In yeast and mammalian cells, the spindle assembly checkpoint proteins Mad1p and Mad2p localize to the nuclear pore complex (NPC) during interphase. Deletion of \textit{MAD1} or \textit{MAD2} did not affect steady-state nucleocytoplasmic distribution of a classical nuclear localization signal-containing reporter, a nuclear export signal-containing reporter, or Ran localization. We utilized cells with conditional mutations in the yeast Ran GTPase pathway to examine the relationship between Ran and targeting of checkpoint regulators to the NPC. Mutations that disrupt the concentration of Ran in the nucleus displaced Mad2p but not Mad1p from the NPC. The displacement of Mad2p in M-phase cells was correlated with activation of the spindle checkpoint. Our observations demonstrate that Mad2p localization at NPCs is sensitive to nuclear levels of Ran and suggest that release of Mad2p from NPCs is closely linked with spindle assembly checkpoint activation in yeast. This is the first evidence indicating that Ran affects the localization of Mad2p to the NPC.

In all eukaryotes, mitosis is controlled through the abundance of B-type cyclins and Pds1p (also known as Securin) (15). B-type cyclins associate with and activate the cdc28 kinase in budding yeast during mitosis. Pds1p forms an inactive complex with the Esp1p protease (also known as Separaese). Upon loss of Pds1p, Esp1p cleaves the cohesin Scc1p, which holds the sister chromatids together, allowing the onset of anaphase. Both B-type cyclins and Pds1 are ubiquitinated at the metaphase-anaphase transition by an E3 ligase called the anaphase-promoting complex (APC) (13). Premature APC activation is prevented in the presence of unattached kinetochores by a set of conserved spindle assembly checkpoint proteins, including Mad1p, Mad2p, Mad3p, Mps1p, Bub1p, and Bub3p (9). Some of these proteins have been localized to unattached kinetochores (Mad1p, Mad2p, Bub1p, and Bub3p), leading to a model in which a diffusible “delay anaphase” signal that prevents the activation of APC is generated by these proteins at unattached kinetochores (11, 22). After all of the chromosomes have achieved bipolar attachment and aligned on the mitotic spindle, the checkpoint is turned off, the APC becomes active, and anaphase ensues (9, 13).

Nuclear pore complexes (NPCs) are large, multiprotein structures that perforate the nuclear envelope (NE) and act as the conduits for nuclear-cytoplasmic trafficking (20). Recent studies have suggested a surprising link between the spindle checkpoint pathway and NPCs (19). In \textit{Saccharomyces cerevisiae}, the Mad1p/Mad2p complex associates with the NPC through the Nup53p-containing complex of nucleoporins (7). In the same study, it was observed that upon checkpoint activation, Mad2p (but not the bulk of Mad1p) is released from the NPC and accumulates on kinetochores. These findings led to the proposal of a model wherein the association between Mad1p and the Nup53p complex sequesters Mad2p at the NPC when the spindle checkpoint is inactive (7). In a separate study, Gillett et al. used chromatin immunoprecipitation analysis to demonstrate that both Mad1p and Mad2p localize to kinetochores upon treatment with microtubule-destabilizing drugs. This finding does not demonstrate that Mad1p is quantitatively removed from NPCs but does suggest that at least some portion of Mad1p relocalizes to the kinetochores upon checkpoint activation (6).

Ran is an abundant, highly conserved Ras-like GTPase that acts as a critical regulator of nuclear-cytoplasmic trafficking (10). Ran has been implicated in the regulation of mitosis (18) and shown to directly control checkpoint activation in \textit{Xenopus} egg extracts (3). We wished to examine the in vivo relationship between Ran and NPC-associated components of the spindle checkpoint pathway. Here we show that Mad2p localization to the NPC is closely linked to proper nuclear-cytoplasmic compartmentalization of the yeast Ran homologue, Gsp1p. The converse relationship was not found, since disruption of the \textit{MAD1} and \textit{MAD2} genes had no apparent effect on steady-state Gsp1p localization or classical nuclear localization signal (NLS) or nuclear export signal (NES) transport. Our data further indicate that there is a close relationship between Mad2p release from NPCs and spindle checkpoint activation.

