A mitotic nuclear envelope tether for Gle1 also affects nuclear and nucleolar architecture

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ABSTRACT During Aspergillus nidulans mitosis, peripheral nuclear pore complex (NPC) proteins (Nups) disperse from the core NPC structure. Unexpectedly, one predicted peripheral Nup, Gle1, remains at the mitotic nuclear envelope (NE) via an unknown mechanism. Gle1 affinity purification identified mitotic tether for Gle1 (MtgA), which tethers Gle1 to the NE during mitosis but not during interphase when Gle1 is at NPCs. MtgA is the orthologue of the Schizosaccharomyces pombe telomere-anchoring inner nuclear membrane protein Bqt4. Like Bqt4, MtgA has meiotic roles, but it is functionally distinct from Bqt4 because MtgA is not required for tethering telomeres to the NE. Domain analyses showed that MtgA targeting to the NE requires its C-terminal transmembrane domain and a nuclear localization signal. Of importance, MtgA functions beyond Gle1 mitotic targeting and meiosis and affects nuclear and nucleolar architecture when deleted or overexpressed. Deleting MtgA generates small, round nuclei, whereas overexpressing MtgA generates larger nuclei with altered nuclear compartmentalization resulting from NE expansion around the nucleolus. The accumulation of MtgA around the nucleolus promotes a similar accumulation of the endoplasmic reticulum (ER) protein Erg24, reducing its levels in the ER. This study extends the functions of Bqt4-like proteins to include mitotic Gle1 targeting and modulation of nuclear and nucleolar architecture.

INTRODUCTION
The nuclear envelope (NE) separates the nucleus and its DNA from the rest of the cell. The NE consists of two juxtaposed membranes—the inner nuclear membrane (INM) facing the nuclear interior, and the outer nuclear membrane (ONM), which is continuous with the endoplasmic reticulum (ER) and faces the cytoplasm (Hetzer, 2010). Alterations of the NE occur during aging and disease (Wolfrer and Wilson, 2001; Worman, 2004; Worman et al., 2009; Walters et al., 2012; Hatch and Hetzer, 2014). For example, aberrant nuclear morphology is a diagnostic marker for cancerous cells (Zink et al., 2004; Chow et al., 2012). It is not only in disease states that the NE can have abnormal structures. In many cell types in both plants and mammals, the NE can form intranuclear invaginations referred to as the nucleoplastic reticulum, although the functional significance and etiology of these structures are not understood (Malhas et al., 2011).

The NE is traversed by nuclear pore complexes (NPCs), which form sites where the INM and ONM are fused (Doucet et al., 2010; Fichtman et al., 2010; Talamas and Hetzer, 2011) and provide regulated conduits to allow selective transport of macromolecules in and out of nuclei. They are composed of ∼30 different proteins called nucleoporins (Nups), which are present in multiple copies per NPC. The structure of NPCs is conserved and contains a core scaffold embedded in the NE, which surrounds the central channel transport pore. Peripheral Nups are anchored to the core scaffold and extend into the central channel, cytoplasm, or nucleoplasm (D’Angelo and Hetzer, 2008).

During mitosis, the NE undergoes dramatic structural changes (Prunuske and Ullman, 2006; Guttinger et al., 2009), and there is great diversity in how organisms modify the NE to ensure successful...
mitosis. Whereas vertebrate cells completely disassemble their NPC and NE (open mitosis), in some organisms, both NPCs and the NE remain intact (closed mitosis). Between these extremes, the NE remains physically intact in the model filamentous fungus *Aspergillus nidulans*, but NPCs partially disassemble, with 14 of the ~30 Nups dispersing from the core NPC scaffold (partially open mitosis; De Souza et al., 2004; Osmani et al., 2006a; Liu et al., 2009). All 14 of the dispersed Nups are peripheral Nups. However, 1 peripheral Nup, Gle1, unexpectedly remains at the NE and does not disperse from the NE during mitosis (Osmani et al., 2006a; Ukil et al., 2009). Gle1 is a conserved, essential RNA export factor with additional roles in translation initiation and termination (Bolger et al., 2008). Shuttling of Gle1 between nucleoplasm and cytoplasm is essential for mRNA export in humans (Kendirgi et al., 2003). Gle1 associates with the cytoplasmic filaments of NPCs and aids in the release of messenger ribonucleoprotein particle (mRNP) complexes into the cytoplasm from the nucleus by activating the ATPase and mRNP remodeling activity of the DEAD-box helicase Dbp5 (Rayala et al., 2004; Kendirgi et al., 2005; Folkman et al., 2011).

In addition to the partial disassembly of its NPCs, the mitotic NE in *A. nidulans* is also dramatically spatially modified. After anaphase, the NE constricts around daughter nuclei on either side of the nucleolus, which becomes positioned between the segregating chromosomes (Ukil et al., 2009). Therefore the NE surrounds three components during anaphase—the central nucleolus and the two forming daughter nuclei. Although all core Nups localize preferentially to the NE around forming daughter nuclei, Gle1 is an exception and locates evenly at the NE around all three of these components during anaphase by an unknown mechanism. The presence of Gle1 at the mitotic NE in *A. nidulans* is therefore intriguing. We report here the requirement for the tail-anchored integral NE protein mitotic tether for Gle1 (MtgA; an orthologue of *Schizosaccharomyces pombe* Bqt4) for targeting Gle1 to the NE specifically during mitosis and maintaining normal nuclear and nucleolar architecture.

