

Asynchronous nuclear division cycles in multinucleated cells

Amy S. Gladfelter, A. Katrin Hungerbuehler, and Peter Philippsen

Department of Molecular Microbiology, Biozentrum University of Basel, 4056 Basel, Switzerland

Synchronous mitosis is common in multinucleated cells. We analyzed a unique asynchronous nuclear division cycle in a multinucleated filamentous fungus, *Ashbya gossypii*. Nuclear pedigree analysis and observation of GFP-labeled spindle pole bodies demonstrated that neighboring nuclei in *A. gossypii* cells are in different cell cycle stages despite close physical proximity. Neighboring nuclei did not differ significantly in their patterns of cyclin protein localization such that both G1 and mitotic cyclins were present regardless of cell cycle stage, suggesting that

the complete destruction of cyclins is not occurring in this system. Indeed, the expression of mitotic cyclin lacking NH₂-terminal destruction box sequences did not block cell cycle progression. Cells lacking AgSic1p, a predicted cyclin-dependent kinase (CDK) inhibitor, however, showed aberrant multipolar spindles and fragmented nuclei that are indicative of flawed mitoses. We hypothesize that the continuous cytoplasm in these cells promoted the evolution of a nuclear division cycle in which CDK inhibitors primarily control CDK activity rather than oscillating mitotic cyclin proteins.

Introduction

Multinucleated cells are found in a variety of organisms and are integral to processes as diverse as the early development of the fruit fly, bone remodeling, placenta formation, and cancer metastasis. However, information on how multinucleated cells regulate nuclear division is limited to relatively few systems. Fusion experiments with mammalian cells demonstrated that multiple nuclei in a shared cytoplasm synchronize their nuclear division cycles (Rao and Johnson, 1970). These experiments suggested that separate nuclei within a common cytoplasmic environment will alter their normal cell cycle kinetics and align temporally with the other nuclei present. Synchronous mitosis has also been observed in binucleate yeast cells (Hoepfner et al., 2002). Furthermore, in naturally multinucleated cells, mitoses often occur either synchronously, as in the slime mold *Physarum polycephalum* (Nygaard et al., 1960), or parasynchronously, with a linear wave of nuclear division spreading across a cell, as in the filamentous fungus *Aspergillus nidulans* (Clutterbuck, 1970).

One hypothesis for the molecular basis for synchronous mitoses in multinucleated cells is that cell cycle regulators shuttle between nuclear and cytoplasmic compartments, which would allow them to diffuse and coordinate multiple nuclei in

the same cytoplasm. Thus, the dynamic localization of the cell cycle machinery may promote continual communication between individual nuclei and the cytoplasm and could explain synchronous nuclear division cycles, although in no instance has this hypothesis been tested in multinucleated cells.

In some cases, nuclei appear to behave independently of one another despite sharing a common cytoplasm. In newly fertilized binucleate sea urchin embryos, the two pronuclei have been shown to initiate nuclear envelope breakdown and mitosis at different times if one nucleus is paused as a result of DNA damage (Hinchcliffe et al., 1999). Signaling for cell cycle progression was also nuclear limited in experiments involving binucleate yeast cells in which a single nucleus received DNA damage (Demeter et al., 2000). Similarly, in binucleated cultured marsupial cells, anaphase entry is asynchronous if chromosome attachment is delayed in one spindle (Rieder et al., 1997). Thus, at least for the case of responding to DNA damage or unattached chromosomes, multinucleated cells can undergo asynchronous mitosis. Furthermore, the filamentous fungi *Neurospora crassa* and *Podospira anserina* are also thought to have asynchronous mitoses, but in neither case has the phenomenon been studied in depth (Minke et al., 1999; unpublished data). Thus, it seems to be possible for the nucleus itself to control the decision to enter mitosis, although the molecular basis for such nuclear autonomy in any system remains unknown.

In this study, we present evidence for and initial molecular analysis of asynchronous nuclear division cycles in the

Correspondence to Amy S. Gladfelter: amy.gladfelter@unibas.ch

A.S. Gladfelter's present address is Department of Biology, Dartmouth College, Hanover, NH 03755.

Abbreviations used in this paper: CDK, cyclin-dependent kinase; NES, nuclear export sequence; SPB, spindle pole body.

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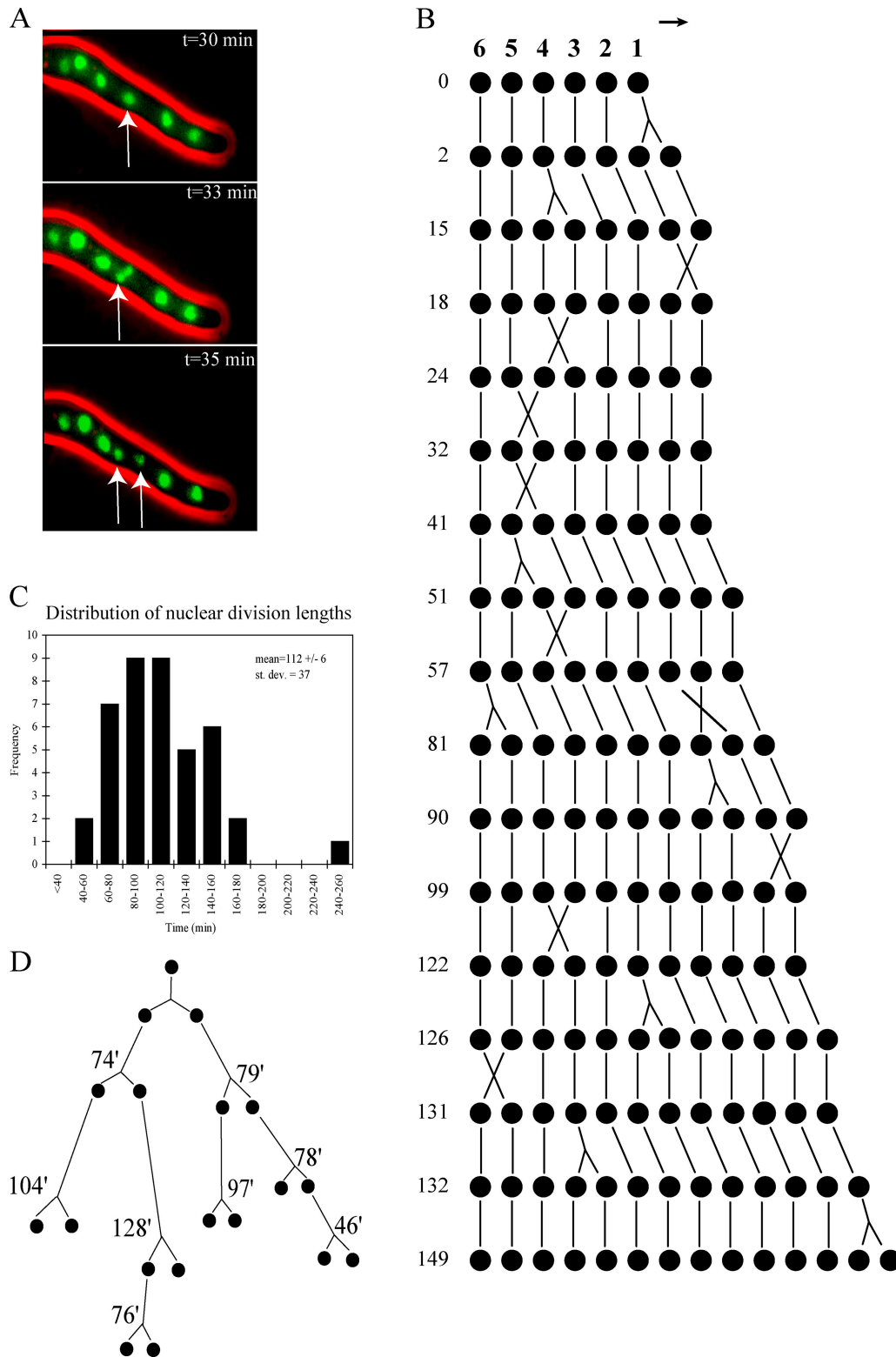


Figure 1. **Mitoses are asynchronous in multinucleated *A. gossypii* cells.** (A) Individual frames from a time-lapse video recording the growth of a strain (*HHF1-GFP*) with histone-GFP-labeled nuclei. The nuclei indicated by the arrows divide independently from their neighbors. Pictures were acquired every 30 s. (B) Example of a nuclear pedigree demonstrating the capacity for all nuclei to divide. Six starting parent nuclei were followed through subsequent nuclear divisions. Numbers are time in minutes from start of time-lapse acquisition. A total of 11 such pedigrees were performed on independent hyphae. In the pedigree diagram, X is a by-passing event, an upside-down Y is a mitosis, and the arrow denotes the direction of growth. (C) Histogram summarizing variability in nuclear division cycle length from 41 mitoses observed in time-lapse videos of *HHF1-GFP* strains. (D) Nuclear lineage demonstrating variability in nuclear division cycle length within related nuclei, where the time in minutes represents the time between nuclear divisions.

multinucleated filamentous fungus *Ashbya gossypii*. This fungus shares a common ancestor with *Saccharomyces cerevisiae* before the duplication of the budding yeast genome (Dietrich et al., 2004; Kellis et al., 2004). Over 90% of the *A. gossypii* genome contains syntenic homologues in budding yeast, and, in particular, all cell cycle control proteins described for budding yeast are present with homology ranging from 20 to 95% identity (Dietrich et al., 2004). Despite this conservation of cell cycle gene sets, unlike budding yeast, *A. gossypii* cells grow exclusively as hyphae containing many nuclei in linear arrays within the same cytoplasm. We predict that these hyphal cells undergo a closed mitosis like many fungi, including *S. cerevisiae*, and there is evidence that the hyphae are compartmentalized with partial septa that contain pores large enough for nuclei to pass between compartments (Alberti-Segui et al., 2001).

We have generated nuclear pedigrees with time-lapse video microscopy showing that nuclei in *A. gossypii* cells divide asynchronously in the same cytoplasm. We analyzed this apparent nuclear autonomous behavior and evaluated how periodic cell cycle proteins act in such a system in which nuclei are not in sync with each other. Our experiments showed that both G1 and mitotic cyclins are present across all cell cycle stages. We present evidence that the continuous cytoplasm in these cells permits proteins to diffuse so that nuclei continually receive proteins from the cytosol, which are expressed from neighboring nuclei. These data have led us to hypothesize that differences in cell architecture and spatial organization may have selected for alternative modes of cell cycle control during the evolution of *S. cerevisiae* and *A. gossypii*. When combined, our experiments suggest that cyclin-dependent kinase (CDK) inhibitors rather than cycles of accumulation and complete destruction of the cyclins provide the primary source of oscillation in the cell cycle machinery of *A. gossypii*.

Results

Mitosis is asynchronous in *A. gossypii* cells

Growth and nuclear dynamics were followed in the growing tip region of *A. gossypii* cells using in vivo fluorescence and time-lapse video microscopy. Nuclei were readily visualized using a GFP-tagged histone (H4) protein, *AgHhf1-GFP*. As can be observed in the accompanying video and has been reported previously, these nuclei oscillated rapidly and tumble through the cytoplasm in three dimensions, frequently exchanging places through by-passing one another (Video 1, available at <http://www.jcb.org/cgi/content/full/jcb.200507003/DC1>; Alberti-Segui et al., 2001). Multiple mitoses were easily observed in the growing tip compartment, and, remarkably, in most cases, nuclei divided asynchronously, independently of neighboring nuclei in close physical proximity (Fig. 1, A and B; and Video 1). The distance between nuclei varied between <1.0 μm , when nuclei pass each other, and 7 μm , with a mean of 3.6 ± 0.1 μm . Nuclear division occurred irrespective of the distance from the hyphal tip, and no spatial patterns of mitosis could be discerned from the time-lapse data.

One way for cells to establish asynchrony would be for some nuclei to be arrested in G_0 and act as “insulators” or

barriers between actively dividing nuclei. To investigate this possibility, we generated a series of nuclear pedigrees in which nuclei in living cells were followed through several generations of division and were also charted as to their relative positions in the cell (Fig. 1 B). In all pedigrees examined, most nuclei divided, yielding daughter nuclei that also divided ($n = 11$ independent pedigrees). The mean time between divisions of a nucleus, which represents the total nuclear division cycle length, was 112 min but varied greatly from 46 to 250 min ($n = 41$ mitoses; Fig. 1 C). Interestingly, variability in cell cycle length was observed within a single lineage starting from one nucleus. Daughter nuclei produced from the same mitosis do not necessarily divide again in the same time interval (Fig. 1 D). Frequently, daughter nuclei by-passed neighboring nuclei soon after mitosis was finished (X on pedigree in Fig. 1 B); however, asynchrony was still observed in the absence of such by-passing events. Furthermore, sudden increases in Hhf1-GFP intensity, which may correlate with S phase, generally occurred at different times in sister nuclei produced from the same mitosis. Thus, most nuclei appear to be capable of division but with variable timing, and, surprisingly, they act independently of neighboring nuclei.

