The Fission Yeast Nup107-120 Complex Functionally Interacts with the Small GTPase Ran/Spi1 and Is Required for mRNA Export, Nuclear Pore Distribution, and Proper Cell Division

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We have characterized *Schizosaccharomyces pombe* **open reading frames encoding potential orthologues of constituents of the evolutionarily conserved** *Saccharomyces cerevisiae* **Nup84 vertebrate Nup107-160 nuclear pore subcomplex, namely Nup133a, Nup133b, Nup120, Nup107, Nup85, and Seh1. In spite of rather weak sequence conservation, in vivo analyses demonstrated that these** *S. pombe* **proteins are localized at the nuclear envelope. Biochemical data confirmed the organization of these nucleoporins within conserved complexes. Although examination of the** *S. cerevisiae* **and** *S. pombe* **deletion mutants revealed different viability phenotypes, functional studies indicated that the involvement of this complex in nuclear pore distribution and mRNA export has been conserved between these highly divergent yeasts. Unexpectedly, microscopic analyses of some of the** *S. pombe* **mutants revealed cell division defects at the restrictive temperature (abnormal septa and mitotic spindles and chromosome missegregation) that were reminiscent of defects occurring in several** *S. pombe* **GTPase Ran (RanSp)/Spi1 cycle mutants. Furthermore, deletion of** *nup120* **moderately altered the** nuclear location of Ran^{Sp}/Spi1, whereas overexpression of a nonfunctional Ran^{Sp}/Spi1-GFP allele was specif**ically toxic in the** *nup120* **and** *nup133b* **mutant strains, indicating a functional and genetic link between** constituents of the *S. pombe* **Nup107-120** complex and of the Ran^{Sp}/Spi1 pathway.

Traffic of macromolecules between the nuclear and cytoplasmic compartments, which is fundamental in eukaryotic cells, occurs through the nuclear pore complexes (NPCs), which are macromolecular assemblies embedded in the nuclear envelope (NE). Active nucleocytoplasmic transport of most macromolecules involves a series of interactions among nuclear pore proteins (nucleoporins), soluble transport factors (karyopherins), and cargos that are modulated by the small Ras-like GTPase Ran (reviewed in references 39 and 55). In contrast, export of most spliced mRNA appears to be independent of the Ran-importin β machinery (12, 33). In addition to regulating nuclear transport, Ran plays additional and independent roles in many other cellular processes, including microtubule dynamics and mitotic-spindle formation, regulation of cell cycle progression, and postmitotic NE and NPC assembly (3, 15, 28, 31, 35, 52, 54, 69).

In recent years, extensive progress has been made in the identification and characterization of all NPC components, using proteomic and genomic approaches. *Saccharomyces cerevisiae* and vertebrate NPCs are composed of roughly 30 nucleoporins, of which about two-thirds have been conserved during evolution (14, 51). While the molecular dissection of NPCs is nearly complete in these organisms, only a few nucleoporins have been definitively identified so far in fission yeast.

As *Schizosaccharomyces pombe* and *S. cerevisiae* are evolutionary distant, interspecies comparisons should allow the dissection of conserved functions with more precision, thereby improving our understanding of NPC function during evolution. Indeed, comparisons of a few *S. cerevisiae* and *S. pombe* nucleoporin orthologues have revealed unexpected functional divergences (71, 73).

In this paper, therefore, we have undertaken the functional analysis of the *S. pombe* orthologues of *S. cerevisiae* Nup84 (ScNup84)/vertebrate Nup107-160 complex constituents. In *S. cerevisiae*, this complex is composed of Nup84, Nup85, Nup120, Nup145-C, Seh1, Sec13, and Nup133 (2, 38, 58, 59, 65). In this organism, single mutations in members of the Nup84 complex are not lethal (although double mutations cause synthetic lethality), but most of them lead to a temperature-sensitive growth defect, to nuclear $poly(A)^+$ RNA accumulation at the restrictive temperature, and to NPC clustering at all temperatures. The vertebrate counterpart of the ScNup84 complex, named the Nup107-160 complex (7, 67), has also been demonstrated to be involved in mRNA export and NPC assembly (9, 28, 67, 68). Our study revealed major differences in the viability phenotypes of the corresponding *S. cerevisiae* and *S. pombe* deletion mutants and some differences in the behaviors of these nucleoporins during biochemical fractionation. Functional studies also suggested that the involvement of the complex in NPC distribution and $poly(A)^+$ RNA export has been at least partly conserved between these highly divergent yeasts. However, our study also revealed cell division defects in some of these *S. pombe* mutant strains. The involve-

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Strain		Genotype	Source
SP826	h^+/h^+	ade6-210/ade6-216 ura4-D18/ura4-D18 leu1-32/leu1-32	D. Beach
FC420	h^+	ade6-216 ura4-D18 leu1-32	F. Chang
FC418	h^-	ade6-210 ura4-D18 leu1-32	F. Chang
SPV21	h^+/h^+	$nup107^{+}/\Delta nup107$::kanMX ade6-210/ade6-216 ura4-D18/ura4-D18 leu1-32/leu1-32	This study
SPV25	h^+/h^+	nup133a ⁺ / Δ nup133a::kanMX ade6-210/ade6-216 ura4-D18/ura4-D18 leu1-32/leu1-32	This study
SPV94	h^+/h^+	$nup85^{+}/\Delta nup85^{+}/\Delta nup85^{+}/\Delta nMX$ ade6-210/ade6-216 ura4-D18/ura4-D18 leu1-32/leu1-32	This study
SPV97	h^+/h^+	$nup120^{+}/\Delta nup120$::kanMX ade6-210/ade6-216 ura4-D18/ura4-D18 leu1-32/leu1-32	This study
SPV560	h^+/h^+	$\frac{1}{2}$ seh1 ⁺ / Δ seh1:: $\frac{1}{2}$ kanMX ade6-210/ade6-216 ura4-D18/ura4-D18 leu1-32/leu1-32	This study
SPV34	h^+	Anup133a::kanMX ade6-210 ura4-D18 leu1-32	This study
SPV53	h^-	Δnup133b::kanMX ade6-210 ura4-D18 leu1-32	This study
SPV56	h^-	∆nup133a::kanMX nup133b∆::kan' ade6-216 ura4-D18 leu1-32	This study
SPV113	h^+	Δ nup120:: k anMX ade6-210 ura4-D18 leu1-32	This study
SPV561	h^+	Aseh1::kanMX ade6-210 ura4-D18 leu1-32	This study
SPV33	h^+	nup107-GFP(S65T)::kanMX ade6-216 ura4-D18 leu1-32	V. Doye
SPV477	h^-	kanMX-P3nmt1-GFP(S65T)-nup85 ade6-210 ura4-D18 leu1-32	This study
SPV639	h^+	kanMX-P3nmtl-GFP(S65T)-seh1 ade6 ⁻ ura4-D18 leu1-32	This study
SPV76	h^2	nup107-GFP(S65T)::kanMX \psqr133a::kanMX ade6 ⁻ ura4-D18 leu1-32	This study
SPV72	$h^?$	nup107-GFP(S65T)::kanMX Δ nup133b::kanMX ade6 ⁻ ura4-D18 leu1-32	This study
SPV584	h^2	$nup107-GFP(S65T):: kanMX \Delta nup120::kanMX + pREP4X-HA3-Nup120 ade6^- leu1-32$	This study
SPV574	h^2	kanMX-P3nmt1-GFP(S65T)-nup85 Δ nup133a::kanMX ade6 ⁻ ura4-D18 leu1-32	This study
SPV572	h^2	kanMX-P3nmt1-GFP(S65T)-nup85 \psqtp133b::kanMX ade6 ⁻ ura4-D18 leu1-32	This study
SPV583	h^2	$kanMX-P3nmt1-GFP(S65T)-nup85 \Delta nup120::kanMX + pREP4X-HA3-Nup120 ade6^{-}leu1-32$	This study
SPV697	h^2	k anMX-P3nmt1-GFP(S65T)-seh1 Δ nup133a:: k anMX ade6 ⁻ ura4-D18 leu1-32	This study
SPV701	h^2	k anMX-P3nmt1-GFP(S65T)-seh1 Δ nup133b:: k anMX ade6 ⁻ ura4-D18 leu1-32	This study
Δ nup184	h^+	Δ nup184::ura4 ⁺ ade6 ⁻ leu1-32	R. Dhar
Δ mad 2	h^-	Δ mad2::ura4 ⁺ ade6-210 ura4-D18 leu1-32	S. Sazer
Δ mph1	h^-	Δ mph1::ura4 ⁺ ade6-216 ura4-D18 leu1-32	S. Sazer
SPV526	h^2	∆nup120::kanMX ∆mad2::ura4 ade6 ⁻ leu1-32 ura4-D18	This study
SPV531	h^2	Δ nup120:: k anMX Δ mph1::ura4 ade6 ⁻ leu1-32 ura4-D18	This study
YP10.22a	h^+	Ch^{16} (ade6-216); ade6-210 ura4-D6 leu1-32	M. Yanagida
SPV323	h^2	$\Delta nup120$::kanMX + pREP4X-HA ₃ -Nup120 Ch ¹⁶ (ade6-216) ade6-210 leu1-32	This study
SPV327	h^2	$\Delta nup120$:: $kanMX + pREP4X-HA_3-Nup120$ ade6-210 leu1-32	This study
$NC102 + p(nuc2^{+})$	h^-	$nuc2-663 + pLEU2-(nuc2^{+})$ leu1-32	M. Yanagida
MA217	h^+	nuf2-CFP:: k anMX ura4-D18 leu1-32 his2 ⁻	T. Toda
SPV830	h^2	kanMX-P3nmt1-GFP(S65T)-nup85 nuc2-663 Nuf2-CFP::kanMX ura4-D18 leu1-32	This study

