Mutagenesis of the putative sterol-sensing domain of yeast Niemann Pick C–related protein reveals a primordial role in subcellular sphingolipid distribution

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Lipid movement between organelles is a critical component of eukaryotic membrane homeostasis. Niemann Pick type C (NP-C) disease is a fatal neurodegenerative disorder typified by lysosomal accumulation of cholesterol and sphingolipids. Expression of yeast NP-C–related gene 1 (NCR1), the orthologue of the human NP-C gene 1 (NPC1) defective in the disease, in Chinese hamster ovary NPC1 mutant cells suppressed lipid accumulation. Deletion of NCR1, encoding a transmembrane glycoprotein predominantly residing in the vacuole of normal yeast, gave no phenotype. However, a dominant mutation in the putative sterol-sensing domain of Ncr1p conferred temperature and polyene antibiotic sensitivity without changes in sterol metabolism. Instead, the mutant cells were resistant to inhibitors of sphingolipid biosynthesis and super sensitive to sphingosine and C2-ceramide. Moreover, plasma membrane sphingolipids accumulated and redistributed to the vacuole and other subcellular membranes of the mutant cells. We propose that the primordial function of these proteins is to recycle sphingolipids and that defects in this process in higher eukaryotes secondarily result in cholesterol accumulation.

Introduction

The function of eukaryotic membranes reflects their lipid composition as much as that of proteins, and yet the mechanisms by which lipid partitioning is maintained are largely unknown. Niemann Pick type C (NP-C) disease is a neurodegenerative lipid storage disorder typified by lysosomal accumulation of cholesterol and sphingolipids. Expression of yeast NP-C–related gene 1 (NCR1), the orthologue of the human NP-C gene 1 (NPC1) defective in the disease, in Chinese hamster ovary NPC1 mutant cells suppressed lipid accumulation. Deletion of NCR1, encoding a transmembrane glycoprotein predominantly residing in the vacuole of normal yeast, gave no phenotype. However, a dominant mutation in the putative sterol-sensing domain of Ncr1p conferred temperature and polyene antibiotic sensitivity without changes in sterol metabolism. Instead, the mutant cells were resistant to inhibitors of sphingolipid biosynthesis and super sensitive to sphingosine and C2-ceramide. Moreover, plasma membrane sphingolipids accumulated and redistributed to the vacuole and other subcellular membranes of the mutant cells. We propose that the primordial function of these proteins is to recycle sphingolipids and that defects in this process in higher eukaryotes secondarily result in cholesterol accumulation.

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Abbreviations used in this paper: ABA, aureobasidin A; DHS, dihydrosphingosine; E–L, endosomal–lysosomal; IPC, inositolphosphorylceramide; LDL, low density lipoprotein; MIPC, mannosyl-IPC; M(IP)2C, mannosyl-dip-IPC; NPC-C, Niemann Pick type C; NPC1, NP-C gene 1; NCR1, NP-C–related gene 1; SCAP, SREBP cleavage-activating protein; SREBP, sterol regulatory element-binding protein; SSD, sterol-sensing domain.
to sterol and fatty acid homeostasis (26% identity over 190 residues; Fig. 1). The latter conservation corresponds to a putative sterol-sensing domain (SSD).

**NPC1** is ubiquitously and constitutively expressed in mammalian tissues and is conserved from yeast to humans (Loftus et al., 1997). In contrast, many aspects of receptor-mediated uptake of sterol are absent over the same period of evolution, suggesting a primordial function of the Npc1 protein family that predates exogenous sterol transport. We demonstrate that the *NP-C–related gene 1* (*NCR1*), the yeast orthologue of mammalian *NPC1*, performs a conserved function that is distinct from trafficking of sterol. We show that in mammalian cells the proteins are interchangeable with respect to cholesterol and ganglioside transport, and yet a null mutation in yeast *NCR1* has no sterol-related phenotype. A dominant mutation in the SSD of yeast *NCR1* confers pleiotropic phenotypes consistent with altered sphingolipid transport. We propose that in this ancestral eukaryote, the role of NPC1-like proteins is to recycle sphingolipids and that in multicellular organisms, one consequence of this is transport of cholesterol.

