Nonessential nucleoporins Nup2, Nup60 and Nup84 are required for normal meiosis in

*Saccharomyces cerevisiae*

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Abstract

The nuclear pore complex (NPC) selectively transports cargo between the nucleus and the cytoplasm. The inner nuclear membrane (INM) face of the NPC also serves as a hub where gene silencing and DNA repair are spatially coordinated. In *Saccharomyces cerevisiae*, partitioning of active and silenced chromatin at subtelomeric regions depends on the boundary activity of nonessential nucleoporin proteins Nup2 and Nup60, along with Htz1, the histone variant H2A.Z. The INM is also important for the chromosome events of meiosis since Ndj1-mediated telomere attachment and clustering at the INM is required for efficient homolog pairing, recombination, and segregation. Here we tested possible meiotic roles for Nup2, Nup60, Htz1, and other nonessential nucleoporins by analyzing the effects of deletion mutations on sporulation, spore viability and possible phenotypic epistasis with *ndj1Δ*. Deleting *NUP2*, *NUP60*, and *HTZ1* reduced spore viability compared to wild-type (WT). A detailed analysis of spore lethality indicated that homolog nondisjunction in these mutants was elevated compared to WT, yet unlike *ndj1Δ*, this was not the predominant cause of spore death. Deleting *NUP84* reduced meiosis I nuclear divisions, while deleting *NUP53*, *NUP100*. and *NUP157* had no effect on sporulation, spore viability, or the kinetics of meiosis I progression. Surprisingly, *nup2Δ ndj1Δ* uniquely failed to undergo meiosis I nuclear divisions, suggesting Nup2 and Ndj1 function in partially redundant pathways or create a poisonous intermediate. The meiosis I division was also delayed by 2 hours in *nup2Δ* compared to WT pointing to a specialized role for Nup2 in the meiotic program.
Introduction

The nuclear pore complex (NPC) is a ring structure, which sits in the nuclear envelope bridging the inner and outer nuclear membranes. The primary function of the NPC is to selectively transport cargo into and out of the nucleus. Aside from its nuclear transport role the NPC has also been implicated in aspects of nuclear architecture that impact processes such as the DNA damage response, transcriptional silencing, boundary activity, transcriptional memory, mRNA processing, and mRNA export (FAHRENKROG et al. 2000; DENNING et al. 2001; DILWORTH et al. 2005; BUKATA et al. 2013; PTAK et al. 2014). The NPC is composed of proteins called nucleoporins (Nups), which fall under several functional components of the NPC. The main structure the NPC is made up of two inner rings which are sandwiched between two outer rings, with one outer ring touching the nucleoplasmic side and the other touching the cytoplasmic side (Figure 1; AITCHISON AND ROUT 2012).

The selective permeability of the NPC is generated by phenylalanine-glycine (FG) Nups which contain an FG-repeat motif made up of FGs separated by ~20 amino acids (AITCHISON AND ROUT 2012). FG Nups predominantly attach to the inner rings and the FG-repeat motif generally results in a long unstructured filament which extends into the central channel of the NPC (ALBER et al. 2007). FG Nups filaments also extend into the cytoplasm as cytoplasmic filaments and into the nucleoplasm as the nuclear basket. FG Nups serve to block passive diffusion and direct transport across the NPC (Figure 1). The cytoplasmic filaments and nuclear basket serve to direct release of cargo following transport through the NPC.

Indirect evidence from non-yeast species links Nups to the meiotic program which is a specialized series of cell divisions required to create haploid gametes from diploid parent cells
A hallmark of meiosis is the pairing, synapsis and recombination between homologous chromosomes that ensures their segregation at the first meiotic division (Zickler and Kleckner 2015). A mutation in Tmem48/NDC1 in mouse causes defects in homologous chromosome pairing and the arrest of cells in late meiotic prophase, and sterility (Akiyama et al. 2013). In Drosophila melanogaster, a P element insertion into Nup154 (yeast NUP157) results in defects in both male and female meiosis with complete prophase arrest in males (Giguotti et al. 1998). The mouse Nup2 homolog, mNup50, and its paralog, n50rel, are enriched in the testis (Fan et al. 1997; Smitherman et al. 2000).

During meiotic prophase, telomeres cluster at the nuclear envelope near the microtubule organizing center in a characteristic bouquet conformation (Zickler and Kleckner 1998). In plants and animals, nuclear pores reorganize to cluster near the bouquet telomeres during meiotic prophase; in yeast, direct interactions between bouquet telomeres and nuclear pores have been observed (Church 1976; Holm 1977; Klein et al. 1992). In yeast, the bouquet is directed by Ndj1 which physically connects the telomeres to the nuclear envelope (Chua and Roeder 1997; Conrad et al. 1997).