TABLE 1. Fission yeast strains used in this study

ment of the *S. pombe* Ran (Ran^{Sp})/Spi1 pathway with respect to the various phenotypes associated with perturbations of the functions of the *S. pombe* Nup107-120 (SpNup107-120) complex is discussed.

MATERIALS AND METHODS

BLAST searches at the Sanger Center *S. pombe* server were used to identify open reading frames (ORFs) encoding proteins similar to *S. cerevisiae* or murine nucleoporins. Sequence comparisons were performed using ClustalW (http: //clustalw.genome.ad.jp/).

S. pombe **strains, media, and genetic techniques.** The strains used in this study are listed in Table 1. Standard cell culture procedures and media were used (44). The strains were grown in rich nonselective yeast extract medium (YE5S) or in synthetic Edinburgh minimal medium (EMM), appropriately supplemented. Sporulation was performed in malt extract medium at 25 or 30°C. Yeast transformation was achieved by the lithium acetate-dimethyl sulfoxide method (6). The thiamine-regulable *nmt1* promoter, in expression vector pREP (43), was repressed by the addition of 5 μ g of thiamine/ml of EMM. Phloxine B solution (Bio 101 Systems, Qbiogen, Carlsbad, Calif.) was used at 0.25 ml/liter. Sensitivity to thiabendazole (TBZ) (Sigma) was monitored at 23°C by 10-fold serial-dilution colony spotting on YE5S plates containing the microtubule drug at the appropriate concentrations.

S. pombe **disruption and genomic green fluorescent protein (GFP)-tagging strategy for** *nup107***,** *nup85***,** *nup120***,** *nup133a***,** *nup133b***, and** *seh1***.** One-step gene disruption (6) was carried out in an h^+/h^+ diploid strain by homologous integration of a PCR fragment corresponding to the KanMX6 module (conferring G418 resistance) in the *nupX* ORFs of interest (*nup107*, *nup85*, *nup120*, *nup133a*, *nup133b*, or *seh1*). YE5S containing 100 or 200 mg of G418/ml was used for selecting G418^R cells, and correct gene targeting was confirmed by PCR analysis. After induction of meiosis, h^+/h^{90} $\Delta nupX$:: $kanMX6/nupX^+$ sporulating colonies were identified by iodine exposure, and tetrads were microdissected. N-terminal tagging of Nup85 and Seh1 expressed under the control of the *nmt1* promoter in their chromosomal locations was accomplished by an analogous method (6) using the pFA6a-kanMX6-P3nmt1-GFP vector.

Plasmids. An XhoI-SmaI fragment corresponding to the *nup133a* ORF was first PCR amplified from genomic DNA using the Sp133a-Xho1F and Sp133a-SmaIR oligonucleotides (sequences of all oligonucleotides used are available upon request) and cloned into pBluescript. The XhoI-SmaI fragment was then subcloned into the pREP3 vector (43), in which the GFPmut2 ORF (13) had been previously inserted as an XhoI/SalI fragment, to generate pREP3- GFP(mut2)-Nup133a and into the pSLF173 vector (21) digested with SalI and SmaI to generate pREP4X-HA₃-Nup133a.

A cDNA encoding the N-terminal domain of Nup133b and obtained by reverse transcription-PCR and a DNA fragment corresponding to the C-terminal domain of Nup133b, PCR amplified from genomic DNA, were ligated in pASΔΔCyHA to generate pASΔΔCyHA-Nup133b (7). For the construction of pREP4X-GFP-Nup133b or pREP4X-HA3-Nup133b, an NcoI-blunt-ended/SmaI fragment of pASCyHA-Nup133b was cloned at the SmaI site of pSGP573 (http://pingu.salk.edu/ \sim forsburg/vectors.html) or pSLF173, respectively.

The *nup120* ORF was PCR amplified from genomic DNA and was cloned in the pSLF173 vector. To construct the pREP4X-GFP-Nup120 plasmid, the NotI-SmaI insert of pREP4X-HA₃-Nup120 was subcloned into pSGP573.

The NLS-GFP-LacZ-containing fragment from the *S. cerevisiae* plasmid pPS817 (36) was subcloned into pREP4X (20). Plasmids pREP82X-GFP-*nsp1* (74), pREP3X-Spi1 (42), and pREP41X-Spi1-GFP (63) were previously described.

Immunoprecipitation experiments. *S. pombe* $\Delta nupX/GFP$ -tagged NupY strains expressing HA_3 -tagged $nupX$ ($nupX$ is $nup133a$, $nup133b$, or $nup120$; $nupY$ is *nup85*, *nup107*, or *seh1*) were constructed by genetic crossing. The strains were grown at 30°C to log phase in supplemented EMM in the presence of thiamine to avoid overexpression of the tagged nucleoporins. For whole-cell lysate prep-

^a The name and predicted mass (in kilodaltons) of each nucleoporin are indicated. To avoid additional nomenclature, these proteins have been named according to the first homologous nucleoporin identified. The *S. pombe* GeneDB systematic names are also specified (http://www.genedb.org/genedb/pombe/index.jsp). To fit the *S. pombe* gene nomenclature, SpNup133a and SpNup133b also a

Percentages of identity and similarity among S. pombe (Sp), S. cerevisiae (Sc), and Homo sapiens (Hs) nucleoporins were calculated using ClustalW (http: //clustalw.genome.ad.jp/). Alignments of *S. cerevisiae*, *S. pombe*, and human Nup107, Nup85, Seh1 (unpublished data), Nup133, and Nup120/160 (7) indicate that the similarity is spread over the entire sequence and does not allow the identification of functional domains within these nucleoporins.

aration, \sim 10 9 cells were washed in stop buffer and resuspended in immunoprecipitation buffer as previously described (60), with the following changes: Triton was replaced by 1% NP-40, and aprotinin and pepstatin were replaced by a protease inhibitor cocktail (Roche). The cells were then broken with glass beads in a Fastprep (Bio 101 Systems) and spun, and the supernatants were recovered. Protein G-Sepharose beads (Amersham Biosciences) were loaded with anti-GFP antibodies (Roche) in A buffer (50 mM HEPES, pH 7.5, 100 mM NaCl, 1 mM EDTA) at 4°C for 2 h. After four washes in A buffer, the beads were incubated with the extracts at 4°C for 3 h, washed five times in A buffer supplemented with 1% NP-40 and then once in A buffer, and resuspended in loading buffer. The resulting samples were analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis followed by Western blotting using anti-GFP or -HA (12CA5) antibodies. The antibodies were detected using horseradish-peroxidase-conjugated goat anti-mouse secondary antibodies. Immunoblots were visualized using enhanced-chemiluminescence reagents (Pierce).