**Results**

**Yeast NCR1, an orthologue of human NPC1**

The predominant sterol of *Saccharomyces cerevisiae* is ergosterol, which is synthesized, esterified, used, and regulated in a similar manner to cholesterol in mammals (Sturley, 2000).
last 64 residues of human NPC1 to Ncr1p (NCR1-Cterm). Expression constructs for NCR1 and NCR1-Cterm were transfected into the CHO cells, and sterol transport was assessed with filipin staining and fluorescence microscopy. Expression of NCR1-Cterm restored cholesterol clearance to levels indistinguishable from cells expressing human NPC1 (Fig. 2 and Table I). A significant decrease in E–L filipin staining was also observed upon expression of NCR1. Although fewer filipin negative cells were observed, most transfected cells displayed “partial” clearance of cholesterol. This diminished suppression likely reflects mislocalization of the native yeast protein; a GFP fusion with Ncr1p localizes to the early endosome, whereas NCR1-Cterm–GFP was detected in the late endosomal compartment (unpublished data).

The subcellular accumulation of sphingolipids, particularly gangliosides, is another characteristic of the NP-C syndrome, and in the case of GM1 has been observed in NPC1-deficient CHO cells (Sugimoto et al., 2001). Expression of NCR1-Cterm or human NPC1 reversed the accumulation of GM1 in the E–L compartment of transfected CT60 cells as detected by fluorescent cholera toxin (subunit B) binding assays (Fig. 2 B). Thus, when provided with the correct localization motif, the yeast and human proteins function interchangeably, suggesting a conserved activity despite several billion years of divergence.

Expression of yeast NCR1 complements loss of NPC1

To test the functional equivalence of the yeast and human proteins, we expressed NCR1 in CHO CT60 and NPC1-trap cells lacking Ncr1p (Cruz et al., 2000; Higaki et al., 2001). Yeast proteins lack targeting information for mammalian endosomes. Because Npc1p localization, which is mediated by COOH-terminal motifs is required for full complementation (Watari et al., 1999; Ioannou, 2000), we fused the

Null mutations in yeast NCR1 have no effect on sterol metabolism

In mammalian cells lacking NPC1, exogenous sterol is sequestered from the esterification reaction and sterol biosynthesis is misregulated. During anaerobic growth (Gollub et al., 1974) or in ncr2-1 mutants (Lewis et al., 1988), sterol is absorbed by yeast in an ABC-transporter–dependent process (Wilcox et al., 2002) and esterified by the acylCoA chole-

Table I. Restoration of cholesterol transport in npc1-deficient CHO cells by expression of yeast NCR1

| Phenotypes tested | NCR1 (normal) | ncr1Δ | Y718D | Y718N | Y717C |
|------------------|---------------|-------|-------|-------|-------|
| Sterol synthesis ([^1]H)acetate, (P vs. NPC1) | 18.2 ± 2.9 | 19.7 ± 1.2 | 13.2 ± 2.3 | 13.2 ± 1.8 | 16.2 ± 1.6 |
| Sterol esterification ([^1]H)cholesterol, (P vs. NPC1) | 11.0 ± 2.8 | 11.9 ± 2.4 | 11.9 ± 0.9 | 10.5 ± 2.7 | 10.7 ± 1.7 |
| Exogenous sterol esterification ([^1]C)cholesterol, (P vs. NPC1) | 74.3 ± 2.0 | 69.2 ± 6.3 | 72.2 ± 3.8 | 77.9 ± 3.6 | 78.1 ± 0.8 |
| 30°C (2% dextrose) | ++ | ++ | + | ++ | ++ |
| 38°C (2% dextrose) | ++ | ++ | – | ++ | ++ |
| Anaerobiosis | ++ | ++ | – | ++ | ++ |
| Nystatin (5 μg/ml) | ++ | ++ | – | ++ | ++ |
| Ethanol (3%) | ++ | ++ | – | ++ | ++ |
| Acetate (2%) | ++ | ++ | – | ++ | ++ |
| Sodium chloride (0.75 M) | ++ | ++ | – | ++ | ++ |
| Sorbitol (1.0 M) | ++ | ++ | – | ++ | ++ |
| Calcium chloride (0.8 M) | ++ | ++ | + | ++ | ++ |
| Hydroxyurea (40 mM) | ++ | ++ | + | ++ | ++ |