**RESULTS**

**MtgA is a conserved protein identified from Gle1 affinity purification**

We previously reported that Gle1 remains at the NE during mitosis, whereas all other peripheral Nups disperse from the NE (Osmani et al., 2006a; Ukil et al., 2009). During anaphase, Gle1 locates at the NE around the centrally positioned nucleolus, whereas core Nup proteins do not (Osmani et al., 2006a; Ukil et al., 2009). We thus reasoned that Gle1 remains at the mitotic NE by associating with a NE protein that is not a core Nup. To test this, we performed Gle1 affinity purifications and identified copurifying proteins by mass spectrometry. Gle1 copurified with two peripheral Nups (SonA and Nup42), the *A. nidulans* importin-α (KapA), and AN0162 (Figure 1A). Because SonA-GFP (De Souza et al., 2004) and KapA-CR (Markina-Inarrairaegui et al., 2011) disperse from nuclei during mitosis (Supplemental Figure S1A), they are unlikely to be the mitotic NE tethers for Gle1. Furthermore, although Nup42 was found to partially remain at the mitotic NE, it is not required for Gle1 targeting to the NE during interphase or mitosis (Supplemental Figure S1B). Therefore we focused on the previously uncharacterized Gle1-copurifying protein encoded from the locus AN0162 as a potential mitotic NE tether for Gle1. From >100 protein affinity purifications, including all known NPC proteins in *A. nidulans*, AN0162 copurified only with Gle1, KapA, and SonA (A.H.O., unpublished data). Of importance, AN0162–S-tag copurified Gle1, as well as Nup42, SonA, and KapA (Supplemental Figure S1C). For reasons discussed later, we named this protein MtgA.

**FIGURE 1:** MtgA is a transmembrane protein that copurifies with Gle1 and is conserved among the Ascomycetes and Basidiomycetes. (A) Identity and sequence coverage of proteins identified from Gle1–S-tag affinity purification (strain SO901) followed by mass spectrometry. SonA and Nup42 are peripheral Nups. KapA is the orthologue of importin-α. AN0162 (MtgA) is a previously uncharacterized protein with a single-pass TMD at its very C-terminus and a conserved APSES KilA-N-like nucleic acid–binding domain (KilA-N). (B) KilA-N proteins in *A. nidulans* and their predicted orthologues in *S. pombe* and *S. cerevisiae*, with e-values given in parentheses. (C) Phylogram representing protein orthologues of MtgA across fungi. Organisms shown are *A. nidulans*, *Sclerotinia sclerotiorum* (locus SS1G_08511), *Neurospora crassa* (locus NCU06560), *S. pombe*, *Schizopyllum commune* (locus SCHCODRAFT_269950), and *Ustilago maydis* (locus UMAG_11055). *S. commune* and *U. maydis* are Basidiomycetes. All other taxa are Ascomycetes. StuA, RgdA, and AN6715 are the three other KilA-N proteins in *A. nidulans*. We were particularly interested in MtgA due to the presence of a single-pass transmembrane domain (TMD) at its very C-terminus (Figure 1A), suggesting that it could integrate into the nuclear membrane. Also of interest was its putative nucleic acid–binding KilA-N domain. The KilA-N-domain family includes the previously defined fungal APSES domain of some fungal transcription factors (Iyer et al., 2002). There are four KilA-N–domain proteins in *A. nidulans*.
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...cifically from Bqt4 with regard to telomere tethering to the NE during the meiotic cell cycle, although it is required for normal levels of meiotic recombination.

Mtga is an integral membrane protein of the nuclear envelope

Green fluorescent protein (GFP) fused to the N-terminus of endogenous Mtga localizes to the NE during interphase and mitosis. Of note, just like Gle1-GFP, during mitosis, GFP-MtgA localizes to the NE surrounding the separating daughter nuclei, as well as the centrally positioned nucleolus during telophase (Figure 3A).

Given that Mtga associates with a NPC protein, it could be a transmembrane Nup. We therefore observed GFP-MtgA localization in the Δnup133 genetic background, which causes NPC clustering (Doye et al., 1994). In this background, proteins associated with NPCs appear in clusters, whereas nuclear membrane proteins adopt a more even distribution around the NE. GFP-MtgA was not seen in NPC clusters (marked by the transmembrane Nup Pom152) and had an even distribution around the NE (100% of 154 nuclei), indicating that it does not associate with NPCs (Figure 3B). This shows that Mtga is a nuclear membrane protein—likely an INM protein like Bqt4 in S. pombe (Chikashige et al., 2009).

Some integral INM proteins are transported to the INM after integrating into the endoplasmic reticulum (ER) membrane via their TMD and subsequent nuclear localization sequence (NLS)-dependent transport through the nuclear pore membrane into the INM (King et al., 2006; Lusk et al., 2007; Meinema et al., 2011). Such proteins that lack their NLS but retain their TMD and therefore localize to the ER. To determine whether Mtga localization is consistent with that of an INM protein, we performed a deletion analysis using mCherry-tagged truncated versions of Mtga to replace the wild type and tracked their localization using confocal microscopy (Figure 3C).

To determine whether Mtga requires its TMD for INM insertion, we tested truncations that lacked the TMD and the TMD and NLS-dependent localization of Mtga was lost, indicating that the TMD is required for INM localization.

Mtga is required for recombination in the NE and...
orthologue, Bqt4. During mitosis, as does Gle1-GFP (strain MC207), in contrast to wild-type cells, in which Gle1 remains at the NE during mitosis, in the ΔmtgA background, Gle1-GFP localizes normally to the NE during interphase but disperses from the NE during mitosis and returns to the NE upon completion of mitosis (Figure 4A, 100% of 71 mitoses). This indicates that Gle1 transitions from NPCs to the nuclear membrane via MtgA at mitosis. Of note, it is difficult to observe transition of Gle1 from NPCs to the mitotic nuclear membrane in WT cells due to the close proximity of NPCs with the nuclear membrane. However, in Δnup133 cells, which have clustered NPCs, this transition was clear. Gle1 located to the NPC clusters (Figure 4B, G2) during interphase but moved from the NPC clusters onto the nuclear membrane during mitosis (Figure 4B, M; 100% of 84 mitoses). If the transition of Gle1 from NPC clusters to the nuclear membrane during mitosis is via MtgA, then deletion of mtgA in the Δnup133 background would be predicted to result in Gle1 localization to NPC clusters during G2, followed by its dispersal from NPCs into the cytoplasm during mitosis due to the absence of MtgA at the nuclear membrane. Subsequently Gle1 would be expected to return to NPCs after mitotic exit. These expectations are fulfilled in the Δnup133ΔmtgA double-deletion background (in 100% of 32 mitoses; Figure 4C). These experiments provide additional lines of evidence that Gle1 associates with NPCs during interphase but dissociates from NPCs during mitosis and localizes to the INM through its interaction with MtgA at mitosis. Of note, it is difficult to observe transition of Gle1 from NPCs to the mitotic nuclear membrane in WT cells due to the close proximity of NPCs with the nuclear membrane. However, in Δnup133 cells, which have clustered NPCs, this transition was clear. Gle1 located to the NPC clusters (Figure 4B, G2) during interphase but moved from the NPC clusters onto the nuclear membrane during mitosis (Figure 4B, M; 100% of 84 mitoses). If the transition of Gle1 from NPC clusters to the nuclear membrane during mitosis is via MtgA, then deletion of mtgA in the Δnup133 background would be predicted to result in Gle1 localization to NPC clusters during G2, followed by its dispersal from NPCs into the cytoplasm during mitosis due to the absence of MtgA at the nuclear membrane. Subsequently Gle1 would be expected to return to NPCs after mitotic exit. These expectations are fulfilled in the Δnup133ΔmtgA double-deletion background (in 100% of 32 mitoses; Figure 4C). These experiments provide additional lines of evidence that Gle1 associates with NPCs during interphase but dissociates from NPCs during mitosis and localizes to the INM through its interaction with MtgA. To our knowledge, the data reveal the first example of a protein targeted to two distinct regions of the NE via MtgA at mitosis. Scale bar, 5 μm.