For mass spectrometry analysis, after coloration with colloidal Coomassie blue (G250; Bio-Rad), the protein spots were excised from the gel, reduced using dithiothreitol, and alkylated using iodoacetamide. The proteins were subjected to digestion with trypsin (Sigma) following the protocol of Shevchenko et al. modified by overnight digestion at 30°C (57). The extracted peptides were analyzed by matrix-assisted laser desorption-ionization–time of flight mass spectrometry (Voyager-DETM PRO; Applied Biosystems). Peptide masses obtained by this analysis were used to search the National Center for Biotechnology Information database to identify the full-length proteins.

Fluorescence and electron microscopy. Fluorescence in situ hybridization (FISH) for poly $(A)^+$ RNA was performed as described by Brown et al. (10), with the following modifications: spheroplasts were obtained by digestion with 0.1 mg of Zymolyase-100T (Seikagaku)/ml in PEM buffer {50 mM PIPES [piperazine-*N*,*N'*-bis(2-ethanesulfonic acid)], pH 6.9, 0.5 mM EGTA, 0.5 mM MgSO₄} plus 1 M sorbitol, and the 50-nucleotide oligo(dT) probe was directly coupled to Cy3 (Amersham Pharmacia Biotech). To detect the distribution of the pre-60S ribosomal particles, FISH was also performed using a probe complementary to the 25S rRNA, conserved from *S. cerevisiae* to *S. pombe* (5' CCCGTTCCCTTGGC TGTGGTTTCGCTAGAT*A 3', where the asterisk represents amino-purified nucleotides coupled with Cy3) (26).

Stainings of the cell walls and septa and the nuclei of formaldehyde-fixed cells were carried out using calcofluor (2 μ g/ml; Sigma) and DAPI (4',6'-diamidino-2-phenylindole; 1 μ g/ml) (48). Tubulin immunofluorescence of formaldehydefixed cells was performed as described previously (48), using a TAT1 monoclonal antibody (72) together (when indicated) with an anti-Mid1 rabbit antibody (48). Spi1p was localized using affinity-purified rabbit anti-Spi1p antibody (41), and NPCs were visualized using mAb414 (BAbCO) as previously described (16).

Images of fixed cells and of live cells expressing GFP- or cyan fluorescent protein (CFP)-tagged proteins were acquired with an upright motorized widefield microscope (DMRA2; Leica), using an oil immersion objective ($100 \times PL$ APO HCX; 1.4 numerical aperture) and a high-sensitivity cooled interlined charge-coupled device camera (CoolSnap HQ; Roper). For some observations, stacks were acquired using a piezo-electric motor (LVDT; Physik Instrument) mounted underneath the objective lens. The whole system was controlled with Metamorph 5 software (Universal Imaging Corp.). Z sections of DAPI signals (see Fig. 7) were converted into single two-dimensional images by taking the maximum signal at each pixel position in the images. The length of the mitotic spindle (see Fig. 8) was determined based on a two-dimensional projection of Z sections, and used to determine approximate classes as follows: premetaphase,

 $\langle 2 \mu m;$ metaphase, 2 to 3 μm ; early anaphase, 3 to 5 μm ; late anaphase, $>$ 5 μm . For deconvolution (see Fig. 5A and 9D), image stacks were acquired without camera binning, with a plane spacing of $0.2 \mu m$, leading to a voxel size of 64.5 by 64.5 by 200 nm³ . Deconvolution was performed by the three-dimensional deconvolution module from Metamorph (Universal Imaging Corp.), using the fast iterative constrained PSF-based algorithm. Images in Fig. 10A were obtained with a Zeiss Axioskop microscope, and digitized images were captured with a DVC 1300 charge-coupled device camera and QED software.

Thin-section electron microscopy was performed as described previously (49), using cells freshly grown overnight on YE5S plates at semipermissive temperature (30°C).

Fluorescence-activated cell-sorting (FACS) analysis. The DNA content was analyzed as described previously (62), using a Becton Dickinson FACSCalibur with the following modifications: buffer A was replaced by 50 mM sodium citrate, pH 7.5, and RNase A was added to a final concentration of 0.5 mg/ml and incubated at 37°C for 3 h. Propidium iodide was then added to a final concentration of 5μ g/ml.

Chromosome loss assays. The YP10.22 strain, containing a 550-kb minichromosome (47), was crossed with a haploid $\Delta nup120$ strain carrying the pREP4X-HA₃-Nup120 plasmid (*ura4*⁺ marker). After dissection, a Δ*nup120* strain carrying pREP4X-HA₃-Nup120 and containing or lacking the minichromosome were isolated and then spotted at 23°C onto EMM plates supplemented with 1 mg of 5-fluoroorotic acid (5-FOA)/ml and 50 mg of uracil/liter to select for cells able to lose the $ura4^+$ -containing plasmid. Chromosome loss was also monitored in *ade6-M210/ade6-M216 h*⁺/h⁺ diploid strains. For scoring Ade⁻ phenotypes, cells from 10 small colonies were plated after dilution on yeast extract medium and cultured at 30 or 36°C for \sim 3 days. The percentage of pink colonies (chromosome loss event) was determined, taking the average of four independent experiments.

RESULTS

The Nup107 complex is conserved in fission yeast. By searching public databases, several *S. pombe* ORFs encoding proteins with moderate sequence similarity to constituents of the ScNup84 and the human Nup107-160 complexes were identified. They included putative homologues of *S. cerevisiae* Nup133 (designated Nup133a and Nup133b), Nup120, Nup84, Nup85, Nup145-C, and Seh1 (Table 2) (7, 14, 66, 67). Because of the low levels of sequence similarity between these *S. pombe* proteins and their *S. cerevisiae* and human counterparts (Table 2), we first asked whether these proteins were bona fide nucleoporins. It was previously reported that Nup107-GFP was localized at the NE (66) (Fig. 1). Similarly, in vivo analysis of GFP-tagged Nup85, Seh1, Nup120, Nup133a, and Nup133b revealed punctate rim-like staining around the nucleus, which is typical of NPC localization (Fig. 1).

To determine whether the organization of these nucleoporins in an NPC-subcomplex has been conserved in *S. pombe*,

FIG. 1. In vivo localization of GFP-tagged Nup107, Nup85, Seh1, Nup120, Nup133a, and Nup133b in *S. pombe*. Nup107-GFP, GFP-Nup85, and GFP-Seh1 strains were grown in YE5S medium. *nup120*, *nup133a*, and *nup133b* strains expressing GFP-Nup120, GFP-Nup133a and GFP-Nup133b fusion proteins, respectively, were first grown in appropriately supplemented EMM containing thiamine and then shifted overnight to EMM without thiamine. The distribution of the various GFP fusions was observed in live cells. All the fusion proteins localized to the nuclear periphery in a ring-like staining pattern characteristic of nucleoporins. Note that under these growth conditions, the level of the nucleoporins expressed under the control of the *nmt1* promoter is similar to that of Nup107 expressed under the control of its own promoter. Bar, $10 \mu m$.

immunoprecipitation experiments were performed using anti-GFP antibodies on total extracts from *S. pombe* cells expressing Nup107-GFP, GFP-Nup85, GFP-Seh1, or no GFP fusion in strains with *nup133a*, *nup133b*, or *nup120* deleted and complemented by plasmids encoding the corresponding HAtagged nucleoporins (Table 1). As shown in Fig. 2A, this study revealed that Nup107-GFP efficiently coprecipitated with HA-Nup133a and HA-Nup133b, confirming previous results based on two-hybrid interactions (7), but did not significantly interact with HA-Nup120. Conversely, GFP-Nup85 and GFP-Seh1 efficiently copurified with HA-Nup120 but not with HA-Nup133a or with HA-Nup133b.

