Identification and Characterization of Major Lipid Particle Proteins of the Yeast Saccharomyces cerevisiae

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Lipid particles of the yeast Saccharomyces cerevisiae were isolated at high purity, and their proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis. Major lipid particle proteins were identified by mass spectrometric analysis, and the corresponding open reading frames (ORFs) were deduced. In silico analysis revealed that all lipid particle proteins contain several hydrophobic domains but none or only few (hypothetical) transmembrane spanning regions. All lipid particle proteins identified by function so far, such as Erglp, Erglp, and Erglp (ergosterol biosynthesis) and Faalp, Faa4p, and Fatlp (fatty acid metabolism), are involved in lipid metabolism. Based on sequence homology, another group of three lipid particle proteins may be involved in lipid degradation. To examine whether lipid particle proteins of unknown function are also involved in lipid synthesis, mutants with deletions of the respective ORFs were constructed and subjected to systematic lipid analysis. Deletion of YDL193w resulted in a lethal phenotype, which could not be suppressed by supplementation with ergosterol or fatty acids. Other deletion mutants were viable under standard conditions. Strains with YBR177c, YMR313c, and YKL140w deleted exhibited phospholipid and/or neutral lipid patterns that were different from the wild-type strain and thus may be further candidate ORFs involved in yeast lipid metabolism.

All types of eukaryotic cells, such as plants (13, 20, 39), mammals (6), and yeasts (8), contain intracellular lipid particles. These particles consist of a highly hydrophobic core formed from neutral lipids (triacylglycerols and steryl esters) surrounded by a phospholipid monolayer in which only a few proteins are embedded. The structure of this cellular compartment is reminiscent of lipoproteins in mammals (24).

Principal studies of lipid particles from the yeast Saccharomyces cerevisiae were carried out by Clausen et al. (8). Triacylglycerols and steryl esters were identified as the main components (approximately 50% each), and it was suggested that lipid particle functions as a storage for components needed for membrane formation. Under sterol depletion, steryl esters of lipid particles are mobilized, and sterols set free through this process are incorporated into cellular membranes (27). Under conditions of fatty acid deficiency, fatty acyl moieties of triacylglycerols and steryl esters can be incorporated into phospholipids (9).

The protein pattern of yeast lipid particles is rather simple. Biochemical studies led to the identification of some of these proteins. One of the major proteins of yeast lipid particles is sterol-Δ24-methyltransferase (Erglp) (27). Localization studies of this protein revealed a 700- to 800-fold enrichment in lipid particles over the homogenate (27). Subsequently, another major lipid particle protein was identified as squalene epoxidase (Erglp) (28). It was demonstrated that Erglp is not exclusively localized to lipid particles but is also present in the endoplasmic reticulum, thus pointing to a relationship between these two compartments. Similar observations were made with Slclp, a 1-acylglycerol-3-phosphate acyltransferase involved in the biosynthesis of phosphatidic acid (3, 34). Slclp was identified as a component of lipid particles by two-dimensional electrophoresis and functional analysis by using an slcl deletion strain. In addition, a glycerol-3-phosphate acyltransferase activity catalyzed by the hypothetical Gatlp was detected in lipid particles (3, 8, 43). (The nomenclature GAT1 is used in this paper for the gene encoding the putative major glycerol-3-phosphate acyltransferase of the yeast. The gene is not identical to GAT1 [also referred to as NIL1; ORF YFL021w] listed in the Proteome Database, encoding a zinc finger transcription factor that plays a supplemental role to Gln3p, which activates genes needed to use nonpreferred nitrogen sources.) The gene encoding this protein and the polypeptide itself, however, have not yet been identified.

In the present paper, we report the identification of the major yeast lipid particle proteins by systematic mass spectrometric analysis. This strategy allowed us to classify some proteins of known function as lipid particle components and to identify additional unassigned open reading frames (ORFs) which code for novel lipid particle proteins. Phenotypic analysis of strains with deletions of the respective ORFs is described, and common features of lipid particle proteins are discussed.