In this study, we explored possible role(s) for nonessential Nups in mediating normal meiotic progression in budding yeast. Through a past genome duplication event in budding yeast, deletions of several NUP genes are viable due, in part, to functional redundancy (Kellis et al. 2004; Aitchison and Rout 2012). We created deletions in genes encoding a component of the inner ring (Nup157), the outer ring (Nup84), and four FG Nups (Nup2, Nup53, Nup60, and Nup100) that localize within the central channel or near the nuclear basket of the NPC (Wente et al. 1992; Loeb et al. 1993; Aitchison et al. 1995; Siniossoglou et al. 1996; Marelli et al. 1998;
ROUT et al. 2000). For this study, each deletion strain was tested for growth under different challenged regimens, sporulation efficiency, spore viability, and the timing of meiotic progression. We also evaluated possible genetic interactions between these mutations and the deletion of NDJ1.
Results

Growth of nup mutants in the SK1 yeast strain background

To evaluate the effects of nup mutations in meiosis, we created homozygous gene knockouts of the nonessential Nups listed above in the SK1 strain background, which is well known for its high efficiency of sporulation and spore viability (Figure 1; KANE AND ROTH 1974; ELROD et al. 2009). Since growth defects could contribute to inefficient meiosis, we analyzed the colony formation of each mutant and wild type (WT) strain at various temperatures on rich (YPD) and minimal solid media. At 20°, 25°, and 30°, all strains gave similarly sized colonies as WT except nup60Δ and nup84Δ, which displayed modest growth defects at these temperatures on YPD (Figure 2A). Surprisingly, nup2Δ and nup60Δ mutants gave larger colonies on YPD at 37° compared to WT (Figure 2A). The nup100Δ mutant also gave larger colonies compared to WT on minimal medium at all temperatures (Figure 2B). These results uncover surprising antagonistic pleiotropic phenotypes of nup2Δ, nup60Δ and nup100Δ mutants on colony growth.

Sporulation efficiency and spore viability of nup mutants

We next assessed sporulation and spore viability in the nup mutants at 30° (Table 1). All nup mutants sporulated with the same efficiency as WT (95%; n = 200) on solid sporulation medium except for nup60Δ (71%; n = 200) and nup84Δ (<1%; n = 200). While 70% of nup84Δ cells could undergo the meiosis I (MI) nuclear division, no cells went on to form tetrads. For the remaining strains, four-spore tetrads were dissected on YPD and growth was assessed after four days of incubation at 30°. Among four-spore tetrads, WT strain gave 97.1% (n = 1152) viable spores from four-spore asci, while nup2Δ and nup60Δ gave 86.1% (n = 1536) and 70.5%
There was no measurable defect in spore viability in nup53Δ (95.8%; n = 288), nup100Δ (98.6%, n = 288) or nup157Δ (98.3%; n = 288; Table 1).

We included the deletion of HTZ1 (histone variant H2A.Z), since Htz1 is associated with the NPC and plays a role in the boundary activity of Nup2 and Nup60 (DILWORTH et al. 2005). The htz1Δ mutation gave 65.9% viable spores among four-spore tetrads (n = 1152; Table 1). Since Nup2, Nup60, and Htz1 function in telomere silencing, we tested their mutant phenotypes in the ndj1Δ mutant strain background, in which telomere attachment to the nuclear envelope is disrupted (TRELLES-STICKEN et al. 1999). The ndj1Δ mutant alone gives 82.0% (n = 768) spore viability in our strain background compared to 75% as reported by Wanat et al., 2008. The nup60Δ ndj1Δ and the htz1Δ ndj1Δ double mutants showed no epistasis with respect to spore inviability compared to the single mutant phenotypes, suggesting that Nup60 and Htz1’s roles during meiosis are independent of Ndj1 (Table 1). Interestingly, the nup2Δ ndj1Δ double mutant failed to undergo the MI nuclear division. A more detailed analysis of this phenotype will be reported separately (D.B.C and S.M.B, unpublished).

