The Yeast Nucleoporin Nup188p Interacts Genetically and Physically with the Core Structures of the Nuclear Pore Complex

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Abstract. We have isolated a major protein constituent from a highly enriched fraction of yeast nuclear pore complexes (NPCs). The gene encoding this protein, Nup188p, was cloned, sequenced, and found to be nonessential upon deletion. Nup188p cofractionates with yeast NPCs and gives an immunofluorescent staining pattern typical of nucleoporins. Using immunoelectron microscopy, Nup188p was shown to localize to both the cytoplasmic and nucleoplasmic faces of the NPC core. There, Nup188p interacts with an integral protein of the pore membrane domain, Pom152p, and another abundant nucleoporin, Nic96p. The effects of various mutations in the NUP188 gene on the structure of the nuclear envelope and the function of the NPC were examined. While null mutants of NUP188 appear normal, other mutants allelic to NUP188 exhibit a dominant effect leading to the formation of NPC-associated nuclear envelope herniations and growth inhibition at 37°C. In addition, depletion of the interacting protein Pom152p in cells lacking Nup188p resulted in severe deformations of the nuclear envelope. We suggest that Nup188p is one of a group of proteins that form the octagonal core structure of the NPC and thus functions in the structural organization of the NPC and nuclear envelope.

NUCLEOPORINS are the molecular constituents of the nuclear pore complex (NPC). Approximately 50 such individual nucleoporins (Rout and Blobel, 1993) constitute the complex 100 MD NPC structure and mediate multiple nuclear translocation processes. The core of the NPC reveals an octagonally symmetric architecture with several repetitious structural elements such as spokes, rings, and radial arms surrounding a central tube (termed "central plug") (Akey and Radermacher, 1993; Hinshaw et al., 1992). Filamentous structures attached to the core appear asymmetric with linear filaments extending into the cytoplasm and a nuclear cage projecting into the nucleoplasm (Unwin and Milligan, 1982; Jarnik and Aebi, 1991; Goldberg et al., 1992; Ris and Malecki, 1993). The nucleoporins so far characterized in yeast fall into two subgroups: those that contain repetitive peptide motifs (GLFG, XFXFG, or FG) and those that do not (for review see Rout and Wente, 1994; Doye et al., 1994; Kraemer et al., 1995; Gorsch et al., 1995; Pemberton et al., 1995; Li et al., 1995; Aitchison et al., 1995a,b; Heath et al., 1995). Among the nonrepeat nucleoporins is a further subgroup of major proteins including Nup170p, Nup157p, Nic96p, and the pore membrane protein Pom152p (Grandi et al., 1993; Wozniak et al., 1994; Aitchison et al., 1995a), which appear more abundant than the other constituents.

Repeat-containing nucleoporins are directly involved in the mechanics of nuclear translocation. In vitro assays have established that the repetitive peptide domains of these nucleoporins are docking sites for soluble transport factors (Rout et al., 1995; Moroi and Brokenbrow, 1995; Kraemer et al., 1995; Rexach and Blobel, 1995; Iovine et al., 1995). This observation is in accordance with the inhibition of specific nuclear transport functions associated with repeat nucleoporin mutants in vivo (reviewed in Rout and Wente, 1994; Fabre and Hurt, 1994). However, mutations in repeat nucleoporins not only affect transport processes, but can also cause structural defects such as NPC clustering and massive distortions of the nuclear morphology. It has therefore not been possible to distinguish between nucleoporins involved in transport and those responsible for the structural organization of the NPC, since both functions are closely intertwined (reviewed in Rout and Wente, 1994; Doye et al., 1994; Pemberton et al., 1994; Gorsch et al., 1995; Aitchison et al., 1995a,b).

With this background, further understanding of NPC function hinges on integrating subgroups of nucleoporins into the context of known NPC substructures. Apart from Pom152p (Wozniak et al., 1994), a major nonrepeat protein located in the nuclear pore-membrane domain of the nuclear envelope (NE) (and thus termed a pom), and Nup159p, a repeat nucleoporin (Kraemer et al., 1995; Gorsch
et al., 1995) that is exposed at the cytoplasmic face of the NPC, as yet no other NPC protein has been sublocalized in yeast. Some progress has been made in defining biochemical subcomplexes within the NPC through nearest neighbor analysis. One such subcomplex containing three repeat- (Nup49p, Nup57p, Nsp1p) and one nonrepeat nucleoporin, Nic96p (Grandi et al., 1995), has been identified in yeast. However, this subcomplex remains to be sublocalized within the NPC.

