Internal lipid synthesis and vesicle growth as a step toward self-reproduction of the minimal cell

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Abstract One of the major properties of the semi-synthetic minimal cell, as a model for early living cells, is the ability to self-reproduce itself, and the reproduction of the boundary layer or vesicle compartment is part of this process. A minimal bio-molecular mechanism based on the activity of one single enzyme, the FAS-B (Fatty Acid Synthase) Type I enzyme from *Brevibacterium ammonigenes*, is encapsulated in 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylcholine (POPC) liposomes to control lipid synthesis. Consequently molecules of palmitic acid released from the FAS catalysis, within the internal lumen, move toward the membrane compartment and become incorporated into the phospholipid bilayer. As a result the vesicle membranes change in lipid composition and liposome growth can be monitored. Here we report the first experiments showing vesicles growth by catalysis of one enzyme only that produces cell boundary from within. This is the prototype of the simplest autopoietic minimal cell.

Keywords Minimal cell · Origin of life · Self-reproduction · Synthetic biology · Vesicle

Introduction

The self-reproduction process in a cell involves both self boundary layer reproduction, which occurs by membrane replication, and core self-reproduction, that is based on genome replication. During the origin of life the initial mechanisms driving prebiotic cell formation are thought to have originated from the spontaneous self-assembly of molecules. The bilayer membrane constituting the first cell compartment may have formed originally through the self assembly of amphiphilic molecules into membrane vesicles (Hargreaves and Deamer 1978; Apel et al. 2002; Monnard and Deamer 2002). Furthermore, the prebiotic boundary layer self-reproduction process may have been the consequence of the vesicle growth, by spontaneous incorporation of lipids, until instability triggered division into smaller stable vesicles (Walde et al. 1994; Berclaz et al. 2001); other suggestive mechanisms of chemical induction of vesicle growth and division have been reported more recently by Sugawara team (Takakura et al. 2003; Takakura and Sugawara 2004). A detailed process was recently reconstructed where vesicle growth was monitored by Fluorescence Resonance Energy Transfer (FRET) assay, directly measuring the fraction of added fatty acids that become incorporated into preexisting vesicle marked with membrane localized fluorescent dyes (Hanczyc et al. 2003).

The major difficulty to propose models describing prebiotic cells, based on bio-molecules and basic metabolisms, is underlined by the absence of a historical record to suggest how the initial cell-biochemistry was organized. To overcome this problem a synthetic biology approach can be followed to build a cell model for early living cells, where extant enzymes and synthetic compartments are used to construct a semi-synthetic minimal cell in the laboratory (Luisi et al. 2006; Murtas 2007).

In this challenge the first milestone is represented by the implementation of a self-maintenance or a basic metabolism, based on protein synthesis, into a lipid or liposome compartment. Complex biochemical mechanisms or protein synthesis systems have been already reconstructed and tested in lipid vesicles (Oberholzer et al. 1995a, b; Ishikawa et al. 2004; Noireaux and Libchaber 2004). Good progress

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was recently achieved introducing protein synthesis in liposomes using a minimal set of enzymes (Murtas et al. 2007).

The introduction of a bio-mechanism to allow boundary layer self-reproduction would certainly deliver the second important milestone: a liposome that builds its own membrane with the help of entrapped enzymes. This represents a prototype of the simplest autopoietic minimal cell (Fig. 1), (Luisi 2006; Bachmann et al. 1992). Initial studies on membrane biogenesis, demonstrating an enzymatic driven membrane reconstitution, have been carried out in Deamer lab (Deamer and Boatman 1980; Deamer and Gavino 1983). The first attempts to propose an autopoietic system based on enzymes that produce cell boundary from within was set up in Luisi’s laboratory when two (Kuruma et al. 2009) or all four enzymes (Schmidli et al. 1991) of the salvage pathway were entrapped in liposomes to control lipid synthesis. Although lipid synthesis was demonstrated, unfortunately no vesicle growth was clearly proven in both cases.

