, but not in any of the known PfaBs of EPA-producing bacteria (Okuyama et al., 2007). The quantities of DHA are limited in native DHA-producing bacteria (DeLong & Yayanos, 1986; Hamamoto et al., 1995; Watanabe et al., 1997) and recombinants carrying their genes, compared with the quantities of EPA produced by native EPA-producing bacteria and recombinants carrying their genes (Table 2) (Orikasa et al., 2004; Satomi et al., 2003; Orikasa et al., 2007), and this must be controlled by pfa gene products other than PfaB and PfaE. In the case of PfaE, pfaE genes from EPA-producing S. pneumatophori SCRC-2738 and DHA-producing M. marina MP-1 resulted in almost the same levels of EPA (10%-12% of total fatty acids) in combined expression in E. coli DH5α containing the pEPA\(\Delta\), 2,3 vector which included four \(pfaA-pfaD\) genes derived from S. \(pneumatophori\)

SCRC-2738 (Nishida *et al.*, 2006a; Orikasa *et al.*, 2006a). Recently Jiang *et al.*, (2008) showed that the total number of ACPs in PfaA controls the levels of polyunsaturated fatty acids.

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Figure legends

Fig. 1. Organization of the five *pfa* genes for biosynthesis of EPA of *S. pneumatophori* SCRC-2738 and pDHA4 for biosynthesis of DHA of *M. marina* MP-1. The functional domain structure of individual *pfa* genes based on analysis by Okuyama *et al.* (2007) is shown in different colors. The relative direction of *pfaE* in pDHA4 has not been determined. Blank arrows are open reading frames unnecessary for EPA or DHA biosynthesis. The structure of the vector is omitted.

Fig. 2. A possible pathway for the bacterial biosynthesis of EPA and DHA. All five *pfa* gene products for EPA [Pfa(EPA)] and those for DHA [Pfa(DHA)] are involved in the biosynthesis of EPA (closed arrow and box of solid line). The KS domain of PfaB(DHA) is not necessary for the production of EPA, but is specifically required for the biosynthesis of DHA (open arrow and box of broken line). Unsaturated fatty acids tentatively considered to be intermediates are abbreviated as 4,7,10,13–16:4 (hexadecatetraenoic acid), where the number before the colon is the number of carbon atoms and the number after the colon is the number of double bonds in the fatty acid molecule. Intermediates before 4,7,10,13–16:4 were omitted. This pathway is slightly modified from that presented previously (Ootaki *et al.*, 2003).

Table 1. Strains and vectors used in this study

Strain/plasmid/cosmid	Relevant characteristics	Source or reference
Strain		
E. coli DH5α	$deoR$, $endA1$, $gyrA96$, $hsdR17(rK^-mK^+)$,	Takara Bio ^a
	$recA1\ phoA,\ relA1,\ thi-1,\ \Delta(lac\ ZYA-argF)$	
	U169 ϕ 80d lac Z Δ M15, F $^-$, λ^- , sup E44	
M. marina MP-1	Wild type	ATTC 15381
S. pneumatophori	Wild type	Hirota et al., 2005
SCRC-2738		
S. marinintestina IK-1	Wild type	Satomi et al., 2003
Plasmids/Cosmids		
pET21a:: <i>pfaE</i>	pET21a carrying <i>pfaE</i> for DHA	Orikasa et al., 2006a
pSTV29	Low-copy-number cloning vector, Cm ^r	Takara Bio
pDHA3	pSTV29 carrying <i>pfaA</i> , <i>B</i> , <i>C</i> , <i>D</i> for DHA	Orikasa et al., 2006a
pDHA4	pSTV29 carrying <i>pfaA,B,C,D,E</i> for DHA	This study
pSTV28	Low-copy-number cloning vector, Cm ^r	Takara Bio
pSTV28:: <i>pfaB</i> (DHA)	pSTV28 carrying <i>pfaB</i> for DHA	This study
pSTV28:: <i>pfaB</i> (EPA)	pSTV28 carrying <i>pfaB</i> for EPA	Yazawa et al., 1998
pSTV28:: pfaD(EPA)	pSTV28 carrying <i>pfaD</i> for EPA	This study
pUC18	High-copy-number expression vector, Amp ^r	Takara Bio
pUC18:: <i>pfaA</i> (EPA)	pUC18 carrying <i>pfaA</i> for EPA	Yazawa et al., 1998
pUC18:: <i>pfaB</i> (EPA)	pUC18 carrying <i>pfaB</i> for EPA	This study