Silver and colloidal-blue staining of the GFP-Nup85, GFP-Seh1, and Nup107-GFP precipitates revealed distinct bands that were not present in the control samples (Fig. 2B). Mass spectrometry analysis of some of these bands identified the \sim 75-kDa polypeptide present solely in the GFP-Seh1 purified fraction as Nup85. The \sim 110-kDa polypeptide, reproducibly detected in both the GFP-Nup85 and GFP-Seh1 fractions, was identified as the C-terminal domain of an *S. pombe* nucleoporin similar to *S. cerevisiae* Nup145 and mammalian Nup98/96 and designated Nup189 in fission yeast (61). Nup189 was previously shown to be cleaved in vivo, most likely by autoproteolysis, as demonstrated in *S. cerevisiae* and mammals (19, 64), leading to a 95-kDa N-terminal product that interacted with Ned1 in a two-hybrid screen (61). In contrast, the protein identified in our complex corresponds to the *S. pombe* orthologue of ScNup145-C.

Therefore, there are two stable subcomplexes in *S. pombe*, one consisting of Nup120, Nup85, Seh1, and Nup145-C and another comprising Nup107 associated with Nup133a and/or Nup133b (Fig. 2C). Our results thus corroborate the structural

model proposed for the *S. cerevisiae* Nup84 complex (38) (Fig. 2C). However, unlike in *S. cerevisiae* or human cells, we were not able to demonstrate a stable interaction between Nup107 or Nup133 and the Nup120-containing subcomplex under the biochemical fractionation employed here. Conversely,

FIG. 2. The *S. pombe* Nup107-120 NPC subcomplex biochemically purifies as two distinct entities. (A) Western blot analysis of immunoprecipitation experiments performed on *S. pombe* whole-cell lysates using an anti-GFP antibody. *S. pombe* strains expressing an HA-Nup133a, -Nup133b, or -Nup120 fusion in a *nup133a*, *nup133b*, or *nup120* background, respectively, and either no GFP fusion or a GFP-tagged nucleoporin (Nup107-GFP, GFP-Nup85, or GFP-Seh1) were grown at 30°C to log phase in supplemented liquid EMM in the presence of thiamine. Immunoprecipitation experiments were performed on these 12 different *S. pombe* whole-cell lysates using an anti-GFP antibody. Equivalent amounts of total protein extracts (T), depleted supernatants (S), and a 10-fold equivalent of the immune pellets (P) were analyzed by Western blotting using anti-GFP (a) or anti-HA $(a', b', and c')$ antibody. The results with the GFP antibody were similar in all experiments and are presented only for the strain expressing HA-Nup133a. Molecular mass markers are on the right in kilodaltons. (B) Silver staining of immunoprecipitates from control, GFP-SpNup85, GFP-Seh1, and Nup107-GFP cell extracts obtained with an anti-GFP antibody. The asterisks indicate the position of the GFP fusion in each lane. The dots indicate additional bands subsequently identified by mass spectrometry as Nup145-C (band 1) and Nup85 (band 2). (C) Schematic model of the Nup107-120 complex in *S. pombe*, based on the structural data obtained from *S. cerevisiae* (38).

FIG. 3. Growth properties of mutant strains with disrupted Nup107-120 complex nucleoporin genes. (A) Tetrads from $nup120^+/2nup120$, *nup107⁺*/ $\Delta nup107$, and $nup85$ ⁺/ $\Delta nup85$ heterozygous diploids were dissected and incubated on YE5S plates at 25°C for 5 days. Highermagnification images of $nup107$ ⁺, $\Delta nup107$, $nup85$ ⁺, and $\Delta nup85$ colonies (boxed) are also shown (insets). (B) Serial dilutions of *S. pombe* cells carrying *nup133a*, *nup133b*, *seh1*, and *nup120* deletions or a combined deletion of both *nup133a* and *nup133b* (*nup133a*/*b*) were spotted onto YE5S plates containing phloxine-B at the indicated temperatures. wt, wild type.

ScNup133 appears to be loosely associated with the ScNup84 complex (2, 38). Despite this difference, we will subsequently designate this subset of nucleoporins as constituents of the *S. pombe* Nup107-120 complex by analogy to the vertebrate Nup107-160 complex.

The *S. pombe* **null mutant strains display different viability phenotypes than their** *S. cerevisiae* **counterparts.** To investigate the functions of constituents of the SpNup107-120 complex, null alleles were generated (see Materials and Methods). As in *S. cerevisiae*, deletion of *S. pombe nup120* leads to a temperature-sensitive phenotype (Fig. 3B, right). In contrast, spores carrying the $\Delta nup107$ or $\Delta nup85$ allele, while able to germinate at 25 $^{\circ}$ C, formed microcolonies of only \sim 20 to 60 cells (equivalent to four to six divisions) (Fig. 3A). This indicated that, unlike their *S. cerevisiae* counterparts, *nup107* and *nup85* are required for vegetative growth in *S. pombe*. Conversely, while deletion of *S. cerevisiae SEH1* leads to a cold-sensitive phenotype (59) , analysis of *S. pombe* Δ *seh1* cells did not reveal any detectable growth defect in a range of temperatures from 16 to 36°C (Fig. 3B and data not shown). Single or combined deletion of the *nup133a* and *nup133b* genes did not lead to any major growth defect at any temperature tested (16 to 36°C). However, phloxine B-containing plates revealed an increased intracellular accumulation of this vital dye in cells disrupted for *nup133b* but not *nup133a*, indicating that deletion of *nup133b* slightly impairs cell viability at all temperatures (Fig. 3B).

FIG. 4. Localization of GFP-tagged nucleoporins in *nup133a*, *nup133b*, and *nup120* mutants reveals NPC aggregation and specific genetic interactions within the Nup107-120 complex. (A) In vivo localization of GFP-Nup85, Nup107-GFP, and GFP-Seh1 in *nup133a* and *nup133b* strains. The cells were grown at 30°C on YE5S plates. Bar, 5 μ m. In contrast to $\Delta nup133a$ cells, $\Delta nup133b$ cells display a heterogeneous distribution of the GFP fusions, which is particularly pronounced in the case of Nup107-GFP. (B) Serial dilutions of wild-type (wt), *nup133b*, *nup107*-GFP, *nup133b*/*nup107*-GFP, and *nup133a*/*nup107*-GFP cells were spotted onto YE5S medium and grown for 4 days at 30°C or for 3 days at 36°C. The growth of $\Delta nup133b/nupl07$ -GFP cells carrying (+) a pREP3-HA-Nup133b, pREP3-HA-Nup133a, or pREP3-HA plasmid further indicates that the temperature-dependent growth defect of the Δ*nup133b*/*nup107*-GFP mutant can be rescued by Nup133b, as well as Nup133a.

FIG. 5. Alterations of NPC distribution in $\Delta nup133b$ and $\Delta nup120$ cells. (A) In vivo distribution of GFP-Nsp1^{Sp} expressed from the *nmt1*^{**} (low-strength) promoter. Wild-type (wt), *seh1*, *nup133a*, *nup133b*, *nup133a*/*nup133b*, and *nup120* cells expressing GFP-Nsp1Sp were grown on thiamine-free EMM plates at 25°C or, when indicated, incubated for 6 h at 36°C, and the localization of the fusion protein was examined in live cells. Representative deconvolved images are shown. The arrowheads point to NPC clusters. Bar, $10 \mu m$. (B) Thin-section electron micrographs of wild-type and *nup120* cells grown at semipermissive temperature (30°C). The small arrows point to NPCs. Nuclear pore clusters in the $\Delta nup120$ mutant are indicated by large arrows. N, nucleus; C, cytoplasm. Bar, 200 nm.