32 amino acids (aa) N-terminal to it (MtgA1-372). Neither was able to integrate into the INM but instead localized to the nuclear interior (in 96.7% of 303 nuclei expressing CR-MtgA1–404, and 98% of 603 nuclei expressing CR-MtgA1–372), indicating that they contained a functional NLS (Figure 3C, 1–404 and 1–372). Further analysis confirmed that MtgA contains a functional NLS between aa 291 and 372 (Figure 3C, 291–372, 97.6% of 206 nuclei). This version and the version lacking only the TMD both localize exclusively to nuclei during mitosis but then disperse from nuclei during interphase like other nuclear proteins, including NLS-DsRed version and the version lacking only the TMD both localize exclusively to nuclei during interphase but then disperse from nuclei during mitosis, in the ΔmtgA background, Gle1-GFP associates with NPC clusters marked by Pom152-CR. However, during mitosis (M), Gle1-GFP adopts a smoother NE distribution and does not associate with NPC clusters. (C) Gle1-GFP localizes to NPC clusters during interphase but disperses from the NE in strain MC167, which lacks both nup133 and mtgA. (D) Model for Gle1 localization during interphase and then mitosis. During interphase, Gle1 is at NPCs, but it transitions from NPCs to the INM during mitosis via MtgA.

During mitosis, Gle1 translocates from NPCs to MtgA

To test whether MtgA tethers Gle1 to the NE during mitosis, we followed Gle1-GFP distribution during the cell cycle in ΔmtgA cells. In contrast to wild-type cells, in which Gle1 remains at the NE during mitosis, in the ΔmtgA background, Gle1-GFP locates normally to the NE during interphase but disperses from the NE during mitosis and returns to the NE upon completion of mitosis (Figure 4A, 100% of 71 mitoses). This indicates that Gle1 transitions from NPCs to the nuclear membrane via MtgA at mitosis. Of note, it is difficult to observe transition of Gle1 from NPCs to the mitotic nuclear membrane in WT cells due to the close proximity of NPCs with the nuclear membrane. However, in Δnup133 cells, which have clustered NPCs, this transition was clear. Gle1 located to the NPC clusters (Figure 4B, G2) during interphase but moved from the NPC clusters onto the nuclear membrane during mitosis (Figure 4B, M; 100% of 84 mitoses). If the transition of Gle1 from NPC clusters to the nuclear membrane during mitosis is via MtgA, then deletion of mtgA in the Δnup133 background would be predicted to result in Gle1 localization to NPC clusters during G2, followed by its dispersal from NPCs into the cytoplasm during mitosis due to the absence of MtgA at the nuclear membrane. Subsequently Gle1 would be expected to return to NPCs after mitotic exit. These expectations are fulfilled in the Δnup133ΔmtgA double-deletion background (in 100% of 32 mitoses; Figure 4C). These experiments provide additional lines of evidence that Gle1 associates with NPCs during interphase but dissociates from NPCs during mitosis and localizes to the INM through its interaction with MtgA. To our knowledge, the data reveal the first example of a protein targeted to two distinct regions of the NE via different targeting proteins that is regulated in a cell cycle–dependent manner (Figure 4D).

MtgA is required to maintain nuclear shape and size

MtgA-deleted strains have nuclei that are noticeably rounder than wild-type cells (Figure 5A), and their nuclear volume is also reduced.
MtgA affects nuclear architecture

MtgA maintains spatial organization of nucleolar proteins

In A. nidulans, as in yeast cells, kinetochores cluster at the spindle pole body (SPB) during interphase (Wigge and Klarmann, 2001; Yang et al., 2004). The SPB is embedded in the NE in a position typically opposite to the nucleolus, which is located adjacent to the NE at the other end of the nucleus. The ribosomal DNA (rDNA) region forming the nucleolar-organizer region (NOR) in A. nidulans is located on the long arm of chromosome V (Brody et al., 1991; Clutterbuck and Farman, 2008). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009). The nucleolus is generated around the NOR, with the single NOR generating one nucleolus per nucleus. The nucleolar structure can be divided into three subregions. The fibrillar center (FC) and dense fibrillar (DF) regions are close to the NOR, whereas the granular component (GC) lies more at the nucleolar periphery (Boisvert et al., 2009).
by Bop1 (Figure 6B, WT). In contrast, Bop1 almost entirely surrounds fibrillarin in mtgA-deleted strains (Figure 6B, ΔmtgA).

Because the nucleolus is built around NORs, we tracked the NOR to investigate its relationship with MtgA. To enable live-cell imaging of the NOR, we imaged endogenously tagged Pol I (Pol I-CR), which associates specifically with the NOR (Boisvert et al., 2007; Ukil et al., 2009). Within WT nuclei, Pol I-CR locates as a focus toward the NE at the nuclear periphery, as does the nucleolus. In 37% of 265 nuclei observed, MtgA was observed to concentrate at distinct regions of the NE, with the majority of these nuclei (79%) having MtgA concentrations at the NE region to which Pol I–CR also associates (Figure 6C, arrowhead). This result suggested that MtgA might be required for the location of Pol I or vice versa. If MtgA is involved in locating Pol I within the nucleus, then deletion of MtgA could influence the location of Pol I. Of interest, Pol I localizes away from the NE at a higher frequency in MtgA-deleted cells than in wild-type cells (Figure 6, D and E). Collectively the data support a model in which wild-type nuclei have their NOR in regions of the NE with higher concentrations of MtgA (Figure 6F). This in turn influences the spatial arrangement of fibrillarin and Bop1. In mtgA-deleted cells, the NOR would dissociate from the NE, thereby shifting fibrillarin distribution more to the nuclear interior and become more evenly surrounded by Bop1.