Neighboring nuclei are in different cell cycle stages

The time-lapse data lead to the prediction that adjacent nuclei are in different phases of their division cycle at the same time. To determine the cell cycle stages of neighboring nuclei, we visualized tubulin (Fig. 2 A) in fixed cells and spindle pole bodies (SPBs; fungal equivalent of the centrosome; Fig. 2 B) in living cells. Overall, $62 \pm 4.8\%$ of nuclei had a single SPB (noted as 1 in figures), $18 \pm 2.6\%$ had duplicated, adjacent SPBs (noted as 2 in figures; scored based on either two distinct dots or a brighter and expanded area of fluorescent signal compared with 1), and $20 \pm 2.7\%$ showed separated SPBs and a spindle (noted as Meta for metaphase and Ana for anaphase in figures; Fig. 2, A and B; $n > 1,000$). Similar proportions were observed by evaluating microtubules in fixed cells with a tubulin antibody (Fig. 2 A) and observing SPBs with *SPC42-GFP* (Fig. 2 B). We found that only 35% of nuclei ($n > 1,000$) were in the same spindle stage as their neighbors as judged by SPB morphology, and these were nearly all nuclei with only a single SPB. Focusing specifically on nuclei with mitotic spindles and separated SPBs, $>80\%$ were beside a neighbor with only a single or just duplicated SPB, showing that even actively dividing nuclei do not seem to influence neighbors in different cell cycle stages.

Four-dimensional time-lapse microscopy of hyphae expressing *SPC42-RFP* and *Hhf4-GFP* revealed that one nucleus frequently underwent SPB duplication, whereas neighboring nuclei remained with a single SPB (Fig. S1 A, available at <http://www.jcb.org/cgi/content/full/jcb.200507003/DC1>). Nuclei spent an extended period with duplicated but not separated SPBs (from 20 min to >1 h, with a mean time of 47 ± 6 min; $n = 11$ nuclei). Thus, nuclei in *A. gossypii* appear to transit the different stages of the cell cycle regardless of the state of their neighbors, and many nuclei pause for extended time periods before or during bipolar spindle assembly.

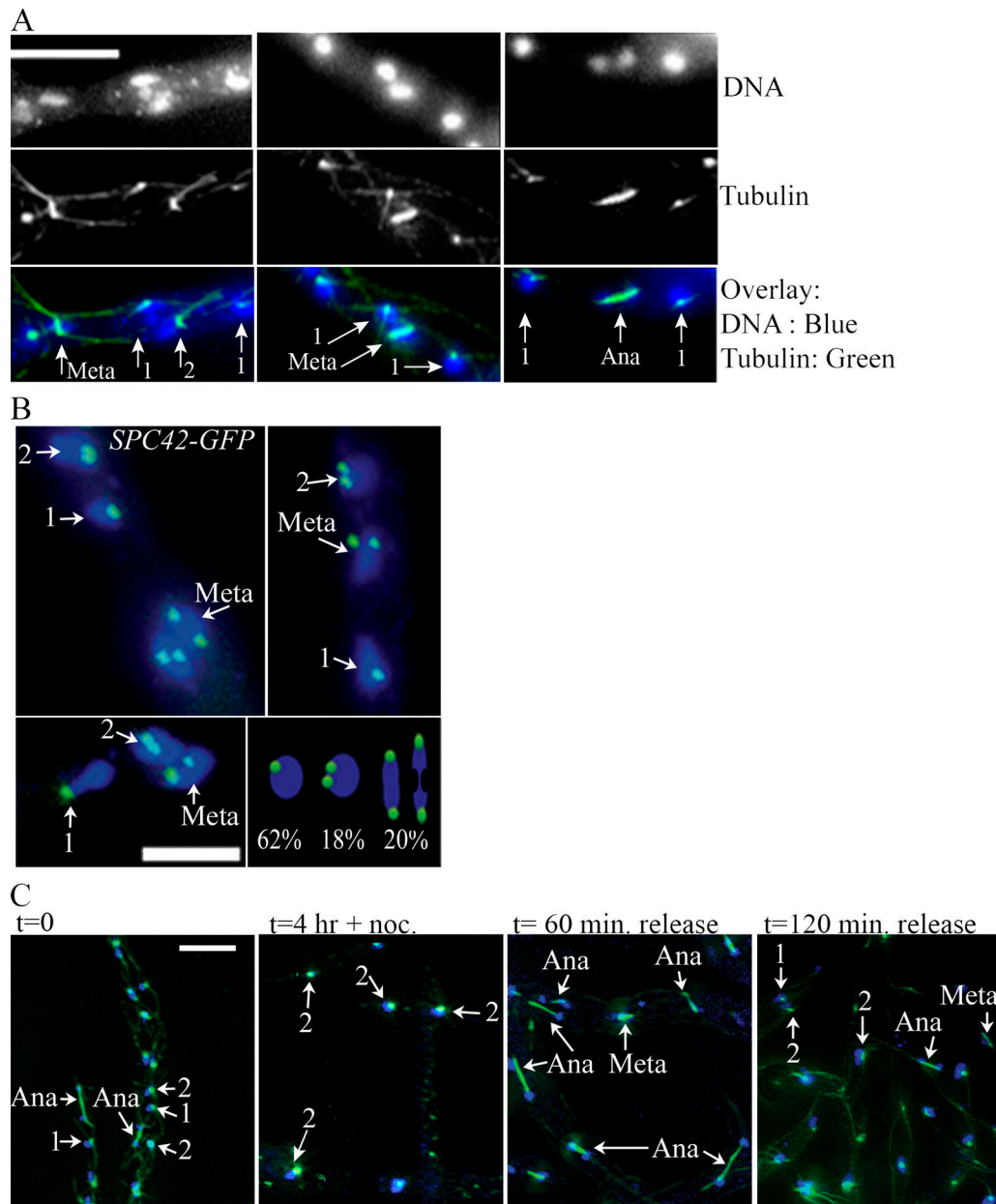


Figure 2. Neighboring nuclei are in different cell cycle stages. Spores were grown overnight at 30°C until young mycelium were formed that contained ~75–100 nuclei. In all panels, arrows highlight neighboring nuclei in different spindle stages. 1 indicates nuclei with a single SPB; 2 indicates nuclei with adjacent, duplicated SPBs or SPBs that are larger in diameter and $\geq 2\times$ brighter than single SPBs; Meta indicates likely metaphase nuclei with a spindle aligned across the middle of the DNA; Ana indicates anaphase nuclei in which the spindle connects two separated DNA signals. (A) Tubulin visualization in wild-type cells by immunofluorescence using antitubulin antibody for spindle observation. (B) *SPC42-GFP* (ASG38) visualization for SPB observation. Proportions are the percentage of nuclei with one, two adjacent, or two separated metaphase/anaphase SPBs ($n > 1,000$). (C) Wild-type spores were grown for 12 h until a bipolar germling stage and incubated with nocodazole for 4 h. Cells were then washed before release into fresh media and processed at the indicated time points for tubulin immunofluorescence. Nuclei are in blue, and microtubules or SPBs are in green. Bars (A and C), 10 μm ; (B) 5 μm .

Asynchrony returns rapidly after artificial synchronization

The asynchrony described in Figs. 1 and 2 could come about in either of two ways: each nucleus may independently modulate the lengths of the division cycle phases as suggested by the pedigrees, or all nuclei could undergo a standard cell cycle, but because the nuclei move around, each nucleus is out of phase with respect to its neighbors. To address these possibilities, we used the microtubule poison nocodazole to arrest nuclei in mitosis and

synchronize this normally asynchronous system. If nuclei have a standard cycle, synchrony should persist after release from the arrest. All cytoplasmic microtubules were depolymerized by nocodazole, producing tubulin staining limited to SPBs that were used to determine cell cycle stage. After a 4-h incubation in the drug, >70% of the nuclei appeared in synchronous arrest with adjacent, bright SPBs, whereas control cells treated with DMSO still displayed asynchrony similar to that of wild-type cells ($n = 200$ nuclei for each condition; Fig. 2 C, Fig. S1 B, and not depicted).

Cells were released from arrest into fresh media, and nuclear division cycle progression was followed. Initially after release (15-, 30-, and 45-min time points), the extensive network of cytoplasmic microtubules was repolymerized before recovery of nuclear microtubules. By 1 h after release, the cytoplasmic networks of microtubules were largely rebuilt, and the majority of nuclei was in synchrony and had extended spindles indicative of anaphase/telophase. However, by 2 h after the release from nocodazole, only 30% of the nuclei were in the same spindle stages as their immediate neighbors, much like the starting cell population, indicating that asynchrony was quickly restored (Fig. 2 C). These data suggest that each nucleus seems to have an autonomous clock that functions independently of nuclei in their vicinity.

In the process of analyzing these arrest–release experiments, it became clear that hyphal morphogenesis proceeded normally even when the nuclear cycle was blocked (Fig. S1 B). This led to a decrease in the nucleocytoplasmic ratio, and the distance between nuclei ranged from 4 to >30 μm , with a mean of $22 \pm 2 \mu\text{m}$. Within ~ 4 h after recovery (roughly two cell cycle lengths), a nearly normal density of nuclei was restored (5.0 μm between nuclei), suggesting that the nucleocytoplasmic ratio is monitored and that the cell cycle length and/or growth rate can respond to alterations in this ratio.

G1 and mitotic B-type cyclins are nuclear and present across all spindle stages

Are there clues in the *A. gossypii* genome to explain the molecular basis for nuclear asynchrony? The *A. gossypii* genome shows striking synteny patterns and similarity in gene set to *S. cerevisiae* despite their different lifestyles and growth form. Whole genome comparisons between the two organisms strongly suggest that they diverged before the duplication of the budding yeast genome, and, thus, the *A. gossypii* genome lacks many gene pairs that are retained in budding yeast after this duplication (Dietrich et al., 2004). *A. gossypii* genes are named based on their syntenic homologue in *S. cerevisiae*, and in the cases where a syntenic gene pair exists in yeast, the single homologue in *A. gossypii* is called by both names separated by a slash. We examined homologues of the core cell cycle machinery in *A. gossypii* and found that the entire gene set is present at syntenic positions, with amino acid identity ranging from 31 to 86% (Table I). Thus, *A. gossypii* and *S. cerevisiae* share similar transcription factors, cyclins, the CDK, inhibitors, activators, and degradation machinery involved in cell cycle control.

How might this conserved network of cell cycle regulators be constructed to direct asynchronous mitosis in a multinucleated cell? Cyclin proteins are integral to the cell cycle clock in eukaryotic cells and, when complexed to a CDK, drive cell cycle progression (Murray, 2004). Periodic transcription and degradation of cyclins is believed to be a central mode of oscillation that is critical for orderly cell cycle progression. Upstream sequences of *AgCLN1/2* and *AgCLB1/2* showed the conservation of transcriptional regulatory elements, including some Swi4/6 cell cycle box-binding factor and Mbp1–Swi6 cell cycle box-binding factor transcription factor–binding sites in the *AgCLN1/2* promoter and Fkh1p, Fkh2p, and Mcm1p sites in the *AgCLB1/2*

promoter (consensus sequences summarized in Kellis et al., 2003). Furthermore, protein sequences have degradation motifs, including possible PEST sequences in *AgCLN1/2p* and the destruction (D)-box and KEN-box motifs in *AgCLB1/2p* (Fig. 3; Glotzer et al., 1991; Holloway et al., 1993; Yamano et al., 1996; Pflieger and Kirschner, 2000). The *AgCLB1/2p* sequence has one additional NH₂-terminal D box, one of two nuclear export sequences (NESs) that is compared with *ScCLB2p*, and one NLS (Hood et al., 2001).

Based on the homology and similar domain composition of the *A. gossypii* cyclins compared with those of budding yeast, we would predict that individual nuclei are expressing and degrading distinct cyclins depending on their cell cycle stage. Given that protein translation occurs in a common cytoplasm, however, it is unclear how independent transcriptional programs and oscillating protein levels are achieved by neighboring nuclei that are out of phase with each other. In an effort to determine whether such periodic expression patterns could be locally established in a syncytial cell, the *A. gossypii* cyclins were localized.

The predicted G1 cyclin *AgCLN1/2* and the B-type cyclin *AgCLB1/2* were epitope tagged at their endogenous loci with 13 copies of the c-myc epitope to evaluate cyclin protein distribution in the cell. These epitope-tagged strains displayed normal growth and wild-type levels of nuclear asynchrony, and proteins of the proper size were recognized on an anti-myc Western blot (Fig. 4 A and not depicted).