MATERIALS AND METHODS

Strains and culture conditions. The haploid wild-type yeast strains S. cerevisiae X2180-1A (MATa SUC2 tet1 gal2 CUl1) and FY1679 (MATa ura3-52 trplΔΔ3 len2Δ1 hisΔ200) and the diploid wild-type strain FY1679 (MATa/α ura3-52/ura3-52 len2Δ1/LEU2 hisΔ200/HIS3 trplΔΔ3/TRPI GAL2/GAL3) were used throughout this study.

Cells were grown aerobically in 2-liter Erlenmeyer flasks to the late logarithmic phase at 30°C in YPD medium (1% yeast extract [Oxoid], 2% peptone [Oxoid], 2% glucose [Merck, Darmstadt, Germany]). Five hundred milliliters of culture medium were inoculated with 0.3 ml of a preculture grown aerobically for 48 h. Growth was monitored by measuring the optical density at 600 nm.

Construction of deletion strains. A dominant resistance marker module, kanMX4, containing the coding sequence of the Kanr gene of the Escherichia coli transposon Tn903 on vector pFA6a (38) was used to replace yeast ORFs. The Kanr gene encodes an aminoglycoside phosphotransferase activity (35) which renders S. cerevisiae resistant to the drug geneticin (G418) (21). A replacement

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The underlined sequences are homologous to the kanMX4 gene. S1, sequence 1; S2, sequence 2 were used for the deletion of ORFs by the short flanking homology method.

(pH 8.85), 25 mM KCl, 5 mM (NH4)2SO4 with 2 mM MgSO4, 0.2 mM (each) nucleotide primers with 70 nucleotides homologous to the target locus at the 5' end followed by 18 to 19 nucleotides homologous to pFA6a-kanMX4 were used for correct integration of the respective deletion cassette.

A 1.65-kb PCR fragment was generated with oligonucleotides with 70 nucleotides homologous to the target locus at the 5' end followed by 18 to 19 nucleotides homologous to pFA6a-kanMX4 were used for correct integration of the respective deletion cassette.

In diploid yeast transformants, the correct integration of the marker resulted in the appearance of one PCR fragment characteristic for the wild-type allele and one fragment characteristic for the mutated allele.

Diploid yeast transformants were sporulated in liquid medium containing 0.5% potassium acetate and 0.02% sodium benzoate for 3 to 5 days. Tetrad dissection was performed on YPD plates. At least nine tetrads were dissected for each ORF and incubated at 30°C for 2 to 3 days prior to phenotypic analysis.

Isolation of lipid particle proteins. Highly purified yeast lipid particles with an enrichment factor of 700 to 800 for triacylglycerols, sterol esters, and Erg6p over the homogenate were prepared from cells grown to the late logarithmic phase as described by Leber et al. (37). Prior to protein analysis, the lipid particle fraction was dialyzed. Nonlipid proteins were extracted with 2 volumes of diethyl ether. The organic phase was withdrawn, residual diethyl ether was removed under a stream of nitrogen, and proteins were precipitated from the aqueous phase with trichloroacetic acid at a final concentration of 10%. The protein pellet was solubilized in 0.1% sodium dodecyl sulfate (SDS)-4% 1NaOH. The protein was quantified by the method of Lowry et al. (30) with bovine serum albumin as a standard.

SDS-polyacrylamide gel electrophoresis (SDS-PAGE) was carried out by the method of Laemmli (36a), with proteins for gel staining excised by using a standard and by visual inspection.