Increased MtgA expression alters nuclear shape and size and modifies compartmentalization of the nucleolus

MtgA deletion results in smaller, rounder nuclei, suggesting that it might play a role in controlling nuclear size. If this were true, excess MtgA would potentially result in larger nuclei. To test this, we expressed extra GFP-MtgA under the transcriptional control of the inducible alcA promoter (Waring et al., 1989; Felenbok, 1991). When grown in glycerol (nonrepressive and noninducive for alcA), GFP-MtgA levels increase over time, with the induced GFP-MtgA initially distributing quite evenly around the NE (Figure 7A). As cells accumulate higher levels of GFP-MtgA, it starts to localize around the NE unevenly, resulting in a more intense accumulation around a specific nuclear compartment (Figure 7B). When the alcA-driven expressed GFP-MtgA displays this localized accumulation, its overall nuclear signal intensity is on average fivefold greater than that of GFP-MtgA expressed from its native locus (Figure 7C). In addition, whereas the null had an 18% reduction in nuclear volume, the overexpression caused a 42% increase in nuclear volume.

Costaining cells with the green fluorescent membrane dye DiOC6(3) (Koning et al., 1993; Saunders et al., 2010) showed that MtgA accumulations correspond to membranous structures (Figure 7D). We found the compartment around which extra MtgA accumulates to be within the nucleus when we observed single confocal Z-sections (Figure 7E). Based on its general size and location to one side of the nucleus, the subnuclear region to which excess MtgA accumulates looked like it could be the nucleolus. This was confirmed by monitoring extra MtgA in a strain expressing fibrillarin-CR, showing that the nucleolus was within the nuclear compartment around which MtgA accumulated in 112 of 137 (~82%) nuclei observed (Figure 7F). The remaining 18% of nuclei had a small portion of either the nucleolus or nucleoplasm within the intranuclear compartment (unpublished data). Of note, the intranuclear MtgA-surrounded compartment was largely devoid of Nup49-CR signal (Figure 7G), indicating that NPCs are excluded from the MtgA membrane within nuclei.

To determine how the nucleolus was surrounded by MtgA, we used live-cell imaging during the transition from general NE accumulation of excess MtgA to its preferential accumulation around the nucleolus. A consistent pattern was observed, starting with the NE restricting in around the nucleolus almost to the point of the nucleolus being budded off from the rest of the nucleus (Figure 8A, 12-min arrowheads). In the second phase, additional MtgA accumulated in the NE region at the restriction point between the nucleolus and the rest of the nucleus (Figure 8A, 12-min arrowheads). In the third phase, additional MtgA accumulated in the NE region at the restriction point between the nucleolus and the rest of the nucleus (Figure 8A, 12-min arrowheads). In the final stage, extra MtgA was seen to expand from the restriction point and move around to engulf the entire nucleolus, often displaying a cup-like intermediary configuration (Figure 8A, times 20–36 min). The process led to extra MtgA surrounding the nucleolus at a higher level than in the rest of the NE (Figures 7, B, D, and E, 8, A and B, 9A, and 10, A and B). The process of NE expansion upon MtgA overexpression is microtubule independent—nuclei treated with the microtubule poison benomyl were able to form enlarged nuclei and accumulated MtgA in the NE (Supplemental Figure S3).
Wild-type interphase nuclei of A. nidulans contain a single nucleolus with a single focus of fibrillarin. In contrast, we observed two or more fibrillarin nucleolar signals in 27 of 114 (∼24%) nuclei with excess MtgA (Figure 8B). We found no correlation between multiple nucleolar signals and preferential accumulation of MtgA around the nucleolus, as multiple nucleolar signals were also observed in nuclei with an even MtgA distribution around the NE (unpublished data). During this process, the parental nucleolar signal for fibrillarin was reduced (Figure 8C), indicating that fragmentation of the parent nucleolus might generate the additional nucleolar signals.

The observed effects on fibrillarin distribution could be the result of altered distribution of the NOR. As described previously, GFP-MtgA concentrates more at the NE region to which Pol I–CR associates, and lack of MtgA alters Pol I location within the nucleus. Consistent with extra MtgA also being able to influence the location of Pol I, additional MtgA was found to dramatically influence the distribution of Pol I (Figure 9A). Instead of Pol I–CR locating to a distinct perinuclear focus as seen in WT cells, in nucleoli with excess GFP-MtgA enriched around them, Pol I is located in curved structures and often as a ring around the periphery of the nucleolus (Figure 9, A and B, at 6 min). In addition to the more circular configurations, in some nuclei, Pol I–CR appears in a more linear manner (Figure 9A, top). The circular and linear Pol I configurations are unique to nuclei accumulating excess MtgA and are absent from WT nuclei. Abnormal Pol I configurations were observed in 62.5% of 128 MtgA-overexpressing nuclei, compared with 0% of 92 wild-type nuclei. The findings suggest that there is an affinity between MtgA and Pol I or its associated proteins and/or nucleic acids. Live-cell imaging provided further evidence for this, as with time, we observed an association between excess accumulation of MtgA and the locations of Pol I–CR as its location within the nucleolus changed over time (Figure 9B; compare Pol I–CR localization to regions of high MtgA demarcated by arrows in the pseudocolor panel at time points 0, 15, and 21 min).

Collectively our findings from MtgA overexpression and deletion experiments suggest an affinity between MtgA and a component(s) of the nucleolus that can affect nucleolar architecture when levels of MtgA are altered.