Yeast strains with the missense mutations Y718D, Y718N, and Y571C in the SSD of NCR (Fig. 1B) were assessed for growth or metabolic incorporation (percent incorporation ± SEM) under the conditions described. The growth of the normal strain is considered ++ and the growth of the other strains is expressed relative to that scale.
sterol acyltransferase orthologues (Yang et al., 1996). Deletion of NCR1 had no effect on acetate incorporation into sterols, oleate incorporation into steryl esters, anaerobic viability, or esterification of exogenous [14C]cholesterol (Table II). Cell sterol composition can be exaggerated by deletion of ERG6 (C-24 sterol methyl transferase; Gaber et al., 1989) or ERG9 (squalene synthase; Fegueur et al., 1991) or treatment with zaragozic acid, an inhibitor of squalene synthase (Bergstrom et al., 1993). This exaggeration results in membrane perturbations and, in the latter two cases, sterol auxotrophy; however, loss of NCR1 in these contexts had no effect on cell viability or sterol uptake (unpublished data).

In contrast to observations with NP-C mutant fibroblasts (Fig. 2 A), the majority of sterol in ncr1Δ and normal yeast strains detected by filipin fluorescence resides at the plasma membrane (Fig. 3 A). In validation of the assay, arv1Δ strains that accumulate sterols in subcellular membranes (Tinkelenberg et al., 2000) displayed significant subcellular filipin staining, whereas an erg6Δ mutant had barely detectable plasma membrane fluorescence. Moreover, ncr1Δ strains displayed

![Figure 3. Sterol distribution in yeast NCR1 mutants.](image)

**Figure 3. Sterol distribution in yeast NCR1 mutants.** (A) Yeast strains (normal, ncr1Δ, and NCR1Y718D) were stained with filipin to detect ergosterol. Deletions in ERG6 (erg6Δ) block synthesis of ergosterol, whereas ARV1 deletions (arv1Δ) accumulate sterol in subcellular membranes. (B) Cell membrane preparations from normal and NCR1 deletion strains were subjected to subcellular fractionation, lipid extraction, and TLC analysis after [14C]oleate or [3H]acetate incorporation. Distributions of sterols relative to phospholipid are given as a ratio of [3H]acetate to [14C]oleate. Total [3H]acetate and [14C]oleate incorporation was 3.06 and 1.79 (normal) versus 2.77 and 1.18 (deletion) x 10^5 dpm/OD600, respectively. Fractions were characterized by immunoblotting with antisera to the plasma membrane ATPase (Pma1p, solid line, squares) and the vacuolar membrane H^+-ATPase (Vph1p, dashed line, diamonds), followed by scanning densitometry (arbitrary units, representative data).

![Figure 4. NCR1 expression in yeast.](image)

**Figure 4. NCR1 expression in yeast.** (A) RNA hybridization of the indicated strains grown in YPD media at 30°C. The loading control of 28S ribosomal RNA is shown. No significant differences in NCR1 transcript levels for normal, NCR1Y718D, NCR1Y718N, or NCR1Y571C (not depicted) strains were detected. (B) Expression of Ncr1-HAp. The chromosomal copy of NCR1 was tagged with HA at the COOH terminus by homologous recombination. Duplicate membrane extracts were solubilized with Triton X-100, deglycosylated (endo-H), and resolved by SDS-PAGE and immunoblotting (αHA 12CA5 mAb). Molecular mass markers (Bio-Rad Laboratories) are shown. (C) Subcellular localization of Ncr1-HAp. Membrane preparations from cells expressing Ncr1-HA were fractionated by ultracentrifugation in Renograffin 60. Fractions 1–14 were characterized by SDS-PAGE and immunoblotting with antibodies to Ncr1-HA (Anti-HA), the plasma membrane ATPase (Anti-Pma1p), and the vacuolar H^+-ATPase (Anti-Vph1p).
no changes in sensitivity to polyene antibiotics (e.g., nystatin) that form lethal complexes with ergosterol. Finally, the relative distribution of sterols and phospholipids across a subcellular fractionation was unchanged in ncr1/H9004 mutants (Fig. 3B), further suggesting that membrane composition, particularly with respect to sterol, was unaffected by loss of NCR1.