Deletion of eight ORFs encoding yeast lipid particle proteins. Primers were designed to construct deletion cassettes by PCR (38, 39). Deletion cassettes containing the ATG codon of the ORF to be deleted, the kanMX4 gene, and the stop codon of the ORF, thus eliminating the entire target ORF. Adjacent ORFs were not affected by this deletion procedure. All deletions were made in the FY1679 strain background. To generate marker DNA flankned by short homology regions, a pair of oligonucleotide primers with 70 nucleotides homologous to the target locus at the 5' end followed by 18 to 19 nucleotides homologous to pFA6a-kanMX4 were used (Fig. 1). A 1.65-kb PCR fragment was generated with Pwo polymerase (Boehringer, Mannheim, Germany) by using approximately 50 ng of gel-purified NotI-digested pFA6a-kanMX4 plasmid or 150 ng of plasmid as a template in a 50-μl reaction mixture. After a denaturation step for 5 min at 94°C, fragments were amplified for 30 cycles of 94°C, 30 s at 54°C, 30 s at 72°C and for 20 cycles of 30 s at 94°C, 30 s at 65°C, and 105 s at 72°C, followed by a final elongation step for 12 min at 72°C. PCR fragments were ethanol precipitated and 400 to 700 ng was used for transformation.

Diploid FY1679 yeast cells were transformed by using the high-efficiency lithium acetate transformation protocol (38). Transformants were grown in YPD at 30°C overnight and then spread on YPD plates containing 200 μg of G418 (Calbiochem, La Jolla, Calif.) per liter. After incubation for 2 to 3 days, large colonies were transferred to fresh YPD-G418 plates. Only those clones that yielded colonies were considered as positive transformants and further checked for correct integration of the respective deletion cassette. Correct replacement of the respective ORFs by the kanMX4 module in G418-resistant transformants was verified by analytical PCR with Dynazyme polymerase and whole yeast genomic DNA as a template (38). Transformants were grown on YPD at 30°C overnight and then spread on YPD plates containing 200 μg of G418 (Calbiochem, La Jolla, Calif.) per liter. After incubation for 2 to 3 days, large colonies were transferred to fresh YPD-G418 plates. Only those clones that yielded colonies were considered as positive transformants and further checked for correct integration of the respective deletion cassette. Correct replacement of the respective ORFs by the kanMX4 module in G418-resistant transformants was verified by analytical PCR with Dynazyme polymerase and whole yeast genomic DNA as a template (38).

In silico sequence analysis. Information about proteins with known and by-identification was retrieved from the Yeast Protein Database (36a), the Saccharomyces Genome Database (37a), and the Munich Information Center for Protein Sequences (37b). Chained proteins were aligned with the aid of a sample applicator (Linomat IV; CAMAG, Muttenz, Switzerland), and homology searches were done by using BLAST Search (2).

Lipid analysis. Lipids of whole yeast cells were extracted after cell disruption by the procedure of Folch et al. (11). Individual phospholipids were resolved by two-dimensional thin-layer chromatography on silica gel 60 plates (Merck) by using chloroform-methanol-water-phosphoric acid (65:45:5:5 vol/vol/vol/vol) as the second developing solvent. Phospholipids were visualized on thin-layer chromatography plates by staining with iodine vapor, scraped off the plate, and quantified by the method of Broekhuyse (5).

For the analysis of neutral lipids, extracts were applied to silica gel 60 plates with the aid of a sample applicator (Linomat IV; CAMAG, Muttenz, Switzerland), and chromatograms were developed in an ascending manner by using the solvent system light petroleum-diethyl ether-acetic acid (25:25:1 vol/vol/vol) for the first third of the total distance. Then, the plates were dried briefly and further developed to the top of the plate by using the solvent system light petroleum-diethyl ether (49:1 vol/vol). Quantification of ergosterol and ergosteryl esters was carried out by densitometric scanning at 275 nm with ergosterol as a standard. Triacylglycerols were visualized by postchromatographic staining with a chromatogram immersion device (CAMAG). Plates were dipped for 6 s into a developing reagent consisting of 0.36 g of MnCl2·4H2O, 4H2O, 60 ml of water, 60 ml of methanol, and 4 ml of concentrated sulfuric acid, briefly dried, and heated at 100°C for 30 min. Quantification of acylglycerols was carried out by densitometric scanning at 400 nm with triolein (NuChek, Inc., Elysian, Maine) as a standard.