**Excess MtgA results in redistribution of an integral membrane ER protein into nuclei**

Nuclear invaginations of the NE can form from the INM alone or from the entire NE containing both INM and ONM proteins and NPCs. We therefore monitored the effects of excess GFP-MtgA on the levels and location of Erg24, a transmembrane protein of the ER containing eight TMDs. Typically, NE and ER marker proteins colocalize around the NE, as the ER is contiguous with the ONM, as can be seen for MtgA and the ER marker Erg24 (Figure 10A, WT, arrows). Neither locates to the nuclear interior, and only Erg24-CR is present in the cytoplasmic ER (Figure 10A, WT, asterisk). However, when the expression of MtgA is increased, the locations of Erg24-CR are markedly modified such that some of it colocalizes with GFP-MtgA within nuclei around the nucleolus (Figure 10A, alcA-MtgA, arrowhead. Note that the exposure for GFP-MtgA was 200 and 80 ms for the alcA-GFP-MtgA cell). Live-cell imaging during GFP-MtgA accumulation around the nucleolus revealed that some Erg24-CR located with MtgA as it first locates to the restriction point.
Elevated MtgA results in asymmetric two-step mitotic NE division

Considering the dramatic changes in nuclear architecture caused by extra MtgA, we wanted to understand what effects, if any, excess MtgA might have on mitosis. Monitoring DNA segregation using

and subsequently spreads around the nucleolus (Figure 10B, 105-min arrowheads). During relocation of Erg24 into nuclei, we noticed that its cytoplasmic levels decrease compared with WT cells (Figure 10C). Quantitation of the cytoplasmic Erg24-CR signal intensity over the course of MtgA up-regulation across a population of cells showed a statistically significant decrease in cytoplasmic Erg24 levels (Figure 10D; \( n = 20 \)). Western blot analysis revealed no difference in overall Erg24-CR levels between WT and MtgA-overexpressing cells (Figure 10E). This analysis indicates that nuclear invaginations generated via excess MtgA cause relocation of a transmembrane protein from the ER into the nuclear interior.

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histone H1 revealed no major defects (Supplemental Figure S4), consistent with the lack of obvious growth defects associated with excess MtgA (unpublished data). We did, however, observe defects in the behavior of the NE during mitosis. As described earlier (Figure 3A), after anaphase, the NE restricts simultaneously on either side of the nucleolus, transiently forming three structures—the centrally located nucleolus and two daughter nuclei (Figure 11A, 8 min). However, in nuclei with extra GFP-MtgA accumulated around their nucleolus, the NE first restricts only around one of the daughter nuclei (Figure 11B, arrowhead) during mitosis. This daughter nucleus was the one located within the NE region having lower levels of GFP-MtgA (Figure 11B). The other daughter nucleus was generated several minutes after the first by a second NE restriction that occurred between itself and the nucleolus (Figure 11B, 12-min arrowhead; see also Figure 11D, 15-min arrowhead). The two sequential NE restrictions seen in nuclei with excess GFP-MtgA suggest that the first generated nucleus might enter G1 before the second. By monitoring the nuclear reimport of NLS-DsRed, a hallmark of entry into G1 (Suelmann et al., 1997; De Souza et al., 2004), we observe similar levels of NLS-DsRed imported into newly formed daughter nuclei in wild-type cells (Figure 11C). In contrast, in MtgA-overexpressing cells, earlier and higher levels of NLS-DsRed import are observed in the first generated daughter nucleus (Figure 11D). Of interest, this discrepancy in nuclear levels of G1 NLS-DsRed was not seen for nucleolar proteins (Figure 11B). This is most likely because an additional limiting factor in G1 nuclear import of nucleolar proteins occurs when they get released into the cytoplasm. Because this occurs after the second NE restriction, both daughter nuclei have similar potential for reimporting the released cytoplasmic nucleolar proteins. Therefore, although the sequence and timing of some mitotic events are different in nuclei in which MtgA has accumulated, mitosis is successfully completed.

DISCUSSION

MtgA is the A. nidulans orthologue of Bqt4 and facilitates meiosis independently of mitotic telomere NE tethering

We identified the A. nidulans orthologue of fission yeast Bqt4, an INM protein that in fission yeast is involved in tethering telomeres to the NE during vegetative growth and also meiotic recombination (Chikashige et al., 2009). The Aspergillus Bqt4 orthologue was identified via affinity purification of the RNA export NPC protein Gle1 and found to play an essential role in tethering Gle1 to the mitotic NE, as discussed later. We named this previously unstudied protein in A. nidulans MtgA for “mitotic tether for Gle1.” Like Bqt4, MtgA contains an APSES-type DNA-binding domain and a C-terminal TM domain. Both Bqt4 and MtgA require an NLS upstream of the TM domain for exclusive targeting to the NE. We therefore suggest that MtgA is inserted into the INM like Bqt4 (Chikashige et al., 2009; Chikashige et al., 2010). Bqt4/MtgA proteins are conserved among both the Ascomycota and Basidiomycota, although S. cerevisiae appears not to have an orthologue.

MtgA, like Bqt4, is required for normal levels of meiotic recombination, but it is not required for mitotic telomere tethering to the

FIGURE 11: Mitotic dynamics in MtgA-overexpressing cells. (A) Time-lapse imaging of GFP-MtgA expressed from its native promoter, showing simultaneous double restriction of the NE on either side of the nucleolus marked by Fib-CR (Strain MC271) during anaphase (time point 8 min). The nucleolus undergoes disassembly in the cytoplasm (10 min; arrow in Fib-inverted points to disassembling nucleolus) and subsequent import into the newly formed nuclei (10, 15 min). (B) Time-lapse imaging of alcA-GFP-MtgA in conjunction with Fib-CR (strain MC232), showing that the NE undergoes a two-step restriction of the NE (first at time point 6 min and then at 12 min, indicated by arrowheads) before the nucleolus is released into the cytoplasm for disassembly and subsequent nuclear import (14–26 min). (C) GFP-MtgA followed over time in conjunction with NLS-dsRed (strain MC313). Mitotic entry is marked by dispersal of NLS-dsRed. At the point of double restriction of the NE, NLS-dsRed is absent from the nucleus (time point 6 min) and is subsequently imported equally into the new G1 nuclei (9 min). (D) In strain MC248 expressing alcA-GFP-MtgA, NLS-dsRed is dispersed upon entry into mitosis as normal. Two daughter nuclei are then generated sequentially, as indicated by the arrowheads (11 and then 13 min). Because the daughter nucleus to the right is generated first, it accumulates more NLS-dsRed. Scale bar, 5 μM.
NE. Mitotic telomere tethering to the NE in fission yeast is required for subsequent telomere clustering to form the bouquet arrangement of chromosomes during meiotic prophase. Our data suggest, however, that the role of MtgA in meiotic recombination does not involve telomere NE tethering before meiosis. Instead, the observed meiotic defects in MtgA-deleted cells might involve the new functions we defined for this class of protein, including mitotic-specific NE tethering of Gle1 and modulation of nuclear and nucleolar architecture.