**MI progression is delayed in nup2Δ compared to WT**

We measured the timing of the MI nuclear division in the nup knockout strains following the transfer of synchronized cell cultures to sporulation medium (t = 0 hours). Post-MI stages are marked by the presence of at least two DAPI-stained nuclear bodies. In nup2Δ, MI was delayed ~two hours compared to WT (Figure 3). By contrast, nup53Δ, nup100Δ, and nup157Δ deletion mutations had no measurable effect on MI timing. The nup60Δ, nup84Δ, and htz1Δ
mutants were excluded from the MI analysis because the cells would be unlikely to enter meiosis synchronously in liquid culture due to their mitotic growth defects at 30°.

**Causes of spore death in nup2Δ, nup60Δ, and htz1Δ**

The distribution of tetrads giving 4, 3, 2, 1, or 0-viable spores can lend insight into the causes of spore death (CHU AND BURGESS 2016). For example, mutants with demonstrated increased levels of MI nondisjunction (MI-ND) also give rise to an increased incidence of two spore and zero spore viable tetrads (SHONN et al. 2000; MARTINI et al. 2006; WANAT et al. 2008). We recently introduced two R-based tools, TetSim and TetFit, for analyzing the causes of mutant spore death based on the distribution of live:dead tetrad classes (File S1; CHU AND BURGESS 2016). Detailed annotated R-scripts and examples of all the generated outputs for each test are presented in File S1.

TetSim simulates the expected distribution of tetrads if spore death was purely random; that is, when one spore dies there is no corresponding death of another spore (CHU AND BURGESS 2016). If spore inviability of a given mutant was due solely to random spore death (RSD), the observed distribution of live:dead tetrads should resemble the output of TetSim (Figure 4A). If spore death was due, in part, to MI-nondisjunction, the observed distribution would show outliers from the 99% confidence limits of the simulated distribution. Applying TetSim to previously published tetrad data sets for WT and ndj1Δ (Wanat et al., 2008) using TetSim showed that not all of the live:dead tetrad classes fit a distribution predicted by RSD alone (Figure 4A, horizontal solid lines; CHU AND BURGESS 2016). Our own tetrad data agreed with these findings. For nup2Δ, nup60Δ, and htz1Δ, at least two of the live:dead tetrad classes fell outside
the 99th percentile for RSD alone (Figure 4A). These results indicate that the increased levels of spore death in these mutants was not solely due to random spore death. The \textit{nup53}\textDelta, \textit{nup100}\textDelta, and \textit{nup157}\textDelta mutants were not analyzed since spore viability in these strains was indistinguishable from WT (Table 1).

We next used the R-script TetFit to test if the decrease in observed spore viability for each genotype (Table 1 and Table S1) could be explained by MI-ND or a combination of MI-ND and RSD (Table S2; Chu AND Burgess 2016). TetFit-A finds the combination of these values that gives the best fit to the observed tetrad distribution (Table 2; Figure 4B). For WT, \textit{ndj1}\textDelta, \textit{nup2}\textDelta, \textit{nup60}\textDelta, and \textit{htz1}\textDelta the calculated best fits were not statistically different from the observed distributions ($P > 0.05$, adjusted for multiple comparisons), suggesting that both MI-ND and RSD could account for all of the observed spore death (Figure 4B).

TetFit also reports on the frequencies of death due to MI nondisjunction (nondisjunction death; NDD), thus allowing for comparison of the relative contributions of MI-ND and RSD to spore inviability. The NDD/RSD ratios for our WT and \textit{ndj1}\textDelta data sets were 2.0%/0.9% and 15.78%/2.7%, respectively (Table 2 and Figure 4B); in both cases the cause of spore death was skewed to NDD. This is consistent with NDD/RSD outcomes of TetFit analysis of previously published data sets for these genotypes (Wanat et al. 2008; Chu and Burgess, 2016), where the calculated NDD/RSD ratios for WT and \textit{ndj1}\textDelta were 1.8%/1.8% and 20.8%/5.8%, respectively. The output of TetFit applied to the observed tetrad distributions of \textit{nup2}\textDelta, \textit{nup60}\textDelta, and \textit{htz1}\textDelta gave greater NDD compared to WT, suggesting that spore death in these mutants can be attributed, in part, to MI-ND. However, the best fit NDD/RSD ratios in these mutants were skewed to greater RSD values: \textit{nup2}\textDelta was 5.2%/9.9%; \textit{nup60}\textDelta was 8.9%/23.1%; and \textit{htz1}\textDelta was
7.5%/29.4% (Table 2 and Figure 4B). Thus, unlike ndj1Δ, the cause(s) of spore death in nup2Δ, nup60Δ, and htz1Δ appear to be predominantly due to RSD rather than MI-ND.