**MATERIALS AND METHODS**

**Yeast strains and plasmids.** All yeast plasmids and strains used in this study are listed in Table 1.

**Localization of GFP fusion proteins.** In all cases, a fully functional Mad2-green fluorescent protein (GFP) was expressed from a low-copy-number plasmid by using its native promoter, and Mad1-GFP was integrated at the \textit{MAD1} locus. All GFP fusion proteins were localized by directly viewing the GFP signal in living cells through a GFP-optimized filter with a Zeiss Axioscope epifluorescence microscope equipped with an Orela digital camera and analyzed with Openlab software (1) (see Fig. 1, 2D, 4, and 5), or they were viewed with a Zeiss LSM510 META confocal microscope equipped with a 100× Plan Apochromat (NA 1.4, oil, differential interference contrast [DIC] objective and a 488-nm line Argon laser (25-mW nominal output: detection long pass [LP], 505 nm), and images were overlaid onto DIC images with Zeiss confocal microscopy software (version 3.2) (2) (see Fig. 2A and 3). To analyze \(G_{1}\)-arrested cells, cells were...
arrested at 25°C with α-factor mating hormone (5 μM) and then shifted to 37°C for 3 h in the presence of α-factor.

**Analysis of tsBN2 cells.** Cells from a hamster cell line that contains a temperature-sensitive allele of the vertebrate Ran nucleotide exchange factor (tSBN2 cells) were grown at 32°C and then shifted to 39.5°C for 3 h. The cells were permeabilized by using 0.2% Triton X-100 in phosphate-buffered saline for 5 min, fixed with 3.7% formaldehyde in phosphate-buffered saline for 5 min, and permeabilized by using 0.2% Triton X-100 in phosphate-buffered saline for 5 min. The cells were incubated with polyclonal anti-MAD2 antibodies (1:200 dilution) (gift from R. Chen) and mAb414 antibodies (1:5,000) followed by appropriate fluorescent secondary antibodies. Slides were examined with a Zeiss Axioskop fluorescence microscope, and images were collected and analyzed with Openlab software.

### RESULTS AND DISCUSSION

We analyzed Mad1p and Mad2p localization, Gsp1p localization, and checkpoint activation in cells with conditional mutations in the yeast Ran GTPase pathway (10): Ntf2p promotes the import of Gsp1p-GTP from the cytosol into the nucleus, whereas Prp20p promotes nucleotide exchange to generate Gsp1p-GDP (15). Gsp1p-GTP binds to transport receptors and exits from the nucleus via NPCs. In the cytosol, Gsp1p's GTPase-activating protein, Rna1p, promotes hydrolysis of the Gsp1p-associated GTP. The Ran pathway mutants that we analyzed included ntf2-1, ntf2-2 (5, 15), prp20-1 (2), and mad1-1 (4).

To assay Mad1p and Mad2p localization in living cells, both proteins were fused to the carboxy terminus to GFP. In all cases, Mad2-GFP was expressed from its native promoter on a low-copy-number plasmid that fully complements the benomyl sensitivity of a mad2Δ strain (data not shown). Mad1-GFP was integrated into the genome at the Mad1 locus and has previously been shown to fully complement endogenous MAD1 function (7). We initially compared the localization of Mad1-GFP and Mad2-GFP in ntf2-1, ntf2-2, and wild-type (WT) cells. The ntf2-1 mutation disrupts dimerization, while the ntf2-2 mutation disrupts binding to Gsp1p (15). Mad1-GFP remained at the NPC under both permissive (25°C) and non-permissive (37°C) conditions in all strains (Fig. 1A), indicating that these mutations did not disrupt its NPC targeting. However, Mad2-GFP was lost from NPCs in an allele-specific fashion: at 25°C, Mad2-GFP was barely visible at NPCs in 75% of WT cells and 66% of ntf2-1 cells, while only 36% of the ntf2-2 cells localized Mad2-GFP to NPCs (Fig. 1A and B). At 37°C, only 2% of ntf2-2 cells retained Mad2-GFP at the NPC, compared to 82% of WT cells and 70% of ntf2-1 cells. Notably, not only was the percentage of ntf2-2 cells that localized Mad2-GFP to the NPC significantly reduced in comparison to WT cells, but also the intensity of Mad2-GFP accumulation on NPCs was visibly lower in the subset of ntf2-2 cells where NPC accumulation could be observed than in WT cells. Mad2-GFP was retained within the nucleoplasm of ntf2-2 cells at 37°C, indicating that its failure to bind the NPC was not a simple result of its exclusion from the nucleus.