**MtgA is required for mitotic-specific tethering of Gle1 to the NE**

Gle1 is an mRNA export factor expected to disperse from NPCs during the partially open mitosis of *A. nidulans*; however, it remains at the NE during mitosis (Osmani et al., 2006a; Ukil et al., 2009). This paradox has been solved: we demonstrate that Gle1 associates with NPCs during interphase but during mitosis, transitions to locate at the INM via MtgA. During mitosis, Gle1 remains tethered to the NE via MtgA and returns to NPCs after mitosis in G1. Thus, although Gle1 remains at the NE throughout the cell cycle, it does so at two distinct locations. To our knowledge, this is the first reported instance of a protein being tethered at two different regions of the NE in a mitotically regulated manner.

The dispersal of Gle1 from the mitotic NE caused by mtgA deletion does not result in obvious mitotic phenotypes, although we have not tested the fidelity of mitotic events in the absence of mtgA. This suggests that Gle1 at the mitotic NE has nonessential functions that are perhaps also not readily visualized. As an example, it could be required to help remove RNA molecules remaining in the nucleus and/or nucleolus during mitosis. However, this work does provide novel insights into the mitotic NE tethering of Gle1, and it will be interesting if similar mitotic-specific locations occur in other species and if this might play roles in the disease states associated with Gle1 mutations (Nousiainen et al., 2008; Glass et al., 2016).

**MtgA levels dramatically modify nuclear architecture**

In addition to its roles in meiosis and mitotic tethering of Gle1 to the NE, MtgA was also found to affect dramatically nuclear architecture when either deleted or in excess. The most obvious defect caused by lack of mtgA was the formation of slightly smaller, rounder nuclei. This suggests that Gle1 at the mitotic NE has nonessential functions that are perhaps also not readily visualized. As an example, it could be required to help remove RNA molecules remaining in the nucleus and/or nucleolus during mitosis. However, this work does provide novel insights into the mitotic NE tethering of Gle1, and it will be interesting if similar mitotic-specific locations occur in other species and if this might play roles in the disease states associated with Gle1 mutations (Nousiainen et al., 2008; Glass et al., 2016).

Truncation gene replacement experiments indicate that integration of MtgA into the INM is involved in its ability to maintain the size and shape of nuclei.

In addition to causing rounder and smaller nuclei, deletion of MtgA also alters the internal organization of the nucleolus. The nucleolus is assembled around NORs and contains a distinctive internal architecture to which specific proteins locate. For example, fibrillarin occupies a centrally located position within the nucleolus and is surrounded unevenly by Bop1. However, in the absence of MtgA, the spatial relationship between fibrillarin and Bop1 is changed such that fibrillarin is more evenly surrounded by Bop1 (Figure 6A). This shows that MtgA is required to maintain the distinctive distribution of proteins within the nucleolus. One potential mechanism is modification of the location of the NOR, an idea further supported by the effects caused by excess MtgA.

Because lack of MtgA generates smaller nuclei and modifies the architecture of the nucleolus, we anticipated that extra MtgA might enlarge nuclei and perhaps change the structure of the nucleolus. These expectations were fulfilled. Nuclei became enlarged as excess MtgA initially accumulated evenly in their NE. However, these nuclei apparently have a limit on the size to which they can enlarge. This is suggested by the fact that, as nuclei accumulated further MtgA, they stopped enlarging and started to accumulate the extra MtgA specifically around their nucleoli. During this process, the nucleolus became largely surrounded by MtgA and its associated membrane (Figure 7). Although several studies showed that induced expression of proteins (Wright et al., 1988; Ralle et al., 2004; Malhas et al., 2011; Masuda et al., 2016), including those of the INM (Ma et al., 2007; Friederichs et al., 2011; Goto et al., 2014), can cause elaboration of intranuclear membranes, none that we know of revealed a specific expansion of intranuclear membrane around the nucleolus.

Several questions arise from the finding that excess MtgA accumulates specifically around the nucleolus. For example, what is the nature of the intranuclear membrane associated with extra MtgA, and how might it form? The intranuclear MtgA membrane is sustained by the lipophilic membrane dye DiOC6(3) and contains a transmembrane ER marker, suggesting that it represents mislocalized NE. This membrane could be generated, at least in part, through redistribution of membrane from the ER, as the cytoplasmic levels of an ER membrane marker decrease during the process. However, the intranuclear MtgA membrane does not represent a complete NE because it lacks NPCs.

There are two possibilities for how the excess MtgA-induced intranuclear membrane might be formed. It could represent an expansion of just the INM to which ONM proteins are now able to locate. Alternatively, the MtgA-promoted intranuclear membrane could represent both the INM and ONMs from which NPCs are excluded. Of further interest, and most notably, accumulation of MtgA induces the NE to invaginate into the nucleoplasm specifically at the interface between the nucleoplasm and nucleolus. This novel yet specific accumulation suggests that MtgA might have a weak affinity for a nucleolar component. Then, as MtgA levels increase within the NE, the net weak nucleolar affinity might accumulate to a level that promotes the NE being pulled in and expanding around the nucleolar surface but not around the bulk DNA. The contact between the INM and nucleolus mediated by excess MtgA potentially could also generate a mechanical barrier to both the movement of NPCs into this NE and the de novo generation of nuclear pores at this NE. This would explain how MtgA accumulation promotes NE invagination around the nucleolus containing ONM and INM proteins but not NPCs.

Further experimentation will be needed to test these working hypotheses, but some findings lend support to the idea that MtgA has weak affinity for the nucleolus. For example, in wild-type cells, MtgA partially concentrates to the NE directly adjacent to the nucleolus, where Pol I also locates (Figure 6C). In addition, without MtgA, the architecture of the nucleolus is modified. Finally, the normal location of Pol I is compromised in both MtgA-lacking and MtgA-overexpressing cells. Our findings therefore collectively indicate that MtgA is required to maintain the normal position of the NOR and nucleolar architecture.