The magnitude of the NDD/RSD ratio defines two classes of mutants (Chu and Burgess, 2016. Class A mutants give NDD/RSD > 3.3 and include those with known defects in proper homolog segregation (e.g. mad1Δ, mad2Δ, hypomorphic alleles of spo11, ndj1Δ, csm4Δ, msh5Δ, mlh1Δ and exo1Δ (Chua and Roeder 1997; Conrad et al. 1997; Kirkpatrick et al. 2000; Shonn et al. 2000; Marston et al. 2004; Cheslock et al. 2005; Martini et al. 2006; Wanat et al. 2008; Keelagher et al. 2011; Thacker et al. 2011). Class B mutants give NDD/RSD < 0.8 and include those with increased levels of precocious sister chromatid separation (PSCS) and/or meiosis II nondisjunction (MII-ND; e.g. sgs1Δ, sgs1Δ795, cbf1Δ, iml3Δ; and cep1Δ; Figure 4C; Masison and Baker 1992; Ghosh et al. 2004; Marston et al. 2004; Jessop et al. 2006; Rockmill et al. 2006; Chu and Burgess 2016). We found that the calculated NDD/RSD ratios for nup2Δ, nup60Δ, and htz1Δ were all less than 0.8 (Figure 4C; Table 5), which would classify them as Class B mutants.

To analyze the variation of the output values from TetFit-A, we ran 50 independent simulations using TetFit.Test using the calculated NDD and RSD values for each genotype and the corresponding number of experimentally dissected tetrads (Figure 4B, E columns; Table S2). The graphical output of the simulations shown in Figure 4C shows that each genotype gives well defined clusters representing NDD and RSD values for 50 simulated tetrad data sets.
Discussion

Here we describe a systematic evaluation of growth and meiosis phenotypes caused by deleting nonessential Nups in the SK1 strain background. Comparisons for growth on rich media and minimal synthetic media at four-different temperatures uncovered growth phenotypes for \textit{nup2Δ, nup60Δ, nup84Δ,} and \textit{nup100Δ}. A growth defect for \textit{nup84Δ} has been reported previously (SINIOSSOGLOU et al. 1996). For \textit{nup60Δ}, however, we found growth phenotypes (both positive and negative) where previous studies found no effect on growth on solid YPD media (DILWORTH et al. 2005; BARYSHNIKOVA et al. 2010). Side-by-side comparisons would be necessary to conclude that the defect we see is due to the SK1 strain background.

The \textit{nup2Δ, nup60Δ} and \textit{nup100Δ} mutants showed increased fitness compared to WT under certain stress conditions. The \textit{nup2Δ} and \textit{nup60Δ} mutants formed larger colonies on YPD at 37° compared to WT, while \textit{nup100Δ} mutant growth surpassed WT on minimal media at 25° and 30°. Increased fitness of a mutant compared to WT is characteristic of antagonistic pleiotropy (AP), where under certain environmental conditions, the WT copy of a gene decreases fitness compared to a mutant (WILLIAMS 2001). Genome-wide analysis of single-gene knockouts in yeast revealed that ~ 5.1% of nonessential genes show differences in the fitness of growth under one or more different challenged growth regimens (QIAN et al. 2012). To our knowledge, the conditions where we found AP (i.e. 37° YPD and minimal media) have not been evaluated in genome-wide analysis (BARYSHNIKOVA et al. 2010; QIAN et al. 2012). Our results indicate that Nup2, Nup60 and Nup100 suppress the efficiency of cell growth under certain conditions in the SK1 strain background.
Our analysis of sporulation and spore viability in the nup mutants uncovered phenotypes consistent with roles for Nup2, Nup60, Htz1 and Nup84 in the yeast meiotic program, but not Nup53, Nup100 and Nup157. We expected that these outcomes would correlate with the results of a genome-wide analysis of the yeast deletion collection that used a visual assay to score the segregation of a GFP-tagged chromosome III in intact tetrads (MARSTON et al. 2004). All of the mutants we analyzed were interrogated in this screen except for nup157Δ. While we found that spore viability of nup53Δ and nup100Δ and the kinetics of meiotic progression were the same as WT, the Marston study reported elevated levels of 2:2 GFP+: GFP- spores in these mutants, where primarily 4:0 would be expected for WT. In principle, the observed missegregation of chromosomes would have resulted in increased spore death, but this was not the case for our analysis.