Individual sterols were identified after alkaline hydrolysis (29) of the lipid extract by gas liquid chromatography (GLC). GLC was performed on a Hewlett-Packard 5890 equipped with a flame ionization detector equipped with a capillary column (Hewlett-Packard 5; 30 m by 0.32 mm by 0.25 μm film thickness). After a 1-min hold at 150°C, the temperature was increased to 310°C.
FIG. 2. Protein pattern of lipid particles. Lipid particle proteins (LP) were separated by SDS-PAGE, reisolated from the gel, and subjected to mass spectrometric analysis as described in Materials and Methods. ST, standard proteins.

RESULTS

To identify proteins of yeast lipid particles, polypeptides of a highly enriched lipid particle fraction were separated by SDS-PAGE (Fig. 2), reisolated from the gel, and subjected to mass spectrometric analysis as described in Materials and Methods. This strategy led to the identification of the major yeast lipid particle proteins as summarized in Table 1. In some of the bands excised from the gel (Fig. 2), more than one protein was detected, namely Fat1p and Faa4p (the amount of Faa4p was greater than Fat1p), Erg7p and Fat1p (the amount of Erg7p was greater than Fat1p), Tgl1p and the YOR059c gene product (approximately equal amounts), Erg6p and the YDL193w gene product (equal amounts), and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and Yju3p (the amount of Yju3p was greater than GAPDH). Since the mass spectrometric method employed did not allow exact quantification of proteins identified in the same band, the molar ratio could be only roughly estimated. Some proteins or their polypeptide fragments were identified in several bands, namely Fat1p, Tgl1p, Erg6p, and the gene product of YIL124w (Fig. 2). The reason for this finding is most likely degradation of proteins during the isolation procedure. Alternatively, different posttranslationally modified forms of proteins might be present on the lipid particle surface, although such modifications could not be demonstrated by the methods used for this study.

Lipid particles contain enzymes involved in lipid metabolism. A number of lipid particle proteins identified in this study had been previously characterized by function, although in some cases the precise subcellular localization of the respective protein was questionable. Among these components, there are several enzymes involved in ergosterol biosynthesis (Erg1p, Erg6p, and Erg7p) and fatty acid activation (Faa1p, Faa4p, and Fat1p). It has to be mentioned, however, that the occurrence of all these proteins is most likely not restricted to lipid particles. For example, Erg1p and Erg6p were also found in the endoplasmic reticulum (28). Also, Fat1p was originally reported to be a fatty acid transporter (10) and assumed to be associated with the plasma membrane. Recent results (40), however, have demonstrated that Fat1p has very-long-chain fatty acid acyl coenzyme A (CoA) synthetase activity and may be a component of peroxisomes. Unspecific cosolubilization of proteins with lipid particles is unlikely because (i) the protein pattern of lipid particles can be obtained in a highly reproducible way and (ii) proteins of other subcellular compartments are not randomly associated with lipid particles. One exception may be GAPDH, whose association with lipid particles (Table 1 and Fig. 2) came as a surprise. Three different forms of GAPDH are known (32). They are encoded by the ORFs YJL052w, YJR009c, and YGR192c (Table 1). Mass spectrometry did not allow us to distinguish between these three forms, because differences in the amino acid sequence of the isoforms are only minor. The amount of GAPDH in lipid particles was very low, and Western blot analysis with monospecific antibodies failed to detect the protein in this fraction (data not shown). Thus, it is likely that the presence of GAPDH, which is a cytosolic enzyme, is due to unspecific interaction with the surface of lipid particles. A tendency of GAPDH to associate with other subcellular membranes in a rather unspecific way has been reported before (42).