Nup133b and Nup120 are required for normal NPC distribution in *S. pombe***.** Since deletions of members of the *S. cerevisiae* Nup84 complex cause severe defects in the distribution and organization of NPCs, we analyzed NPC distribution in *S. pombe* strains carrying deletions of the nonessential constituents of the Nup107 complex by combining deletion of the *nup133a* or *nup133b* gene with an integrated copy of GFP-Nup85, GFP-Seh1, or Nup107-GFP. As shown in Fig. 4A, a homogeneous punctate perinuclear distribution of these nucleoporins, similar to that observed in wild-type cells (Fig. 1), was observed in a Δnup133a strain. In contrast, a clear clustering phenotype was observed with Nup107-GFP, GFP-Nup85, or GFP-Seh1 in Δ*nup133b* cells, which was more pronounced in the Δnup133b strain expressing Nup107-GFP. This $\Delta nup133b/nup107$ -GFP strain also displays growth defects at 36°C, a temperature at which the individual *nup107*- GFP and Δ*nup133b* strains grow very well (Fig. 4B). In contrast, the $\Delta nup133a/nup107$ -GFP, $\Delta nup133b/GFP-nup85$, and *nup133b*/GFP-*seh1* strains are not temperature sensitive (Fig. 4B and data not shown). It thus appears that GFP tagging of Nup107 at its C terminus, although capable of complementing the lethal phenotype of $\Delta nup107$ and allowing biochemical interactions with at least Nup133a and Nup133b, may slightly impair its function. The temperature-sensitive phenotype of the $\Delta nup133b/nup107$ -GFP strain could be complemented by HA- or GFP-tagged Nup133b (Fig. 4B and data not shown), but surprisingly, also by HA- or GFP-tagged Nup133a expressed under the control of the *nmt1* promoter in the presence of thiamine, which does not completely eliminate expression (i.e., conditions similar to those used for Fig. 2). This suggests that endogenous Nup133a is expressed at very low levels in wild-type cells and indicates that its mild overexpression can overcome some of the defects caused by the *nup133b* deletion (see Discussion).

Attempts to obtain $\Delta nup120$ strains carrying an integrated copy of Nup107-GFP or GFP-Nup85 revealed mild and severe synthetic lethality, respectively. As an alternative approach to follow NPC distribution, mutant strains were transformed with a plasmid encoding a fusion between GFP and the *S. pombe* homologue of ScNsp1p (Nsp1^{Sp}) (74). As shown in Fig. 5A, a homogeneous ring-like staining of the nuclear periphery was observed in wild-type, Δseh1, and Δnup133a strains. In contrast, *nup133b* and *nup133a*/*nup133b* cells gave a discontinuous, patchy ring around the NE, and an even stronger phenotype was observed in $\Delta nup120$ cells, in which fewer bright spots were observed at the nuclear periphery in most cells grown at 25°C, and to a lesser extent at 36°C (Fig. 5A). In these strains, a more pronounced phenotype was observed when cells were grown on plates, suggesting that, as previously reported for several *S. cerevisiae* mutants (8, 27, 45), the extent of NPC clustering depends on growth conditions. Finally, thin-section electron microscopy confirmed the nonhomogeneous NPC distribution in $\Delta nup120$ cells and revealed, in some cases, minor alterations of NE structure (Fig. 5B).

Mutations in the SpNup107-120 complex affect the export of $poly(A)^+$ **RNAs.** To analyze the subcellular distribution of poly $(A)^+$ RNAs in the various mutant strains, FISH exper-

FIG. 6. Mutations in the Nup107-120 complex cause mRNA export defect. (A) Rapid accumulation of poly(A)⁺ RNA in $\Delta nup120$ and *nup133b*/*nup107*-GFP mutants at restrictive temperature. Wild-type (wt), *nup133b*, *nup120*, and *nup133b*/*nup107*-GFP strains grown at 25°C or shifted to 36°C were fixed and subjected to in situ hybridization with a Cy3-labeled oligo(dT)₅₀ probe. DNA was stained with DAPI. Bar, 10 μ m. (B) Distribution of the pre-60S ribosomal particles in Nup107-120 complex mutants. Strains grown at 25°C or shifted to 36°C for 3 h were processed for FISH using a Cy3-labeled probe complementary to the 25S rRNA.

iments were performed. In wild-type and $\Delta nup133b$ cells, $poly(A)^+$ RNAs were uniformly distributed throughout the cells at both 25 and 36°C (Fig. 6A). In contrast, *nup120* cells grown at 25 $\rm ^{\circ}C$ already accumulate poly $(A)^+$ RNA inside their nuclei. In these cells, a clear nuclear $poly(A)^+$ RNA signal was observed in \sim 30% of the cells within 10 min at 36°C (data not shown) and in most cells by 30 min (Fig. 6A). A similar, albeit less severe, phenotype was also observed in Δ*nup133b/nup107*-GFP cells, in which nuclear accumulation of $poly(A)^+$ RNA could be observed in $~10\%$ of the cells following a 1-h shift to 36°C (data not shown) and in most of these cells after 4 h at 36°C (Fig. 6A). Therefore, both *nup120* and *nup133b*/ *nup107*-GFP strains exhibit a poly $(A)^+$ RNA export defect.

We also analyzed the nuclear export of the pre-60S ribosomal particles using probes complementary to the 25S rRNA. As a control, we used the *pim1-d1* strain that carries a mutation in the nucleotide exchange factor for the small GTPase Ran^{Sp}/ Spi1 (56). In agreement with data obtained from *S. cerevisiae* (34), nuclear retention of 25S rRNA was observed in the temperature-sensitive *pim1-d1* strain (Fig. 6B). In contrast, no detectable intranuclear accumulation of pre-60S particles could be detected in the *nup120*, *nup133b*/*nup107*-GFP, or $\Delta \Delta nup133a/b$ strain (Fig. 6B). Although one cannot formally exclude a minor defect in pre-60S rRNA export, this result suggests that the poly $(A)^+$ RNA export defect observed in the *S. pombe* Nup107-120 mutant strains is specific and does not reflect a general impairment of nucleocytoplasmic transport in these strains.

nup120 **and** *nup133b***/***nup107***-GFP mutant strains exhibit severe cell division defects at restrictive temperature.** Unexpectedly, microscopic analyses of $\Delta nup120$ cells, and to a lesser extent $\Delta nup133b/nup107$ -GFP cells, revealed cell division defects at the restrictive temperature. In these cells, calcofluor staining of asynchronous cultures showed an increase of the septation index upon incubation at 36°C. In $\Delta nup120$ cells, \sim 40% had a septum after a 3- or 6-h shift to 36°C compared to \sim 10% of wild-type cells or $\Delta nup120$ cells grown at 25°C (Fig. 7A). Moreover, $\Delta nup120$ cells exhibited septation defects (i.e., widened, misplaced, or occasionally, multiple septa) even at 25°C (\sim 10% of the septa) and more frequently at 36°C (\sim 20%) of the septa at 3 h and \sim 40% at 6 h) (Fig. 7A). To further characterize this cytokinesis defect, wild-type or $\Delta nup120$ cells synchronized in G_1/S at 25 $°C$ using hydroxyurea were released at 36°C upon hydroxyurea washout. Under these conditions, we observed an \sim 1-h delay of septation onset in $\Delta nup120$ cells compared to wild-type cells and the aberrant persistence of septated cells (data not shown).