For nup84Δ, yet we found that less than 1% of these cells sporulated, while no decrease in sporulation or increased nondisjunction was reported in the Marston study. Dramatic differences like this could be due to the presence of a suppressor mutation that could have been introduced from the yeast deletion collection in the Marston study, for which there is precedence (WINZELER et al. 1999; HUGHES et al. 2000). A more in depth evaluation of this phenotype could potentially uncover a functional link between these processes. For example, the meiotic defect we see in nup84Δ may reflect the defects of the nup84Δ mutant in telomere localization at the NE and DSB repair in vegetatively dividing cells as reported previously (LOEILLET et al. 2005; BUKATA et al. 2013).

For nup60Δ and htz1Δ we found defects in spore viability. While spore inviability was not measured in the Marston study, they found no evidence for chromosome missegregation.
This would be consistent with the increased RSD we found for these mutants, however, TetFit estimated a modest contribution of MI-ND to spore death. On the other hand, Marston et al found that \textit{nup2Δ} gave increased levels of 3:1 chromosome segregation errors, which is characteristic of defects leading to precocious sister chromatid separation (PSCS) or MI-ND segregation errors. This phenotype would appear as RSD using TetFit analysis and could possibly explain the increased spore death phenotype we saw for \textit{nup2Δ}. Together, these results suggest that spore death in \textit{htz1Δ} and \textit{nup60Δ} is due to different causes than for \textit{nup2Δ}.

The most surprising outcome of the study was that the \textit{nup2Δ ndj1Δ} double mutant uniquely gave a synthetic block to completing the MI nuclear division compared to all of the other \textit{nup ndj1Δ} double mutants. The \textit{nup2Δ} mutation alone delayed progression through MI unlike \textit{nup53Δ}, \textit{nup100Δ}, and \textit{nup157Δ}. These results suggest functional link between Nup2 and the chromosome events of meiosis. A more detailed analysis of Nup2’s role in meiosis will be described in a future publication (D.B.C and S.M.B., unpublished).
Materials and Methods

Strains

All strains in this study are derivatives of SK1 and listed in Table 3. All media were generated as previously described (TRECO AND LUNDBLAD 2001; HO AND BURGESS 2011; LUI et al. 2013). All experiments were performed with at least two independent trials and in all cases similar results were obtained. Gene knockouts were constructed using standard tailed PCR based gene replacement and tagging techniques (LONGTINE et al. 1998; GOLDSTEIN AND MCCUSKER 1999).

Statistical analysis

All statistical tests for significance are indicated in the text. Whenever multiple comparisons were performed the p values were corrected using the Holm method (HOlm 1979).

Spotting assay for vegetative growth

Diploid cells were prepared by first patching and mating haploid cells taken from glycerol stocks stored at –80° on YPG plates (2% peptone, 1% yeast extract, 3% glycerol, 2% bacto agar, 0.01% adenine sulphate, 0.004% tryptophan, 0.002% uracil) for 15 hours at 30°. Cells were then streaked onto YPD plates (2% peptone, 1% yeast extract, 2% glucose, 2% bacto agar, 0.01% adenine sulphate, 0.004% tryptophan, 0.002% uracil) for 2 days at 30°. Well-isolated diploid single colonies were inoculated into 5ml liquid YPD (2% peptone, 1% yeast extract, 2% glucose, 0.01% adenine sulphate, 0.004% tryptophan, 0.002% uracil) for 24 hours at 30° on a roller drum. The cells were diluted from their 5 ml culture to OD600 = 1.0. Suspended
cells were serially diluted 1:10 in a 96-well plate and 2.5 μl of the diluted cultures were spotted onto YPD and minimal medium plates (1% Difco yeast nitrogen base without amino acids, 2% glucose, 2% bacto agar) using a multichannel pipettor, followed by incubation at 20°, 25°, 30°, and 37°. YPD plates were imaged after incubation for 48 and 72 hours and minimal medium plates were imaged after incubation for 72 and 120 hours using an Epson Perfection 1200 U scanner.

**Time course protocol**

The time course protocol was followed as previously reported (Ho and Burgess 2011; Lui et al. 2013). Cells were synchronized for progression through meiosis by the same regimen used for assaying vegetative growth (above), except that the inoculated YPD liquid culture was grown for 30 hours instead of 24 hours and was not supplemented with uracil. Following the 30-hour growth period, cells were added to YPA (2% peptone, 1% yeast extract, 1% KOAc, 0.01% adenine sulphate, 0.004% tryptophan) to a final OD600 of 0.23 and grown for 14.5 hours at 30° on a roller drum. Cells were pelleted at 1000 x g for 3 minutes, washed with SPM (1% KOAc, 0.02% raffinose, 0.009% synthetic complete dropout powder) and resuspended in SPM to a final OD600 of 3.0. Cells were removed from the culture at various time points thereafter (starting at t = 0 hours), fixed in 40% ethanol and stained with DAPI to follow the kinetics of MI progression. The MI division was marked by the formation of cells with two well-differentiated DAPI stained foci. At least 200 cells were analyzed for each time point.