Normal and ncr1/H9004 strains grew comparably on a variety of carbon sources and at different temperatures (12°C, 30°C, 38°C, and 40°C). Similarly, there was no growth differential due to supplementation with 1 M sorbitol or 0.75 M NaCl (Table II). A transmembrane permease activity with specificity for oleic acid and acriflavine has been suggested for Npc1p (Davies et al., 2000). However, we could detect no defect in uptake of [3H]oleate or in accumulation of acriflavine in ncr1 mutants (unpublished data). In a further attempt to reveal a phenotype due to loss of NCR1, we undertook a transcriptional profiling approach. Oligonucleotide arrays (Affymetrix) were used to compare normal and ncr1/H9004 strains. Consistent with the redundancy of NCR1, there were no significant (i.e., greater than twofold) transcriptional differences between these strains (unpublished data). In summation, the deletion of NCR1 had no detectable physiological consequences.

Expression and localization of Ncr1p

The NCR1 gene is transcribed (Fig. 4A) and expressed in normal strains and growth conditions as evidenced by tagging the endogenous protein with the HA epitope NCR1-HA. The biological activity of this allele was confirmed in the NPC1 mutant CHO cells (unpublished data). Ncr1-HAp was detected as a membrane-associated, endoglycosidase H-sensitive glycoprotein (∼130 kD; Fig. 4B), indicating its passage through the ER. In a further example of concordance between the yeast and human proteins, Ncr1-HAp cofractionated predominantly with normal yeast vacuoles, the equivalent organelle to the E–L system of mammalian cells (Fig. 4C). Furthermore, a GFP-Ncr1 fusion protein localizes to the vacuolar membrane (Huh et al., 2003; unpublished data).

A dominant mutation in the SSD of yeast NCR1

Ncr1p is biologically active when expressed in mammalian cells, and yet loss of Ncr1p is of no apparent consequence in yeast. To bypass this redundancy, we introduced mutations in NCR1 (Fig. 1B) analogous to dominant mutations in the SSD of SCAP (Nohturfft et al., 1998). The presence of asparagine instead of aspartic acid at residue 443 in SCAP confers sterol-resistant SREBP cleavage; thus, we mutated the analogous residue (Tyr718) of Ncr1p to either aspartic acid (NCR1Y718D) or asparagine (NCR1Y718N). In addition, we changed an invariant tyrosine of the Ncr1p SSD (Tyr571) to cysteine (NCR1Y571C). The mutations were introduced at the NCR1 locus by homologous recombination and did not affect NCR1 transcript levels (Fig. 4A).

Because the NCR1Y571C, NCR1Y718D, and NCR1Y718N mutations reside in a putative SSD, we anticipated an alter-
ation in sterol metabolism. NCR1Y718D strains, unlike the others, were hypersensitive to nystatin, suggestive of perturbed plasma membrane properties (Fig. 5 A). However, anaerobic growth, an indicator of exogenous sterol transport, was unaltered relative to aerobic growth (Table II). Furthermore, sterol synthesis, esterification (Table II), and localization (Fig. 3 A; and see Fig. 7 C) were not significantly different in these strains compared with normal strains. The NCR1Y718D allele conferred poor growth at 30°C, inviability at 38°C (Fig. 5 A), and salt sensitivity and poor growth on carbon sources such as acetate and ethanol (Table II). The NCR1Y718N and NCR1Y77C, strains were modestly impaired at elevated temperatures (Fig. 5 A), but were otherwise indistinguishable from normal strains. The NCR1Y718D strains were not uniformly sick; responses to sorbitol, calcium, hydroxyurea, and low pH were indistinguishable from growth on YPD at 30°C (Table II).

Yeast strains with the NCR1Y718D mutation exhibit pleiotropic phenotypes, whereas the ncr1Δ strain showed no defects, predicting that this variant would act as a dominant allele. Normal, ncr1Δ, and NCR1Y718D strains were transformed with vector control or a plasmid carrying NCR1. NCR1 expression was confirmed by RT-PCR assays (not depicted) but failed to rescue inviability at 38°C of the NCR1Y718D strains (Fig. 5 B). In addition, heterozygous NCR1/NCR1Y718D diploids retain the temperature sensitivity of the NCR1Y718D allele, further indicating dominant inheritance (unpublished data).