Here, we report the identification and characterization of a novel major nucleoporin, Nup188p. We demonstrate that Nup188p is located to both the nucleo- and the cytoplasmic faces of the NPC core, where it interacts genetically and physically with the pore membrane protein Pom152p. We show that Nup188p function is required for a colony-sectoring assay described in Aitchison et al. (1995a) with the following exceptions: p33, pSB32 (LEU2)

Materials and Methods

Yeast Manipulations and Plasmids

The strains used in this work are listed in Table I. All yeast manipulations were performed as outlined in Aitchison et al. (1995a). The plasmids used in this study have been previously described in Wozniak et al. (1994) and Aitchison et al. (1995a) with the following exceptions: p33, pSB32 (LEU2) containing the NUP188 locus; YepH and YepW (Wimmer et al., 1992); pNUP188, pUN100 that contains a genomic HindIII/BamHI fragment including the NUP188 gene; pNUP188CB, pUN100 (CEN/LEU2) containing NUP188 in which the STOP codon has been substituted by A, and restriction site (see below); pNUP188PA and pNUP188BPA, pUN100 (CEN/LEU2) containing protein A-tailed NUP188 (PA) or a truncated version of NUP188 tagged with protein A (BPA; see below); pPOM152-PA, the plasmid pPM1-HA to which the coding sequence of six histidine residues was added to the 3' end of the POM152 open reading frame (ORF); pPOM152-H, the plasmid pPOM152 to which the coding sequence of six histidine residues was added to the 3' end of the POM152 ORF; pBlueNUP188; pBluescript (Stratagene Cloning Systems, La Jolla, CA) containing a genomic HindIII/BamHI fragment including NUP188.

Table I. Yeast Strain Genotype

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Derivation</th>
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<tr>
<td>ps14</td>
<td>Mata ade2 ade3 ura3 his3 trpl leu2 con1 pom152-2::HIS3 nup188-4 pCH1122-POM152(ADE3-URA3)</td>
<td>Synthetic lethal derived from PM152-CP (Aitchison et al., 1995a)</td>
</tr>
<tr>
<td>ps144</td>
<td>Mata ade2 ade3 ura3 his3 trpl leu2 con1 pom152-2::HIS3 nup188-44 pCH1122-POM152(ADE3-URA3)</td>
<td>Synthetic lethal from PM152-CP</td>
</tr>
<tr>
<td>NP188Δ</td>
<td>Mata/Mata ade 2-1ade2-1 ura3-1ura3-1 his3-11,15/his3-11,15 trpl-1/hrpl-1 leu2-3,112/leu2-3,112 can1-100/can1-100 nup188-1::HIS3/+</td>
<td>Integrative transformation of W303 with linearized p33::HIS3</td>
</tr>
<tr>
<td>Y28</td>
<td>Mata ade2-1 ura3-1 trpl-1 leu 2-3 con1-100 nup188-2::HIS3</td>
<td>see Materials and Methods</td>
</tr>
<tr>
<td>CSL5</td>
<td>Mata ade2-1 ura3-1 leu2-3 con1 pom152-2::HIS3 nup 188-3::TRP1 pYESUra3-POM152 (URA3)</td>
<td>see Materials and Methods</td>
</tr>
<tr>
<td>NP188PA</td>
<td>Mata ade2-1 ura3-1 trpl-1 leu 2-3 con1-100 nup188-2::HIS3 pNUP188PA (LEU2)</td>
<td>Y28 transformed with pNUP188PA</td>
</tr>
<tr>
<td>NP188BPA</td>
<td>Mata ade 2-1 ura3-1 trpl-1 leu 2-3 con1-100 nup188-2::HIS3 pNUP188BPA (LEU2)</td>
<td>Y28 transformed with pNUP188BPA</td>
</tr>
<tr>
<td>P7AB-1</td>
<td>Mata/Mata ade2-1ade2-1 ura3-1ura3-1 his3-11,15/his3-11,15 trpl-1/hrpl-1 leu2-3,112/leu2-3,112 can1-100/can1-100 pom152-2::HIS3/pom152-2::HIS3 pom152-2 (LEU2)</td>
<td>Segregant of sporulated NP188Δ</td>
</tr>
<tr>
<td>NP188-2.4</td>
<td>Mata ade2-1 ura3-1 his3-11,15 trpl-1 leu2-3,112 can1-100 nup188-1::HIS3</td>
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<td>P7AB-B61H</td>
<td>Mata/Mata ade2-1ade2-1 ura3-1ura3-1 his3-11,15/his3-11,15 trpl-1/hrpl-1 leu2-3,112/leu2-3,112 can1-100/can1-100 pom152-2::HIS3/pom152-2::HIS3 pPOM152-HAH (LEU2)</td>
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diploid strains, including those derived from ps14 and ps144, failed to grow on 5-FOA-containing plates, suggesting the mutants are allelic to NUP188.