Characterization of mutants with deletions of ORFs encoding lipid particle proteins with unknown functions. The fact that almost all lipid particle proteins of known function are enzymes of lipid biosynthetic pathways led us to speculate that the other hypothetical proteins of lipid particles might also be involved in lipid metabolism. To obtain some general information about these uncharacterized proteins, strains with deletions of the respective ORFs were constructed and subjected to phenotypic analysis. Lack of the hypothetical proteins resulted in only one case in lethality (YDL193w), whereas all other mutants were viable under standard conditions (Table 2). The YDL193w deletion strain could not be rescued by supplementation with ergosterol or long-chain fatty acids under aerobic or anaerobic conditions (data not shown). Thus, involvement of this gene in sterol or fatty acid biosynthesis appears unlikely. Since the YDL193w gene product shows weak homology to Ca²⁺ channel proteins in different organisms (Table 1), complementation of the deletion defect by addition of various ions to the culture medium was tested. Supplementation of the mutant with neither Ca²⁺ nor other divalent or monovalent cations restored growth. The other deletion strains were viable and grew like the wild type on YPD medium at 15, 30, and 37°C. The only exception was the mutant with a deletion of YBR177c, which exhibited a slight temperature sensitivity at 37°C but grew better than the wild type at 15°C.

Haploid strains with deletions of unassigned ORFs were subjected to systematic lipid analysis. In the mutant with a deletion of YBR177c, the phospholipid pattern was slightly but significantly different from that of the wild-type strain. In this mutant, cellular levels of phosphatidylinositol (PtIns) and phosphatidic acid were increased at the expense of phosphatidylethanolamine and phosphatidyltrimethylethanolamine (Table 3). The strain with a deletion of YMR313c contained significantly higher amounts of triacylglycerols and the strains...
with deletions of YBR177c and YKL140w had higher amounts of ergosterol esters than the wild type (Table 4). The pattern of individual sterols (Table 5) was similar in all deletion strains, with the exception of the mutant having a deletion of YBR177c. This strain contained a smaller amount of fecosterol and a higher level of lanosterol than the wild type. Phospholipid, sterol, and neutral lipid compositions of the other deletion strains were like those of the wild type. The fatty acid composition of all deletion strains was similar to that of FY1679 (data not shown). In summary, these results suggest that the gene products of YBR177c, YMR313c, and YKL140w are candidates for involvement in lipid metabolism through secondary effects. Since we did not observe all-or-nothing effects, it is unlikely that major enzymes of lipid metabolism are directly affected by the respective mutations.

Most lipid particle proteins lack transmembrane spanning domains. Computer analysis of functionally characterized and uncharacterized yeast lipid particle proteins did not unveil obvious common motifs that could be regarded as targeting sequences of these polypeptides to lipid particles. The observation, however, that most lipid particle proteins of yeast lack transmembrane spanning (TM) domains or contain only one TM domain (Table 2) deserves our attention. The only exceptions are Fat1p and Erg1p, with two or three putative TM segments. Preferential association of proteins lacking TM domains with lipid particles is most likely because the surface membrane of lipid particles is a phospholipid monolayer (27). Thus, proteins containing hydrophobic stretches different from TM domains may preferentially interact with the surface of lipid particles.

DISCUSSION

Mass spectrometric analysis was used to identify major proteins of lipid particles of the yeast S. cerevisiae. This approach supported previous findings concerning the localization of Erg6p and Erg1p to lipid particles (27, 28). In addition, several proteins with known function could be attributed to the lipid particle fraction during this study, namely Erg7p, Faa1p, Faa4p, and Fat1p (4, 10, 22, 23, 40). Furthermore, some novel gene products encoded by unassigned ORFs were identified as lipid particle components.
TABLE 2. Characterization of gene products present in yeast lipid particles