**Spore viability and nondisjunction analysis**
Viability was determined as the percent viable spores from dissected tetrads. Single colonies from YPD plates were patched onto solid SPM media (1% KOAc, 0.02% raffinose, 0.009% synthetic complete dropout powder, 2% bacto agar), incubated for three days at 30°, and then dissected on YPD plates. The YPD plates were incubated for four days at 30° and colony formation was evaluated by unaided visual inspection. Any visible growth was considered to be a viable spore. The calculated MI-ND frequencies were determined using TetFit (CHU AND BURGESS 2016). Expected live:dead tetrad frequencies were generated for all genotypes using TetFit with the following conditions: the number of nondisjunction intervals (ndint) was 500, the number of random spore death intervals (rsdint) was 500, aneuploidy induced death (anid) was 0.035, and the MI-ND multiplier (ndm) was 10. Detailed methods are described in File S1.

Data availability

All strains are available upon request. File S1 contains a detailed description of the R-scripts used in this study and example data input and output formats associated with TetSim, TetFit and TetFit.Test.

Acknowledgements

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Abbreviations List

WT – Wild type
MI – Meiosis I
MII – Meiosis II
ND - Nondisjunction
NPC – Nuclear pore complex
Nup – Nucleoporin
References


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Figure Legends

Figure 1

Schematic of the position(s) of each nonessential Nup protein examined in this study

Z-axis slice schematic of the nuclear pore complex showing the relative position of the Nups analyzed in this study (adapted from Alber et al., 2007). The cytoplasmic side is on the top of the outer of the nuclear membrane (ONM) and the nucleoplasmic side in below the inner nuclear membrane (INM; thick horizontal lines). Nup84 is located in the outer ring (green); Nup157 is located in the inner ring (black); Nup100 is an FG-Nup found on the cytoplasmic side of the pore channel (light gray); Nup53 is an FG-Nup found on both sides of the channel (dark gray); Nup60 (orange) and Nup2 (red) are FG-Nups is primarily on the nucleoplasmic side of the channel near the nuclear basket.

Figure 2

Growth of nup mutants on rich media (YPD) and minimal media at 20°, 25°, 30°, and 37°.

(A) Growth of WT and nup mutants at on YPD after 48 and 72 hours at the indicated temperatures. Shown are spotted 1:10 serial dilutions. (B) Same as in (A) except cells were spotted on minimal medium and imaged after 72 and 120 hours at the indicated temperatures.

Figure 3
Meiotic phenotypes of nonessential nucleoporins

(A) Percent viable spores are based on growth of spore clones from dissected 4-spore tetrads for the indicated deletion mutants (n = number of total spores dissected) after four days at 37°C.

(B) Representative meiotic time course of WT (blue), nup2Δ (red), nup53Δ (dark gray), nup100Δ (light grey), and nup157Δ (black). Multinucleate cells are those with 2 or more DAPI staining bodies. At least 200 cells were scored for each data point. (C) Same as in part (A) for the indicated deletion strains.

Figure 4

Estimation of nondisjunction death and random spore death caused by nup2Δ, nup60Δ, and htz1Δ.

(A) TetSim simulates a distribution of 4:0, 3:1, 2:2, 1:3, and 0:4 tetrads based on the observed fraction of viable cells (CHU AND BURGESS 2016) and compares the 99% cutoff values for the simulated distribution to the observed distribution for each genotype. The number of simulations for each panel is equal to the number of tetrads dissected experimentally. A distribution of dissected tetrad simulations is shown (in red, green, blue, teal, and purple) with the upper and lower dashed lines represent 99% and 1% percentiles, respectively. The observed frequencies of live:dead tetrad distributions for individual strains are shown as solid lines.