Cyclin protein localization in the cell was determined using indirect immunofluorescence. The myc antibody did not show any signal in untagged wild-type strains (Fig. 4 B). As would be predicted, *AgCLN1/2p* was concentrated in nuclei with a single SPB and was diffusely present in the cytoplasm (Fig. 4, C and D; left). Surprisingly, however, this G1 cyclin homologue is also readily visible in nuclei with mitotic spindles. Additionally, *AgCLN1/2p* was observed at many hyphal tips, which are sites of polarized growth (Fig. 4 D, left). Similarly, the B-type cyclin homologue *AgCLB1/2p* was concentrated in nuclei with two SPBs and metaphase spindles as expected but, remarkably, was also clearly present in nuclei with a single SPB and in anaphase nuclei (Fig. 4, C and D; right).

Both types of cyclins seem to be present throughout all cell cycle stages (Fig. 4, C and D). *AgCLB1/2p* was found in 91% of nuclei in a mycelium, and *AgCLN1/2p* was observed in 74% of nuclei ($n > 200$ nuclei). Thus, the proportion of nuclei with a given cyclin is greater than the proportion of nuclei in the corresponding spindle stage (on average only 20% nuclei are in mitosis, yet >90% have mitotic cyclin). Furthermore, nuclei lacking either type of cyclin signal were not in any specific cell cycle stage as determined by spindle appearance, suggesting that these unstained nuclei are results of technical limitations in resolving all proteins by immunofluorescence rather than a large oscillation in cyclin levels in specific stages of the cell cycle (Fig. S2, available at <http://www.jcb.org/cgi/content/full/jcb.200507003/DC1>). There was some variability in the intensity of signals between nuclei, but this also did not correlate with cell cycle stage, and we predict this is caused by variability inherent in immunofluorescence in fungal cells where there is

Table 1. Amino acid sequence comparison of a subset of cell cycle control proteins

<i>S. cerevisiae</i> cell cycle protein	<i>A. gossypii</i> homologous protein	Percent identity
Cyclins and CDK		
Cdc28	Cdc28	86
Cln3	Cln3	34
Cln1	Cln1/2	63
Cln2	Cln1/2	60
Clb5	Clb5/6	44
Clb6	Clb5/6	39
Clb3	Clb3/4	51
Clb4	Clb3/4	52
Clb1	Clb1/2	56
Clb2	Clb1/2	65
Degradation machinery		
Cdc4	Cdc4	52
Cdc34	Cdc34	82
Skp1	Skp1	87
Cdc53	Cdc53	64
Grr1	Grr1	50
Cdh1	Cdh1	66
Cdc20	Cdc20	59
Cdc16	Cdc16	53
Apc1	Apc1	35
CDK inhibitors and regulators		
Sic1	Sic1	37
Far1	Far1	31
Hsl7	Hsl7	45
Hsl1	Hsl1	43
Mih1	Mih1	34
Swe1	Swe1	52
Cdc5	Cdc5	73
Transcription factors		
Swi6	Swi6	49
Swi4	Swi4	43
Mbp1	Mbp1	53
Mcm1	Mcm1	54
Swi5	Swi5/Ace2	35
Ace2	Swi5/Ace2	35
Fkh1	Fkh1/2	46
Fkh2	Fkh1/2	50

Percentage of amino acid sequence identity between selected *S. cerevisiae* cell cycle genes and the syntenic *A. gossypii* homologues. Genes that are duplicated in *S. cerevisiae* but present in a single copy in *A. gossypii* are named using both yeast homologues (e.g., AgClb1/2p).

irregularity in digestion of the cell wall. Thus, the cyclins in *A. gossypii* do not display complete degradation in sync with cell cycle progression but, instead, are found in most nuclei regardless of their cell cycle stage, suggesting that many nuclei have different types of cyclins present at the same time.

Mitotic cyclins are present in nuclei where they were not transcribed

Synchrony in multinucleated systems is hypothesized to depend upon the free diffusion of mitotic regulators in the cytoplasm. Could asynchrony be established simply by limiting diffusion and concentrating cell cycle factors in nuclei? If so, the nuclear limited localization of mitotic cyclins in *A. gossypii* cells could compartmentalize the cell and prevent communication between individual nuclei leading to asynchrony. It is unclear, however, if nascent cyclin proteins expressed from a single nucleus and translated in the common cytoplasm are free to diffuse and enter

neighboring nuclei of different cell cycle stages. To evaluate the basis of the ubiquitous nuclear cyclin localization, we generated heterokaryon cells in which only a subset of nuclei express a tagged version of the cyclin. We then asked whether the nuclei that do not themselves contain an epitope-tagged cyclin gene have detectable levels of tagged cyclin proteins expressed from neighboring nuclei.

AgCLB1/2-13myc was expressed from an autonomous replicating sequence plasmid that also contained lac operator repeats (lacO) and was put into an *A. gossypii* strain that had GFP-lacI-NLS integrated in the genome (Fig. 5 A; Belmont and Straight, 1998). Autonomous replicating sequence-containing plasmids are maintained by *A. gossypii* in a heterokaryon state such that only a subset of nuclei ever contain a plasmid under limited selection pressure, and plasmids are readily lost from nuclei after the removal of all selection (Fig. 5 A). Nuclei containing the plasmid and, thus, also the tagged version of

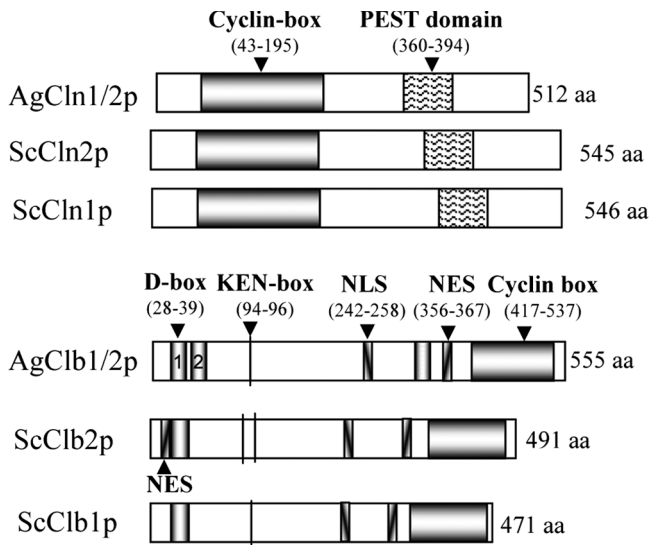


Figure 3. **Comparison of *A. gossypii* and *S. cerevisiae* cyclins.** Domain comparison between AgCln1/2p, ScCln1p, and ScCln2p and, below, between AgClb1/2p, ScClb1p, and ScClb2p. The amino acid positions noted for motifs are for the *A. gossypii* homologue. The cyclin box was identified by a ProSite scan, and the PEST motif is based on Salama et al. (1994). The identity within the PEST domain is 30%, although all key Cdc28p phosphorylation sites are conserved. The D box [db1 (indicated by 1); Ag aa 28–39; Glotzer et al., 1991], the COOH-terminal cyclin box [ProSite scan], one of two KEN boxes (Pfleger and Kirschner, 2000; Hendrickson et al., 2001), the NLS, and one of two NESs (Hood et al., 2001) are conserved in the *A. gossypii* *CLB1/2* sequence. *AgCLB1/2* appears to have an additional putative D-box sequence at aa 84–96 (db2; indicated by 2) and a more NH₂-terminal KEN box at aa 6–8.

AgCLB1/2 were visible by a bright “dot” of GFP-lacI-NLS where it concentrates as a result of the lacO repeats present on the plasmid (Fig. 5 B, left). When all selection for the plasmid was removed, the bright dots of GFP signal were lost from many nuclei so that only a fraction (28%; $n = 300$ nuclei) of nuclei retained a plasmid (5 h after release from selection; Fig. 5 B, right). We then asked whether the *AgClb1/2-13myc* protein was visible in nuclei that did not appear to contain a plasmid. Tagged cyclin proteins were detected in nearly all nuclei regardless of whether they contained the plasmid expressing the epitope-tagged gene. In hyphae in which very few GFP dots were visible, *AgClb1/2-13myc* protein was still apparent in many nuclei. Thus, nuclei that clearly lacked the plasmid containing the tagged gene had tagged protein product. This product most likely derived from neighboring nuclei that still possessed a plasmid or, in some cases, may still contain a reservoir of stable protein. This experiment suggests that mitotic cyclins are freely diffusing, taken up by, and maintained in neighboring nuclei.

Asynchrony persists when cyclins are displaced from nuclei with an exogenous NES

To further address how mitotic cyclin nuclear localization contributes to asynchrony, we shifted a proportion of the cyclin protein out of the nucleus with the addition of a NES to the *AgClb1/2-13myc* protein (Edgington and Futcher, 2001). If nuclear sequestration of newly made cyclin protein is important

for nuclear autonomy, we would predict that the displacement of more mitotic cyclin into the cytoplasm, where it could potentially more freely diffuse, would lead to an increase in synchrony of nuclear division. The additional NES led to an increase in the cytoplasmic levels of *AgClb1/2-13myc* protein as visible by immunofluorescence but still left sufficient protein in the nucleus to promote normal growth and nuclear division. There was no increase in cell cycle synchrony observed despite the increase in cytoplasmic levels of cyclin proteins between cells with an inactive NES or active NES fusion protein (Fig. 6 A). Interestingly, however, these cells did show a higher proportion of nuclei with duplicated SPBs or mitotic spindles, suggesting that cyclin protein concentration gradients were altered by this displacement, leading to an increase in the frequency of mitosis (Fig. 6 B). Thus, nuclear asynchrony or autonomy seems to be established independently of the origins, levels, and localization of mitotic cyclins.

Mitotic cyclin degradation does not correlate with mitotic exit

Cyclin proteins appear to be present in all cell cycle stages, which raises the question of how nuclei cycle with accuracy. To further examine the possible persistence of a pool of mitotic cyclin protein across all cell cycle stages, B-type cyclins were evaluated during a nocodazole arrest and release experiment. This allowed *AgClb1/2p* localization and protein levels to be followed through a synchronous mitotic exit, the time period when mitotic cyclins are predicted to be degraded. Mitotic cyclins were readily observed both by immunofluorescence and by Western blots on whole cell extracts even in cells where the majority (73%) of nuclei had extended spindles and separated DNA indicative of late anaphase/telophase (Fig. 7, A and B). This suggests that B-type cyclin degradation does not correlate with mitotic exit and that any regulative degradation, if present, must occur on only a small fraction of the total protein. Thus, mitotic cyclins seem to persist across the mitosis–G1 transition and potentially may be inactivated for G1 by direct inhibition rather than degradation. When *AgCLB1/2* was expressed in yeast cells either from its own promoter or from a GAL promoter, yeast cells showed a delay in telophase, further suggesting that the *A. gossypii* protein may be somewhat intrinsically stable compared with the yeast cyclins (Fig. S3, available at <http://www.jcb.org/cgi/content/full/jcb.200507003/DC1>).