DAPI staining also revealed aberrant mitoses, such as unequally segregated chromosomes and lagging chromosomes, in *nup120* cells shifted for 3 or 6 h to 36°C (Fig. 7A and 8; see below). We also detected many mononucleated Δ*nup120* cells showing hypercondensed chromatin, suggesting a lag at metaphase in the $\Delta nup120$ mutant. In binucleated and septated *nup120* cells that accumulate at 36°C, atypical DAPI staining patterns were frequently observed (Fig. 7A). Flow cytometry analyses showed that while $\Delta nup120$ cells mainly have a 2C DNA content at 25°C, 4C-containing cells accumulated at 36°C (Fig. 7B), indicating that DNA replication had occurred. Finally, the NLS–GFP–β-galactosidase reporter accumulated efficiently in the nuclei of $\Delta nup120$ cells at both permissive and

FIG. 7. $\Delta nup120$ strain accumulates binucleated cells with abnormal septa and hypercondensed chromatin at 36°C. (A) Samples from wild-type (wt) and $\Delta nup120$ cells grown at 25°C or shifted for 3 or 6 h to 36°C were fixed and stained with calcofluor or DAPI as indicated. The differential interference contrast (DIC) image of the DAPI-stained cells revealing the presence of the septum is also shown. The arrows indicate septated *nup120* cells with condensed or unequally segregated chromosomes. The arrowheads point to nuclei with atypical DAPI staining patterns. Bar, 10 μ m. (B) Samples from the same cultures were stained with propidium iodide and analyzed by FACS. The fluorescence (the DNA content on an arbitrary scale) and frequency (relative cell number) are plotted along the *x* and *y* axes, respectively. Relative fluorescences corresponding to 2C and 4C DNA contents are indicated. (C) The localization of the NLS–GFP–β-galactosidase fusion was monitored in living Δ*nup120* cells grown at 25 \degree C or incubated for 3 h at 36 \degree C. Bars, 10 μ m.

restrictive temperatures (Fig. 7C), indicating that unlike several mutant alleles defective in the $\text{Ran}^{\text{Sp}}\text{/Spilpathway}$ (53), NE integrity is not impaired in $\Delta nup120$ cells.

Altogether, our results indicate that $\Delta nup120$ cells shifted to 36°C are at least transiently blocked at metaphase, leading to a delay in septation onset. After the first nuclear division, these cells can proceed to septum formation, replicate their DNA, and accumulate as septated cells with an intact NE but an atypical chromatin structure.

To further characterize these mitotic defects, *nup120* and

wild-type cells grown at 25°C or shifted to 36°C for 3 or 6 h were analyzed by immunofluorescence using an anti-tubulin antibody and DAPI. Analysis of the various stages of mitosis, assessed by the length of the mitotic spindle, revealed an increased proportion of $\Delta nup120$ cells in premetaphase and metaphase after 3 h at 36°C compared to wild-type cells or *nup120* cells grown at permissive temperature (Fig. 8A and C). In addition, this study revealed that even in cells grown at 25°C, \sim 20% of the mitotic $\Delta nup120$ cells had undergone aberrant mitoses. At 36 \degree C, this proportion reached \sim 70% of the

FIG. 8. *nup120* cells display abnormal microtubule spindles and aberrant mitosis. Wild-type (wt) and *nup120* cells grown asynchronously at 25°C or shifted to 36°C for 3 or 6 h were processed for immunofluorescence using the TAT1 antibody to visualize microtubules and DAPI to observe the DNA morphology. (A) The frequencies of the different stages of mitosis were determined according to the lengths of the mitotic spindles (panel C). An average of 40 mitoses were examined for each sample, and a χ^2 procedure was applied for statistical analysis (wt/ $\Delta nup120$ at 0 h and 36°C, $P =$ not significant; wt/ $\Delta nup120$ at 3 h and 36°C, $P \le 0.001$; wt/ $\Delta nup120$ at 6 h and 36°C, $P \le 0.001$). Note that the increase in the early anaphase stage observed in $\Delta nup120$ cells after a 6-h shift to 36°C may reflect the accumulation of abnormal metaphases with aberrant spindle lengths. (B) The percentage of abnormal mitoses was also quantified. (C) Prometaphase (a), metaphase (b), and early (c) and late (d and d) anaphase in *nup120* cells grown at 25°C. (D) Examples of abnormal mitoses exhibited by *nup120* cells shifted for 6 h to 36°C. Rows: a, misplaced nucleus or spindle; b, chromosome missegregation; c, lagging chromosomes; d, chromosomes unattached to the spindle; e, monopolar spindle; f, malformed mitotic spindle. For each example, images of DIC combined with DAPI (DIC+DAPI), DAPI, tubulin staining, and DAPI plus tubulin staining (merge) are shown. In row a, the medial plane of septation was visualized using anti-Mid1 immunostaining (arrowhead). Bar, $10 \mu m$.

mitotic cells after a 6-h shift to 36°C (Fig. 8B and D) ($P \le 0.01$). Aberrant mitoses included displaced positioning of the nucleus and microtubule spindle relative to the long axis of the cell, with the three condensed chromosomes displaced from the center of the cell (Fig. 8D, a and f); unequal sister chromatid segregation; lagging or unattached chromosomes (Fig. 8D, b, c, and d); and compromised microtubule structures (monopolar spindles and misshaped spindles) (Fig. 8D, e and f).

Accurate chromosome transmission is defective in a *nup120* **strain.** As many mutants with altered spindle assembly and function show hypersensitivity to microtubule-depolymerizing agents, such as TBZ, we performed serial cell dilutions on rich medium containing various concentrations of TBZ. As shown in Fig. 9A, 10 μ g of TBZ/ml severely inhibited the growth of the $\Delta nup120$ mutant at permissive temperature (23 $^{\circ}$ C) compared to wild-type cells, indicating that *nup120* deletion affects, most likely indirectly, microtubule integrity or spindle function. We then determined if the spindle checkpoint, which monitors mitotic spindle integrity and attachment of the kinetochores to the spindle, was required for the viability of the *nup120* strain. Strains with both *nup120* and either of the spindle checkpoint genes, *mph1* (29) or *mad2* (30), deleted were viable. Unlike single mutants, Δ*nup120*/Δ*mph1* cells were sensitive to 5 μ g of TBZ/ml at 23°C, indicating a synergistic effect of the two mutations (Fig. 9A). In contrast, sensitivity to TBZ was indistinguishable in $\Delta nup120$ and $\Delta nup120/\Delta mad2$ mutants. This pattern of synthetic interaction with *mph1* but not with Δ*mad2* is similar to that seen with a point mutant of *spi1* (18) and likely reflects differences in the severities of the spindle checkpoint defects in the two mutant strains.

To analyze chromosome segregation in $\Delta nup120$ cells in more detail, we monitored the presence of a nonessential minichromosome (47) whose loss results in adenine auxotrophy and the subsequent appearance of pink colonies on plates containing low adenine. However, attempts to isolate an *S. pombe* $\Delta nup120$ strain containing this minichromosome were not successful. Furthermore, while the $\Delta nup120$ strain transformed by a complementing pUra4-Nup120 plasmid was able to lose the plasmid and thus to grow on 5-FOA-containing plates at the permissive temperature, $\Delta nup120$ cells carrying pUra4-Nup120 and a minichromosome failed to grow on 5-FOA (Fig. 9B). Accordingly, the presence of an additional chromosome in a $\Delta nup120$ mutant severely impairs cell viability. Similar results were previously reported in the case of the *alp14* mutant and were interpreted as resulting from extreme instability of the minichromosome (25).

As an alternative approach, we followed chromosome segregation in a heterozygous $\Delta nup120/nup120$ ⁺ h^+/h^+ ade6- $M210/ade6-M216$ (Ade⁺) diploid by monitoring the loss of one *ade6* mutant allele that leads to the appearance of pink colonies. This diploid strain is homozygous at the mating-type locus and consequently is incompetent for meiosis. Interestingly, a significant increase in the number of pink (Ade^-) colonies was observed in a $\Delta nup120/nup120^+$ diploid grown at 36°C compared to a wild-type $nup120^+/nup120^+$ strain or a $\Delta nup120/$ *nup120* diploid grown at 30°C (Fig. 9C). Flow cytometric analysis of the DNA content revealed that the pink-colony cells had mainly a 2C DNA content, in contrast to the white-colony cells, which were 4C (data not shown). Such a "haploidization" phenotype (loss of the disomic state for all three chromosomes

