C Terminus of Nce102 Determines the Structure and Function of Microdomains in the Saccharomyces cerevisiae Plasma Membrane†

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Received 8 January 2010/Accepted 21 June 2010

The plasma membrane of the yeast Saccharomyces cerevisiae contains stably distributed lateral domains of specific composition and structure, termed MCC (membrane compartment of arginine permease Can1). Accumulation of Can1 and other specific proton symporters within MCC is known to regulate the turnover of these transporters and is controlled by the presence of another MCC protein, Nce102. We show that in an NCE102 deletion strain the function of Nce102 in directing the specific permeases into MCC can be complemented by overexpression of the NCE102 close homolog FHN1 (the previously uncharacterized YGR131W) as well as by distant Schizosaccharomyces pombe homolog fhn1 (SPBC1685.13). We conclude that this mechanism of plasma membrane organization is conserved through the phylum Ascomycota. We used a hemagglutinin (HA)/Suc2/His4C reporter to determine the membrane topology of Nce102. In contrast to predictions, its N and C termini are oriented toward the cytosol. Deletion of the C terminus or even of its last 6 amino acids does not disturb protein trafficking, but it seriously affects the formation of MCC. We show that the C-terminal part of the Nce102 protein is necessary for localization of both Nce102 itself and Can1 to MCC and also for the formation of furrow-like membrane invaginations, the characteristic ultrastructural feature of MCC domains.

Stable lateral domains coexist within the plasma membrane of the yeast Saccharomyces cerevisiae. Nce102, a protein originally thought to be involved in nonclassical export (6) and more recently in sensing sphingolipids (10), is the major organizer of one type of these domains, termed MCC (membrane compartment of Can1) (25). MCC consists of evenly distributed, isolated patches enriched in sterols and specific proteins (15, 16, 25, 26). We showed that MCC-specific proton symporters accumulate in these patches in a reversible, membrane potential-dependent manner. This Nce102-mediated transient MCC accumulation plays a key role in the turnover of the transporters (16). Each MCC patch is accompanied by an eisosome, a cytosolic complex located directly beneath the membrane (36).

In an early freeze-etching study, Moor and Mühlethaler (28) demonstrated that the yeast plasma membrane contains numerous furrow-like invaginations. Recently, MCC patches were identified with these plasma membrane structures, and Nce102 was shown to be necessary for furrow formation. On the ultrastructural level, the MCC patches of nce102Δ cells appeared as flat, smooth, elongated areas within an otherwise particle-rich plasma membrane (32).

There is now increasing evidence that cytosolic Pil1, a primary component of eisosomes, is a prerequisite for MCC patch formation. It marks the sites where Nce102 and the MCC-specific transporters will subsequently accumulate (16, 23, 29). Data published so far do not indicate a direct involvement of cytoskeletal components in this process (26). Accordingly, markers of classical endocytosis, which are coupled to the cortical patches of actin, were localized outside the MCC (16). In this paper we examine the contribution of Nce102 to the organization of MCC patches and of furrow-like invaginations. Our results indicate that, in contrast to the prediction of four transmembrane domains (TMDs), the Nce102 molecule might span the plasma membrane only twice, the C and N termini being oriented toward the cytoplasm. We find that the C-terminal 6 amino acids of Nce102 are essential for MCC patch formation as well as for the formation of the furrow-like membrane invaginations. In addition it is shown that this Nce102 function is phylogenetically conserved among Ascomycota.

MATERIALS AND METHODS

Yeast strains and growth conditions. Yeast strains used in this study are listed in Table 1 (4, 15, 19, 33). If not stated otherwise, cells were grown in a rich medium (YPD [1% yeast extract, 2% peptone, 2% glucose]) at 30°C on a shaker. Synthetic defined (SD) medium contained 0.67% Difco yeast nitrogen base without amino acids and 2% glucose supplemented with, depending on which marker was used to select for transformed cells, uracil and adenine (both 20 μg/ml). All yeast transformations were carried out by the high-efficiency method described by Gietz and Woods (14).