Although we are unaware of other instances of induced intranuclear membranes locating specifically around the nucleolus, studies using *S. cerevisiae* identified the NE adjacent to the nucleolus as a site for induced NE membrane accumulation. In these studies, under conditions that promote extra NE formation (including mitotic delay and induced imbalance in lipid synthesis), the NE expands out of the nucleolus, forming “flares” that extend into the cytoplasm. The expandable NE adjacent to the nucleolus is believed to act as a “membrane sink” allowing the nucleus to accommodate increased NE levels without dramatically increasing nuclear volume (Campbell et al., 2006; Witkin et al., 2012; Meseroll and Cohen-Fix, 2016). We propose that *A. nidulans* deals with the excess NE problem caused
by extra MtgA by incorporating it within nuclei around the nucleolus rather than out into the cytoplasm, such that the extra NE does not uncontrollably affect overall nuclear size, and so it helps to maintain an acceptable nuclear/cell volume ratio (Huber and Gerace, 2007; Meseroll and Cohen-Fix, 2016). By exclusion of NPCPs from this extra membrane, normal levels and direction of nuclear transport presumably can also be maintained.

It is surprising that turning the nucleolus into a largely membrane-bound organelle, as occurs upon MtgA accumulation, does not cause clear growth defects. In part, this is probably because the excess MtgA nuclear membrane does not prevent completion of successful mitosis. In spite of this, the mitotic process itself is modified, as daughter nuclei are generated sequentially rather than in unison. This is accompanied by a delay in the normal process of mitotic nucleolar disassembly into the cytoplasm. Because nucleolar disassembly still occurs after daughter nuclei have formed in G1 and reestablished nuclear transport, both subsequently get a relatively equal portion of nucleolar proteins, ensuring faithful nucleolar protein segregation. If indeed the nucleolus is surrounded by membrane, it would be expected that transport of nucleolar products out into the cytoplasm would be hindered. However, because the nucleolus is disassembled and its contents released into the cytoplasm during mitosis, this gives the nucleolus the opportunity to release its products, even if it is membrane bound, once per cell cycle.

The large diversity of nuclear morphologies exhibited across both healthy and diseased cells demonstrates that there are important regulated mechanisms used by organisms to maintain nuclear size and architecture (Zink et al., 2004; Webster et al., 2009; Malhas et al., 2011; Edens et al., 2013; Vukovic et al., 2016). Collectively our work furthers our understanding of how organisms can maintain nuclear architecture and cope with excess nuclear membrane. This study uncovered a novel INM protein–dependent mechanism that impeded further growth (unpublished data), and protoplasts were collected as per standard techniques (Osmani et al., 2006b).

Alternatively, pyrG" strains were protoplasted from mycelia as described previously (Szewczyk et al., 2006) with minor modifications. The GFP-MtgA tagged strain was generated as described using three-way fusion PCR (Szewczyk et al., 2006) and primers listed in Supplemental Table S2. Upstream and downstream mtgA sequences were generated using primers AO390/AO388 and AO391/AO359, respectively. The gfp sequence was amplified from plasmid pFN03 (Yang et al., 2004) using primers HP108 and KF043. The construct derived from fusion of gfp and upstream and downstream mtgA sequences was generated with primers AO358/AO389, transformed into SO451, and PCR confirmed using primers AO359/AO390. To generate aca-GFP-MtgA strains, we used the pAL5 plasmid vector system (Doonan et al., 1991), which allows for in-frame insertion of the gene of interest under the control of the inducible aca promoter (Waring et al., 1989; Felenbok, 1991). pAL5 was modified by cloning the fragment KpnI-NsiI-gfp-BglII-Nhel into the KpnI and Nhel sites of the pAL5 backbone that are situated downstream of the aca promoter such that GFP is under the control of the aca promoter. This new plasmid, pAL5-S'5'GFP, was generated by Colin De Souza and kindly gifted for use in this study. The gene of interest can be inserted in-frame into a unique polylinker after the initial in yeast extract/glucose medium at 32°C with shaking until emergence of germ tubes was observed (shmoo cells) and their cell wall digested as follows. The cells were centrifuged and washed with 0.55 M KCl and 50 mM citric acid, pH 6.5, and transferred to a solution consisting of 10 mg/ml VinoFlow FCE (Novozymes, Bagsvaerd, Denmark) or 15 mg/ml Vinotaste (Novozymes) in 0.55 M KCl and 50 mM citric acid. Cells were protoplasted under conditions that impeded further growth (unpublished data), and protoplasts were collected as per standard techniques (Osmani et al., 2006b).