FIG. 9. Hypersensitivity to TBZ and chromosome segregation defects in $\Delta nup120$ cells. (A) Deletion of $nup120$ induces hypersensitivity to the microtubule-destabilizing drug TBZ. Serial dilutions of wildtype (wt) and mutant cells were spotted on YE5S medium containing or lacking $(-)$ TBZ (5 or 10 μ g/ml) and incubated at 23°C. A TBZ-free plate was also incubated at 36°C. Note that the TBZ sensitivity of the $\Delta nup120$ cells is further enhanced in a $\Delta mph1$ background. (B) The presence of a minichromosome is toxic in $\Delta nup120$ haploid cells. *nup120* strains complemented by the pUra-Nup120 plasmid and containing $(+mc)$ or lacking $(-mc)$ a minichromosome were spotted on supplemented EMM plates in the presence or absence of 5-FOA at 23°C. $\Delta nup120$ cells that carry both pUra-Nup120 and a minichromosome cannot grow on 5-FOA, indicating that the pUra-Nup120 plasmid cannot be easily lost by these cells. (C) $\Delta nup120$ /wt heterozygous diploid cells undergo chromosome loss at 36°C. wt/wt h^+/h^+ ade6-*M210*/*ade6-M216* (wt/wt) and $\Delta nup120/wt$ h^+/h^+ ade6-M210/ade6- $M216$ (Δ nup120/wt) diploids were plated on low-adenine yeast extract medium at 30 or 36°C. The percentage of pink [Ade⁻] colonies that had lost one of the two *ade6* alleles was determined. The average of four independent experiments is presented. The error bars represent standard deviations. (D) Unlike Nuf2-CFP, GFP-Nup85 is not detectable at kinetochores in metaphase-arrested cells. *nuc2-663* cells with integrated copies of both GFP-Nup85 and Nuf2-CFP were arrested in metaphase by a 3-h shift to 36°C. Deconvolved images of two Z sections ($0.6 \mu m$ apart) of a live cell are presented. The perinuclear labeling observed in the CFP channel is due to the bleedthrough of the GFP-Nup85 signal. The DIC image of the cell is also shown. Bar, $10 \mu m$.

as a consequence of initial loss of one chromosome in *S. pombe* diploids) was previously described (46) and is very similar to that previously reported in the case of a $\Delta spil/spil^+$ diploid (40, 53). Our data thus demonstrate that Nup120 is essential for genome stability and integrity during mitosis.

Because several constituents of the human Nup107 complex are localized at kinetochores during mitosis (7, 28), it was tempting to link these defects in chromosome segregation to a possible function of this complex at kinetochores. To determine whether the *S. pombe* Nup107 complex is similarly localized at kinetochores during mitosis, we analyzed the

FIG. 10. Localization of endogenous Spi1 and toxicity of Spi1-GFP in Nup107-120 complex nucleoporin mutants. (A) *nup120* cells grown at 25°C or shifted for 4 h to 36°C were processed for immunofluorescence using affinity-purified anti-Spi1 antibody, mAb414 antibody to visualize the NPCs, and DAPI to visualize the DNA. Note that in the Δ*nup120* strain at both 25 and 36°C, the nuclear pores surround the DNA and Spi1 is predominantly but not exclusively localized to the nucleus. Despite the overall weaker Spi1 and mAb414 signals at 36°C, moderate changes in the intracellular distribution of Spi1 can be observed in *nup120* cells shifted to the restrictive temperature. (B) Wild-type (lanes 1), *nup133a* (lanes 2), *nup133b* (lanes 3), and *nup120* (lanes 4) cells expressing the Spi1-GFP fusion under the control of the *nmt1** promoter were grown at 25°C in liquid EMM supplemented with thiamine (repressed [R] *nmt1* promoter, R. 25°C) and shifted either for 24 h at 25°C to thiamine-free medium (induced [I] *nmt1* promoter, I. 25°C) or for 4 h to 36°C in thiamine-containing medium (R. 36°C). Equivalent amounts of whole-cell lysates were analyzed by Western blotting using anti-Spi1 antibody. The same blot was reprobed with the mAb414 antibody that mainly recognized a protein
most likely corresponding to Nsp1^{Sp} (theoretical mass, 61 kDa; GeneDB database). (lanes 1) and *nup120* (lanes 4) cells even when shifted to 36°C (R. 36°C), whereas the expression levels of Spi1-GFP and its degradation products (*) are lower in *nup133b* (lanes 3) and *nup120* (lanes 4) strains grown at 25°C in thiamine-free liquid medium (I. 25°C). (C) Spi1-GFP expression is toxic in *nup120* and *nup133b* mutant cells. Wild-type (wt), *nup133a*, *nup133b*, and *nup120* cells expressing the Spi1-GFP fusion under the control of the *nmt1*^{*} promoter were grown to mid-log phase in liquid EMM supplemented with thiamine. The cells were then either spotted onto EMM-plus-thiamine plates (repressed *nmt1* promoter) or shifted for 24 h at 25°C to thiamine-free liquid medium and spotted on thiamine-free EMM (induced *nmt1* promoter). The plates were incubated at 25, 30, or 36°C.

localization of GFP-Nup85 in metaphase-arrested cells, taking advantage of the *nuc2*-*663* mutation (32). In addition, we used the Nuf2-CFP marker (24) to visualize kinetochore positioning. In the resulting GFP-*nup85*/*nuf2*-CFP/*nuc2*-*663* cells, intranuclear Nuf2-CFP dots were clearly detected in metaphase. However, while GFP-Nup85 gave a bright localization at the nuclear rim, no specific labeling could be observed at kinetochores (Fig. 9B). Although the presence of undetectable amounts of GFP-Nup85 at kinetochores cannot be excluded, this result suggests that kinetochore localization of the Nup107 complex might be restricted to higher eukaryotes.

The SpNup107-120 complex is functionally and genetically linked to the Ran^{Sp}/Spi1 pathway. While this study was in progress, Gao and coworkers reported that nuclear accumulation of the small GTPase $\text{Ran}^{\text{Sc}}/\text{Gsp1p}$ was impaired in *S. cerevisiae* Δ*nup133, nup120-1/rat2-1, and Δnup85* mutant strains (23). These results, together with the similarity of cell cycle defects observed in the *S. pombe* $\Delta nup120$ strain and several mutants defective in the Ran^{Sp}/Spi1 pathway, prompted us to analyze whether alteration of the Spi1 pathway could be responsible for some of the defects observed in $\Delta nuv120$ mutant cells. We therefore investigated the expression level and distribution of Spi1 in $\Delta nup120$ strains grown at permissive

temperature or shifted for 4 h to 36°C. Western blot analysis did not reveal any significant alteration of the expression level of endogenous Spi1 in Δnup120 cells grown at 36°C (Fig. 10B, compare R. 25°C and R. 36°C, lanes 1 and 4). Upon fluorescence analysis, both the Spi1 and mAb414 signals were reproducibly fainter in the $\Delta nup120$ strain at 36°C (Fig. 10A). However, since (i) NE targeting of GFP-Nsp 1^{Sp} is not significantly altered in $\Delta nup120$ cells at 36°C (Fig. 5A) and (ii) the expression level of neither endogenous Spi1 nor the major $65-kDa$ protein (most likely $Nsp1^{Sp}$) recognized by the mAb414 antibody is affected (Fig. 10B), this decreased staining most likely reflects technical limitations associated with the efficiency of fixing or spheroplasting these cells once they are grown at 36°C. Despite the weaker overall staining, fluorescence analysis revealed moderate changes in the distribution of Spi1 in $\Delta nup120$ cells shifted to the restrictive temperature: although a nuclear signal remained detectable in most cells, the relative signal for Spi1 in the nucleus relative to the cytoplasm was lower in cells grown at the restrictive temperature (Fig. 10A).

It has been reported that mutations in several genes of the RanSp/Spi1 pathway, such as *pim1* and *mog1*, are rescued by overexpression of wild-type Spi1 (40, 56, 63). Using the

pREP3X-Spi1 plasmid (42), we could not observe any rescue of either cell growth or septation defects of $\Delta nup120$ cells at 36°C (data not shown). However, this study revealed that expression of Spi1-GFP under the control of the mediumstrength *nmt1* (*nmt1**) promoter (Fig. 10B) enhanced the growth defect of $\Delta nup120$ cells and decreased the viability of *nup133b* mutant cells (Fig. 10C). In contrast, the growth properties of wild-type and *nup133a* cells were not altered by Spi1-GFP expression. As the Spi1-GFP fusion is not functional (63), its expression, although not affecting wild-type cell growth, may lead to a competition with endogenous Spi1 in *nup120* and *nup133b* mutants. Most likely correlated with this toxicity, we reproducibly found that the expression level of Spi1-GFP was lower in both Δnup133b and Δnup120 cells (Fig. 10B). These data thus strengthen the functional link between the Nup107-120 complex and the Spi1/Ran pathway.