**Construction of NUP188 Null Mutations and the CSL5 Strain**

We have deleted and disrupted the NUP188 gene by integrative transformation using the procedure of Rothstein (1991). The majority of the NUP188 ORF was deleted by removing a NotI/Xbal fragment from the plasmid p33 and replacing it with the HIS3 marker gene derived from pGAL. The resulting plasmid (p33::HIS3) was linearized and transformed into the diploid strain W303 (Aitchison et al., 1995a). His+ transformants were selected on synthetic media (SM)-histidine plates, and heterozygous diploids carrying the p33::HIS3 disruption were identified by Southern blotting. A null strain, NP188A, was sporulated and tetrads were dissected. All of the resulting spores were viable and displayed a 2:2 segregation of the HIS3 marker. Southern blotting confirmed that the His+ haploids contained only the disrupted copy of NUP188.

Two additional disruptions of the NUP188 gene were also performed using procedures similar to those described for NP188A. The TRP1 and HIS3 marker genes were PCR amplified from the plasmids YepW and YepH, respectively, using primers containing 5' EcoRI sites. The resulting products were used to replace an internal 4.5-kb EcoRI fragment, the NUP188 ORF encoding to amino acid residues 99-1612 in the plasmid pBlueNUP188. The nup188::HIS3 construct was used to disrupt a chromosomal copy of NUP188 in the diploid strain W303. From the resulting cells, a haploid strain (Y28) containing the nup188::HIS3 was isolated. To create the CSL5 strain, the nup188::TRP1 disruption construct was transformed into a haploid strain PM152-75 (pom152Δ; Aitchison et al., 1995a) carrying POM152 under the control of the GAL1 promoter in the plasmid pYEur3-PM152 (Wozniak et al., 1994). Transformants were selected on galactose medium lacking tryptophan, and homologous integrations verified by Southern blotting.

**Isolation and Peptide Sequence of Nup188p**

Fractions derived from the SDS-hydroxyapatite chromatography of isolated yeast NPCs and containing the 170-kd species were pooled (corresponding to fraction 50-54 in Fig. 3 of Wozniak et al., 1994), and the polypeptide constituents were separated by SDS-PAGE. This gel revealed a closely spaced doublet at ~170 kD (see Fig. 1A). Polypeptides were electroblotted transferred to polyvinylidifluoride membrane, visualized with 0.1% Pierce red (in 1% acetic acid), and the faster migrating species was excised. Several internal peptides were sequenced using previously described procedures (Wozniak et al., 1994).

**Isolation and Sequencing of the NUP188 Gene**

The sequence of a peptide fragment derived from the faster migrating species of a prominent 170-kd double band on SDS-PAGE was used to design a pair of degenerate sense and antisense oligonucleotide primers encoding amino acid residues YLNSRI (residues 1611-1616 of Nup188p) and QQQLED (residues 1624-1629 of Nup188p), respectively. Following the method of Lee et al. (1988), these primers and Saccharomyces cerevisiae genomic DNA (Promega, Madison, WI) were used to PCR amplify the cDNA segment encoding the 19-amino acid residue peptide. From the sequence of the PCR product an oligonucleotide was designed corresponding to the nucleotide sequence between the primers. The oligonucleotide was labeled with υ[γ32P]ATP using polynucleotide kinase (Boehringer Mannheim, Mannheim, Germany) and used to screen a S. cerevisiae genomic library in plasmid Ycp50 (Rose et al., 1987) as described (Ausabel et al., 1992). Two clones were isolated. Plasmids derived from these clones contained overlapping inserts encoding a deduced ORF of 1655 amino-acid residues.

**Production of Nup188p- and Nup170p-specific Antiserum**

A cDNA fragment encoding 59- and 56-kD segments of Nup188p (amino acid residues 1041-1613) and Nup170p (amino-acid residues 697-1207), respectively, were synthesized using PCR and each fused to the 3' end of the IgG binding domains of Staphylococcus aureus protein A in the prokaryotic expression vector pRIT2T (Pharmacia Biotech Inc., Piscataway, NJ). Expression and purification of the resulting fusion proteins on IgG-Sepharose 6 fast flow (Pharmacia), were performed as described by the manufacturer. The purified protein A fusion proteins were used to elicit antibodies in rabbits, and the specificity of the resulting antiserum was examined by immunoblotting. Procedures for the blocking, probing (with anti-Nup188p), and the detection of antibody binding with the ECL detection system were performed essentially as instructed by the manufacturer (Amersham, Arlington Heights, IL).