<table>
<thead>
<tr>
<th>No.</th>
<th>Gene or ORF designation</th>
<th>Gene product</th>
<th>TM</th>
<th>Hydrophobicity domain</th>
<th>Deletion strain phenotype</th>
<th>Deletion strain phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>FAA1</td>
<td>1 2 (middle and C terminal)</td>
<td>1</td>
<td>2 (middle and C terminal)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>2</td>
<td>FAT1</td>
<td>3 2 (N terminal and middle)</td>
<td>1</td>
<td>2 (N terminal and middle)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>3</td>
<td>FAA4</td>
<td>0 2 (middle and C terminal)</td>
<td>1</td>
<td>2 (middle and C terminal)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>4</td>
<td>ERG7</td>
<td>0 1 (C terminal)</td>
<td>1</td>
<td>1 (C terminal)</td>
<td>Sterol auxotroph</td>
<td>Sterol auxotroph</td>
</tr>
<tr>
<td>5</td>
<td>YMR313c</td>
<td>0 1 (C terminal)</td>
<td>1</td>
<td>1 (C terminal)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>6</td>
<td>TGL1</td>
<td>1 1 (N terminal)</td>
<td>1</td>
<td>1 (N terminal)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>7</td>
<td>ERG1</td>
<td>2 3 (N and C terminal)</td>
<td>1</td>
<td>3 (N and C terminal)</td>
<td>Sterol auxotroph</td>
<td>Sterol auxotroph</td>
</tr>
<tr>
<td>8</td>
<td>YOR059c</td>
<td>0 1 (middle)</td>
<td>1</td>
<td>1 (middle)</td>
<td>Viable (temperature sensitive)</td>
<td>Viable (temperature sensitive)</td>
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<tr>
<td>9</td>
<td>YMR313c</td>
<td>0 1 (middle)</td>
<td>1</td>
<td>1 (middle)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>10</td>
<td>YIM1</td>
<td>0 0</td>
<td>0</td>
<td>0</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>11</td>
<td>ERG6</td>
<td>0 0</td>
<td>0</td>
<td>0</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>12</td>
<td>YDL193w</td>
<td>1 1 (middle)</td>
<td>1</td>
<td>1 (middle)</td>
<td>Lethal</td>
<td>Lethal</td>
</tr>
<tr>
<td>13</td>
<td>TDH1</td>
<td>0 0</td>
<td>0</td>
<td>0</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>14</td>
<td>YJU3</td>
<td>0 0</td>
<td>0</td>
<td>0</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>15</td>
<td>YIM4</td>
<td>0 2 (middle and C terminal)</td>
<td>0</td>
<td>2 (middle and C terminal)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td>16</td>
<td>YKR0456</td>
<td>1 0–1</td>
<td>1</td>
<td>0–1</td>
<td>Viable</td>
<td>Viable</td>
</tr>
<tr>
<td></td>
<td>SLC1</td>
<td>1 2 (N terminal)</td>
<td>1</td>
<td>2 (N terminal)</td>
<td>Viable</td>
<td>Viable</td>
</tr>
</tbody>
</table>

aData Hydrophobicity index from Kyte and Doolittle plots (25); the positions of the hydrophobic domains within the polypeptide are shown in parentheses.

What is the physiological role of lipid particle proteins? Since several enzymes of ergosterol synthesis are located on lipid particles, it is tempting to speculate that these proteins actively participate in cellular sterol formation. This is very likely for Erg6p (27) and Erg7p (44), which are enzymatically active in lipid particle preparations. Similarly, a 3-phosphate acetyltransferase encoded by the unidentified GAT gene and the 1-acylgllycerol-3-phosphate acyltransferase SlcIp, which contribute to phosphatidic acid biosynthesis, were previously identified as lipid particle components (3, 7, 43). It has to be mentioned, however, that SlcIp escaped detection by the mass spectrometric approach most likely due to its low abundance. None of the unassigned ORFs could be identified as GAT since lipid particles of all deletion strains tested contained wild-type levels of glycerol-3-phosphate acyltransferase activity (2a).