pUC18:: <i>pfaC</i> (EPA)	pUC18 carrying <i>pfaC</i> for EPA	Yazawa et al., 1998
pUC18:: <i>pfaD</i> (EPA)	pUC18 carrying pfaD for EPA	This study
pWE15	Cosmid expression vector, Km ^r , Amp ^r	Stratagene ^c
pEPAΔ1	pWE15 carrying <i>pfaA,B,C,D,E</i> for EPA	Orikasa et al., 2004
pEPAΔ6	pWE15 carrying <i>pfaA</i> , <i>C</i> , <i>D</i> , <i>E</i> for EPA	Orikasa et al., 2006a;
		Yazawa et al., 1998

^a Takara Bio Inc., Tokyo, Japan.

^b Abbreviations of antibiotics: Km, kanamycin; Amp, ampicillin; and Cm, chloramphenicol.

^c Stratagene, La Jolla, CA, USA.

Table 2. Combined expression of *pfa* genes from *S. pneumatophori* SCRC-2738 and from *M. marina* MP-1 in *E. coli* DH5α

Combination ^a	EPA or DHA produced			
	(% of total fat			
Vector carrying a <i>pfa</i> gene(s)	Vector carrying a pfa	EDA	DIIA	(n =)
from SCRC-2738 ^b	gene(s) from MP-1 ^c	EPA	DHA	
None	pDHA4 ^d	ND ^e	3.7 ± 0.1	(5)
pUC18:: <i>pfaA</i> (EPA)	pDHA4	ND	2.1 ± 0.5	(3)
pUC18:: <i>pfaB</i> (EPA)	pDHA4	3.4 ± 0.8	0.8 ± 0.2	(5)
pUC18:: <i>pfaC</i> (EPA)	pDHA4	ND	5.0 ± 1.5	(3)
pUC18:: <i>pfaD</i> (EPA)	pDHA4	ND	3.5 ± 0.4	(3)
pEPAΔ1	pSTV28:: <i>pfaB</i> (DHA)	12.8 ± 0.2	ND	(3)
рЕРАΔ6	pSTV28::pfaB(DHA)	9.2 ± 0.8	0.2 ± 0.1	(6)
pEPAΔ6	pSTV28:: <i>pfaB</i> (DHA)	$13.8 \pm 0.9^{\rm f}$	$0.6 \pm 0.1^{\rm f}$	(3)

^a *E. coli* DH5α cells carrying pDHA4 and any pUC18 vector were cultivated in LB medium containing chloramphenicol at 30 μg mL⁻¹ and ampicillin at 50 μg mL⁻¹ at 15°C for 72 h. Cells carrying the cosmid vector (pEPAΔ6 or pEPAΔ1) and pSTV28::*pfaB*(DHA) were cultivated in LB medium containing ampicillin at 50 μg mL⁻¹ and chloramphenicol at 50 μg mL⁻¹ at 15°C for 96 h. In the combination of pEPAΔ6 and pSTV28::*pfaB*(DHA), recombinants were cultivated in the presence and in the absence of cerulenin at 5 μg mL⁻¹.

^b S. pneumatophori SCRC-2738

^c M. marina MP-1

^d Details for vectors, see Table 1.

^e ND, not detected.

 $^{^{\}mathrm{f}}$ These values are those from the recombinant cells cultivated in the presence of cerulenin.

Fig. 1

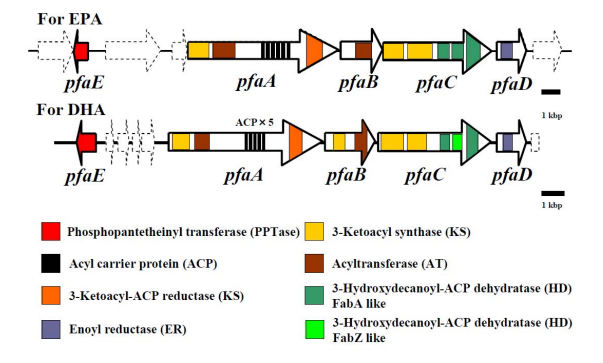


Fig. 2.