Mitotic cyclin D-box mutants do not show altered cell cycle progression

These localization and protein level data suggest that cyclin levels do not dramatically fluctuate in a cell cycle–dependent manner in these syncytial cells. The *AgClb1/2p* homologue, which is an essential gene in *A. gossypii*, does contain two NH₂-terminal D-box sequences that, in other systems, have been shown to lead to anaphase-promoting complex-mediated degradation by the proteasome (Glotzer et al., 1991). In *Xenopus laevis*, yeast, and *Drosophila melanogaster* cells, the expression of cyclins lacking the D boxes leads to cell cycle arrest as a result of an inability to exit mitosis (Vodermaier, 2001). If some amount of cyclin destruction is required in vegetatively growing

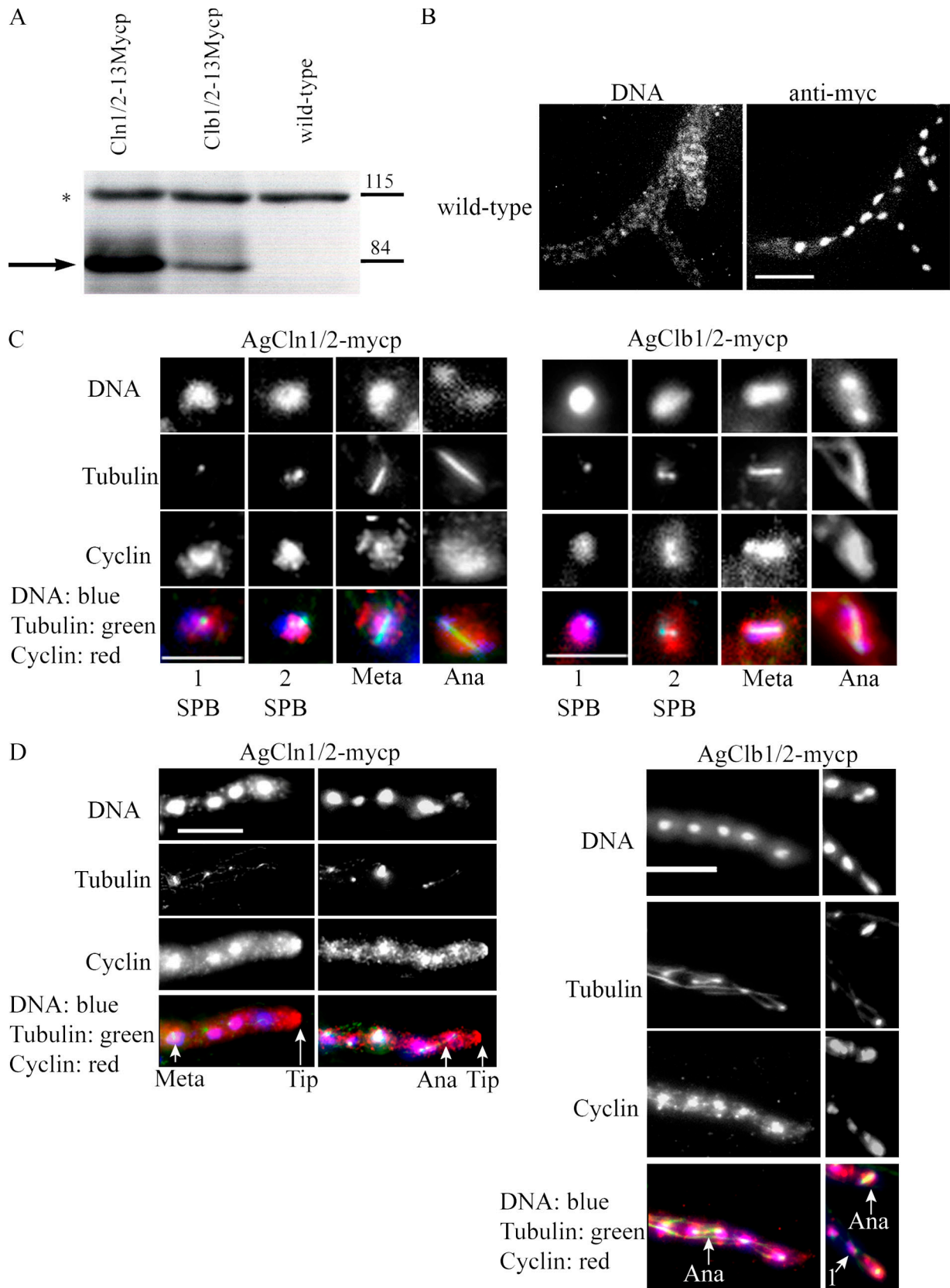


Figure 4. **Cyclins are nuclear and present throughout all nuclear division stages.** (A) Western blot on whole cell extracts showing recognition of epitope-tagged cyclins at 80 kD (arrow) with anti-myc antibody (ASG43 and ASG60) and the absence of a band in an untagged wild-type control strain. The load control is a nonspecific cross-reacting band marked by an asterisk, and the molecular mass marker is given in kilodaltons. All images in B–D are derived from stacks of images acquired in the Z axis and compressed to capture all possible signals in the hyphae. (B) Wild-type, untagged *A. gossypii* cells were processed for immunofluorescence and stained with the myc antibody to show the level of background fluorescence with this antibody. (C) Cyclins

A. gossypii cells, *Agclb1/2* D-box mutants should display a clear cell cycle block or delay. The expression of mutants in which either D-box was deleted individually (*Agclb1/2-db1* and *Agclb1/2-db2*) or both were deleted (*Agclb1/2-dbd*) led to no growth defect or alterations in nuclear division based on the proportion of nuclei in different cell cycle stages and levels of asynchrony (Fig. 7, C [top] and D; $n > 200$ cells scored for each strain). To determine whether these mutant strains were delayed in exiting mitosis, cells were synchronized with nocodazole and released to observe a synchronous mitosis in the population. (As with Fig. 2, in the first hour after release from nocodazole, cells rebuild cytoplasmic microtubules before restoring nuclear microtubules). When compared with cells with only wild-type *AgCLB1/2*, both the single and double D-box mutants showed similar rates of return to asynchrony and similar proportions of nuclei in mitosis after 120 min, suggesting no significant delay in mitotic exit (Fig. 7 C, $t = 120$ min). Surprisingly, protein levels of *AgClb1/2-db1p*, *-db2p*, and *-dbdp* were comparable with wild-type *AgClb1/2p* (Fig. 7 E). We predict that the D-box sequences may be required in a different growth stage but are dispensable for normal nuclear division during vegetative growth.

AgSic1p contributes to accurate nuclear division

Normal mitotic progression does not appear to require sharp degradation of the mitotic cyclin pools, yet nuclear division is apparently accurate and faithful in *A. gossypii*. What prevents premature entry into mitosis given the presence of mitotic phase-promoting cyclin early in the cell cycle? One possibility is that mitotic CDK inhibitors such as *AgSwe1p* and *AgSic1p* are acting in a nuclear-autonomous fashion and repress any CDK-*Clb1/2p* complexes present in G1 nuclei. Although mutants lacking *AgSWE1* showed minimal mitotic defects (unpublished data), 35% of *Agsic1Δ* cells stopped growth with 10–20 nuclei, aberrant multipolar spindles, and irregular nuclear distribution and density (Fig. 8, A [middle] and B). A proportion of *Agsic1Δ* cells (55%) formed a developed mycelium, but, out of these, the majority ultimately arrested growth with up to 100 densely packed nuclei (Fig. 8 A, bottom). About 10% of mutant spores produced viable and sporulating mycelia, potentially as a result of the acquisition of a suppressing mutation or the production of multinucleated spores during sporulation that contain a wild-type nucleus. Many cells in an *Agsic1Δ* mutant population show defects in nuclear division, suggesting that *AgSic1p* may help limit the premature activity of mitotic CDK complexes.

AgSic1p localization changes across the nuclear division cycle

If *AgSic1p* is a source of *AgCDK/Clb1/2p* oscillation, the protein may vary in localization and/or abundance across different stages of the cell cycle. We localized *AgSic1p-13myc* protein

by immunofluorescence and evaluated its behavior during different nuclear division cycle stages to examine whether this predicted CDK inhibitor may oscillate in any way. We found that *AgSic1p* undergoes notable changes to its sub-nuclear localization, correlating with different stages of nuclear division (Fig. 9). In nuclei with a single SPB, *AgSic1p* was present throughout the nucleoplasm. Interestingly, the *AgSic1p* was not uniformly distributed; rather, there were foci of intense signal in addition to a more diffuse signal across the nucleus. In nuclei with duplicated SPBs, the nucleoplasm signal weakened, and some protein was also observed on short spindles. When metaphase spindles were present, *AgSic1p* was barely detectable in the nucleoplasm, and, occasionally, a low level of *AgSic1p* was apparent on spindles. In anaphase, as soon as any separation was apparent between the stained DNA, *AgSic1p* reappeared on elongating spindles as an intense signal. *AgSic1p* was observed on spindles throughout late anaphase/telophase as well as in the nucleoplasm of the daughter nuclei. These dynamics would suggest that *AgSic1p*'s subnuclear localization is regulated to alter either its own activity or its contact with interacting proteins through different stages of the nuclear cycle.

Discussion

In this study, we provide evidence for nuclear-autonomous division in multinucleated fungal cells. Nuclear pedigrees and observations of spindle morphology in neighboring nuclei demonstrate that nuclei divide asynchronously in these cells and display independence in a common cytoplasm. Cyclin proteins appear to be highly enriched in nuclei but able to diffuse in the cytoplasm without disturbing asynchrony. Remarkably, the mitotic cyclin protein is present in all stages of the nuclear division cycle, suggesting that oscillation of CDK activity is not primarily generated by periodic expression and complete degradation of cyclins (Fig. 9 B). From our experiments, we propose that the syncytial nature of these cells has favored the evolution of a cell cycle oscillator built primarily upon the action of CDK inhibitors rather than control of cyclin protein abundance. These data raise two unique problems for these cells: how does cell cycle progression occur accurately given the apparent lack of coordinated oscillation in cyclin proteins, and how do nuclei behave independently in the same cytoplasm?

Current models of cell cycle control are rooted in the principle of a biochemical oscillator built through the intertwined regulation of synthesis and destruction of many proteins (Murray, 2004). In yeast, ~800 genes display periodic expression over the cell cycle, including cyclins, CDK inhibitors, degradation machinery, and transcription factors, and many of these proteins similarly have periodic destruction patterns (Spellman et al., 1998). Extensive experimental data combined with recent mathematical models have shown that this choreographed synthesis and destruction program enables the cell cycle to function

AgCln1/2-13myc (ASG60) and *AgClb1/2-13myc* (ASG43) were visualized along with tubulin by immunofluorescence. Individual nuclei are shown representing different spindle stages. All images were acquired and processed under identical conditions. (D) Images showing intact hyphae of *AgCln1/2-13myc* and *AgClb1/2-13myc* strains prepared as in C. Arrows highlight nuclei of different cell cycle stages as described for Fig. 2. In overlays, DNA is blue, tubulin is green, and cyclins are red, leading to purple nuclei containing cyclins. Bars (B and D), 10 μm ; (C) 5 μm .

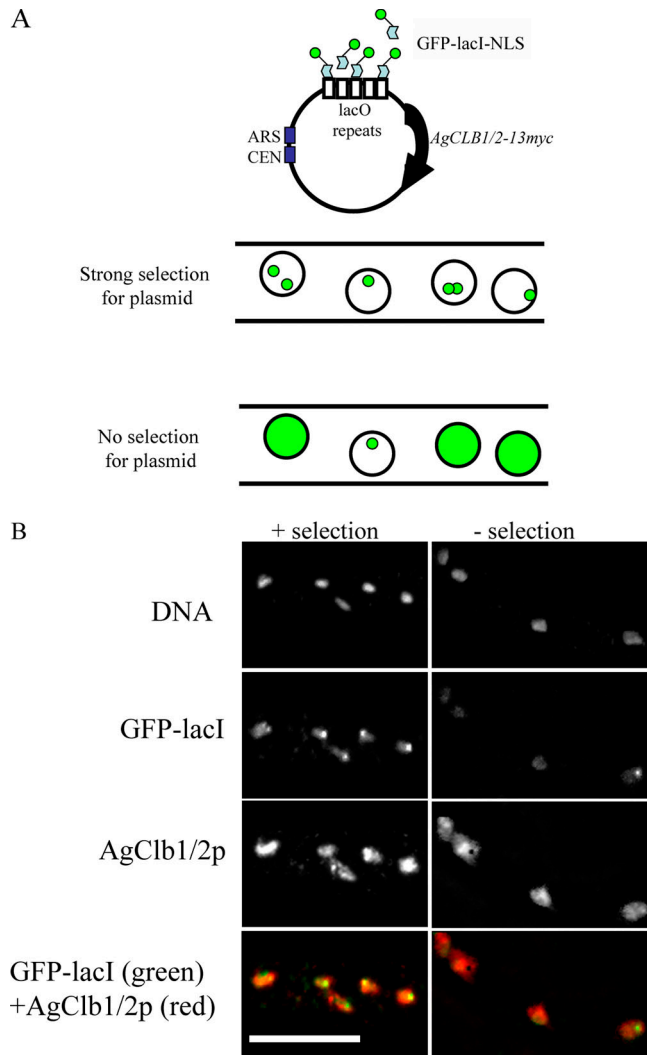


Figure 5. Cyclins are present in nuclei that are distinct from the site of expression. (A, top) Centromere-based plasmids containing an epitope-tagged *AgCLB1/2* gene, *lacO* repeats, and a dominant drug resistance marker (*GEN3*) were constructed (pAKH38) that can replicate and segregate in *A. gossypii* nuclei. (bottom) *A. gossypii* GFP-lacI-NLS cell strains were transformed with the plasmid containing the *AgCLB1/2-13myc* and the *lacO* repeats. In the presence of the G418 drug, many nuclei show one or two tight dots of GFP signal that are indicative of plasmid. When the selective drug pressure is removed, within 5 h, the GFP signal is lost in the majority of nuclei, and GFP-lacI appears as a diffuse signal in the nucleoplasm, which is indicative of no plasmid. CEN, centromeric. (B) GFP-lacI-NLS + *placO-CLB1/2-13myc* (AKHAg28) were grown overnight for 16 h in 200 $\mu\text{g/ml}$ G418 drug. Half of the cells were fixed before washing out the drug (left), and the remaining half of the cells were resuspended in fresh media lacking the selective drug and allowed to grow for 5 h before fixation (right). Cells were processed for immunofluorescence to visualize GFP-lacI-NLS and *AgClb1/2-13myc*. Bar, 10 μm .

as a bistable system that alternates between two stable but mutually exclusive and irreversible states, G1 and S/G2/mitosis phases (Chen et al., 2000; Cross et al., 2002). In yeast, the basis of this bistability is thought to originate from essentially two redundant, although distinct, biochemical oscillators: one rooted in a negative feedback loop that triggers mitotic cyclin (ScClb2p) degradation and the other, which is a “relaxation oscillator” that ensures alternating states of high Clb2p/low Sic1p and low Clb2p/high Sic1 activity (Cross, 2003). Only one of the two

oscillators needs to be present for yeast to divide. Importantly, however, both oscillators require fluctuating protein levels of either Clb2p or Sic1p to properly function and maintain the bistable nature of the system (Cross, 2003).