In order to introduce a biological mechanism to control boundary layer self-reproduction in liposomes, biochemical synthesis of fatty acids based on a FAS type I enzyme catalysis (Schweizer and Hofmann 2004) was selected for this work. This was the mechanism of choice because fatty acid synthesis is simply controlled by only one enzyme, in agreement with the low complexity of our minimal cell model system. In particular we used the FAS-B type I enzyme from Brevibacterium ammoniagenes already biochemically characterized, expressed and purified from the heterologous system Escherichia coli (Stuible et al. 1997).

Moreover the main product of this catalysis, palmitate, is one of the surfactant synthesized in the laboratory under presumably prebiotic conditions (Walde 2006; Óró 1994; Simoneit 2004). It is chemically less reactive, therefore stable in comparison to unsaturated fatty acids such as oleate, due to the absence of double bonds between the carbon atoms of the fatty acid chains. We propose that a similar and minimal enzymatic catalysis may have occurred in vesicle compartments to control the first biological boundary layer self-reproduction process (Fig. 1).

**Results**

Palmitate synthesis by FAS type I catalysis in liposomes

The biochemical components for the synthesis of lipids are the bacterial FAS Type I enzyme FAS-B from B. ammoniagenes (Stuible et al. 1996) as the catalyst and the precursors or reagents required for the fatty acid synthesis of palmitate as shown in Fig. 2a. This FAS type I is a water soluble multifunctional enzyme consisting of a dimer with a single long multifunctional polypeptide encoded by a single c-DNA. The FAS-B protein has already been expressed in heterologous system, purified and characterized by Stuible et al. (1997) with an E. coli transformant based on the fasB gene isolated from B. ammoniagenes. The same E. coli clone was generously provided from the Schweizer team (University of Erlangen-Nürnberg, Germany) and used to express and purify the FAS-B enzyme following the methods for FAS-B purification reported by Stuible et al. (1997). Since endogenous E. coli FAS is a non aggregated type II system, based on the activity of 6 independent genes, protein contamination of the heterologous type I FAS by any of the endogenous type II FAS is considered unlikely.

After purification, an SDS–PAGE gel was loaded with aliquots of the purified FAS-B protein showing only one

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**Fig. 1** A liposome that builds its own membrane based on entrapped enzymes is the prototype of the simplest autopoietic minimal cell. 

P Precursors are the reagents required for an enzymatic catalysis, E enzyme/s controlling lipid catalysis, L lipids are released within the liposome lumen and spontaneously become incorporated into the lipid bilayer and this results in vesicle growth. The enzymatic pathway produces the palmitate represented as the dark shaded lipids.
high molecular weight band of 325 kDa and corresponding
to the expected size of the FAS-B enzyme, as reported by
Stuible et al. (1997) (data not shown). The FAS-B enzyme
activity, based on the biochemical reaction shown in
Fig. 2a, was first assayed in vitro monitoring the NADPH
oxidation, by spectrophotometric analysis, in the presence
or in the absence of the prime precursor malonyl-CoA
(Fig. 2b). The FAS activity reported in Fig. 2b was assayed
according to the method of Lynen (1980), and corre-
sponding to 0.37 \( \mu \text{mol/min/mg} \), and described in “Mate-
rials and Methods”.

Once the activity of the lipid synthesis pathway was
confirmed in vitro, the system (Fig. 2a) was introduced into
POPC liposomes. The protocol followed the FAS-B assay
conditions described in detail in “Materials and Methods”,
with the reaction incubated at 30°C for 3 h. In order to
examine the products of the FAS-B catalysis either in bulk
solution or in liposomes, the synthesis was performed with
in vitro incorporation of \(^{14}\text{C}\) radioactively labeled malonyl-
CoA, and the resulting fatty acids analyzed by Thin Layer
Chromatography (TLC). As reported in Fig. 3, the auto-
radiography of the TLC chromatogram indicates that with
the FAS-B reaction the radioactivity is exclusively incor-
porated into long chain fatty acids and palmitic acid was
produced under these conditions.