DISCUSSION

The evolutionarily conserved ScNup84/vertebrate Nup107- 160/SpNup107-120 complex: conservation and divergences. In this study, we have identified and functionally characterized *S. pombe* nucleoporins homologous to components of the ScNup84/vertebrate Nup107-160 complex. Unexpectedly, the *S. pombe* genome contains two ORFs encoding Nup133 orthologues, which we designated Nup133a and Nup133b. Sequence analysis indicated that the corresponding proteins were nearly as divergent from each other (19.5% homology between SpNup133a and SpNup133b) as they are from their *S. cerevisiae* or human counterparts (Table 2), yet the two nucleoporins were both targeted to the NPCs and able to interact with Nup107. Although single or combined deletion of *nup133a* and *nup133b* did not affect cell viability, Nup133b appeared to have a function that was predominant over Nup133a: unlike deletion of *nup133a*, deletion of *nup133b* led to (i) minor growth defects; (ii) synergistic growth defects when combined with $nup107$ -GFP, $\Delta nup120$, or $\Delta nup184$ (whose product is the *S. pombe* homologue of ScNup188p [71]) (Fig. 4 and data not shown); and (iii) toxicity of Spi1-GFP expression. Despite the lack of any detectable function of the *nup133a* gene, its overexpression was able to complement the phenotypes induced by *nup133b* deletion. Although Nup133a may have a specific function that was not revealed by our assays, a more likely explanation is that, under some conditions, the protein may provide a backup pool for Nup133 function. In this respect, it is noteworthy that Nup133a appears to be mildly induced (2- to 2.5-fold) under several stress conditions $(H₂O₂)$, cadmium, or heat), a feature that is not shared by any other constituent of the Nup107-120 complex (11; http://www.sanger.ac.uk /PostGenomics/S_pombe/projects/stress/).

Another unexpected result arising from this study was the lack of a major growth defect induced by *Spnup133a/b* deletions that differs from the temperature-sensitive phenotype induced by *NUP133* deletion in *S. cerevisiae* (17, 37, 50) and the essential function of Nup133 in vertebrate NPC assembly (68). Yet, Nup133 inactivation in *Caenorhabditis elegans* does not lead to any major defect (reference 22 and references therein). More generally, comparison of the phenotypes obtained upon deletion of *S. cerevisiae* and *S. pombe* genes, or induced by RNA interference in *C. elegans* (reference 22 and references

therein) or in human cells (9, 28, 68), revealed major divergences among these organisms. The fact that the viability phenotypes were as divergent between the two yeasts as with either *C. elegans* or vertebrates suggests that these differences reflect the evolution of individual nucleoporins within the ScNup84, SpNup107-120, and metazoan Nup107-160 complexes rather than specific features of evolutionarily divergent organisms (i.e., both of the yeasts that undergo a closed mitosis versus metazoans with an open mitosis).

In summary, although in certain instances some of its individual constituents might be dispensable for cell viability or less tightly associated with other subunits, the evolutionarily conserved ScNup84/SpNup107-120/metazoan Nup107-160 complex appears to be required, as an entity, for cell viability in all organisms tested to date.

The major functions of the ScNup84/SpNup107-120 complexes have been evolutionary conserved. Despite the divergence observed at the level of individual nucleoporins, functional analysis of several of the *S. pombe* mutant strains revealed a disturbed NPC distribution, similar to the phenotypes previously reported for *S. cerevisiae* constituents of the Nup84 complex. Although the NPC clustering phenotype is not as pronounced in *S. pombe* as in several *S. cerevisiae* mutants, these newly identified *S. pombe* mutants provide a useful tool to discriminate NE- from NPC-associated proteins. In addition, FISH studies revealed nuclear accumulation of $poly(A)^+$ RNA at the restrictive temperature in the $\Delta nup120$ and *nup133b*/*nup107*-GFP mutants. Our study also revealed major mitotic defects associated with mutations within the Nup107-120 complex. Although not extensively characterized, mitotic defects were also previously reported in *S. cerevisiae* $\Delta nup120$ (1) and $\Delta nup84$ strains (59). It is noteworthy that the dual function of the SpNup107-120/ScNup84 complex in mRNA export and the cell cycle represents an additional example of potential linkage between these two pathways: while Rae1, initially identified as an mRNA export factor, was also shown to be required for proper mitosis in both *S. pombe* (10, 70) and higher eukaryotes (5), mutations in the *cut1* and *cut2* genes, involved in sister chromatid separation, were recently reported to impair mRNA export in *S. pombe* (4). How these different pathways are connected thus remains to be elucidated.

Mitotic defects of $\Delta nup120$ mutants are consistent with a **mild alteration of the Ran pathway.** In *S. pombe*, the multiple Ran-dependent cellular processes are differentially sensitive to the level of active Ran (53). Among these processes, those most sensitive to the level of functional Ran (i.e., microtubule dynamics and chromosome transmission) were also impaired in the $\Delta nup120$ mutant grown at restrictive temperature. In contrast, phenotypes associated with more severe defects in the Ran pathway (accumulation of septated cells in G_1 with hypercondensed unreplicated chromosomes and a fragmented NE) were not detected in this mutant. Accordingly, the *nup120* mutant behaves in *S. pombe* like strains carrying weak loss-of-function alleles of the Ran pathway.

Gao et al. (23) recently reported that several members of the *S. cerevisiae* Nup84 complex are required for the nuclear accumulation of $\text{Ran}^{\text{Sc}}/\text{Gsp1}$, a phenotype that could be correlated with an altered association of Ntf2 (the nuclear transport factor of Ran) with the NE. Immunofluorescence analyses

performed in *S. pombe* $\Delta nup120$ cells suggest that the nuclear accumulation of Spi1 might be slightly diminished at 36°C. Consistently, we also found that expression of nonfunctional Spi1-GFP was specifically toxic in $\Delta nup120$ and in $\Delta nup133b$ cells, but not in wild-type or $\Delta nup133a$ cells. These data suggest that some of the phenotypes of Nup107-120 complex mutants may reflect a disturbance of the Ran GTPase system. Whether an alteration of nuclear import of Ran^{Sp}/Spi1 in these mutant cells is primarily responsible for the subsequent mitotic-spindle and chromosome segregation defects remains to be determined, and further experimentation will be necessary to establish the mechanism by which the Ran^{Sp}/Spi1 pathway and cell cycle progression are affected.

While $poly(A)^+$ RNAs strongly accumulate within the entire nucleus in Δnup120 and Δnup133b/nup107-GFP mutants, export of most mRNAs appears to be independent of the Ran-GTPase (12, 63). Accordingly, defects in mRNA export occurring in the Nup107-120 complex mutants are likely to be uncoupled from those induced by alterations of the Ran pathway. In contrast, it is tempting to speculate that the clustering defects observed upon alteration of the ScNup84 and SpNup107-120 complex could somehow be linked to the previously reported requirement for Ran in NE and NPC integrity in both *S. pombe* (16, 41) and *S. cerevisiae* (52).

In conclusion, the data presented here provide functional and genetic links between the *S. pombe* Nup107-120 subcomplex and Ran-dependent mitotic processes. In vertebrates, recent studies suggest additional links between the Nup107-160 complex and Ran: this GTPase was shown to regulate interactions between Nup107 and other nucleoporins that may be required for NPC formation (69). In addition, a fraction of mammalian Nup107, Nup133, and Nup85 localizes at kinetochores in mitosis (7, 28), and Ran was recently shown to regulate kinetochore function (3). Whether these independent studies reflect a general function of the Nup107 complex during mitosis in both yeasts and vertebrates remains quite speculative.

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