At 37°C, ntf2-2 cells arrest in G2/M phase as large budded cells with duplicated DNA, whereas ntf2-1 cells show a loss of viability that is not associated with arrest in any particular phase of the cell cycle (15). It was therefore plausible that Mad2p may be released from Mad1p at the NPC near the G2/M transition and that the loss of Mad2-GFP from NPCs

### TABLE 1. Strains and plasmids

| Strain or plasmid | Relevant genotype | Source |
|-------------------|------------------|--------|
| PSY 580          | Wild type        | 5      |
| ACY109           | prp20-1          | 15     |
| ACY60            | mad1-1           | 15     |
| BQY392           | MAD1-GFP::HIS5   | This study |
| BQY393           | ntf2::HIS3 (+pPS920) | This study |
| KH141            | mad2::URA3       | 15     |
| YMB1299          | MAD1-GFP::HIS5   | 7      |
| BQY141           | nup53Δ KanMX     |         |
| BQY142           | Wild type        | Research Genetics |
| BQY157           | mad1Δ KanMX      | Research Genetics |
| BQY381           | mad2Δ KanMX      | Research Genetics |
| BQY238           | ntf2::HIS3 (+pPS882) | This study |
| BQY239           | ntf2::HIS3 (+pPS920) | This study |
| BQY240           | ntf2::HIS3 (+pPS919) | This study |
| BQY413           | mad2Δ KanMX ntf2::HIS3 (+pPS920) | This study |
| BQY412           | ntf2::HIS3 (+pPS920) | This study |
| BQY414           | prp20-1 mad2Δ KanMX | This study |

| Plasmid          | Relevant genotype | Source |
|------------------|------------------|--------|
| pPS882           | CEN LEU2 NTF2    | 5      |
| pPS919           | CEN LEU2 ntf2-1  | 5      |
| pPS920           | CEN LEU2 ntf2-2  | 5      |
| pAC212           | 2μm URA3 pADH1 SV40-NLS PKI-NES 2×GFP | 8      |
| pAC410           | 2μm URA3 GSPI-GFP | 15     |
| pAC256           | 2μm URA3 PDS1-GFP | This study |
| pBQ181           | CEN TRP1 MAD2-GFP | This study |
| pBQ184           | 2μm URA3 pGAL SV40-NLS 2×GFP | This study |
| pBQ288           | CEN URA MAD2-GFP | This study |
ntf2-2 cells might result indirectly from cell cycle arrest. To test this possibility, we arrested WT and ntf2-2 cells in G1 phase with /H9251-factor, followed by a shift to 37°C for the continued presence of /H9251-factor (Fig. 1C). Both WT and ntf2-2 cells retained a shmoo morphology throughout the experiment, confirming that they remained appropriately arrested. Under these circumstances, Mad2-GFP remained at the NPC in the WT cells, but it was displaced from the NPC in the ntf2-2 cells (Fig. 1C). This finding shows that Mad2p is released from the NPC in ntf2-2 cells at the restrictive temperature, regardless of cell cycle stage.

We next analyzed Mad2-GFP localization at the permissive (25°C) and restrictive (37°C) temperatures in prp20-1 and rna1-1 cells. There were fewer prp20-1 cells with Mad2-GFP at the NPC than both WT and rna1-1 cells at 25 and 37°C (Fig. 2). At 37°C, the residual Mad2-GFP was completely lost from the NPC in prp20-1 cells. In rna1-1 cells, the reduction of Mad2-GFP association with NPCs at 37°C was less dramatic: we found no NPC-associated Mad2-GFP within 72% of cells, while 28% of cells still showed a concentration of Mad2-GFP at their NEs. As with ntf2-2 cells, Mad2-GFP still concentrated within the nucleoplasm in the majority of prp20-1 and rna1-1 cells at 37°C, indicating that a loss of NPC association was not an indirect result of reduced levels of Mad2-GFP or nuclear exclusion. Together with observations of ntf2-2 cells, these observations suggested that disruption of the Ran pathway profoundly alters the localization of Mad2p to the NPC.