This experiment shows that the fatty acid palmitate can
be synthesized by the FAS-B enzyme activity within ves-
cicles. In particular, to demonstrate that this reaction can
take place exclusively within the liposome lumen, there-
fore in the absence of any external FAS activity, Proteinase
K was added to the outside of liposomes after palmitate
pathway entrapment. As expected, when Proteinase K is
added palmitate synthesis is weaker than the signal pro-
duced by palmitate synthesis in both the liposome lumen
and the external volume (Fig. 3). In the same experiment,
the specific activity of the FAS-B enzyme and the essential requirement of the reagents in FAS (Fig. 2a) were tested. Clearly, only in the absence of precursors such as acetyl-CoA or in the absence of the enzyme FAS-B there is no lipid synthesis, confirming the lack of any contaminant that promote palmitate synthesis. Finally, the proteolytic activity of the Proteinase K on the FAS enzyme was tested by adding Proteinase K to the FAS-B reaction in bulk solution; as a result no palmitate is detected on TLC.

Liposome growth by internal FAS type I palmitate synthesis and incorporation into membrane

We initially examined how the surfactant palmitic acid behaves in the chemical induction of vesicle membrane growth by fatty acids incorporation as spontaneous fraction of added palmitate micelles into membrane of preformed POPC vesicles. Lipid bilayer growth was monitored by FRET assay based on membrane-localized lipid fluorescent dyes (Struck et al. 1981; Malinin et al. 2001). For this purpose pre-formed vesicles were prepared with POPC and 1% of the donor fluorescent dye, the NBD-PE (N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-1,2-dihexadecanoyl-sn-glycero-3 phosphoethanolamine, triethylammonium salt), and the acceptor fluorescent dye, rhodamine DHPE (Lissamine rhodamine B 1,2 dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt) that become integrated within the bilayer. The vesicles with both donor and acceptor dyes, integrated within the bilayer, grow by incorporating palmitic acid chains and the average distance between donor and acceptor molecules increases with a consequent decrease in FRET efficiency. The FRET test was carried out by fluorometric analysis measuring the donor/acceptor fluorescence ratio \( F_{\text{don}}/F_{\text{acc}} \), normalized to the \( F_{\text{don}}/F_{\text{acc}} \) after detergent lysis and was set up on four samples of pre-formed vesicles diluted 300 times (10 ul sample diluted in 3 ml buffer), where palmitic acid was added in different concentration as reported in “Materials and Methods”. The dyes used for these experiments exchange extremely slowly between phospholipid vesicles (Hanczyc et al. 2003) and consequently even less after 300 X dilution. Therefore, a decrease of FRET efficiency in this experiment can only result from the growth of the preformed vesicle membranes due to palmitic acid incorporation (See “FRET analysis” in “Materials and Methods”). In the same way, as already demonstrated for the oleate addition and incorporation into membrane of preformed vesicles (Hanczyc et al. 2003), palmitate can induce vesicle membrane growth as shown in Fig. 4a. The preformed vesicles used for this experiment were extruded again after rehydration in order to make them more homogenous (Fig. 4b), and the FRET analysis shows an increase in \( F_{\text{don}}/F_{\text{acc}} \) ratio within the range of values reported in Fig. 4a.

To test whether the FAS Type I catalysis of palmitate synthesis within liposomes can induce vesicle growth by
fatty acid insertion to the lipid bilayer, the biochemical components of the reaction reported in Fig. 2a were entrapped in POPC liposomes. A film of POPC mixed with 1% of the donor fluorescent dye NBD-PE and the acceptor fluorescent dye DHPE was prepared in eppendorf tubes. Precursors for palmitate synthesis were mixed together with the FAS-B enzyme (Fig. 2a) and added to the lipid film for rehydration and vesicle formation. In order to prevent lipid synthesis outside of the liposomes, the Proteinase K enzyme was added externally.