The NCR1Y718D allele alters sphingolipid metabolism

NP-C disease is characterized by sphingolipid accumulation. Furthermore, the sensitivity of yeast to temperature, high salt, or polyns can result from modulations in sphingolipids (Dickson and Lester, 2002). Thus, in the absence of a sterol-related phenotype, we questioned whether or not sphingolipid metabolism was disturbed by mutations in NCR1. In S. cerevisiae, ceramide is synthesized from phytosphingosine and is the precursor for synthesis of inositolphosphoglycerides (Nickels and Broach, 1996; Chung et al., 2001). The NCR1Y718D mutant strain was super sensitive to 10 μM C2-ceramide (Fig. 5 A) and 5 μM sphingosine (Fig. 5 C) compared with the other strains. This phenotype was evident as a prolonged lag phase, which is consistent with retarded nutrient uptake, a trait common to alterations in sphingolipids (Chung et al., 2001).

Changes in sphingolipid status can also be monitored using antifungals such as aureobasidin A (ABA), which blocks synthesis of IPC causing ceramide accumulation and inviability (Nagiec et al., 1997). NCR1Y718D strains were resistant to 500 ng/ml ABA, whereas normal and ncr1Δ strains were sensitive (Fig. 5 C). ABA resistance was associated with increased incorporation of [3H]dihydrosphingosine (DHS) into IPC at the expense of ceramide over a range of drug concentrations (Fig. 5 D), which is consistent with increased flux through the sphingolipid biosynthetic pathway of NCR1Y718D strains.

The sensitivity of the NCR1Y718D strain to sphingosine and C2-ceramide, and resistance to ABA, led us to assess metabolic incorporation of [3H]DHS into sphingolipids in the mutant strains over a 2-h labeling period (Fig. 6). The
levels of ceramide were comparable between the strains. However, \( NCR1Y_{718} \) strains accumulated 2.8-fold more MIPC (23.3% vs. 8.1%, \( P < 0.01 \)) and significantly less IPC and M(IP)2C (Fig. 6 A). No changes in sphingolipids were observed in the other \( NCR1 \) mutants (unpublished data). Given the elevated MIPC levels in \( NCR1Y_{718} \) strains, we assessed the expression of \( CSG1 \), \( CSG2 \) (encoding components of MIPC synthase), and \( IPT1 \) (mediating the synthesis of M(IP)2C). \( CSG2 \) transcript levels were elevated in \( NCR1Y_{718} \) strains (Fig. 6 B).

Upon heat shock, yeast elevate ceramide levels (Jenkins et al., 1997). To assess if the temperature sensitivity of the \( NCR1Y_{718} \) strains was related to sphingolipid metabolism, we monitored the changes in ceramide levels of normal, \( ncr1 \), and \( NCR1Y_{718} \) strains upon shifting to 38°C and labeling with \( [\text{H}] \)DHS. Ceramide levels increased fourfold in normal and \( ncr1 \) strains upon temperature shift, whereas \( NCR1Y_{718} \) strains showed less than a twofold increase (Fig. 6 C). Complex sphingolipids were not altered by heat shock relative to growth at 30°C. The molecular mechanism or purpose for increased de novo synthesis of ceramide after heat stress is not known, however, the diminished response by the \( NCR1Y_{718} \) strains may explain their compromised growth rates.