Comparison of simulated tetrad distributions with mutant data from previously published data sets for WT and ndj1Δ (WANAT et al. 2008). Note the fraction of viable spore in the ndj1Δ
mutant was higher than previously reported by Wanat et al., 2008. (B) Best fitting distributions of 4 (red), 3 (green), 2 (blue), 1 (teal), and 0 (purple)-viable spore tetrads in nup2Δ, nup60Δ, htz1Δ, and ndj1Δ mutants generated by TetFit (Chu and Burgess, 2016). The observed tetrad distributions of genotype (O, middle bar), the expected tetrad distributions with the best fitting MI-ND and RSD (E, right bar), and the expected tetrad distributions using the WT MI-nondisjunction rate and the best fitting RSD (W, left bar). Significance is noted as * P < 0.05; *** P < 0.001 (chi-squared, Holm corrected). (B) Output of TetFit, which finds the calculated estimated distributions of each live:dead tetrad class. The observed tetrad distributions of genotype (O, middle bar), the expected tetrad distributions with the best fitting MI-ND and RSD (E, right bar), and the expected tetrad distributions using WT MI-ND and the best fitting RSD (W, left bar). (C) 50 simulations of TetFit using dissected tetrad simulations corresponding to the estimated tetrad distributions shown in part (B) and the number of experimental tetrads shown in part (A). The solid line represents a NDD/RSD slope of 1, which indicates an equal contribution of NDD and RSD on spore inviability. The green shaded area covers the Class A mutants that have an NDD/RSD ratio above 3.3, and the pink shaded area covers Class B mutants that have an NDD/RSD ratio below 0.8 (CHU AND BURGESS 2016).

Table 1

Sporulation and spore viability in nup mutants
<table>
<thead>
<tr>
<th>Strain</th>
<th>% Sporulation efficiency (n)</th>
<th>Tetrads dissected</th>
<th>4-sv</th>
<th>3-sv</th>
<th>2-sv</th>
<th>1-sv</th>
<th>0-sv</th>
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</tr>
<tr>
<td>nup53Δ</td>
<td>94% (200)</td>
<td>72</td>
<td>66</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>95.8</td>
</tr>
<tr>
<td>nup100Δ</td>
<td>93% (200)</td>
<td>72</td>
<td>69</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>95.9</td>
</tr>
<tr>
<td>nup157Δ</td>
<td>94% (200)</td>
<td>72</td>
<td>69</td>
<td>1</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>98.2</td>
</tr>
<tr>
<td>htz1Δ</td>
<td>67% (200)</td>
<td>288</td>
<td>77</td>
<td>91</td>
<td>69</td>
<td>40</td>
<td>11</td>
<td>66.0</td>
</tr>
<tr>
<td>ndj1Δ</td>
<td>75% (200)</td>
<td>192</td>
<td>136</td>
<td>14</td>
<td>19</td>
<td>6</td>
<td>17</td>
<td>82.0</td>
</tr>
<tr>
<td>ndj1Δ&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td>851</td>
<td>482</td>
<td>112</td>
<td>123</td>
<td>38</td>
<td>96</td>
<td>74.9</td>
</tr>
<tr>
<td>nup2Δ ndj1Δ</td>
<td>1% (200)</td>
<td>NA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>nup60Δ ndj1Δ</td>
<td>46% (200)</td>
<td>48</td>
<td>16</td>
<td>4</td>
<td>8</td>
<td>3</td>
<td>7</td>
<td>65.1</td>
</tr>
<tr>
<td>nup53Δ ndj1Δ</td>
<td>75% (200)</td>
<td>144</td>
<td>103</td>
<td>10</td>
<td>12</td>
<td>6</td>
<td>13</td>
<td>82.0</td>
</tr>
<tr>
<td>nup100Δ ndj1Δ</td>
<td></td>
<td>72</td>
<td>48</td>
<td>3</td>
<td>14</td>
<td>1</td>
<td>6</td>
<td>79.9</td>
</tr>
<tr>
<td>nup157Δ ndj1Δ</td>
<td></td>
<td>72</td>
<td>39</td>
<td>5</td>
<td>14</td>
<td>4</td>
<td>10</td>
<td>70.5</td>
</tr>
<tr>
<td>htz1Δ ndj1Δ</td>
<td>41% (200)</td>
<td>72</td>
<td>15</td>
<td>18</td>
<td>17</td>
<td>13</td>
<td>9</td>
<td>55.9</td>
</tr>
</tbody>
</table>