Construction of overexpression and tagging plasmids. S. cerevisiae genes NCE102 and FHN1 (YGR131W) and S. pombe gene fhn1 (SPBC1685.13) were amplified by PCR from genomic DNA using the following primers (F and R for each of the three genes): NCE102-F (ATATAAGCTTATAATGCTAGC), NCE102-R (ATTACTCGAGACTTGGGAAATGGTT), FHN1-F (A

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1 Published ahead of print on 25 June 2010.
NCE102 forms membrane domains

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**TABLE 1. Yeast strains used in this study**

**RESULTS**

Nce102 homologs share Nce102 function in plasma membrane organization. Phylogenetically, furrow-like invaginations of the yeast plasma membrane represent a highly conserved structure (reviewed by Stradalova et al. [32]). Proteins sharing a significant homology with Nce102, a protein shown to be necessary for the final step of the formation of furrows as well as of MCC patches in *S. cerevisiae*, are found in various species of *Ascomycota*. BLAST (Basic Local Alignment Search Tool) analysis (2) reveals more than 40 different Nce102 orthologs. We tested whether the molecular function of promoting the localization of specific transporters in the MCC/furrows is conserved among these Nce102-like proteins.

In the genome of *S. cerevisiae*, open reading frame (ORF) YGR131W codes for one of the closest Nce102 homologs (Fig. 1A). Ygr131w shows 55% identity (68% similarity) to Nce102, and its overexpression complements the Nce102 deletion. In the plasma membrane of the nce102Δ strain, Can1-GFP is distributed homogeneously (16). When overexpressed in nce102Δ cells, Ygr131w-mRFP not only accumulates in MCC patches (16) but also induces the accumulation of Can1-GFP in this plasma membrane compartment. Due to the ability of YGR131W to function in this Nce102 function, we named the gene *FHN1* (functional homolog of NCE102) (Fig. 1B to D). Since *FHN1* is not able to compensate for the NCE102 dele-
tion when expressed under the control of its own promoter, we conclude that its inherent expression level is not sufficient.

The replacement of Nce102 function in the plasma membrane organization could also be observed when a heterologous NCE102-like gene of Schizosaccharomyces pombe, the uncharacterized ORF SPBC1685.13 (31% identity and 52% similarity to *S. cerevisiae* NCE102) (Fig. 1A) was expressed in the *S. cerevisiae* nce102Δ mutant (Fig. 1E). Similarly to Fhn1, the gene product of SPBC1685.13 was targeted to MCC and induced significant accumulation of Can1-GFP in this compartment. Therefore, we named the gene *fhn1* (functional homolog of *S. cerevisiae* NCE102). Even though the primary structure of Fhn1 shows a relatively low degree of homology with Nce102 (see alignment of NCE102, FHN1, and *fhn1*), it complements the Nce102 deletion. We concluded therefore that the function of Nce102 in plasma membrane organization is probably widely conserved throughout *Ascomycota*.

**Membrane topology of Nce102.** Large conserved parts of Nce102 correspond to highly hydrophobic regions of the protein molecule. According to hydropathy analysis tools (e.g., TMHMM2.0; http://www.cbs.dtu.dk/services/TMHMM-2.0/) (21), the protein is predicted to possess 4 transmembrane helices, with N and C termini oriented toward the lumen of the endoplasmic reticulum (ER) during protein synthesis (Fig. 2A, model I) and outside the cell once it reaches the plasma membrane.

To determine the topology of Nce102 experimentally, we employed a technology of a hemagglutinin (HA)/Suc2/His4C chimeric protein tag as a topology reporter (7, 18), a technique widely used for membrane protein topology determination. We constructed vectors coding for Nce102 C-terminally truncated after the 1st, 2nd, 3rd, or 4th predicted transmembrane domain (at amino acid L28, E63, R94, or I146, respectively) (Fig. 2A, model I). The full-length Nce102 and the individual truncated versions described above were C-terminally tagged with HA/Suc2/His4C and expressed in an auxotrophic his4 strain (7, 9, 18). The ability of the histidinol dehydrogenase (His4C) to convert histidinol to histidine enables a histidine-auxotrophic strain with the reporter located in the cytosol to grow on media lacking histidine but supplemented with histidinol. Invertase (Suc2) and His4C contain eight and four, respectively, consensus acceptor sites for N-linked glycosylation that could be glycosylated only if the reporter is translocated to the lumen of the endoplasmic reticulum. The HA tag is included to allow identification of the expressed fusion proteins by Western blotting.