It appears as though the cell cycle oscillators in *A. gossypii* may not require temporally regulated synthesis and complete degradation of at least some key cell cycle control proteins. Conceivably, there is highly localized degradation of a small but functionally significant fraction of cyclin proteins that has escaped our detection in these studies. However, lack of a dominant phenotype in cells expressing cyclin D-box mutants suggests that regulated proteolysis of cyclins is not critical for regulating mitosis during normal vegetative growth. The continuous cytoplasmic streaming of a syncytial cell complicates the establishment of an oscillating protein gradient through degradation alone because new protein is always supplied by the cytoplasm. Thus, multinucleated cells like *A. gossypii* may have evolved to favor the modulation of CDK activity by inhibitors to generate bistability in the cell cycle circuit. Although at least the *AgCln1/2p* and *AgClb1/2p* cyclins are consistently present in nuclei throughout the cell cycle, we predict that they cannot be simultaneously active. In these cells, the feedback loops that run the cell cycle oscillator would lead to changes in activity instead of changes in protein synthesis and degradation. In fact, a bistable cell cycle oscillator has been shown to function with constant levels of mitotic cyclins in *Xenopus* egg extracts simply through modulating the activity of Cdc2 (CDK; Pomeroy et al., 2003). Our data suggest that Sic1p acts in *A. gossypii* cells to ensure alternating states of CDK activity even in the presence of relatively constant *AgClb1/2p* levels.

Nuclear-autonomous regulation of a single factor such as the inhibitor *AgSic1p* could be sufficient to provide the necessary oscillation in CDK activity for DNA synthesis, spindle construction, and chromosome segregation without great changes in cyclin levels. Thornton and Toczyski (2003) recently presented evidence in *S. cerevisiae* that oscillation in CDK activity could be generated without the degradation of mitotic cyclins simply by the overexpression of ScSic1p (Thornton et al., 2004). In these cells, ScClb2p was modified to be present in G1 phase, as is observed in the normal *A. gossypii* cycle, but was held inactive by an overabundance of Sic1p. Furthermore, ScSic1p becomes essential in yeast cells overexpressing ScClb2p (Cross et al., 2005). Clearly, *AgSic1p* makes a vital contribution to accurate division in these multinucleated cells and may prevent G1 nuclei with *AgClb1/2p* from entering mitosis before the completion of S phase. The *AgSic1p* homologue shows relatively low identity to the yeast homologue, suggesting it could have diverged to be more stable and active and may act as a more potent CDK inhibitor. Additionally, *AgSic1p* subnuclear localization dynamics present the possibility that it is tightly regulated in the nuclear cycle. In the middle of mitosis, when CDK/Clb activity is at a peak, *AgSic1p* levels diminish in the nucleus potentially as a result of either nuclear autonomous degradation or transient nuclear export. The rapid accumulation of *AgSic1p* shortly thereafter in early anaphase, however, argues that nuclear import rather than new synthesis would replenish the nuclear protein pool. Thus, we hypothesize that multinucleated

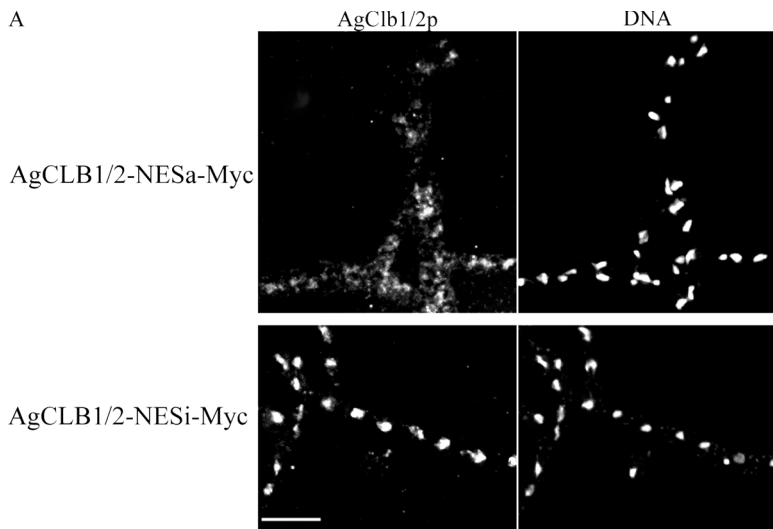
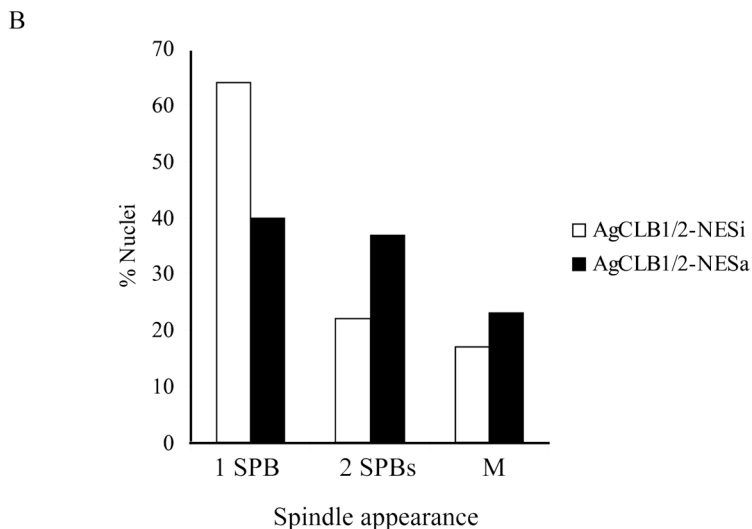


Figure 6. **Displacement of AgCLB1/2p to the cytoplasm with an exogenous NES.** (A) AgCLB1/2-NESa-Myc (AKHAg19, active NES) or AgCLB1/2-NESi-Myc (AKHAg22, inactive NES control) were grown for 16 h and processed for antitubulin and anti-myc immunofluorescence. Images were captured and processed identically to compare protein intensities within the figure. Bar, 10 μ m. (B) Percentage of nuclei in each spindle stage based on tubulin staining. $n > 200$ nuclei scored for each strain. M, mitotic.



cells such as *A. gossypii* may generate oscillation in CDK activity through posttranslational regulation of inhibitor and cyclin protein activity rather than through tightly controlled abundance. It is conceivable that other cyclins in the system, such as AgClb3/4p or AgClb5/6p, may fluctuate, but this will be tested in future experiments.

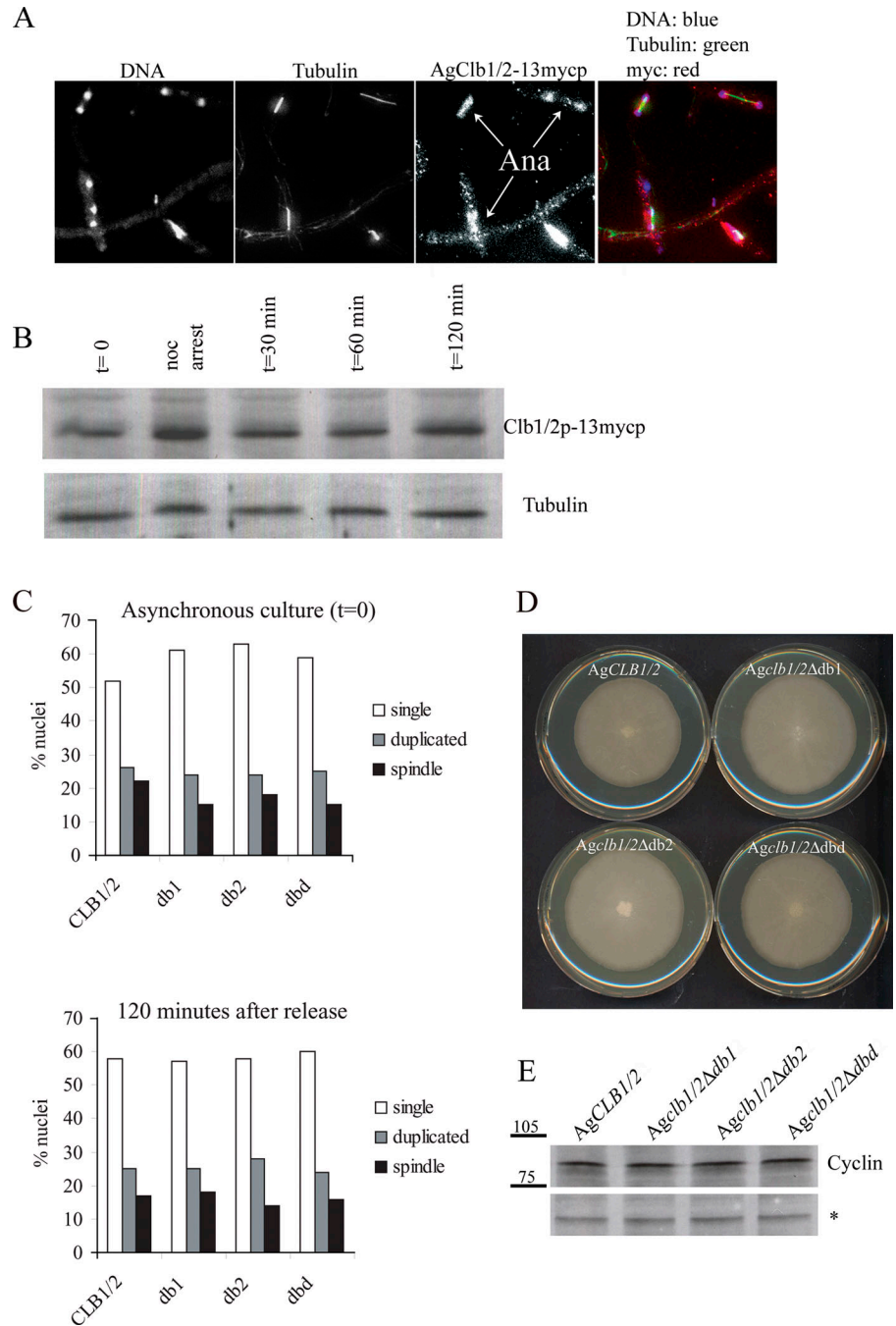
How does each nucleus cycle independently? One simple possibility is that asynchrony or nuclear autonomy could be generated as a result of the nature of the cell cycle circuits in *A. gossypii*, which, as discussed above, may be subtly different than those used in uninucleated cells. Interestingly, recent modeling and experimental work by Pomerening et al. (2005) in *Xenopus* egg extracts in which a positive feedback loop in the Cdc2/anaphase-promoting complex circuit is blocked (producing damped oscillations in CDK activity) led to asynchrony among nuclei in the extracts. In *A. gossypii*, the core circuit regulating CDK oscillations may be wired in such a way that transitions between division stages are gradual (which happens when positive feedback loops are repressed in the *Xenopus* model), making nuclei divide out of sync. Alternatively, cell cycle stage identity is given by periodically transcribed genes that may produce proteins that are translated

in a common cytoplasm but, unlike the cyclins, could be directed back to their transcriptional “mother” nucleus. This factor could be a direct regulator of Sic1p that ensures temporal regulation of the localization and, potentially, activity of the CDK inhibitor.