The samples were kept at 30°C for 3, 5, and 24 h to allow lipid synthesis to occur within vesicles (see “Materials and Methods” : “Synthesis of F.A. in liposomes”). To test vesicle membrane growth, by lipid synthesis and insertion, a FRET assay, based on membrane-localized lipid fluorescent dyes, was set up for those samples diluted 300 times (10 ul sample diluted in 3 ml buffer). As reported for the chemical induction experiment, the FRET test was carried out by fluorometric analysis measuring the $F_{\text{don}}/F_{\text{acc}}$ ratio, normalized to the $F_{\text{don}}/F_{\text{acc}}$ after detergent lysis.

The vesicles with both donor and acceptor dyes, integrated within the bilayer, grow by incorporating palmitic acid chains soon after FAS-B catalysis and the average distance between donor and acceptor molecules increases with a consequent decrease in FRET efficiency. As shown in Fig. 4c, when FASB enzyme, together with precursors, is present within vesicles the FRET efficiency clearly decreases. On the contrary when malonyl-CoA or the FASB enzyme is omitted from the reaction mix the FRET performance is higher demonstrating that in the absence of palmitate synthesis no fatty acid molecule is inserted in the liposome membrane and no liposome growth is detected. When the vesicles used for this experiment were extruded again after the FAS reaction entrapment, in order to make them homogenous in size (Fig. 4d), the FRET analysis reported an increase in $F_{\text{don}}/F_{\text{acc}}$ ratio within the range of values reported in Fig. 4c.

The palmitate synthesis was analyzed in detail, measuring the integrated density of each radioactive signal reported on the TLC autoradiograph (Fig. 3), and estimating the ratio in between these values and the concentration of $^{14}$C radioactively labeled malonyl-CoA incorporated into the radioactive palmitate. The palmitate synthesized within liposome is limited to 30% in yield in comparison to the palmitate released in and out of liposomes (100%). As a result the palmitate produced within liposomes, after 24 h of FAS catalysis, corresponds to an estimated 2 nanomoles of palmitate released in the internal lumen of the liposomes for 200 µl of FAS catalysis assay (see “Synthesis of fatty acids in liposomes” in “Materials and Methods”), that is, on average, one molecule of palmitic acid for one thousand POPC molecules or one palmitate chain every ten molecules of fluorescent dyes present in the vesicles.

**Discussion**

In order to reconstruct a boundary layer self-reproduction process based on enzyme catalysis as a model for the first prebiotic cells, a FAS Type I catalysis was introduced in POPC vesicles. POPC are the vesicles of choice because of their stability, robustness and capability of hosting complex biochemical reactions although the vesicle forming surfactant cannot be considered prebiotic compound. As palmitate is one of the surfactant synthesized in the laboratory under presumably prebiotic conditions, palmitate vesicles could have been the lipid compartments for our experiments, on the other hand those vesicles are not easily manipulated due to their high melting temperature and stability in a limited pH range. However, it is believed that the prebiotic cell compartments were likely to be of heterogeneous lipid composition (Apel et al. 2002; Luisi 2006; Pereto et al. 2004; Walde 2006).

The FAS-B type I enzyme, from *B. ammoniagenes*, was chosen for our semi-synthetic protocell and introduced in liposomes together with the chemical precursors to produce palmitate. Although the FAS-B enzyme, as all the FAS Type I enzymes, is a multienzymes complex, characterized by a very large polypeptide chain, is the only extant enzyme that works autonomously and free in the cytoplasm. Moreover the fatty acids synthesized are released free from catalysis and ready to be integrated within the vesicle bilayer spontaneously. It is reasonable to imagine that during the origin of early cells, the boundary layer self-reproduction process may have been controlled by a simpler ancient FAS enzyme.