The \( NCR1Y_{718} \) mutation causes sphingolipid mislocalization

In NPC1-deficient mammalian cells, the primary defect in sphingolipid metabolism relates to transport out of the E–L system (Choudhury et al., 2002). To assess whether the recycling of complex sphingolipids through the yeast endosomal pathway might be disturbed by mutations in \( NCR1 \), we labeled sphingolipids to steady state with \( [\text{H}] \)DHS, \([\text{C}] \)oleate (phospholipids), or \([\text{H}] \)acetate (sterols) at 30°C for 18 h. (A) Total cellular incorporation into sphingolipid pools (percentage of total extraction, means ± SEM; *, \( P < 0.01 \)) to normal). Total incorporation of \( [\text{H}] \)DHS for normal and \( NCR1Y_{718} \) strains was 1.826 and 2.152 \( \times 10^3 \) cpm per OD600, respectively. (B) Cell membrane preparations of identical cultures to those in A were subjected to subcellular fractionation after \( [\text{H}] \)DHS incorporation. The distributions of IPC, MIPC, and M(IP)2C are presented as the percentage of total incorporation in normal (gray bars) and \( NCR1Y_{718} \) (black bars) strains (representative data). (C) Membrane preparations after \( [\text{H}] \)acetate incorporation were subjected to subcellular fractionation. Total \( [\text{H}] \)acetate incorporation was 3.06 and 1.79 (normal) versus 6.09 and 2.43 (\( NCR1Y_{718} \) strains) \( \times 10^3 \) dpm/OD600, respectively. The distributions of sterol and phospholipids are presented as the percentage of total incorporation in normal (gray bars) and \( NCR1Y_{718} \) (black bars) cells per fraction. (D) Immunoblotting of fractions with antibodies to plasma membrane (Pma1p), dolichol phosphate mannose synthase (Dpm1p, ER), and vacuoles (Vph1). Peak fractions for these markers were coincident between the strains, despite apparent differences in protein expression.
MIPC, and to a lesser extent M(IP)2C, redistributed to subcellular organelles, particularly the vacuolar compartment as inferred by colocalization with the vacuolar H\(^+\)-ATPase (Figs. 7, B and D, Vph1p). Indeed, if we consider fractions 4 and 5 (Fig. 7), in which Vph1p is predominant but Pma1p or Dpm1p are low, NCR1Y718D causes a 6.5- and 7.2-fold accumulation of MIPC relative to controls. This lipid rearrangement in the NCR1Y718D strains was specific to complex sphingolipids. The subcellular distribution of ceramide (Fig. 7 B), phospholipids, and sterols (Fig. 7 C) was similar between strains.

**Discussion**

Niemann Pick disease type C is a neurodegenerative disorder with a significant defect in subcellular transport of exogenous sterols. However, NP-C disease is not solely a cholesterol storage disorder, and the nature of the offending metabolite in this syndrome is contentious, particularly in the brain. Sterol accumulation is detectable in NPC1-deficient neurons, but is not as striking as at extra-neuronal sites. Several works in genetically modified murine models of this disease have concluded that sterol uptake has no role in symptom development (Xie et al., 2000a,b). In contrast, the metabolism and transport of sphingolipids is aberrant in most npc1−/− cells, including neurons (Sugimoto et al., 2001; Zervas et al., 2001a; Zhang et al., 2001). Finally, pharmacological intervention in sphingolipid synthesis alleviates symptoms in animal models of NP-C disease, whereas reductions in plasma cholesterol give no benefit (Patterson et al., 1993; Zervas et al., 2001b).

What does the yeast system inform regarding the human syndrome? Despite evolutionary divergence, the yeast and mammalian proteins are functionally equivalent for sterol and ganglioside transport in mammalian cells. However, we failed to identify any defect in sterol metabolism conferred by mutations in NCR1. Thus, we hypothesize that Ncr1p performs a primordial function that underlies the sterol transport defects associated with NP-C disease. However, a quandary remains: what is the physiological role of this protein in yeast? The similarities of yeast Ncr1p and human Npc1p to the SREBP cleavage-activating protein led us to create constitutive mutations in the SSD of Ncr1p. One such allele, NCR1Y718D, functioned as a dominant neomorph, its most salient phenotype being the errant subcellular accumulation of MIPC. The normal pathways of sphingolipid distribution (Fig. 8) commence with synthesis of ceramide in the ER and transport to the Golgi compartment where complex sphingolipid synthesis progresses (Funato et al., 2002). An asymmetry is then created; MIPC and M(IP)2C accumulate in the plasma membrane, whereas IPC localizes to the vacuole. In the NCR1Y718D strains, the sequestration of MIPC in the vacuole limits its retrograde movement to the Golgi compartment resulting in reduced M(IP)2C synthesis. The same event likely results in up-regulation of biosynthesis, in part at the level of transcription of the CSG2 gene. This increased flux explains resistance to pathway inhibitors, sensitivity to substrates such as C2-ceramide and sphingosine, and inadequate response to heat shock in terms of ceramide accumulation.