**Epitope Tagging of NUP188**

DNA encoding five IgG-binding repeat units of S. aureus protein A were PCR amplified as described in Schnell and Blobel (1993), with the modification that the PCR primers contained 5' BamHI restriction sites. The STOP codon of the NUP188 gene in the plasmid pNUP188 was replaced by a BamHI site. This was achieved by PCR amplifying the 3' 1030-bp of NUP188 using the sense primer 5' AGAGATGACTTTATAAA 3' and the antisense primer 5' TTAGGATCAAGCTTGTAAATTGTA 3' containing a 5' BamHI site. This PCR amplified fragment was digested with restriction enzymes BglII and BamHI, and cloned into vector pNUP188, which had been digested with BglII and BamHI, thus releasing a 1-kb region of the NUP188 ORF including the STOP codon plus the 2.2-kb sequence downstream of NUP188. The resulting plasmid carrying NUP188 with a BamHI site instead of the STOP codon was termed pNUP188BC. Plasmid pNUP188BCB was then digested with BamHI and a PCR-amplified, BamHI-digested DNA fragment coding for the IgG-binding repeats of protein A was cloned into this novel BamHI site at the 3' end of NUP188 to create plasmid pNUP188PA. Alternatively, the IgG repeats were cloned into the plasmid pNUP188BC after digestion with BglII and BamHI, thus truncating the NUP188 at the BglII site and inserting protein A. The resulting plasmid was termed pNUP188BPA. Plasmids were transformed into a haploid nup188Δ yeast strain (Y28) for expression.

**Preparation of Yeast Extracts and Western Blot Analysis**

Whole-cell yeast extracts were prepared as described by Wozniak et al. (1994). Yeast subcellular fraction leading to the isolation of NPCs was performed as described by Rout and Blobel (1993). For Western blot analysis, proteins were separated by SDS-PAGE and then transferred electrophoretically onto nitrocellulose. Blots were probed with mAb118C3 (anti-Pom152p) as described (Strambio-de-Castillia et al., 1995).

**Immunoprecipitation of Pom152p-HA**

NEs were isolated from two yeast strains, P7AB-B61H (synthesizing hemagglutinin antigen [HA]-tagged Pom152p; Pom152p-HA) and P7AB-1 (lacking the HA tag), using the procedure of Strambio-de-Castillia et al. (1995). All of the following procedures were conducted at 4°C. NEs derived from 50 OD660 U of isolated nuclei were suspended in 0.5 ml of 10 mM bisTris, pH 6.5, 0.1 mM MgCl2, 0.2 mM PMSF. NEs were then disrupted with an equal volume of 2x solubilization buffer (10 mM NaCl, 2% Triton X-100, 1 mM EDTA, 25 mM Tris, pH 8.0, 0.01 mM DTT, 0.5 mM PMSF). After a 15-min incubation on ice, samples were centrifuged (90,000 g for 5 min), and the supernatant fraction was then preabsorbed with 1 μl mouse serum for 1 h followed by the addition of 50 μl of protein G-Sepharose 4 beads (Pharmacia Biotech Inc., Piscataway, NJ) for an additional hour to reduce nonspecific binding. As mAb (12CA5) directed against the HA tag (1 μl ascites fluid; Berkeley Antibody Co., Richmond, CA) was then added to the preclaraed NE extract and incubated for 3 h. Immune complexes were absorbed on 50 μl of protein G-Sepharose 4 beads, washed extensively with solubilization buffer, and eluted with 0.5 M acetic acid, pH 3.4. Aliquots of the starting nuclear extract, the eluted fraction, and the unbound fraction were then analyzed by SDS-PAGE, and Western blotting was performed as described above.

**Indirect Immunofluorescence and Electron Microscopy**

Immunofluorescence was performed as described by Kilmartin and Adams (1984) with modifications (Wente et al., 1992; Aitchison et al., 1995a).

For electron microscopy, cells were harvested in midlog phase, washed, and then fixed as described in Aitchison et al. (1995a). For the Pomp12p depletion experiments, CSL5 cells were grown in raffinose-containing medium (1% yeast extract, 2% bactopeptone, 2% raffinose) to midlogarithmic phase before washing and incubation in YPGal orYPD medium. Preembedding labeling immunoelectron microscopy was performed on...
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from the faster migrating band (Fig. 1 A), we cloned the
necessary for the localization to the NPC.

These data suggest that the carboxyl-terminal domain of Nup188p is
fully functional (data not shown). The subcellular distribu-
tions of both fusion proteins were examined by indirect
immunofluorescence. Nup188PAp (full-length Nup188p-
prot A) was visible in a punctate pattern on the surface of
prot A) was visible in a punctate pattern on the surface of
immunofluorescence microscopy [Kilmartin and Ad-
ams, 1984]). After washing, the bound antibody was visualized with an
overnight incubation of a 1:10 dilution of 10-nm gold anti-rabbit IgG
(Sigma Chemical Co., St. Louis, MO) in IEM buffer.