We examined the localization of Mad2p by immunofluorescence in a hamster cell line that contains a temperature-sensitive allele of the vertebrate Ran nucleotide exchange factor, RCC1 (tsBN2 cells) (12). A subpopulation of Mad2 clearly localizes to the nuclear rim in tsBN2 cells at the permissive temperature (32°C) (Fig. 2C). This population of Mad2p was no longer localized to NPCs 3 h after the shift to the restrictive temperature (39.5°C). As in yeast, a large fraction of Mad2p remained within the tsBN2 nucleus at the restrictive temperature. This finding strongly suggests that the Ran pathway is required for the retention of the Mad2 protein on the NPC during interphase in vertebrate cells, suggesting that the role of Ran in Mad2p localization may be highly conserved.

We considered two possible explanations of our findings in yeast. First, localization of Mad2p to the NPC may require ongoing nuclear transport. This idea is not consistent with the presence of Mad2-GFP at the NPC in ntf2-1 cells at 37°C, because it has been previously demonstrated that ntf2-1 cells have lower transport capacity than ntf2-2 cells under nonpermissive conditions (15) (see Fig. 4A). Alternatively, the localization of Mad2p may be directly sensitive to the localization of Gsp1p or its nucleotide binding state. To examine the behavior of Gsp1p in the mutant strains, we analyzed the localization of a Gsp1-GFP fusion protein in each of the mutants at 25 and 37°C (Fig. 3).

As previously reported (15), ntf2-1 cells concentrate Gsp1-GFP within their nuclei at both temperatures (Fig. 3A), al-
though there is an increase in cytoplasmic Gsp1-GFP in these cells compared to WT cells. This may account for the decreased level of Mad2-GFP at the pore observed in this mutant (Fig. 1). By contrast, ntf2-2 cells exhibit extensive loss of Gsp1-GFP compartmentalization at 37°C. In the prp20-1 cells, Gsp1-GFP localization was abnormal even at 25°C: Gsp1-GFP localized to the nuclear periphery instead of filling the entire nuclear compartment (Fig. 3B). The localization of Gsp1-GFP was further disrupted in prp20-1 cells at 37°C, with a loss of Gsp1-GFP compartmentalization similar to that of ntf2-2 cells. Gsp1-GFP localization was also abnormal at 25°C in rna1-1 cells, although this defect was not as severe as that in prp20-1 cells. Approximately 25% of rna1-1 cells showed some punctuate nuclear rim staining at 25°C (Fig. 3B). For reasons that are unclear, Gsp1-GFP protein levels were consistently lower in rna1-1 cells at 37°C than at 25°C (Fig. 3C). The residual Gsp1-GFP was no longer concentrated within the nucleus of rna1-1 cells at 37°C, although it was enriched on the nuclear envelope in 35% of cells (Fig. 3).

In summary, the two strains with mutations in the Ran pathway that showed complete displacement of Mad2-GFP from the NPC (the ntf2-2 and prp20-1 mutants) at the restrictive temperature also showed a loss of Gsp1p compartmentalization under the same conditions. rna1-1 cells exhibited an intermediate phenotype, with a residual population of Gsp1p at the NE and incomplete dissociation of Mad2-GFP from NPCs. By contrast, the single mutant that maintained nuclear Gsp1-GFP localization at 37°C (ntf2-1) also maintained Mad2p on NPCs. Together, these findings suggest that Gsp1p concentration within the nucleus is closely correlated with Mad2p targeting to the NPC. Interphase tsBN2 cells similarly showed both a loss of Ran compartmentalization (16) and release of Mad2 from the NPC at 39.5°C (Fig. 2C), suggesting that similar mechanisms may be operative in metazoan cells.