![Diagram](image)

Based on these observations, we propose that the primordial role of the Npc1 protein family lies in sphingolipid recycling with sterol movement as a consequence, which is consistent with studies of NPC1 in mammalian cells (Coxey et al., 1993; Sugimoto et al., 2001) and with hypotheses that neurodegeneration in NP-C disease results from ganglioside accumulation. There are many precedents for a link between sphingolipid and sterol homeostasis. Multiple sphingolipid storage disorders are associated with perturbations in cholesterol homeostasis (Pagano et al., 2000), perhaps because the two molecules readily form membrane microdomains (Bagan et al., 2000; Simons and Ehehalt, 2002). Indeed, cholesterol accumulation was strikingly reduced when NPC1 deficiency was combined with loss of the \( \beta \)-1,4GalNAc transferase responsible for complex ganglioside synthesis (Liu et al., 2000; Gondre-Lewis et al., 2003).

The SCAPs of mammals and insects via the SSDs are proposed to respond to membrane perturbations rather than changes in a specific molecule (Seegmiller et al., 2002). It is clear that residue 443 of SCAP is a critical position in the SSD (Nothurft et al., 1996). Our work confirms this in a different protein (Ncr1p) and illustrates a clear species and molecule specificity. In mammalian SCAP, insect SCAP (Seegmiller et al., 2002), or yeast Ncr1p, the normal residue is aspartic acid, asparagine, or tyrosine, respectively. In mammalian SCAP or yeast Ncr1p, substitution with asparagine or aspartic acid produce dominant phenotypes, which is likely due to the loss of specific protein–protein interactions. A critical component of this homeostasis is the interaction of SCAP with ER retention proteins such as INSIG1 and INSIG2. Although there is no SCAP orthologue in yeast, INSIG-like proteins are conserved (Hampton, R., personal communication). Although they have no reported mu-
tant phenotype, it is tempting to speculate they may interact with Ncr1p.

One model for the failure to identify an ncr1 null phenotype could be that the protein functions in a redundant pathway for sphingolipid recycling. Accordingly, NCR1-Y718 acts as a dominant mutant that corrupts mutually redundant pathways of sphingolipid recycling, thus revealing a phenotype. The manipulation of these bypass pathways would be obvious targets to circumvent the transport defects and thus treat NP-C disease. Interestingly, overexpression of certain Rab proteins implicated in vesicular transport suppresses loss of NPC1 (Choudhury et al., 2002; Walter et al., 2003). Whether or not this is an avenue to therapy remains to be determined; however, the identification of such bypass pathways in yeast and humans will provide a significant step in this direction.

Materials and methods

General

Standard methods were used for yeast manipulation (Ausubel et al., 1998). Yeast strains are isogenic with W303-1A (Thomas and Rothstein, 1989). Cholesterol, ergosterol, and nystatin were added from stock solutions (5 and 2 mg/ml in 1:1 ethanol/txoloxapol [Sigma-Aldrich] and 10 mg/ml in propylene glycol, respectively). Zaragoasic acid (provided by Y.S. Chao, Merck & Co., Rahway, Nj), sphingosine, C2-ceramide (BIOMOL Research Laboratories, Inc.), and ABA (Panvera) were used at the concentrations indicated.

The BLAST alignment tool was used for sequence comparisons (Altschul et al., 1997).

Mutagenesis of NCR1

NCR1 deletions were constructed by allele replacement (Erdeniz et al., 1997) using a PCR product (oligos: 5'KONP, 3'KONP) homologous to NCR1 at its 5' and 3' ends and flanking the Kluveromyces lactis URA3 gene. The locus was characterized by PCR with 5' and 3' NCR1 or URA3-specific oligos (5'NPC5, 006NSFC, 3'NP-internals, and 3'NPC5). Misseis alleles were created similarly with oligos N25, D25, and C25 for the NCR1200, NCR12002, and NCR1200 alleles, respectively. Ncr1p was tagged at the COOH terminus with a triple HA epitope (18 residues) using a PCR-generated template (Schneider et al., 1995). All alleles were sequenced.