What are possible mechanisms to limit the presence of such an “identity” factor to a specific stage nucleus? Potentially, mRNAs could be targeted to a domain of the ER adjacent to nuclear pores to facilitate rapid uptake into nuclei after translation, or the newly translated proteins could be immediately complexed with nuclear import factors. Conceivably, such a protein could actually be translated within the nuclei, thereby avoiding post-translational diffusion within the cytoplasm (Iborra et al., 2001). Alternatively, perhaps such a factor is constitutively transcribed but would be selectively blocked in certain cell cycle stages by small regulatory RNA molecules that exist only within the nucleus. The nuclear pore complex itself could be remodeled in a cell cycle-dependent manner such that certain proteins are only allowed to enter nuclei of a particular cell cycle phase (Makhnevych et al., 2003; De Souza et al., 2004). Finally, this could be a factor that is associated with some cell cycle event, such as DNA replication, and could be activated in a nuclear-autonomous

Figure 7. **AgClb1/2p levels do not diminish at mitotic exit, and cyclin mutants lacking D-box sequences have normal nuclear division.**

(A) Mitotic AgClb1/2-13myc was visualized in young germling cells (ASG43) that were arrested and released from nocodazole. Clb1/2-13myc protein was evaluated by immunofluorescence and (B) on a Western blot, where tubulin was used as a loading control. Arrows highlight anaphase/telophase nuclei with mitotic cyclin. In overlays, DNA is blue, tubulin is green, and cyclins are red, leading to purple nuclei containing cyclins. (C) Wild-type strains were transformed with plasmids containing either *AgCLB1/2*, *Agclb1/2-db1* (pAKH47), *Agclb1/2-db2* (pAKH45), or *Agclb1/2-dbd* (pAKH46) and grown in liquid media containing G418 to maintain the plasmid in most nuclei. A subset of asynchronous cultures grown for 16 h were fixed and processed for antitubulin immunofluorescence, and nuclei were evaluated according to spindle stage ($t = 0$). The remaining cultures were then arrested and released from nocodazole, and percent nuclei in mitosis were evaluated at 120 min, the point at which wild-type cultures have just returned to normal asynchronous levels of mitosis. (D) Spores from cells containing *AgCLB1/2*, *Agclb1/2-db1*, *Agclb1/2-db2*, or *Agclb1/2-dbd* were inoculated on plates, and radial growth was evaluated after 5 d at 30°C. Similar growth rates were also observed when alleles are integrated in the genome at the *ADE2* locus (not depicted). (E) Spores from cells containing *AgCLB1/2-13myc*, *Agclb1/2-db1-13myc*, *Agclb1/2-db2-13myc*, or *Agclb1/2-dbd-13myc* were inoculated into liquid media containing 150 $\mu\text{g/ml}$ G418 for selection, grown for 16 h at 30°C, and lysed and processed for an anti-myc Western blot. The asterisk is a nonspecific cross-reacting band for a loading control. Molecular mass is shown in E in kilodaltons.



manner only upon completion of the event in an individual nucleus. Thus, there are many ways to conceive of how the nuclei could maintain independent cell cycle stage identity within the same space.

Why would a multinucleated cell evolve mechanisms to ensure asynchronous nuclear division? What advantage does a nuclear-autonomous cell cycle bring to a multinucleated cell? Such cells may limit mitosis so that it does not occur synchronously to prevent a sudden doubling of the nucleocytoplasmic ratio. A nuclear-autonomous cell cycle is a possible way to maintain the balance of nuclei in the cell, ensuring that large fluctuations in the nuclear number do not occur without concur-

rent growth. A nuclear-autonomous cycle also enables nuclei in a specific subcellular location to divide, such as near a branch point or at a site rich in nutrients, without having to duplicate or transport more distant nuclei. Thus, asynchronous mitosis enables conservative and more spatially directed control of mitosis. When combined, these experiments demonstrate that the basic eukaryotic cell cycle control network may have evolved in diverse ways to accommodate the unique geometry of a multinucleated cell. Given the evolutionary relationship between budding yeast and *A. gossypii*, it is clear that dramatic differences in cell behavior can be achieved through modulating a very similar set of proteins.

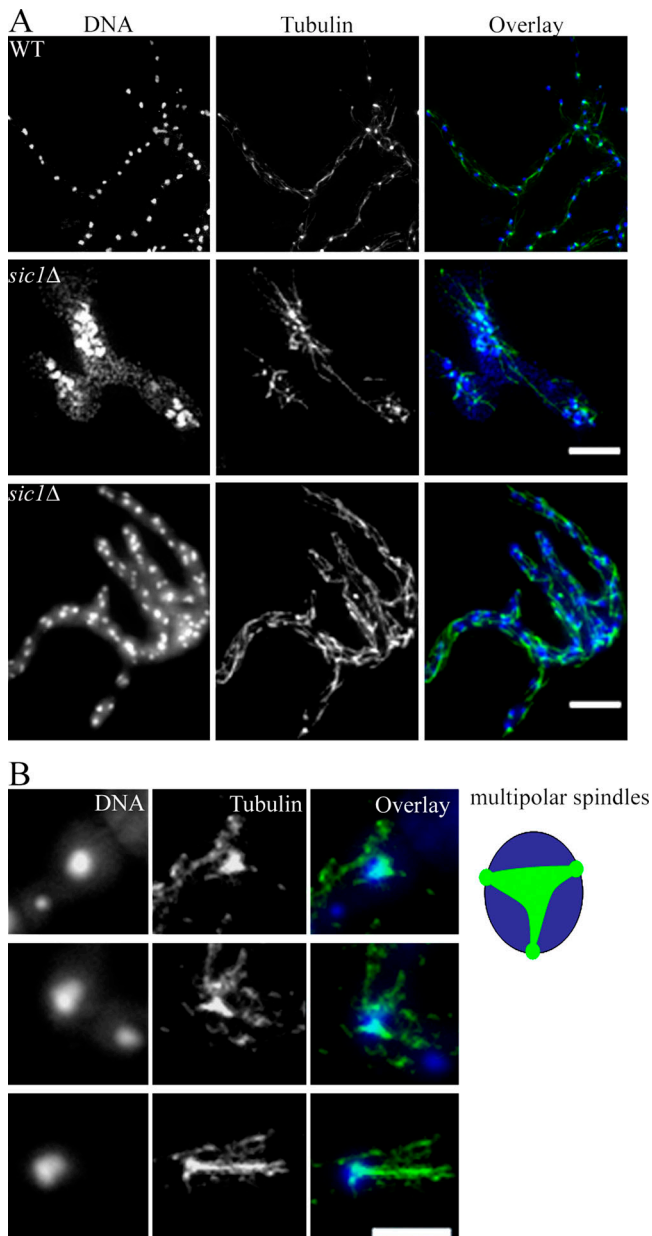


Figure 8. *Agsic1*Δ mutants arrest growth with abnormal nuclear distribution and spindle defects. (A) *Agsic1*Δ mutants (ASG65) were germinated in the presence of 50 μg/ml CloNAT to select for cells with the gene disrupted, were fixed and processed for antitubulin immunofluorescence, and compared with wild-type cells. The middle panels show the typical arrest phenotype of bipolar germlings, and the bottom panels show cells that arrest as a small mycelia and high nuclear density. Bars, 10 μm. (B) Higher magnification of nuclei showing multipolar spindle defects observed in *Agsic1*Δ cells. Nuclei are in blue, and microtubules are in green. Bar, 5 μm.

Materials and methods

Media, strains, and plasmid construction

A. gossypii media, culturing, and transformation protocols are described in Wendland et al. (2000) and Ayad-Durieux et al. (2000), and strains are listed in Table II. Nocodazole stocks were 3 mg/ml in DMSO and used at a final concentration of 15 μg/μl (Sigma-Aldrich). Standard methods were used for culturing *S. cerevisiae* and plasmid manipulations (Sambrook, 2001). Plasmids generated and used in this study are listed in Table S1 (available at <http://www.jcb.org/cgi/content/full/jcb.200507003/DC1>). PCR was performed using standard methods with Taq polymerase from Roche,

oligonucleotides were synthesized at MWG, and all restriction enzymes came from New England Biolabs, Inc. or Roche. Oligonucleotide primers are listed in Table S2.

The wild-type *A. gossypii* strain was transformed with the pHHF4-GFP to make the AgHHF1 strain (Alberti-Segui et al., 2001). This strain was always grown under selective conditions with 200 μg/ml G418 (Calbiochem/Merck) to maintain the plasmid during time-lapse video microscopy for pedigree analyses. *SPC42* was tagged at the COOH terminus with GFP at its genomic locus using PCR-mediated gene targeting with oligonucleotides *SPC42-GFPS1* and *SPC42-GFPS2* and the pGUG template to generate ASG38, which was verified using oligonucleotides *SPC42 I* and *GG2*.

The 13-myc-GEN cassette for *A. gossypii* was generated by digesting pFA6-13myc with BglII and ligating the gel-purified 3.2-kb fragment to the 1.7-kb fragment generated from digesting pGEN3 with EcoRV-BamHI. To COOH-terminally tag the cyclin genes and AgSic1p, this 13-myc cassette was amplified with oligonucleotides *Clb2-MycF* and *Clb2-MycR* for *AgCLB1/2*, *Cln2-MycF* and *Cln2-MycR* for *AgCLN1/2*, and *Sic1-MycF* and *Sic1-MycR* for *AgSIC1*. The resulting PCR products were cotransformed in yeast with the plasmid pAG7578 (for *CLB1/2*) or pAG5016 (for *CLN1/2*) to produce pCLB2-13myc and pCLN2-13myc. *AgSIC1* was tagged at its endogenous locus by direct transformation with the PCR fragment. pCLB2-13myc was digested with BglII and NcoI, and pCLN2-13myc was digested with Apal before the transformation of *A. gossypii* cells to tag the endogenous cyclin genes, creating ASG43 and ASG60, which were verified with oligonucleotides *Clb1-Myc-I* or *Cln2-Myc-I* and *MycI*.

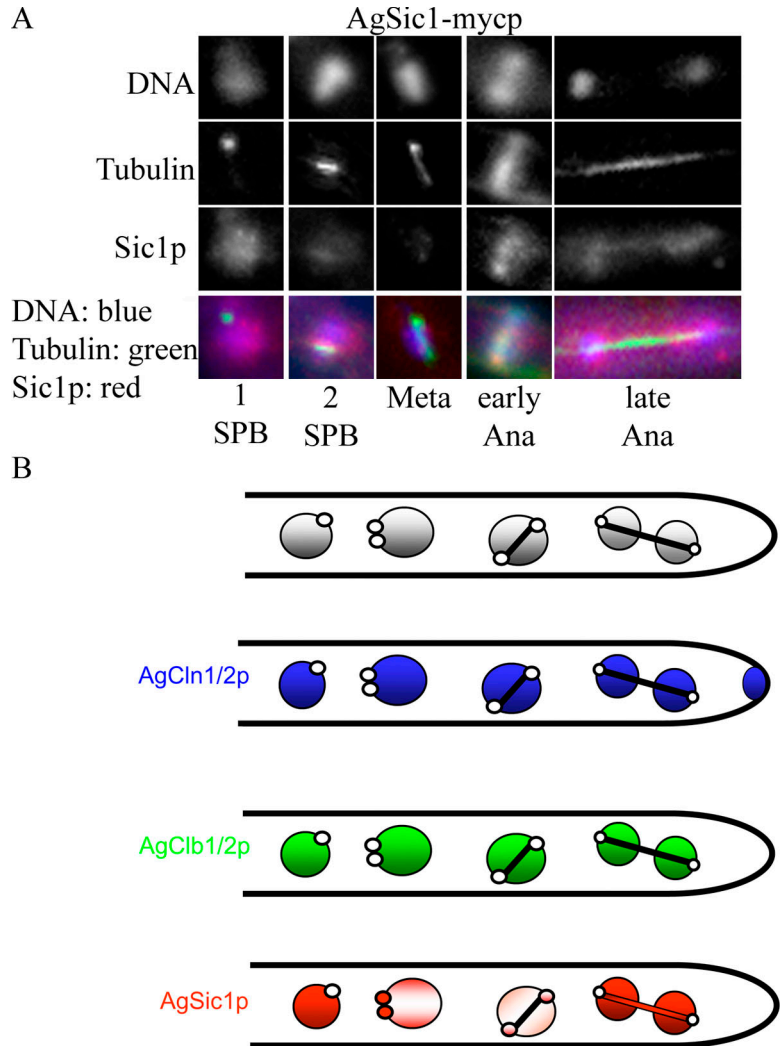
To generate the ASG46 strain, the plasmid pHHF1 [29] was digested with BglII and SacI. Blunt ends were generated using the 5'–3' exonuclease activity of Vent polymerase. The 1,747-bp fragment was subcloned into pAIC opened at the *Scal* site (Knechtle, 2002). The new plasmid pHPH001 carried a COOH-terminal fusion of GFP (S65T) to the ORF of histone H4 (*AgHHF1*) with flanking homologies to the *AgADE2* locus. HHF1-GFP was under the control of the *AgHHF1* promoter and was terminated by *ScADH1-T*. 5 μg pHPH001 was isolated from *Escherichia coli*, digested with EcoRI-HindIII, and transformed into the partially deleted *AgADE2* locus of the *Agade2Δ1* strain (Knechtle, 2002). Transformants were obtained on minimal medium lacking adenine (ASG46). This strain was then used as the basis for constructing ASG50, which contains *HHF4-GFP* and *SPC42-RFP*.