<sup>a</sup> Data reported in Wanat et al., 2008

Table 2

Mean and standard deviation of 50 simulations of TetFit
### Table 3

#### Yeast strains used in this study

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBY1903</td>
<td>MATa/Matα ho::hisG/&quot; leu2::hisG/&quot; ura3/&quot; his4-x::LEU2-(NBam)-URA3/HIS4::LEU2-(NBam)</td>
</tr>
<tr>
<td>SBY4102</td>
<td>nup2Δ/&quot; SBY1903</td>
</tr>
<tr>
<td>SBY5216</td>
<td>nup60Δ/&quot; SBY1903</td>
</tr>
<tr>
<td>SBY5437</td>
<td>nup53Δ/&quot; SBY1903</td>
</tr>
<tr>
<td>SYB5586</td>
<td>nup84Δ/&quot; SBY1903</td>
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<tr>
<td>SBY5549</td>
<td>nup100Δ/&quot; SBY1903</td>
</tr>
<tr>
<td>SBY5256</td>
<td>nup157Δ/&quot; SBY1903</td>
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<tr>
<td>SBY2030</td>
<td>ndj1Δ/&quot; SBY1903</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Strain</th>
<th>RSD(^a)</th>
<th>NDD(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>0.010 ± 0.003</td>
<td>0.024 ± 0.009</td>
</tr>
<tr>
<td>ndj1Δ</td>
<td>0.0262 ± 0.005</td>
<td>0.160 ± 0.018</td>
</tr>
<tr>
<td>nup2Δ</td>
<td>0.104 ± 0.011</td>
<td>0.054 ± 0.011</td>
</tr>
<tr>
<td>nup60Δ</td>
<td>0.229 ± 0.019</td>
<td>0.087 ± 0.023</td>
</tr>
<tr>
<td>htz1Δ</td>
<td>0.292 ± 0.017</td>
<td>0.072 ± 0.025</td>
</tr>
</tbody>
</table>

\(^a\) Mean and standard deviation of estimated fraction of spores that die due to RSD (random spore death) and NDD (death due to nondisjunction) for each genotype.
<table>
<thead>
<tr>
<th>Strain</th>
<th>Mutations</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBY4116</td>
<td>nup2Δ/&quot; ndj1Δ/&quot;</td>
<td>SBY1903</td>
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<tr>
<td>SBY5268</td>
<td>htz1Δ/&quot;</td>
<td>SBY1903</td>
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<tr>
<td>SBY5222</td>
<td>nup60Δ/&quot; ndj1Δ/&quot;</td>
<td>SBY1903</td>
</tr>
<tr>
<td>SBY5278</td>
<td>htz1Δ/&quot; ndj1Δ/&quot;</td>
<td>SBY1903</td>
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A.  

<table>
<thead>
<tr>
<th></th>
<th>20°C</th>
<th>25°C</th>
<th>30°C</th>
<th>37°C</th>
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<tbody>
<tr>
<td>WT</td>
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<td></td>
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</tr>
<tr>
<td>nup2</td>
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<tr>
<td>nup53</td>
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<tr>
<td>nup60</td>
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</tr>
<tr>
<td>nup84</td>
<td></td>
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<tr>
<td>nup100</td>
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</tr>
<tr>
<td>nup157</td>
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</table>

B.  

<table>
<thead>
<tr>
<th></th>
<th>20°C</th>
<th>25°C</th>
<th>Minimal</th>
<th>30°C</th>
<th>37°C</th>
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<tbody>
<tr>
<td>WT</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>nup2</td>
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<td>nup53</td>
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<tr>
<td>nup60</td>
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<td></td>
</tr>
<tr>
<td>nup84</td>
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<td></td>
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<tr>
<td>nup100</td>
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<td></td>
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</tr>
<tr>
<td>nup157</td>
<td></td>
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</tr>
</tbody>
</table>

48 hr:

- 20°C
- 25°C
- 30°C
- 37°C

72 hr:

- 20°C
- 25°C
- 30°C
- 37°C

120 hr:

- 20°C
- 25°C
- 30°C
- 37°C

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A.  

- **WT**  
  Tetrads = 288  
  SV = 0.97  

- **WT**  
  Tetrads = 1199  
  SV = 0.97  

- **ndj1**  
  Tetrads = 192  
  SV = 0.82  

- **ndj1**  
  Tetrads = 851  
  SV = 0.75  

B.  

- **WT**  
  Tetrads = 384  
  SV = 0.86  

- **nup2**  
  Tetrads = 264  
  SV = 0.71  

- **nup60**  
  Tetrads = 264  
  SV = 0.66  

- **htz1**  
  Tetrads = 264  
  SV = 0.66  

C.  

- **ndj1**  
  Nondisjunction Death (NDD)  

- **nup60**  
  Random Spore Death (RSD)  

- **htz1**  
  Nondisjunction Death (NDD)  

- **nup2**  
  Random Spore Death (RSD)