In contrast to the above-mentioned enzymes, it was shown that squalene epoxidase (Erglp) of isolated lipid particles is not enzymatically active in vitro, whereas Erglp present in the endoplasmic reticulum fraction exhibits enzymatic activity (28). It was argued that a component, probably a reductase, that is present in the endoplasmic reticulum but absent from lipid particles may be the missing cofactor. Interaction of the endoplasmic reticulum with lipid particles may activate Erglp of the latter compartment.

The presence of enzymatically inactive proteins on the surface of lipid particles, as described for Erglp, may also be interpreted as a regulatory phenomenon. If Erglp of lipid particles does not contribute to ergosterol synthesis in vivo, this protein might be put on hold on the surface of this compartment for a situation which requires enhancement of lipid biosynthesis. Under these conditions, enzymes could be immediately mobilized from lipid particles and translocated to their site of activation, e.g., the endoplasmic reticulum, thus providing lipids within a short time without new polypeptide synthesis. The idea of depositing proteins in lipid particles during formation of this compartment has been previously advocated by Lum and Wright (31) when studying overexpression of 3-hydroxy-3-methylglutaryl CoA reductase in Schizosaccharomyces pombe. This enzyme, which accumulated first in so-called karmellae, was deposited in lipid particles upon degradation of the former organelle.

The function of several gene products located on yeast lipid particles remains to be demonstrated. With one exception, YDL193w, a deletion of ORFs encoding lipid particle proteins affected neither cell viability nor the formation of lipid particles in a significant way. All these deletion strains contain lipid particles of normal size and physical properties as shown by microscopic inspection and isolation of the respective fractions (44). We can only speculate at present that some proteins located on the surface of lipid particles may be involved in the deposition of triacylglycerols and/or steryl esters in or mobilization of these lipids from this compartment. The gene product of YMR313c may be a candidate for such a function because the strain having a deletion of this ORF accumulates triacylglycerols (44).

An intriguing question concerns targeting and transfer of proteins to lipid particles. Through localization studies of particles, it is tempting to speculate that some proteins to lipid particles. Through localization studies of...
Erglp (28) and of Scllp and Gatlp (3), it was demonstrated that lipid particles and the endoplasmic reticulum share a certain set of proteins and thus appear to be related compartments. At present, we can only speculate about the relationship of these two subcellular fractions. The fact that (i) all lipid particle proteins characterized so far are involved in lipid metabolism and (ii) most lipid particle proteins contain either none or only a small number of TM domains may serve as the basis for the following hypothesis (Fig. 3A). Several enzymes involved in lipid synthesis may be located in specific domains of the endoplasmic reticulum. Clustering of these enzymes might cause local accumulation of newly formed lipids, especially those which are unable to integrate into a phospholipid bilayer, namely triacylglycerols and steryl esters. These neutral lipids may form microdroplets (preforms of lipid particles) between the two leaflets of the endoplasmic reticulum membrane bilayer which bud off after reaching a certain size. The presence of steryl ester synthases Are1lp and Are2lp in the endoplasmic reticulum (45) is in line with this model. Recent findings from our laboratory (3a) suggest that the endoplasmic reticulum is also the major site of triacylglycerol synthesis, indicating that both neutral lipid species of lipid particles are formed in the same compartment. It has to be noted, however, that triacylglycerol synthase activity has previously been attributed to the lipid particle fraction by Christiansen (7). The assay used by this author, however, did not allow for distinguishing between acylation of the substrate diacylglycerol and transacylation reactions.