MATERIALS AND METHODS

General techniques

The genotypes of A. nidulans strains used in this work are listed in Supplemental Table S1 and were generated using standard techniques (Pontecorvo et al., 1953; Yang et al., 2004; Nayak et al., 2006). mtgA was deleted using the pyrG" marker. The deletion construct was generated using fusion PCR followed by transformation of a ΔnuAΔulo/Δulo strain as described previously (Yang et al., 2004; Nayak et al., 2006; Szewczyk et al., 2006) using primers listed in Supplemental Table S2. Deletion of nup42 and nup132 was described previously (Osmani et al., 2006a). To generate Rap1-GFP-S-tag, three-way fusion PCR was performed. The GFP-S-tag:pyrG" cassette was first amplified from plasmid pCD565 (De Souza et al., 2014) using C-terminal tagging primers HP116 and FN01-pyrG (Yang et al., 2004; Liu et al., 2010). The upstream Rap1 targeting sequence was amplified from wild-type DNA using MaC59 and MaC38, whereas the downstream Rap1 sequence, including the stop codon and 3′ untranslated region, was amplified using primers MaC36 and MaC39. MaC38 has an overhang that complements the sequence to primer HP116, whereas MaC39 has an overhang complementing primer FN01-pyrG to enable PCR-directed fusion of the three products. The fusion construct was obtained by amplifying the three PCR products with primers MaC59 and MaC36. The fusion construct was used for strain SO451 transformation. Site-specific integration was confirmed via diagnostic PCR with primers MC34 and MC37. Because MtgA has a transmembrane domain at its C-termi nus, fluorescent protein tagging of MtgA was done at the N-termi nus to avoid possible disruption of localization and folding of MtgA and/or the fluorescent protein arising from the membrane topology of MtgA. The exception to this is MtgAΔTM-GFP-S-tag (see later discussion), which lacked its transmembrane domain and was tagged C-terminally. N-terminal fluorescent protein fusion constructs lacked selection markers and were used for transforming ΔmtgA strains marked by the wild-type pyrG allele from Aspergillus fumigatus (pyrG"), which complements the mutant pyrG" allele in the genetic background. After transformation of ΔmtgA strains, counterselection was done using 1 mg/ml 5-fluoroorotic acid (5-FOA; toxic to pyrG" strains; US Biological, Salem, MA) added to yeast extract/glucose agar medium containing uridine and uracil to select for transformants that incorporated the N-terminal fusion constructs and evicted pyrG". Primers AO359 and AO390 were used to check for site-specific integration. For generation of protoplasts from pyrG" strains to enable transformation, cells were grown initially in yeast extract/glucose medium at 32°C with shaking until emergence of germ tubes was observed (shmoo cells) and their cell wall digested as follows. The cells were centrifuged and washed with 0.55 M KCl and 50 mM citric acid, pH 6.5, and transferred to a solution consisting of 10 mg/ml VinoFlow FCE (Novozymes, Bagsvaerd, Denmark) or 15 mg/ml Vinotaste (Novozymes) in 0.55 M KCl and 50 mM citric acid. Cells were protoplasted under conditions that impeded further growth (unpublished data), and protoplasts were collected as per standard techniques (Osmani et al., 2006b).
with the NheI plus KpnI–digested mCherry sequence to obtain pHMC2. Plasmids pHMC1 and pHMC2 were then used for transformation of SO451 and selection for complementation of pyrG89.

Affinity purification and liquid chromatography–tandem mass spectrometry
Gle1–S-tag (strain SO901) and MtgAΔTMD-GFP–S-tag (strain MC107) affinity purifications followed by mass spectrometry were done as previously described (Liu et al., 2010). Full-length MtgA affinity purifications were unsuccessful (unpublished data), and hence a strain expressing a version of MtgA lacking its TMD and C-terminally fused to GFP–S-tag was used for affinity purifications. The MtgAΔTMD-GFP–S-tag construct was generated as described earlier for Rap1-GFP–S-tag. The upstream MtgA sequence was PCR amplified using primers MaC01/MaC24, and the downstream MtgA sequence was amplified using MaC23/MaC04. Primers MaC23 and MaC24 have overhangs priming FN01-pyrG and HP116, respectively. The fusion construct was obtained by amplifying the three PCR products with primers MaC1 and MaC4. The fusion construct was used for SO451 transformation. Site-specific integration was confirmed with primers AO359 and AO390. Expression of S-tag was confirmed by Western blotting using anti–S-tag antibody (Immunology Consultants Laboratory, Portland, OR).

Live-cell imaging
Live-cell imaging and induction of genes regulated by the alcA promoter was done as described previously (Govindaraghavan et al., 2014) and additionally using a 100×/1.4 numerical aperture total internal reflection fluorescence objective lens on an Eclipse TE 2000-U (Nikon, Tokyo, Japan) microscope equipped with an UltraView ERS spinning-disc confocal system. Benomyl at 2.4 μg/ml was used to depolymerize microtubules. Cell membranes were visualized by adding 1 μl of 10 mM DIOC6(3) (Sigma-Aldrich, St. Louis, MO) to hyphae grown overnight in 3 ml of medium containing glycerol as a carbon source immediately followed by microscopy.

Bioinformatics
Orthologues of A. nidulans Kila-N proteins present in other fungi were identified by BLAST search (blast.ncbi.nlm.nih.gov/Blast.cgi) against species in different fungal families in either the Ascomycota or Basidiomycota. The phylogenetic tree was constructed using MEGA version 7 (Kumar et al., 2016) by aligning sequences using ClustalW with a BLOSUM62 matrix and then generating a maximum-likelihood tree of the alignment.

MtgA domain analysis
mCherry-MtgA and mCherry-MtgA truncations were generated by N-terminal tagging as described. mCherry-MtgA was generated by amplifying mCherry from plasmid pHLB5 (Liu et al., 2009) using primers MaC07 and MaC08. This product was fused to flanking products amplified from a wild-type strain using primers MaC01/MaC24, and the downstream MtgA sequence was amplified using MaC23/MaC04. Primers MaC23 and MaC24 have overhangs priming FN01-pyrG and HP116, respectively. The fusion construct was obtained by amplifying the three PCR products with primers MaC1 and MaC4. The fusion construct was used for SO451 transformation. Site-specific integration was confirmed with primers AO359 and AO390. Expression of S-tag was confirmed by Western blotting using anti–S-tag antibody (Immunology Consultants Laboratory, Portland, OR).

Western blotting
Western analysis was performed as described previously (Osmani et al., 2006a) with the modification that strains were grown in minimal medium using either 55 mM glucose or 1% (vol/vol) ethanol as a carbon source.

Data analyses
Image analyses were carried out using ImageJ software (National Institutes of Health, Bethesda, MD) or Volocity Image Analysis Software (PerkinElmer, Waltham, MA). For Figure 2B, Rap1-GFP nuclear localization was determined for the central focal plane of each nucleus as determined by Nup49-CR distribution. For Figure 5B, nuclear circularity values were calculated during G2 (defined as 6–10 min before onset of nuclear division) for all mutants tested. Three-dimensional renderings in Figure 6B were generated using isosurface renderings with identical settings for each condition using Volocity Image Analysis Software. For Figure 7C, nuclear GFP intensities were calculated after background subtractions for each condition. Erg24-CR cytoplasmic signal intensities in Figure 10 were calculated after background subtraction. Fluorescence intensities and volumes were calculated by identifying objects for desired channels using Volocity Image Analysis Software. Pixel intensity profiles were obtained by drawing indicated regions of interest in ImageJ. Plotting of graphs and statistics was done using Microsoft Excel. Unpaired, two-tailed t tests were performed to calculate statistical significance.

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