Anti-HA antibody detected the fusion proteins in crude membranes prepared from all five strains, each expressing one of the truncated versions of NCE102 or the full-length NCE102 fused to the reporter. The shift to higher molecular masses of the fusion proteins containing amino acids 1 to 28, 1 to 63, and 1 to 94, which can be abolished by Endo H treatment, proved their glycosylation (Fig. 2C). Therefore, the C termini of these proteins was coexpressed either with an empty plasmid (B) or with vectors coding for *S. cerevisiae* Nce102-mRFP (C) and Fhn1-mRFP (D) and *S. pombe* fhn1-mRFP (E). Note the homogenous distribution of Can1-GFP in panel B and its focal appearance in panels C to E. Bar, 5 μm.
three constructs were exposed to the ER lumen during their biosynthesis, and consequently, in the plasma membrane, amino acids 28, 63, and 94 should be oriented outside the cell. No molecular mass shift or effect of Endo H treatment was detected in strains bearing the full-length (amino acids 1 to 173) version or the version consisting of amino acids 1 to 146, indicating thus that the stretch of amino acids 146 to 173 was oriented to the cytoplasm (Fig. 2C). This conclusion was also confirmed by the growth on histidinol of the histidine-auxotrophic strains bearing fusion constructs with the full-length or truncated (1 to 146) protein, indicating the cytoplasmic orientation of the C terminus (Fig. 2B).

These results indicate that the Nce102 protein contains only two transmembrane helices (Fig. 2A, model II), corresponding to the first (N-terminal) and last hydrophobic regions in the Nce102 molecule. The middle large hydrophobic region in this interpretation does not span the membrane. A precaution has to be taken is to consider the possibility that the bulky (125-kDa) topology reporter attached to this small (19-kDa) protein and its even smaller fragments could interfere with both their trafficking and folding. However, as shown in Fig. 3, at least the largest Nce102-derived chimeras are targeted properly to the plasma membrane. To minimize the danger of artificial misfolding, we further checked the Nce102 topology by inserting a shortened Suc2 fragment, D81 to Y133 or V67 to D148, into the intact protein. They were inserted after P64Q; including short additional amino acid stretches was thought to possibly prevent potential interference with the putative TMD 2. In
another construct, L70 in Nce102 was exchanged for N to generate a glycosylation site within the loop exposed to the ER lumen in both the computer-predicted topology (model I) and the Suc2/His4C-based experimentally assessed topology above (model II; in this case, however, the orientation of the whole molecule is inverted [Fig. 2A]). The proteins were immunodetected with specific anti-Nce102 antibody. ER protein Wbp1 was used as a positive control for Endo H digestion. As shown in Fig. 2D, the Endo H digestion was not accompanied by a shift in molecular weight with either of the tested proteins, while a clear shift was detected in the control (Wbp1). These results show that the Suc2 reporter was not glycosylated in the ER lumen. The putative model of 4 TMDs with N and C termini inside the cell has, therefore, not definitely been excluded, and model III of Fig. 2A has still to be considered as possibly correct.

The C terminus of Nce102 is necessary to target MCC-specific transporters into MCC. Besides the hydrophobic regions, also the C terminus is highly conserved among the Nce102 homologs. We tested the physiological significance of this highly conserved C terminus in targeting Nce102 to MCC and/or in the gathering (concentrating) of other MCC residents.

The total C-terminal part following the last predicted transmembrane helix, consisting of 27 amino acids, was removed from Nce102, and this truncated version (Nce102\(_{1-146}\)) was expressed in a wild-type background under the control of an endogenous promoter. In the resulting strain, Can1-GFP was

FIG. 3. The C-terminally truncated version of Nce102 is unable to sequester Can1 into MCC. Plasma membrane distributions of Can1-GFP in the wild type (A) and NCE102 deletion mutant (B) were compared to that in the cells expressing the C-terminally truncated versions of Nce102, Nce102\(_{1-146}\) (C) and Nce102\(_{1-167}\) (D), under the control of the natural promoter. Surface confocal sections and central confocal cross sections are presented. Bar, 5 \(\mu\)m.