Results

Nup188p Is a Major Yeast Nucleoporin

Nucleoporins that form the repeating subunits of the NPC rings and spokes are expected to be abundant constituents of the isolated NPC fraction. SDS-PAGE analysis of the protein constituents of isolated yeast nuclear pore complexes (Rout and Blobel, 1993) reveals several major polypep-
tides (Rout and Blobel, 1993; Wozniak et al., 1994; Aitchison et al., 1995a). Four of these—Pom152p, Nup170p, Nup157p, and Nic96p—have been previously characterized (Grandi et al., 1993; Wozniak et al., 1994; Aitchison et al., 1995a). In addition to these polypeptides, two major protein species were observed, both with an apparent molecular mass of ~170 kD. Using internal peptide sequence information from the faster migrating band (Fig. 1 A), we cloned the NUP188 gene from a yeast genomic library as two overlapping clones that defined a 1655-amino acid residue ORF with a deduced mass 188,459 D. We have termed this gene NUP188, using conventional nucleoporin nomenclature. Deletion of the entire NUP188 ORF in a yeast haploid strain did not affect the growth of this strain at temperatures ranging from 16 to 37°C (see Materials and Methods; data not shown) or its nuclear morphology (see Fig. 5). This phenotype is similar to that observed in null mutations of POM152, NUP170, and NUP157.

A fusion protein containing amino-acid residues 1041–1613 of Nup188p was used to produce antibodies in rabbits which were found to react specifically with Nup188p (Fig. 1 A). A Western blot of sucrose gradient fractions derived from the yeast NPC isolation procedure revealed that Nup188p cofractionated absolutely with isolated NPCs (Fig. 1 B). We epitope tagged either full-length NUP188 or a truncated form of NUP188 ending at residue 1313 by fusing a DNA fragment encoding the IgG-binding domain of S. aureus protein A to the NUP188 carboxy-terminus. When analyzed for their ability to complement an inactive nup188 gene in a synthetically lethal background (see below), the full-length fusion protein (Nup188PAp), but not the truncated version (Nup188BPAp), was found to be fully functional (data not shown). The subcellular distributions of both fusion proteins were examined by indirect immuno-fluorescence. Nup188PAp (full-length Nup188p-prot A) was visible in a punctate pattern on the surface of the nucleus, characteristic of its localization to the NPC (Fig. 1 C). However, Nup188BPAp (truncated Nup188p) was primarily visible in the cytosol (Fig. 1 C). These data suggest that the carboxyl-terminal domain of Nup188p is necessary for the localization to the NPC.

Nup188p Is Associated with the NPC Core

Nuclear envelopes isolated from the protein A–tagged Nup188p strain (NP188PA) were probed with purified

![Figure 1. Nup188p is a nucleoporin. (A and B) Identification of Nup188p as a major constituent of isolated yeast NPCs. In A, polypeptides in a fraction containing a doublet of ~170 kD (derived from SDS-hydroxylapatite chromatography of isolated yeast NPC, see Wozniak et al. [1994], Fig. 3, fractions 43–45) were separated on an SDS 8–5% inverse polyacrylamide gradient gel and visualized by silver staining (Silver). The faster migrating species of the doublet (arrow) was microsequenced, its gene was cloned, and the protein was termed Nup188p. Western blots were performed on identical fractions using an antibody directed against Nup188p (α-Nup188p). In B, protein constituents of fractions derived from the sequential sucrose gradients used in the enrichment of yeast NPCs were separated by SDS-PAGE and Western blotted with antibodies directed against Nup188p. The number of starting cell equivalents per lane is exactly as defined in Fig. 1 of Wozniak et al. [1994]. Nup188p coenriched with the NPCs. Molecular mass markers are indicated in kilodaltons. (C) Localization of Nup188p and Nup188p tagged with protein A. Full-length Nup188p (Nup188PAp) and a carboxyl-terminal truncated Nup188p (Nup188BPAp) fused to the IgG-binding repeats of Staphylococcus protein A were localized by indirect immunofluorescence of whole yeast cells using rabbit anti–mouse IgG, followed by donkey anti–rabbit coupled to the fluorescent tag Cy3. Nup188PAp (full-length Nup188p) gives punctate NE staining, while truncated Nup188p (Nup188BPAp) localizes mainly to the cytoplasm. Corresponding 4',6-diamidino-2-phenylindole (DAPI) staining of DNA is shown below.]
rabbit antibody, and the binding was visualized with gold-labeled secondary antibody by electron microscopy. As has been previously demonstrated with Nup159p (Kraemer et al., 1995), the use of a preembedding immunolabeling technique made it possible to sublocalize Nup188p on the yeast NPC with greater accuracy than the previously used postembedding labeling methods (Fig. 2 A). The distance from the center of each gold particle (165 particles total) to the midplane of its associated NPC was measured for 104 NPCs clearly sectioned perpendicular to that plane (and hence to the plane of the NE) (Fig. 2 B). The gold was distributed equally on both the nucleoplasmic and cytoplasmic faces (Fig. 2, A and B). Most particles were found within 20 nm of the midplane of the NPC (20 ± 13 nm, n = 80, on the nucleoplasmic side; 18 ± 12 nm, n = 85, on the cytoplasmic side) (Fig. 2 B). Many NPCs were multiply labeled, both on the same face and on opposite faces (Fig. 2 A). Unfortunately, without three-dimensional reconstruction, the radial distribution of the gold cannot be as reliably quantified; however, many particles were apparently located close to the cylindrical axis of the NPC (Fig. 2 A).