We considered two possible roles for the regulated localization of Mad2p to the NPC: Mad2p might promote nuclear-cytoplasmic trafficking or its sequestration to NPCs might control its activity in the mitotic checkpoint. It has previously been shown that deletion of MAD1 inhibits import by the Kap121p
transport receptor (7). However, deletion of MAD2 had no apparent effect in the same assay. To determine more generally whether the deletion of MAD1 or MAD2 alters classical nucleocytoplasmic transport, we examined whether cells with deletions in the MAD1 (mad1Δ) or MAD2 (mad2Δ) genes maintain correct localization of a model substrate bearing a classical nuclear localization sequence (cNLS-GFP) (Fig. 4A). For comparison, the localization of the reporter was also examined in ntf2-1 and ntf2-2 cells at the nonpermissive temperature (37°C). ntf2-1 and ntf2-2 cells showed extensive and partial loss of cNLS-GFP nuclear localization, respectively. By contrast, both the mad1Δ and mad2Δ cells were indistinguishable from the WT controls, arguing that the deletion of MAD1 or MAD2 has an insignificant effect on NLS-mediated protein import.

To investigate whether Mad1p or Mad2p has a role in protein export, we examined a similar model substrate bearing a leucine-rich nuclear export sequence (NLS-NES-GFP) in mad1Δ and mad2Δ cells (Fig. 4B). When nuclear export is compromised, this reporter is redistributed to the nucleus due to the presence of a functional NLS within the protein (8). The steady-state distribution of NLS-NES-GFP between the cytoplasm and nucleus in mad1Δ and mad2Δ cells is essentially indistinguishable from the distribution in WT cells, arguing that these cells retain high levels of nuclear export capacity. Together with the observations from cNLS-GFP import, these data suggest that steady-state nucleocytoplasmic trafficking is not altered in the absence of Mad1p and Mad2p. Moreover, GFP-Gsp1p remained localized within the nucleus of both mad1Δ and mad2Δ cells (Fig. 4C), arguing against the notion that Mad2p or Mad1p is required for the maintenance of Gsp1p-GTP within the nucleus. For all of these reasons, we feel that it is very unlikely that Mad1p or Mad2p plays a direct role in regulation of bulk trafficking through the NPC.

Alternatively, it has previously been shown that deletion of the NUP53 gene renders cells resistant to the microtubule (MT)-destabilizing drug benomyl (7). These observations would be consistent with the possibility that Mad2p release from NPC potentiates its activity in the spindle checkpoint, thereby rendering the cells better able to cope with MT destabilization. This hypothesis predicts that Mad2p release by other means should also promote checkpoint activation. To test this prediction, we examined whether ntf2-2, prp20-1, and rna1-1 mutants showed resistance to benomyl (Fig. 5A), as nup53Δ cells do (7). It was previously found that the ntf2-2 mutation conferred benomyl sensitivity (15). In the course of performing these experiments, we determined that the benomyl sensitivity observed in those experiments was due to an additional background mutation present in the ntf2-2 strain used at that time. We grew each of the yeast strains at 25°C on plates containing 20 μg of benomyl/ml. ntf2-1 and prp20-1 cells exhibited benomyl resistance, although ntf2-2 cells did not, consistent with the idea that NPC release potentiates spindle checkpoint activation under conditions where Gsp1p-GTP levels are low (Fig. 5A).