Expression of NCR1 in mammalian cells

NCR1 or NCR1-HA was amplified from yeast genomic DNA (oligos 5'NPC5 and 3'NPC5) and incorporated into pcR3.1 digested with EcoRV (pcR3.1-NCR1 and pcR3.1-NCR1-HA). The COOH-terminal 95 residues of NCR1 were replaced with amino acids 1214–1278 of human NPC1 in pcR3.1-NCR1-Cterm after PCR amplification of the human CDNA/SNCR1tail-BsmI and 3'NPC1tail digestion with BsmI and XhoI, and ligation at the same sites in pcR3.1-NCR1.

CHO lines C660 and NPC1trap were cultured 3 d in Ham's F12 and 10% lipoprotein-deficient serum and transfected (FuGENE 6; Roche Molecular Biochemicals) with pcR3.1, pcR3.1-NCR1, or pcR3.1-NCR1-Cterm and a mammalian nuclear targeting-EGFP vector (Wataria et al., 1999) to identify transfected cells. Cells were incubated with 10% lipoprotein-deficient serum and 50 μg/ml of human LDL for 24 h, fixed with 3% PFA, and stained with 50 μg/ml of filipin (Sigma-Aldrich). Cells were imaged by confocal fluorescence microscopy (model LSM510; Carl Zeiss Micromaging, Inc.); excitation 488 nm for EGFP. For quantitative analysis, filipin-corrupted cells were counted from multiple fields of EGFP-positive cells. For detection of GM1, cells were washed with Ham's F12, 25 mM Hepes, pH 7.4, and 0.01% BSA, incubated with 20 mM of Alexa 555-conjugated CtxB (Molecular Probes, Inc.) for 1 h at 37°C, and fixed with 4% PFA. Fluorescence images were collected after excitation at 568 nm using an microscope (model Axiamat 100; Carl Zeiss Micromaging, Inc.).

Phenotypic analyses

The expression of NCR1-HA was confirmed in Triton X-100-soluble protein extracts (Guo et al., 2001) by immunoblotting with the 12CAS mAb (Roche Molecular Biochemicals). Protein deglycosylation with endoglycosidase H was accomplished as directed (Roche Molecular Biochemicals). For growth in the presence of sphingosine (in ethanol) or ABA (in DMSO), cells were continuously monitored at 25°C at 0.1% ethanol/0.05% tergitol, or 40 mM hydroxyurea were performed on solid YP media. Total RNA was isolated from independent isolates of the ncr1-trap strain grown to OD600 = 0.5–0.6 in YEPD and analyzed in microarray or northern hybridizations (Wilcox et al., 2002).

Cell labeling (10 μCi [3H]acetic acid [NEN Life Science Products], 5.0 μCi [3H]olic acid [NEN Life Science Products], or 2 μCi/ml [3H]DHS [American Radiolabeled Chemicals]; for 2–5 h), lysis, release of lipids, TLC, and data collection were performed as described previously (Yang et al., 1996). Sphingolipids were isolated from cells treated with 5% TCA (ice, 15 min) by extraction with diethylther/ethanol/water/pyridine/ammonia (5:15:15:1:0.18, vol/vol) for 60 min at 60°C, resuspension was performed in chloroform/methanol/water (16:16:5) and TLC was performed on silica gel 60 (EM Science) plates resolved with chloroform/methanol/2.4N ammonia (9:7:2). Ergosterol localization was visualized by filipin staining (20 μg/ml) as above.

Membrane preparations from cells grown for 18 h in the presence of [3H]DHS, [3H]acetic acid, or [3H]olic acid were subjected to discontinuous gradient ultracentrifugation in Renografin-60 (Braaco Diagnostics: Tinkenberg et al., 2000). 250-μl fractions were diluted 10-fold with Tris EDTA buffer and centrifuged at 100,000 g. Radiolabeled sphingolipids were extracted from the pellet and resolved by TLC as before. Marker proteins for the ER (DPM1 [dolichol phosphate mannose synthase]), vacuoles (Vph1p [vacuolar H+-ATPase]), and plasma membrane (Ipm1p [plasma membrane ATPase]) antisera provided by A. Chang, University of Michigan, Ann Arbor, MI indicated peak fractions for these organelles.

Online supplemental material

All oligonucleotides were synthesized by Invitrogen and are described in Table S1. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200310046/DC1.

The work is dedicated to all NP-C-afflicted families and to the memory of Mrs. A.M.R. Sturley.

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