Figure 2. Nup188p is symmetrically distributed on both faces of the NPC. (A) Localization of protein A–tagged Nup188p in isolated NEs, as visualized by the binding of rabbit IgG and gold-labeled secondary antibodies. The nuclear side of the envelope (N) is on the concave face with attached chromatin (Chr); the cytoplasmic face (C) reveals associated ribosomes (Rib). Nup188p appeared to be present in multiple copies and accessible from both sides of the NE, close to the central cylindrical axis of the NPCs. Bar, 0.1 μm. (B) Histogram quantification of the distribution of 165 gold particles on the NEs (count), measured as the distance (Distance (nm)) from the center of each gold particle to the midplane of its associated NPC. Particles to the cytoplasmic side of this midplane have positive distances, while particles to the nucleoplasmic side have negative distances. Most particles were found within 20 nm of either face of the NPC. A diagrammatic silhouette of a yeast NPC (drawn to scale) has been placed behind the histogram for reference.

Figure 3. Nup188p interacts genetically and physically with Pom152p. (A) The mutant strains psl4 and ps144 and parental strain PM152-CP (Aitchison et al., 1995a; PM75CP) were grown on 5-FOA-containing or YPD plates for 3 d at 30°C. The growth of each strain on 5-FOA was evaluated after the introduction of the plasmids containing POM152 (pPOM152), NUP188 (p33), or a control plasmid without insert (pRS315). Both psl4 and ps144 containing pRS315 fail to grow on 5-FOA. Growth on 5-FOA can be restored to both mutants by POM152 or NUP188. (B) NEs isolated from a strain expressing HA-tagged Pom152p (P7AB-B61H; +HA) and a strain expressing untagged Pom152p (P7AB-1; −HA) were extracted with Triton X-100-containing buffer and centrifuged. The supernatant fractions derived from both strains were immunoprecipitated with the anti-HA mAb 12CA5. Polypeptides in the initial supernatant fraction (Sp; 1 equivalent), the immunoadsorbed fraction (Ab; derived from 25 Eq of Sp), and the unbound fraction (Un; derived from 7.5 Eq of Sp) were separated by SDS-PAGE, transferred to nitrocellulose, and probed with antibodies directed against the HA tag (α-Pom152p-HA), Nup188p (α-Nup188p), Nic96p (α-Nic96p), and Nup170p (α-Nup170p). In each case the relevant region of the blot is shown. Nup188p and Nic96p, but not Nup170p, coimmunoprecipitated with Pom152p. None of these proteins were immunoprecipitated from the control strain lacking the HA tag (−HA).
Figure 4. Psl4 cells reveal a temperature-sensitive growth defect and NPC-associated herniations. (A) The psl4 strain (psl4), the psl4 strain containing the plasmid p33 (NUP188)(psl4+NUP188) and parental strain PM152-CP (P75CH) were grown on YPD plates for 3 d at 30°C (left). Only slight differences in the growth rates of the respective strains can be observed. Shown on the right is an electron micrograph of a thin-sectioned psl4 cell grown at 30°C in YPD. NPC-associated NE herniations are marked by arrowheads. Arrows point to fenestrations often visible in the herniations. (B) Shown on the left are the strains described in A grown on YPD plates for 3 d at 37°C. A significant delay in the growth rate of the psl4 strain can be observed at 37°C (psl4). This ts phenotype could be complemented by wild-type NUP188 (psl4+NUP188). Shown on the right is an electron micrograph of a thin-sectioned psl4 cell grown at 37°C for 9 h. NPC-associated NE herniations are marked by arrowheads. At this temperature, some herniations appear filled with electron-dense material. Bar, 0.5 μm.
Taken together, these data indicate that Nup188p-protein A is present in multiple copies and accessible on both faces of the NPC. Assuming that this localization reflects the position of the protein bulk, these data suggest that Nup188p is associated with the NPC core, perhaps as a component of the inner-spoke domain or the central transporter.

**Nup188p Interacts Genetically and Physically with Pom152p**

The NUP188 gene was also identified in a colony-sectoring assay designed to identify genes that are synthetically lethal with POM152 (for details see Materials and Methods and Aitchison et al., 1995a). Of the 32 recessive, non-sectoring mutants (termed psl mutants) obtained in this screen, 20 were allelic to NUP188. As shown in Fig. 3 A, two synthetically lethal mutants, psl4 and psl44, regained the ability to grow on 5-FOA medium and to sector on YPD medium after transformation with either the POM152 or NUP188 gene. Consistent with this observed synthetic lethality, a haploid strain lacking both the POM152 and the NUP188 genes was not viable (data not shown). The remaining 12 mutants originally obtained in the screen, which were not complemented by NUP188, fell into three other complementation groups; one set allelic to the major nucleoporin gene NUP170 (Aitchison et al., 1995a), a second set could be rescued by the nucleoporin Nic96p, and a third by an as yet uncharacterized plasmid termed p40 (M. Marelli and R.W. Wozniak, unpublished results).