FIG. 4. The C-terminally truncated Nce102 does not localize to MCC. The plasma membrane distribution of C-terminally truncated Nce102\(_{1-146}\) (A) tagged with GFP and that of the full-length Nce102-GFP (B) are shown on surface confocal sections (left) and central confocal cross sections (right). Note the absence of a characteristic MCC pattern in panel A. Bar, 5 \(\mu\)m.
distributed homogenously as in the nce102Δ background, which means that Nce102Δ1-146 did not substitute for the Can1 accumulation-related function of Nce102 (Fig. 3A to C). A deletion of the 6 terminal amino acids only (Nce102Δ1-167) resulted in an identical effect (Fig. 3D). Expression levels of the full-length protein and its C-terminally truncated proteins were mutually comparable (data not shown). As shown by localization of GFP fusion proteins, the two truncated Nce102 versions reached the plasma membrane but, in contrast to the full-length molecule, were distributed homogenously (Fig. 4). These results document that the stretch of the very last 6 amino acids of Nce102 is responsible for directing this protein to MCC patches.

The C terminus of Nce102 is necessary for the formation of the plasma membrane invaginations. As previously observed in freeze-etch preparations of plasma membrane replicas, the full deletion of NCE102 results in a lack of furrow-like invaginations (32). We therefore tested whether the C-terminal truncation of the Nce102 protein has a similar effect. For direct comparability with the results obtained by confocal microscopy we used the strains expressing NCE102-GFP and NCE102Δ1-146-GFP for replica preparation. In total, 111 cells expressing full-length NCE102-GFP and 33 cells expressing NCE102Δ1-146-GFP were analyzed. While all cells expressing the full-length NCE102 gene formed furrows, only in 3 out of 33 analyzed Nce102Δ1-146-containing cells could shallow, premature furrow-like invaginations be identified. The remaining 90% of cells exhibited flat plasma membranes devoid of the typical furrow-like invagination pattern. Flat, smooth, elongated areas with surface densities comparable to that observed in furrows of wild-type cells were detected instead (Fig. 5). This remarkable change in the plasma membrane morphology was virtually identical to the phenotype previously observed for the nce102Δ strain (32). We conclude that, after its C-terminal part had been deleted, Nce102 lost its function in formation of furrows.

**DISCUSSION**

So far, we have documented nine proteins sharing the MCC localization: four members of the Sur7 family (Sur7, Ynl149c, Fmp45, and Ylr414c), three proton symporters (Can1, Fur4, and Tat2), and Nce102, with its close homolog Ftih1 (15, 16, 25, 26). Among those, no general MCC targeting sequence was revealed. Our findings rather support a step-by-step mechanism for MCC formation as suggested previously: (i) Pil1-driven assembly of planar membrane domains (primary MCC patches) containing Sur7 family proteins; (ii) recruitment of specific lipids, Nce102, and transporters into these domains; and (iii) lipid- and/or protein-mediated invagination of the MCC membrane (for details see references 27 and 32). Previously we showed that Nce102 directs the proton transporters into MCC patches, where they are protected against untimely turnover (16). In this study we document that the mechanism of plasma membrane domain formation and this MCC function is probably conserved through the largest phylum of fungi, the Ascomycota. This conclusion is based on the observation that, by expressing one of the most distant Nce102 homologs coded by *S. pombe* ftn1, we were able to reconstitute the Can1-enriched MCC patches in the nce102Δ strain of *S. cerevisiae*.