FRET experiments, where FAS catalysis is the only metabolism within our protocell system, have detected a surface area change of the liposome membranes due to palmitate inclusion in between POPC amphiphilic chains. Considering that Proteinase K was added to eliminate external lipid synthesis, we can presume that internal palmitate synthesis has contributed toward liposome growth. In fact, the only source of palmitate is represented by the FAS-B activity within liposomes. On the other hand the FAS-B activity is the only one responsible for palmitate synthesis, because no synthesis is detected if FAS-B enzyme is absent or degraded by Proteinase K (Fig. 3) and if any of the chemical precursors, such as malonyl-CoA or acetyl-CoA (Figs. 3, 4c), are missing.

In order to measure the vesicle growth we examined the liposome sizes by dynamic light scattering (DLS), before and after FAS catalysis and palmitate incorporation, but a change in size is not detectable or too small to be detected.
(data not shown). This result is not surprising if we consider the small amount of palmitate synthesized within liposomes. We therefore argue that while a liposome surface area change, induced by FAS synthesis, is detectable by a sensitive FRET assay, the change in vesicle size is too small to be detected in aggregate by DLS.

In particular, the FAS-B enzyme has a very short lifetime (around 3 h as shown in Fig. 2b) likely due to oxidation mechanisms and, as experienced for many enzymatic catalysis and/or cell-free protein synthesis, activity stops after a few hours because of energy and nutrient consumption (Kawarasaki et al. 1994; Noireaux et al. 2003; Noireaux and Libchaber 2004). The synthesis of palmitate is little within liposomes in comparison to the FAS-B synthesis in solution (Fig. 3) and this is in part due to the reduced inner water pools produced during liposome preparation (lipid aggregates are formed), and to the Proteinase K inhibitory activity in the outer water pools. Therefore, the resulting lipid membrane growth by FAS-B catalysis is definitely reduced in comparison to the vesicle growth reported by the FRET analysis after palmitic acid chemical induction (Fig. 4a, c). During the chemical induction process the palmitic acid is released onto preformed POPC vesicles at a temperature above the melting point of palmitate (63–64°C), where the palmitic acid synthesized by FAS-B within the liposome lumen is released at 30°C and this may cause molecular aggregation and affect the efficiency of lipid incorporation into membrane.

In order to overcome some of these limitations and demonstrate a complete process of boundary layer self-reproduction, a trans-membrane/channel pore system should be introduced in the liposome lipid bilayer. As an example the α-hemolysin toxin is a selective permeability system that allows exchange of nutrients, energy and anti-oxidants and can be used as a trans-membrane protein channel in lipid vesicles. This type of pore can create a selective portal for nutrients, energy and anti-oxidants and is capable of boosting protein expression up to 4 days through repeated feeding of the system externally (Noireaux and Libchaber 2004).

For the first time we monitor liposome vesicle growth based on the entrapment and activity of only one enzyme capable of building its own membrane from within boundary. This is a prototype of the simplest autopoietic minimal cell (Fig. 1). (Luisi 2006).

Materials and methods

Materials

Palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylcholine (POPC) was purchased from Avanti Pola Lipids (Canada). The two membrane localized fluorescent dyes used for the FRET analysis, the Lissamine rhodamine B 1,2 dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (rhodamine DHPE) and the N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (NBD-PE) were purchased from invitrogen. The E. coli strain DH5α (Gibco) was used in this study for the FAS-B heterologous expression. The plasmid pGM44 is a pBlueScript derivative containing the fasB gene together with a 0.8 kb of both its 5′-flanking and 3′-flanking DNA sequences and was kindly provided by Eckhart Shweizer, Universitat Erlangen, Nurnberg, Germany. The B. ammoniagenes DNA inserted into this vector is limited by Clal and NotI restriction sites. For heterologous expression of the B. ammoniagenes FAS proteins, the E. coli DH5α transformants were grown at 30°C in a medium containing 20 g/l casamino acids, 10 g/l yeast extract and 10 g/l NaCl, together with 50 mg/l ampicillin at pH 7.5.