The genetic interactions between POM152 and NUP188 suggested an overlapping role for these proteins in NPC function and/or their physical interaction. In an effort to determine whether in fact Pom152p and Nup188p physically interact, we performed immunoprecipitations on nuclear envelope extracts derived from strains synthesizing HA-tagged Pom152p (Pom152p-HA). Nuclear envelopes were disrupted under conditions that release the majority of Pom152p-HA (~80%) into a soluble supernatant fraction after centrifugation (data not shown). Pom152p-HA was immunoprecipitated from this fraction using an mAb (12CA5) specific for the HA epitope. Analysis of the immunoprecipitated fraction by Western blotting revealed that Nup188p and Nic96p, but not Nup170p, were coimmunoprecipitated with Pom152p-HA (Fig. 3 B, +HA). None of these proteins were detected in parallel immunoprecipitates derived from the strain lacking HA-tagged Pom152p (Fig. 3 B, −HA). These data suggest that Pom152p is physically associated, either directly or indirectly, with Nup188p and Nic96p in an NPC subcomplex lacking Nup170p. We note that a major portion of Pom152p-HA, Nup188p, and Nic96p are not immunoprecipitated with the 12CA5 mAb, suggesting that this complex may be largely inaccessible to the antibody and/or unstable under the conditions we have used to dissociate it from the NE.

**A nup188Δ Mutant, psl4, Reveals Structural Abnormalities of the NE without Affecting Nuclear Transport**

We have examined the growth of a nup188 mutant strain, psl4 (see Fig. 3 A), at various temperatures ranging from 23°C to 37°C. While the doubling time of the psl4 cells at 30°C (2.4 h) was only slightly slower than the PM152-75 parent strain (2.1 h), growth was inhibited at 37°C (Fig. 4 B, left panel). This growth defect could be rescued by introducing wild-type NUP188 (Fig. 4 B, left panel). Mutant cells (psl4) were either maintained at 30°C or shifted to 37°C for 9 h and then processed for thin-section electron microscopy. At both temperatures, the morphology of the nucleus and NE appeared essentially identical (Fig. 4, A and B, right panels), with many NPCs being enclosed on their cytoplasmic face by NE herniations similar to structures previously described by Wente and Blobel (1993). No apparent increase in the frequency of the NPC-associated herniations was observed in cells grown at 37°C. Cells grown at 37°C, but not those grown at 30°C, contained electron-dense material within several of these herniations (Fig. 4, A and B, right panels). This would suggest that at 37°C, the herniations are accumulating material being exported from the nucleus. However, when mRNA export was analyzed at both 30°C and 37°C by in situ hybridization using oligo-dT, no apparent accumulation of mRNA in the nucleus or at the nuclear periphery could be detected at either temperature (data not shown). Also, nuclear import, as monitored by the subcellular distribution of the nuclear protein NSR1 (Lee et al., 1992), was not affected at either temperature (data not shown).

**Pom152p Depletion in a nup188Δ Genetic Background Leads to Structural Distortions of the Nucleus**

The effects of the conditional lethality observed between the NUP188 and POM152 null mutants on the structure of the nucleus were examined in a genetic background lacking the chromosomal copies of POM152 and NUP188, but carrying the POM152 gene, under the control of the regulatable GAL1 promoter in the plasmid pYEUra3 (URA3). The viability of the resulting GAL1::pom152, nup188Δ mutant strain, CSL5, was expected to be dependent on GAL1-controlled POM152 expression. When we analyzed the growth rate of the CSL5 strain, we observed a significant increase in the doubling time upon a shift to glucose-containing medium (2.2 h in galactose to 5 h in glucose). Surprisingly, even after 30 h of growth in glucose, CSL5 cells were still viable. The survival of the CSL5 strain in glucose medium after 30 h was most likely caused by incomplete repression of the GAL1 promoter. This is supported by Western blots of CSL5 cells grown for up to 30 h in glucose, in which detectable amounts of Pom152p were visible (Fig. 5 A). The low level of Pom152p detectable after a 30-h depletion was apparently sufficient to support slow growth.