Ran-GTP has been shown to have roles in MT dynamics in fission yeast (17) and to promote MT polymerization in Xenopus egg extracts (14). While some contribution of MT dynamics toward the benomyl resistance phenotype cannot be fully eliminated, we do not believe that benomyl resistance of ntf2-2 and prp20-1 cells resulted simply from MT stabilization for
several reasons. First, *ntf2*-2 cells arrest with morphologically normal short metaphase spindles, suggesting that their MT dynamics are relatively normal (15). Second, *ntf2*-2 *mad2*/*H9004* double mutants are viable and undergo normal mitosis with WT spindle morphology in the absence of benomyl (15). If the *ntf2*-2 mutation altered spindle assembly, *ntf2*-2 *mad2*/*H9004* cells should show inviability and increased chromosome missegregation. Moreover, similar to the benomyl resistance observed for *nup53*/*H9004* cells (7), the benomyl resistance of *ntf2*-2 and *prp20*-1 cells was entirely dependent upon the presence of Mad2p, since the *ntf2*-2 *mad2*Δ cells showed inviability and increased chromosome missegregation. Moreover, similar to the benomyl resistance observed for *nup53Δ* cells (7), the benomyl resistance of *ntf2*-2 and *prp20*-1 cells was entirely dependent upon the presence of Mad2p, since the *ntf2*-2 *mad2*Δ and *prp20*-1 *mad2*Δ double mutants were as sensitive to benomyl as *mad2*Δ cells (Fig. 5B). This finding suggests that changes in Mad2p activity in these cells at the permissive temperature underlie the benomyl resistance observed.

Interestingly, *mal1*-1 cells were not benomyl resistant at 25°C (Fig. 5A). It is possible that the NPC association of Mad2p was not as perturbed in this mutant as in *ntf2*-2 or *prp20*-1 cells. This idea is consistent with the frequent persistence of Mad2-GFP on NPCs of *mal1*-1 cells at 37°C (Fig. 2B and C). It is also possible that the nucleotide binding state of Gsp1p is important for checkpoint activation: defects in Ntf2p or Prp20p should cause accumulation of GDP-bound Gsp1p, whereas loss of Rna1p function should cause accumulation of GTP-bound Gsp1p. Increased Ran-GTP levels disrupt the localization of Mad2p and other spindle checkpoint regulators to kinetochores in *Xenopus* egg extracts (3), and Gsp1p-GTP might act similarly in yeast. These two possibilities are obviously not mutually exclusive, and it is imaginable that the nucleotide binding state of Gsp1p may influence both the release of Mad2p from the NPC and its activity in the spindle checkpoint.

Finally, we noted that *ntf2*-2 cells arrest at the G₂/M transition when shifted to 37°C. To determine whether this arrest is related to activation of the spindle checkpoint, we analyzed the stability of Pds1p in living WT and *ntf2*-2 cells at 37°C by using a Pds1-GFP fusion protein (Pds1-GFP) (21). Pds1-GFP fluorescence was reduced or completely lost in large budded cells (Fig. 5B, arrows), and a smaller percentage of cells exhibited Pds1-GFP fluorescence. However, more *ntf2*-2 cells retain Pds1-GFP fluorescence at 37°C, with 33% of cells arresting as large budded cells. Localization of Gsp1-GFP (pAC410) in WT (BQY142), *mal1*Δ (BQY157), and *mal2*Δ (BQY381) cells is shown.
results argue that Ran interacts with the spindle checkpoint pathway in yeast as well as in vertebrates, suggesting that Ran’s function as a mitotic regulator may be conserved across all eukaryotes, and this report is the first time that Ran has been shown to affect the localization of Mad2p to the nuclear pore complex.

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FIG. 5. Displacement of Mad2p from the NPC correlates with checkpoint activation. (A) WT (BQY142), mad2Δ (BQY381), nup53Δ (BQY141), prp20-1 (ACY109), rna1-1 (ACY60), ntf2-1 (BQY240), ntf2-2 (top, BQY239; bottom, BQY412), ntf2-2 mad2Δ (BQY413), and prp20-1 mad2Δ (BQY414) cells grown on plates containing benomyl. Cells were grown overnight at 25°C, and 105, 104, 103, 100, and 10 cells were spotted onto yeast extract-peptone-dextrose (YPED) medium and YEPD containing 20 μg of benomyl/ml. Plates were incubated at 25°C for 4 days. (B) Pds1p destruction is delayed in ntf2-2 cells at the restrictive temperature. Shown are micrographs of asynchronously growing WT (BQY142) and ntf2-2 (BQY412) cells expressing Pds1-GFP (pAC256) after shift to 37°C for 3 h. Arrowheads indicate cells in late mitosis. Shown below is the percentage of cells expressing Pds1-GFP at the point diagramed.