The *RFP*-tagging cassette for *A. gossypii* was generated by digesting pRedStar2 (provided by M. Knop, European Molecular Biology Laboratory, Heidelberg, Germany; Maekawa et al., 2003) with BamHI and SphI and ligating the purified 3-kb fragment to the 1.5-kb fragment generated by digesting pGEN3 with BamHI and SphI. This was used as a template to amplify *RFP-GEN3* using oligonucleotides with homology to the COOH terminus of *AgSPC42*, *SPC42RedS1*, and *SPC42RedGS2*. The resulting PCR product was cotransformed into yeast cells together with pAG19228 to generate pAG-*SPC42RFP*, which was then digested with XbaI and MscI to transform strain ASG46 and tag the endogenous *SPC42* gene with *RFP* to produce ASG50, which was verified with oligonucleotides *Red I* and *SPC42 I*.

A. gossypii deletion mutants were made using pairs of gene name S1/S2 oligonucleotides for each locus, which contained 45-bp homology upstream and downstream of the ORFs. Wild-type *A. gossypii* cells were transformed with PCR products amplified off the pGEN3 template with the S1/S2 oligonucleotide pairs. The primary transformation produces heterokaryon cells, which have a mixture of wild-type and transformed nuclei; thus, even mutations in essential genes produce viable transformants. Transformants were verified with oligonucleotide gene name *G1/G4/I*, and three independent transformants were characterized for each mutant. To evaluate phenotypes of lethal mutants, the heterokaryon was sporulated to produce uninucleated spores, which were then germinated under selective conditions.

The *clb1/2Δdb* mutants were made using an overlap PCR approach, which deleted amino acids 27–36 (db1) and/or amino acids 84–96 (db2) in *AgClb1/2*. To generate pAKH47 (db1), pCLB2-13myc was used as a template for a PCR reaction with *clb2dbA* and *clb2dbB* to produce a 643-bp product and a second reaction with *clb2dbC* and *clb2dbD* to create a 537-bp product. Oligonucleotide *clb2dbA* was homologous to a region 500 bp upstream of the D box, and *clb2dbD* was homologous to a region 600 bp downstream of the D box. Oligonucleotide *clb2dbB* contained homology to *CLB2* immediately upstream of the D box but lacked the D-box sequences between nt 81 and 108. Additionally, the 3' end of *clb2dbB* has homology to the region immediately downstream of the D box, which is also present in the 5' end of oligonucleotide *clb2dbC*. This overlapping homology was then used in a second PCR

Figure 9. **AgSic1p localization varies according to nuclear division cycle stage, and summary of *A. gossypii* nuclear division cycle.** (A) AgSic1-13mycp (ASG61) cells were grown for 18 h and processed for antitubulin and anti-myc immunofluorescence. All images in the AgSic1p panel were taken with identical exposure times and processed in parallel with identical processing conditions. 82% of one-SPB nuclei, 76% of two-SPB nuclei, 75% of metaphase nuclei, 80% of early anaphase nuclei, and 79% of late anaphase/telophase nuclei showed signal as depicted by these representative nuclei ($n = 400$ total nuclei). (B) Summary of asynchronous nuclear division and localization of cell cycle regulators.



reaction involving the two products of A + B and C + D and additional clb2dbA and clb2dbD primers to create a 1.1-kb fragment containing an in-frame deletion of the D box (db1) and part of the AgCLB1/2 ORF. To generate a full-length db mutant, this 1.1-kb fragment was cotransformed into yeast strains with pCLB2-13myc, which had been digested with NheI located just upstream of the D box. The gap-repaired plasmid containing the deleted D box was confirmed by sequencing and was called pclb2Δdb1-13myc.

Similarly, to generate pAKH45 (db2), oligonucleotides clb2dbA and clb2db2B were used with pCLB2-13myc as a template in the first reaction, and clb2db2C and clb2dbD were used in the second reaction. Clb2db2B contains sequence just upstream of D box 2 and lacks the sequence from nt 248–287 corresponding to db2. Additionally, the 3' end of clb2db2B has homology to the region immediately downstream of the D box 2, which is also present in the 5' end of oligonucleotide clb2db2C. The full-length mutant lacking db2 was generated as for db1, but the pCLB2-13myc plasmid was digested with AarI for cotransformation in yeast. To delete both D boxes (pAKH46), a similar attempt was chosen, but pAKH45 was used as template for both PCR reactions, using clb2dbA and clb2db1B for the first and clb2db1C and clb2dbD as primers for the second PCR reaction, as described for constructing the db1 mutant.

The GFP-lacI-NLS integration cassette for *A. gossypii* was generated with pKL55Y, which was provided by C. Pearson and K. Bloom (University of North Carolina, Chapel Hill, NC). pKL55Y was digested with SapI and HindIII to generate a 2-kb fragment, and blunt ends were generated with Vent polymerase and ligated into the ScaI site of the pAIC plasmid. This produced two plasmids, pAKH35 and pAKH36, which were verified by sequencing and differ only in the orientation of lacI-NLS-GFP. pAKH35

and pAKH36 were digested with HindIII-NotI and transformed into the partially deleted *AgADE2* locus of the *Agade2Δ1* *A. gossypii* strain (Knechtle, 2002) to generate AKHAg26 and AKHAg27, which contain the lacI-NLS-GFP integrated at the ADE2 locus. These strains were verified by PCR and showed diffuse, nuclear GFP signal.

To generate the plasmid containing the lacO repeats and AgCLB1/2-13myc (pAKH38), pKL60Y (provided by C. Pearson and K. Bloom) was cut with PacI and BamHI to generate a 1.5-kb fragment containing 32 lacO repeats, and blunt ends were generated using Vent polymerase. pUC21 was linearized with the blunt end cutter HincII and ligated with the lacO fragment, resulting in pAKH37. pAKH38 was generated by digesting pAKH37 with KpnI and NdeI and ligating this 1.6-kb fragment to the 9,535-bp fragment generated from digesting pCLB2-13myc with KpnI-NdeI. pAKH38 was sequenced and transformed as plasmid into AKHAg26 and AKHAg27, resulting in AKHAg28 and AKHAg29, respectively.

The forced localization cassettes were built using p306-NESa and p306-NESi, which were provided by N.P. Edgington and B. Futcher (SUNY Stony Brook, Stony Brook, NY). The NESa and NESi of the plasmids p306-NESa and p306-NESi were PCR amplified with primers NES-P1-NdeI and NES-α-P2-SmaI or NES-i-P2-SmaI containing the restriction sites NdeI and SmaI. The 156-bp PCR product was cut with NdeI and SmaI, resulting in a 130-bp fragment, and was ligated into the NdeI- and SmaI-cut pAG13-myc. This gave rise to pAKH20 (containing NESa) and pAKH21 (containing NESi), which were both verified by sequencing.

For construction of the forced localization cassettes of the mitotic cyclin *AgCLB1/2*, the pAKH20 and pAKH21 cassettes were amplified with the oligonucleotides Clb-NES-P1 and Clb-Myc-P2, each giving rise to a 2,800-bp fragment. These PCR fragments were cotransformed into yeast

Table II. *A. gossypii* strains used in this study

Strain	Relevant genotype	Source
WT	<i>leu2Δ thr4Δ</i>	(Altmann-Johl and Philippsen, 1996)
HHF1-GFP	<i>pHHF1-GFP::GEN3 leu2Δ thr4Δ</i>	(Alberti-Segui et al., 2001)
ASG38	<i>SPC42-GFP::GEN3/SPC42 leu2Δ thr4Δ</i>	This study
ASG43	<i>CLB1/2-13myc::GEN3 leu2Δ thr4Δ</i>	This study
ASG46	<i>ade2::HHF1-GFP::ADE2 leu2Δ thr4Δ</i>	This study
ASG50	<i>ade2::HHF1-GFP::ADE2 leu2Δ thr4Δ SPC42-RFP::GEN3/SPC42</i>	This study
ASG60	<i>CLN1/2-13myc::GEN3 leu2Δ thr4Δ</i>	This study
ASG61	<i>SIC1-13myc::GEN3 leu2Δ thr4Δ</i>	This study
ASG65	<i>sic1ΔNAT1 leu2Δ thr4Δ</i>	This study
AKHAg19	<i>CLB1/2-NESa-13myc-GEN3</i>	This study
AKHAg22	<i>CLB1/2-NESi-13myc-GEN3</i>	This study
AKHAg26	<i>ade2::GFP-lacI-NLS::ADE2</i>	This study
AKHAg28	<i>ade2::GFP-lacI-NLS::ADE2+placO-CLB1/2-13myc</i>	This study
AKHAg29	<i>ade2::GFP-lacI-NLS::ADE2+placO-CLB1/2-13myc</i>	This study

All ASG and AKHAg strains are *A. gossypii* derived from the wild-type ($\Delta\Delta\Delta$) strain. Individual nuclei are haploid, and most strains are homokaryotic (all nuclei have same genotype). In cases of heterokaryons (strains with mixed genotypes for a given locus), genotypes of the two different types of nuclei are written, separated by a plus sign.

cells together with pAG7578 to generate pAKH25 and pAKH26. pAKH25 and pAKH26 were digested with Stu1, NcoI, and BsrG1, and the resulting DNA was transformed into *A. gossypii* cells, resulting in the strains AKHAg19 and AKHAg22. These new strains were verified using oligonucleotide CLB1/2-11 and G3.

Immunofluorescence, Hoechst staining, and Western blotting

A. gossypii cells were processed for immunofluorescence as described for yeast cells (Pringle et al., 1991) with slight modifications. Young mycelium containing ~75–100 nuclei were fixed for 1.5 h in 3.7% formaldehyde (Fluka) and digested in 1.0 mg/ml zymolyase + 1% β -mercaptoethanol for 30–45 min before antibody incubation. Anti-myc and tubulin stainings were performed sequentially so as to limit cross reactivity, beginning with mouse anti-myc (Santa Cruz Biotechnology, Inc.) at 1:100, AlexaFluor488 anti-mouse (Invitrogen) at 1:200, rat antitubulin (YOL34; Serotec) at 1:50, and AlexaFluor568 anti-rat (Invitrogen) at 1:200 with Hoechst (Invitrogen) dye to visualize nuclei at 5 μ g/ml. Rabbit anti-GFP (Invitrogen) was used to detect GFP-lacI-NLS at a 1:100 dilution, and AlexaFluor488 anti-rabbit was used at a 1:200 dilution (Invitrogen). Antibody dilutions and washes were made with PBS + 1.0 mg/ml BSA.

When only nuclei were visualized, cells were fixed for 1 h in 3.7% formaldehyde in growth media at growth temperature, washed once in 1 \times PBS, and resuspended in a 5.0- μ g/ml solution of Hoechst dye. Cells were incubated with dye for 30 min at room temperature and washed three times in 1 \times PBS.

Western blotting was performed using standard conditions and nitrocellulose membranes, blocking with 3% milk in 0.1% PBS-Tween (Sambrook, 2001). Mouse anti-myc was used at 1:2,000, and HRP anti-mouse (Jackson ImmunoResearch Laboratories) was used at 1:10,000. ECL chemiluminescence solution (GE Healthcare) was used to develop the Western blots.

Microscopy

The microscope used for all fixed cell images (immunofluorescence and Hoechst stainings with cells mounted in standard fluorescent mounting medium containing 1 mg/ml p-phenylenediamine in 90% glycerol) was essentially as described in Hoepfner et al. (2000) and consisted of an imaging microscope (Axioplan 2; Carl Zeiss MicroImaging, Inc.) with plan Neofluar 100 \times Ph3 NA 1.3 objectives. It was equipped with a 75 WXBO and a 100 WHBO illumination source controlled by a shutter and filter wheel system (MAC2000; Ludl Electronics). The camera was a back-

illuminated cooled CCD camera (TE/CCD-1000PB; Princeton Instruments). The following filter sets for different fluorophores were used: No. 10 for AlexaFluor488 and No. 20 for rhodamine/AlexaFluor568 (Carl Zeiss MicroImaging, Inc.); and No. 41018 for GFP (Chroma Technology Corp.). The excitation intensity was controlled with different neutral density filters (Chroma Technology Corp.). MetaMorph 4.6r9 software (Universal Imaging Corp.) controlled the microscope, camera, and Ludl controller and was used for processing images. The microscope used for time-lapse images was a similarly equipped microscope (Axioplan 2; Carl Zeiss MicroImaging, Inc.) with a plan Neofluar 63 \times NA 1.25 or 100 \times (same as above) objective. The 100 \times objective was driven by a PIFOC P721.10 objective nanopositioner (Piezo device) to move quickly through multiple planes in the Z axis (Physik Instrumente). A Micromax back-illuminated CCD camera (512BFT; Princeton Instruments) was used for rapid acquisition and transfer of images. Stacks of tubulin images were processed using the “no-neighbors” deconvolution within MetaMorph, and, like all other fixed specimens, the Z-stacks were compressed using stack arithmetic, brightness and contrast were adjusted, and images were overlaid using color align.