When examined for morphological changes, CSL5 cells grown in galactose appeared normal (Fig. 5 C, +Galactose). However, after 30 h depletion of Pom152p in glucose medium, the shape of the nucleus became severely distorted, losing its rounded shape (Fig. 5 D, +Glucose). Projections of the NE were visible extending into the cytosol. In addition, the NE appeared to form extensive invaginations into the nucleus. These distortions in the shape of the nuclear envelope are similar to those that were previously observed upon depletion of Nup170p in cells lacking Pom152p (Aitchison et al., 1995a). Unlike the Nup170p depleted strain, however, we observed no apparent decrease in the number of NPCs.
Discussion

Nup188p belongs to a subgroup of NPC constituents that do not contain peptide repeat motifs and together constitute a significant percentage of the NPC mass. We refer to this subgroup of nucleoporins as "major nonrepeat nucleoporins." This classification goes beyond a descriptive level, since the other members of this subgroup, including Nup170p, Nup157p, and Nic96p, have previously been shown to genetically interact with each other and Pom152p (Aitchison et al., 1995a; Zabel et al., 1996). This genetic interaction has its basis in either a physical interaction and/or an overlap in the respective functions of these proteins. Accordingly, Nup188p was identified both as a major protein in a highly enriched pore complex preparation and as a mutant that is synthetically lethal with Pom152p.

The localization of Pom152p to the pore membrane domain established it as a constituent of the NPC core (Wozniak et al., 1994). Nup188p can be assigned to the same NPC substructure. Using immunoelectron microscopy, Nup188p was found symmetrically distributed at both cytoplasmic and nucleoplasmic faces of the NPC. The close proximity of the gold label to the central plane of the NPC strongly suggests that Nup188p is a component of the NPC core, most likely the spokes or the central transporter. Moreover, using HA-tagged Pom152p, Nup188p could be specifically immunoprecipitated from NEs using HA antibodies, demonstrating that these proteins are physically associated. Thus, Pom152p and Nup188p are two major constituents of the NPC core domain that interact both physically and genetically. Interestingly, these interactions also extend to other major nonrepeat nucleoporins. As can be seen from Fig. 3B, Pom152p also appears associated with Nic96p, and in an accompanying study by Zabel et al. (1996), Nup188p was shown to interact with Nic96p. A physical interaction between Pom152p and Nic96p is consistent with previous data showing their genetic interactions (Aitchison et al., 1995a). These results strongly suggest that the genetic interactions between major nonrepeat nucleoporins are based on their physical interaction. It is therefore conceivable that this entire subgroup of major
nucleoporins is involved in forming a common NPC substructure, such as the octagonal core, with its repetitive spoke, ring, and radial arm structures.

A role of Nup188p in the structural organization of the NPC core should ideally correlate with structural NPC defects of nup188 mutants in vivo. One nup188 mutant allele analyzed for morphological abnormalities is psl4. It exhibited a temperature-sensitive phenotype that could be rescued by wild-type NUP188. This growth defect is in contrast to a nup188 null mutation that showed no growth abnormalities. Thus, it is likely that the product of the psl4 allele is more detrimental to NPC function than the complete removal of the wild-type protein. An examination of the morphology of the NE in the psl4 strain at 30° and 37°C revealed the presence of NPC-associated NE herniations similar to those observed in nup116 null mutations at restrictive temperatures (Wente and Blobel, 1993). In contrast to the NUP116 mutants, however, the herniations seen in Fig. 4 were free of electron-dense material at 30°C, and revealed such accumulation only at 37°C. Although this material in the herniations may represent some trapped nuclear transport cargo, no significant inhibition of nuclear transport could be detected in psl4 cells at either temperature (data not shown). The nup188 mutant in psl4 therefore appears to primarily affect the structural organization of the NPC rather than having a direct impact on nuclear transport mechanisms.

Upon depletion of a second NPC core component, Pom152p, in cells already lacking Nup188p, structural distortions of the nucleus became more profound. They were no longer confined to the area of the NPC, but rather affected the structural integrity of the whole nuclear envelope. As shown in Fig. 5, the gradual depletion of Pom152p in cells lacking Nup188p led to severe distortions in the shape of the nucleus which are visible as extended projections and invaginations of the NE. It is unlikely that these structural distortions of the NE are the pleiotropic result of dying cells, since the depletion of Pom152p was not complete and the cells continued to grow. Again, the persistence of growth implies a continuation of nuclear transport mechanisms despite a highly compromised NE structure.

Taken together, these experiments suggest that Nup188p is a component of the NPC core and has an important role in the structural organization of the NPC and the NE. We hypothesize that Nup188p, as well as the other major non-repeat nucleoporins and Pom152p, represent a subgroup of NPC proteins that help to constitute the core of the NPC. This core may represent the structural framework that is required for the correct positioning of repeat nucleoporins and other NPC components directly involved in nuclear transport mechanisms. The only major nonrepeat nucleoporin shown to undergo genetic and physical interactions with repeat nucleoporins is Nic96p (interacting with Nsp1p, Nup49p, and Nup57p [Grandi et al., 1995]). In the context of data presented in this paper and Zabel et al. (1996), Nic96p could therefore link this repeat nucleoporin subcomplex to the NPC core, effectively connecting the elements of the transport machinery to the structural framework of the NPC and, via Pom152p, to the pore membrane domain.

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