DNA manipulation techniques

Transformants were routinely checked by pGM44 plasmid isolation and restriction digestion based on its restriction map. Plasmid isolation was carried out based on the method of Birnboim and Doly (1979). The DNA manipulation methods were performed as described elsewhere (Sambrook et al. 1989).

FAS-B enzyme purification and assay

Escherichia coli heterologously expressed B. ammoniagenes FAS-B enzyme was purified according to the procedure described by Stuible et al. (1997). FAS-B enzyme assay, FAS activity was assayed according to the method of Lynen (1980) with modifications (Stuible et al. 1997): NADPH disappearance, oxidation, was measured spectrophotometrically and carried out in a total volume of 1 ml containing 0.4 M potassium phosphate at 25°C, pH 7.3, 3 mM dithiothreitol, 0.5 mg Fatty acid free bovine serum albumin (SIGMA), 700 μM NADPH, 200 μM malonyl-CoA (Sigma) and 150 μM acetyl-CoA (Sigma) adding 2 μg of purified enzyme FAS-B. The absorbance of samples was recorded every 1-h interval at 340 and 600 nm and the values reported on the graph equal the delta between the values at 340 and 600 nm.

Vesicle preparation

Liposomes were prepared by the dehydration and rehydration method as described by Yomo and coworkers (Ishikawa et al. 2004). Briefly, POPC liposomes were prepared by hydrating a thin POPC film with ultra-pure...
water (Milli-Q, Millipore), followed by extrusion (10 passages) through two stacked 400 nm polycarbonate membranes (Nucleopore Track-Etch Membrane, Whatman) and freeze-drying. The resultant freeze-dried liposomes were therefore hydrated with Bicine buffer (200 mM pH 8.5) to prepare 100 mM POPC preformed vesicles. The vesicles for FRET analysis were modified by dissolving in chloroform two membrane localized fluorescent dyes, rhodamine DHPE and NBD-PE, together with freeze dried POPC liposomes in a 1% ratio (fluorescent dyes/POPC) and dried under vacuum centrifuge for 2 h. The same FRET experiments were carried out with freeze dried POPC liposomes and a 1% ratio (fluorescent dyes/POPC) have been repeated with the vesicles extruded through 0.2 μm filters (four times) after rehydration with Bicine buffer or after entrapment of the 200 μl FAS-B catalysis solution at 4°C.

Palmitate supplied as micelles on preformed POPC vesicles

Palmitic acid 20 mM was dissolved in a solution of 20 mM NaOH to form sodium palmitate at 60°C for 15 min. Palmitate was added on POPC preformed vesicles, marked with membrane localized fluorescent dyes (see vesicle preparation), in a set of four samples at varying concentration. Each sample was characterized by 3 ml of buffered solution (200 mM bicine pH 8.5) where the preformed vesicles were diluted 300 times resulting in 0.3 mM POPC vesicles (10 μl of the preformed vesicles in 3 ml, final volume, of Bicine buffer). Palmitate was added soon after melting treatment at 60°C in a four set of samples as 0, 100, 200, 300 and 400 μl of 20 mM palmitic acid. The sample were analysed 30 min after palmitate addition.

Dynamic light scattering (DLS)

Measurements were carried out with an ALV home-assembled light scattering photometer made of a 25 mW He–Ne laser (Model 127, Spectra-Physics Lasers, Mountain View, Canada), an ALV DLS/SL5000 Compact Goniometer System (ALV, Langen, Germany), two SPCM-AQR avalanche photodiodes (PerkinElmer Optoelectronics, Vaudreuil, Canada) and an ALV-5000 Multiple-tau Digital Correlator (ALV, Langen, Germany). The cylindrical scattering cells were immersed in a fuzzy-thermostated decaline bath (ALV), which was kept at 25.0°C. All experiments were performed at the scattering angle of 90°; other settings were: solvent viscosity 0.899 MPas, solvent refractive index 1.330. Great care was taken at every step to avoid the presence of dust in the liposome preparations. Liposomes (after the incubation period) were diluted (tenfold) in isotonic buffer and measured without any pretreatment. Particle-size distribution was computed by using two different algorithms (ILT and CONTIN), which gave similar results. Number-weighted size distributions were calculated within the Rayleigh–Gans–Debye approximation.