For time-lapse videos, spores were grown 12–16 h in liquid media at 30 $^\circ$ under selective conditions to maintain the HHF1-GFP plasmid. Cells were washed and resuspended in 1/4 \times AFM + G418 + 1 \times glucose to limit fluorescent background from the media during time-lapse acquisition. Glass slides containing a concave well in the middle were filled with hot 1/4 \times AFM + 1 \times glucose + 1% agarose, which solidified and provided a growth surface for the prepared cells. 8 μ l of washed cells were placed on the agarose pad, and a coverslip was gently placed on top of the cell suspension. Slides were then placed in a moist chamber and incubated at 30 $^\circ$ C for 30 min to allow cells to adapt to the solid substrate.

Time-lapse conditions used XBO fluorescence at 5–10% intensity and exposures of 100 ms that were acquired every 30 s. For HHF1-GFP videos, a 63 \times objective was used, and a single-phase image with transmitted light was taken followed by 5 \times 0.5- μ m images in the Z axis with the EGFP filter and fluorescent light. For HHF1-GFP SPC42-RFP videos, a single Hhf1-GFP image was acquired followed by 16 \times 0.5- μ m images in the Z axis with the rhodamine filter to capture the dynamics of Spc42-RFP using the 100 \times objective. Focusing was automatically performed using the find focus function in MetaMorph software, and for HHF1-GFP SPC42-RFP videos, a Piezo motor directed the objective to allow more rapid movement through the Z axis than with the motorized stage. Images were acquired at a constant ambient temperature of 25–27 $^\circ$ C.

Time-lapse images were processed in MetaMorph to construct videos. Phase images were false colored red to provide an outline of the cell, and the GFP signal was colored green. Fluorescent images were flattened into a single plane, and phase and fluorescent images were assembled into separate stacks, which were then aligned using the color align function into a single stack. This then compressed the phase and fluorescent images into a single plane for each time point. This stack was then made into a video in which each image plays for 1/6 s.

Online supplemental material

Figs. S1 shows still images from a time-lapse video observing SPBs and nuclei. Fig. S2 shows the categorization of nuclei that lack detectable cyclin signal according to nuclear cycle stage. Fig. S3 shows that *AgCLB1/2* causes a transient delay in the budding yeast cell cycle. Table S1 presents the plasmids that were used in this study. Table S2 presents the oligonucleotide primers used in this study. Video 1 shows the dynamics and divisions of nuclei in a growing *A. gossypii* hyphal tip. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.200507003/DC1>.

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References

- Alberti-Segui, C., F. Dietrich, R. Altmann-Johl, D. Hoepfner, and P. Philippsen. 2001. Cytoplasmic dynein is required to oppose the force that moves nuclei towards the hyphal tip in the filamentous ascomycete *Ashbya gossypii*. *J. Cell Sci.* 114: 975–986.
- Altmann-Johl, R., and P. Philippsen. 1996. AgTHR4, a new selection marker for transformation of the filamentous fungus *Ashbya gossypii*, maps in a four-gene cluster that is conserved between *A. gossypii* and *Saccharomyces cerevisiae*. *Mol. Gen. Genet.* 250:69–80.
- Ayad-Durieux, Y., P. Knechtle, S. Goff, F. Dietrich, and P. Philippsen. 2000. A PAK-like protein kinase is required for maturation of young hyphae and septation in the filamentous ascomycete *Ashbya gossypii*. *J. Cell Sci.* 113:4563–4575.
- Belmont, A.S., and A.F. Straight. 1998. In vivo visualization of chromosomes using lac operator-repressor binding. *Trends Cell Biol.* 8:121–124.
- Chen, K.C., A. Csikasz-Nagy, B. Gyorfy, J. Val, B. Novak, and J.J. Tyson. 2000. Kinetic analysis of a molecular model of the budding yeast cell cycle. *Mol. Biol. Cell.* 11:369–391.
- Clutterbuck, A.J. 1970. Synchronous nuclear division and septation in *Aspergillus nidulans*. *J. Gen. Microbiol.* 60:133–135.
- Cross, F.R. 2003. Two redundant oscillatory mechanisms in the yeast cell cycle. *Dev. Cell.* 4:741–752.
- Cross, F.R., V. Archambault, M. Miller, and M. Klövstad. 2002. Testing a mathematical model of the yeast cell cycle. *Mol. Biol. Cell.* 13:52–70.
- Cross, F.R., L. Schroeder, M. Kruse, and K.C. Chen. 2005. Quantitative characterization of a mitotic cyclin threshold regulating exit from mitosis. *Mol. Biol. Cell.* 16:2129–2138.
- De Souza, C.P., A.H. Osmani, S.B. Hashmi, and S.A. Osmani. 2004. Partial nuclear pore complex disassembly during closed mitosis in *Aspergillus nidulans*. *Curr. Biol.* 14:1973–1984.
- Demeter, J., S.E. Lee, J.E. Haber, and T. Stearns. 2000. The DNA damage checkpoint signal in budding yeast is nuclear limited. *Mol. Cell.* 6:487–492.
- Dietrich, F.S., S. Voegeli, S. Brachat, A. Lerch, K. Gates, S. Steiner, C. Mohr, R. Pohlmann, P. Luedi, S. Choi, et al. 2004. The *Ashbya gossypii* genome as a tool for mapping the ancient *Saccharomyces cerevisiae* genome. *Science.* 304:304–307.
- Edgington, N.P., and B. Futcher. 2001. Relationship between the function and the location of G1 cyclins in *S. cerevisiae*. *J. Cell Sci.* 114:4599–4611.
- Glotzer, M., A. Murray, and M. Kirschner. 1991. Cyclin is degraded by the ubiquitin pathway. *Nature.* 349:132–138.
- Hendrickson, C., M.A. Meyn III, L. Morabito, and S.L. Holloway. 2001. The KEN box regulates Clb2 proteolysis in G1 and at the metaphase-to-anaphase transition. *Curr. Biol.* 11:1781–1787.
- Hinchliffe, E.H., E.A. Thompson, F.J. Miller, J. Yang, and G. Sluder. 1999. Nucleo-cytoplasmic interactions that control nuclear envelope breakdown and entry into mitosis in the sea urchin zygote. *J. Cell Sci.* 112:1139–1148.
- Hoepfner, D., A. Brachat, and P. Philippsen. 2000. Time-lapse video microscopy analysis reveals astral microtubule detachment in the yeast spindle pole mutant *cnm67*. *Mol. Biol. Cell.* 11:1197–1211.
- Hoepfner, D., F. Schaerer, A. Brachat, A. Wach, and P. Philippsen. 2002. Reorientation of mispositioned spindles in short astral microtubule mutant *spc72Delta* is dependent on spindle pole body outer plaque and Kar3 motor protein. *Mol. Biol. Cell.* 13:1366–1380.
- Holloway, S.L., M. Glotzer, R.W. King, and A.W. Murray. 1993. Anaphase is initiated by proteolysis rather than by the inactivation of maturation-promoting factor. *Cell.* 73:1393–1402.
- Hood, J.K., W.W. Hwang, and P.A. Silver. 2001. The *Saccharomyces cerevisiae* cyclin Clb2p is targeted to multiple subcellular locations by cis- and trans-acting determinants. *J. Cell Sci.* 114:589–597.
- Iborra, F.J., D.A. Jackson, and P.R. Cook. 2001. Coupled transcription and translation within nuclei of mammalian cells. *Science.* 293:1139–1142.
- Kellis, M., N. Patterson, M. Endrizzi, B. Birren, and E.S. Lander. 2003. Sequencing and comparison of yeast species to identify genes and regulatory elements. *Nature.* 423:241–254.
- Kellis, M., B.W. Birren, and E.S. Lander. 2004. Proof and evolutionary analysis of ancient genome duplication in the yeast *Saccharomyces cerevisiae*. *Nature.* 428:617–624.
- Knechtle, P. 2002. AgSPA2 and AgBOI control landmarks of filamentous growth in the filamentous ascomycete *Ashbya gossypii*. Ph.D. thesis. Biozentrum Universität Basel, Basel, Switzerland. 69 pp.
- Maekawa, H., T. Usui, M. Knop, and E. Schiebel. 2003. Yeast Cdk1 translocates to the plus end of cytoplasmic microtubules to regulate bud cortex interactions. *EMBO J.* 22:438–449.
- Makhnevych, T., C.P. Lusk, A.M. Anderson, J.D. Aitchison, and R.W. Wozniak. 2003. Cell cycle regulated transport controlled by alterations in the nuclear pore complex. *Cell.* 115:813–823.
- Minke, P.F., I.H. Lee, and M. Plamann. 1999. Microscopic analysis of *Neurospora* roopy mutants defective in nuclear distribution. *Fungal Genet. Biol.* 28:55–67.
- Murray, A.W. 2004. Recycling the cell cycle: cyclins revisited. *Cell.* 116:221–234.
- Nygaard, O.F., S. Guttes, and H.P. Rusch. 1960. Nucleic acid metabolism in a slime mold with synchronous mitosis. *Biochim. Biophys. Acta.* 38:298–306.
- Pfleger, C.M., and M.W. Kirschner. 2000. The KEN box: an APC recognition signal distinct from the D box targeted by Cdh1. *Genes Dev.* 14:655–665.
- Pomerening, J.R., E.D. Sontag, and J.E. Ferrell Jr. 2003. Building a cell cycle oscillator: hysteresis and bistability in the activation of Cdc2. *Nat. Cell Biol.* 5:346–351.
- Pomerening, J.R., S.Y. Kim, and J.E. Ferrell Jr. 2005. Systems-level dissection of the cell-cycle oscillator: bypassing positive feedback produces damped oscillations. *Cell.* 122:565–578.
- Pringle, J.R., A.E.M. Adams, D.G. Drubin, and B.K. Haarer. 1991. Immunofluorescence methods for yeast. *Methods Enzymol.* 194:565–602.
- Rao, P.N., and R.T. Johnson. 1970. Mammalian cell fusion: studies on the regulation of DNA synthesis and mitosis. *Nature.* 225:159–162.
- Rieder, C.L., A. Khodjakov, L.V. Paliulis, T.M. Fortier, R.W. Cole, and G. Sluder. 1997. Mitosis in vertebrate somatic cells with two spindles: implications for the metaphase/anaphase transition checkpoint and cleavage. *Proc. Natl. Acad. Sci. USA.* 94:5107–5112.
- Salama, S.R., K.B. Hendricks, and J. Thorner. 1994. G1 cyclin degradation: the PEST motif of yeast CLN2 is necessary, but not sufficient, for rapid protein turnover. *Mol. Cell. Biol.* 14:7953–7966.
- Sambrook, J., and D.W. Russell. 2001. *Molecular Cloning: A Laboratory Manual*. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
- Spellman, P.T., G. Sherlock, M.Q. Zhang, V.R. Iyer, K. Anders, M.B. Eisen, P.O. Brown, D. Botstein, and B. Futcher. 1998. Comprehensive identification of cell cycle-regulated genes of the yeast *Saccharomyces cerevisiae* by microarray hybridization. *Mol. Biol. Cell.* 9:3273–3297.
- Thornton, B.R., and D.P. Toczyski. 2003. Securin and B-cyclin/CDK are the only essential targets of the APC. *Nat. Cell Biol.* 5:1090–1094.
- Thornton, B.R., K.C. Chen, F.R. Cross, J.J. Tyson, and D.P. Toczyski. 2004. Cycling without the cytosome: modeling a yeast strain lacking the APC. *Cell Cycle.* 3: 629–633.
- Vodermaier, H.C. 2001. Cell cycle: waiters serving the destruction machinery. *Curr. Biol.* 11:R834–R837.
- Wendland, J., Y. Ayad-Durieux, P. Knechtle, C. Rebuschung, and P. Philippsen. 2000. PCR-based gene targeting in the filamentous fungus *Ashbya gossypii*. *Gene.* 242:381–391.
- Yamano, H., J. Gannon, and T. Hunt. 1996. The role of proteolysis in cell cycle progression in *Schizosaccharomyces pombe*. *EMBO J.* 15:5268–5279.