Fluorescence microscopic evidence obtained recently in our laboratory (23a) demonstrated the appearance of small, newly formed lipid particles in proximity to the endoplasmic reticulum, thus supporting the model presented in Fig. 3A. According to the molecular shape concept (18), PtdIns-rich domains in the endoplasmic reticulum might facilitate the budding process. This hypothesis is in line with the finding that PtdIns comprises approximately 30% of total lipid particle phospho-

<table>
<thead>
<tr>
<th>ORF deleted</th>
<th>Squalene</th>
<th>Zymosterol</th>
<th>Fecosterol</th>
<th>Ergosta-5,7,4(28)-triene</th>
<th>Episterol</th>
<th>Lanosterol</th>
<th>4,4-Dimethylsterol</th>
<th>Ergosterol</th>
</tr>
</thead>
<tbody>
<tr>
<td>None (FY1679)</td>
<td>6.7</td>
<td>7.1</td>
<td>6.0</td>
<td>8.2</td>
<td>6.3</td>
<td>7.8</td>
<td>1.6</td>
<td>56.3</td>
</tr>
<tr>
<td>YBR177c</td>
<td>7.0</td>
<td>8.0</td>
<td>2.2</td>
<td>8.2</td>
<td>6.5</td>
<td>10.2</td>
<td>1.0</td>
<td>56.9</td>
</tr>
<tr>
<td>YKL124w</td>
<td>5.9</td>
<td>8.3</td>
<td>5.5</td>
<td>9.7</td>
<td>4.4</td>
<td>6.9</td>
<td>1.3</td>
<td>57.7</td>
</tr>
<tr>
<td>YKL094w</td>
<td>6.1</td>
<td>8.1</td>
<td>5.9</td>
<td>8.0</td>
<td>6.0</td>
<td>7.4</td>
<td>1.9</td>
<td>56.6</td>
</tr>
<tr>
<td>YKR046c</td>
<td>7.1</td>
<td>7.1</td>
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* Mean values, with a standard deviation of ±3%, are from three independent experiments.

FIG. 3. Hypothetical models describing targeting and transfer of proteins to lipid particles. (A) Budding of a neutral lipid-rich domain from the endoplasmic reticulum is accompanied by association of proteins with the phospholipid monolayer of newly formed lipid particles. According to this model, polypeptides without TM regions would preferentially associate with lipid particles. Nasc-LP, nascent lipid particles. (B) Association of proteins with preexisting lipid particles though a targeting signal on the polypeptide. (C) Transport of proteins to preexisting lipid particles by vesicle (V) flow. LP, lipid particles; ER, endoplasmic reticulum.
lipids (27), whereas PtdIns is only a minor component among endoplasmic reticulum bulk phospholipids. During the budding process, newly formed lipid particles may be wrapped by an endoplasmic reticulum-derived phospholipid monolayer, which indeed forms the surface membrane of lipid particles (27). Proteins with none or only a low number of TM domains initially present in the endoplasmic reticulum may remain associated with the phospholipid monolayer of lipid particles during the budding process, whereas proteins with typical TM regions may be largely excluded.

Although the above-mentioned hypothesis of lipid particle biosynthesis is consistent with experimental evidence obtained during our studies and compatible with the theory of oil body formation in plants (16), alternative possibilities of lipid particle formation should be considered. As an example, proteins could be directed to the surface of preformed lipid particles by a targeting signal (Fig. 3B). Although no such typical motifs were found in lipid particle proteins, signals based on conformational properties that escaped our attention may be important in that respect. As a further possible mechanism for assembly of proteins to lipid particles, transport of proteins through vesicle flux might be considered (Fig. 3C). Experimental evidence for such a mechanism, however, is also missing.

The protein encoded by YBR177c, which is slightly homologous to a probable human membrane receptor (HPS1) (Table 1), might be regarded as a candidate for facilitating such a vesicle docking process. The fact that the mutant with a deletion of YBR177c contains lipid particles with a slightly different protein pattern than the wild type could be an argument for this hypothesis. A detailed analysis of this mutant and characterization of the YBR177c gene product will be required to address this question.

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