The hydrophathy plot of Nce102 revealed four potential membrane-spanning domains. This prediction of the membrane topology, however, does not seem to be correct, at least in the sense of the protein orientation. Using a combination of glycosylation and growth assays, we determined that the C terminus of Nce102 is oriented toward the cytoplasm, while in the predicted protein structure it is exposed to the cell exterior. Taking this observation into account, we consider three putative models of Nce102 accommodation in the plasma membrane (Fig. 2A, models II to IV). The first one (model II), assuming that the protein across the membrane only twice, is based on a set of experiments where the topology was determined using the entire HA/Suc2/His4C reporter. However, this tag is quite bulky (125 kDa) compared to the tested Nce102 fragments, and thus it is possible that, during the membrane insertion, the fusion proteins do not fold correctly. On the other hand, it is worthwhile mentioning that, using the same strategy as that described above, we confirmed unambiguously the tetraspan topology of another protein residing in MCC, Sur7 (20; V. Stradalova, unpublished results). To obviate the interference of the bulky HA/Suc2/His4C reporter with the membrane insertion of Nce102-derived peptides, much smaller Suc2 fragments (6 and 9.3 kDa) were tested as glycosylation reporters. As evident from Fig. 2D, in this case the C-terminal reporter on fragment Nce102Δ1-63 is not glycosylated, which indicates that the hydrophilic loop following amino acid 63 is not exposed to the ER lumen and therefore probably faces the cytoplasm when the Nce102 protein is incorporated into the plasma membrane. The same holds for a newly created glycosylation site generated at amino acid 70. Conclusions from these results are summarized in topology model III in Fig. 2A. Neither of the two models, however, might be fully fitting. It is noteworthy that the first predicted TMD in Nce102 consists of 18 amino acids only. Usually, at least 23 amino acid residues are required for spanning the yeast plasma membrane (31, 37). However, 7 of 10 amino acid residues of the cytoplasmic N terminus are of positive hydrophathy index, and the possibility that they might be involved in protein embedding cannot be excluded. The first extracellular stretch in model III consists of 12 highly polar amino acids, while the second loop, the internal one, contains only 9 amino acids, 7 of which are nonpolar. The other two amino acids are prolines, which are well documented to influence the conformation of, e.g., helical regions (1, 24, 39). Helices containing proline have a pronounced kink, with
bending of the helical axis of approximately 30° away from the side with the proline residue (38). Regarding all our results and the specific features of Nce102, we suggest topology model IV, where the proposed TMDs 2 and 3 do not span the plasma membrane but are inserted into the outer leaflet of the membrane. Due to its high hydrophobicity, the short 9-amino-acid stretch located between potential TMDs 2 and 3 might be incorporated into the lipid bilayer as well. In this case, the whole stretch of amino acids 40 to 95 could be embedded in the plasma membrane and the presence of the prolines might be involved in bending this helix.

It is plausible to speculate that these specific features of the Nce102 primary structure endow Nce102 with a certain topological flexibility, which might be of physiological relevance. In this context it is interesting to refer to the recent studies of Fröhlich et al. (10), where Nce102 has been proposed to act as a sensor for sphingolipids, regulating the appearance of the MCC pattern via phosphorylation of Pil1, a cytosolic protein accumulated beneath the MCC patches. Indeed, the involvement of Nce102 in processes including changes of the plasma membrane lipid composition is indicated by its increased expression reported in several screening studies looking for responses to environmental changes, like heat stress, oxidative and osmotic shocks, nitrogen source depletion, diauxic shift, transition to invasive growth, and introduction of various toxic substances (8, 11, 12, 34). The monitoring and transferring of Nce102 primary structure endow Nce102 with a certain topological flexibility, which might be of physiological relevance.

We identified a short C-terminal motif (PTISQV), the absence of which resulted in a loss of Nce102 function in the plasma membrane (Fig. 2A, model IV). It is known that a shallow insertion of small hydrophobic inclusions effectively induces membrane curvatures (5). A critical amount of Nce102 at the patches could, in this case, induce the membrane bending. Many other proteins involved in the organization of membrane microdomains, for example, caveolins, reticulins, and flotillins (reggies), contain at least one membrane-integrated, but not membrane-spanning domain, which, it has been suggested, adopts a hairpin structure (3).

Based on our ultrastructural studies, we suggest that Nce102 operates also as a structural protein in bending the plasma membrane, which, of course, involves also the effects of specific lipids in the plasma membrane region. The role of Nce102 in specific structure formation does not exclude the possibility that Nce102 has additional functions.

ACKNOWLEDGMENTS

We are very grateful to Ingrid Fuchs for technical assistance. We also thank Katerina Malinska from the Institute of Experimental Botany, Academy of Sciences of the Czech Republic (Prague), for the construction of the pVTU100-mRFP plasmid and Sabine Strahl, University of Heidelberg, for technical advice.

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