Synthesis of fatty acids in vitro and in liposomes

The FAS-B catalysis (in vitro) as reported in Fig. 2a was set up in 200 μl solution with the following stochiometry: 0.4 M potassium phosphate pH 7.3, 3 mM dithiothreitol, 0.5 mg Fatty acid free bovine serum albumin (SIGMA), 700 μM NADPH, 200 μM malonyl-CoA (Sigma), 150 μM acetyl-CoA (Sigma) adding 2 μg of purified enzyme FAS-B. When the FAS-B catalysis was entrapped in liposomes, the 200 ul of the FAS-B stoichiometry was resuspended in eppendorfs with the POPC freeze-dried liposomes and the liposomes formed by vortexing and pipetting for a few seconds to form 25 mM liposomes. For FRET analysis, the same method was used to encapsulate a FAS-B reaction but using freeze dried POPC liposomes modified with fluorescent dyes as prepared and reported in vesicle preparation. Proteinase K (5 μl, 5 mg/ml) was added soon after liposome entrapment to prevent FAS-B activity at the outside of ready formed vesicles unless explicitly stated. The reactions were in all cases incubated at 30°C for 3 h.

TLC experiment: Synthesis of 14C-labeled fatty acids was performed as reported for the FASB catalysis in vitro with the following modifications: 0.5 μCi of [2-14C] malonyl-CoA (Amersham) was added to the radioactive malonyl-CoA. The fatty acyl-CoA derivatives synthesized were hydrolysed using 10 μl 5 M NaOH at 50°C for 3 h followed by the addition of 15 μl 6 M H2SO4. When synthesis of 14C-labeled fatty acids was performed in liposomes the same conditions were applied as described above entrapping the solution reaction in 25 mM POPC vesicles and, after incubation 3 h at 30°C, the liposomes were broken by freeze and thaw, 2 min at −70°C three times. After addition of 10 μg of palmitic acid the free fatty acids were extracted twice with 80 μl ethyl acetate. The ethyl acetate extracts were dried under vacuum and resuspended in 50 μl of acetone. For fatty acid separation Silica-gel 60 thin layer plates were used. The mobile phase used for fatty acids separation was a 90:24:1 (by vol.) mixture of toluene, dioxane and acetic acid. The TLC plates were spotted on one edge with 10 μl out of 50 ul of the acetone resuspension for each sample, dried and the same edge exposed onto the mobile phase soaking only 1 mm of the silica-gel side. The chromatogram was exposed to Kodak BioMax XAR Film O/N (−70°C) and the film developed the day after.
FRET analysis

About 10 µl out of 200 µl FAS-B catalysis in liposomes were diluted 300 times in a 3 ml final volume of 0.4 M potassium phosphate buffer pH 7.3 and analyzed. Fluorescence measurements were made on a Jasco.

Spectrofluorometer FP-6200 Fluorescence resonance energy transfer (FRET) experiments were done using NBD-PE as the donor and rhodamine DHPE as the acceptor using an excitation wavelength of 430 nm, and emission wavelength of 530 nm for NBD-PE and 586 for DHPE. Fluorescence intensities for the donor and the acceptor dyes were normalized to the fluorescence intensity of DHPE. Fluorescence intensities for the donor and the acceptor using an excitation wavelength of 430 nm, and emission wavelength of 530 nm for NBD-PE and 586 for DHPE. Fluorescence intensities for the donor and the acceptor dyes were normalized to the fluorescence intensities after sample lysis with 1% triton X-100. All measurements were taken at sample turbidities of <0.1, and dye absorbance of <0.1 to minimize multiple scattering and re-absorption effects. All FRET experiments reported here were performed in triplicate.

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Conflict of interest statement The authors declare that they have no conflict of interest.

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