

CENP-A exceeds microtubule attachment sites in centromere clusters of both budding and fission yeast

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The stoichiometries of kinetochores and their constituent proteins in yeast and vertebrate cells were determined using the histone H3 variant CENP-A, known as Cse4 in budding yeast, as a counting standard. One Cse4-containing nucleosome exists in the centromere (CEN) of each chromosome, so it has been assumed that each anaphase CEN/kinetochore cluster contains 32 Cse4 molecules. We report that anaphase CEN clusters instead contained approximately fourfold more Cse4 in *Saccharomyces cerevisiae* and ~40-fold more CENP-A

(Cnp1) in *Schizosaccharomyces pombe* than predicted. These results suggest that the number of CENP-A molecules exceeds the number of kinetochore-microtubule (MT) attachment sites on each chromosome and that CENP-A is not the sole determinant of kinetochore assembly sites in either yeast. In addition, we show that fission yeast has enough Dam1–DASH complex for ring formation around attached MTs. The results of this study suggest the need for significant revision of existing CEN/kinetochore architectural models.

Introduction

Accurate protein counts are crucial for building structural and mathematical models and reconstituting multiprotein complexes. Quantitative fluorescence microscopy is used to determine stoichiometries in live cells (Hirschberg et al., 2000; Dunder et al., 2002; Wu and Pollard, 2005; Joglekar et al., 2006, 2008). For example, Wu and Pollard (2005) used immunoblotting to calibrate fluorescence intensity measurements of 28 cytoskeletal and signaling proteins in fission yeast. Joglekar et al. (2006) used an internal protein standard to count molecules in budding yeast kinetochores.

Centromeres (CENs) specify the sites to assemble kinetochores that mediate attachment of chromosomes to the spindle microtubules (MTs). Centromeric nucleosomes contain the histone H3 variant CENP-A/CenH3/Cse4 (Palmer et al., 1987; Sullivan et al., 1994; Stoler et al., 1995). In budding yeast, one MT attaches to each kinetochore (Winey et al., 1995) assembled at the point CEN (~120 bp). Cse4 is essential for kinetochore assembly (Collins et al., 2005). Chromatin immunoprecipitation (ChIP) shows that each CEN contains

only one Cse4 nucleosome (Furuyama and Biggins, 2007), whereas another study suggests a wider range of Cse4 binding (Riedel et al., 2006).

The CENs and kinetochores of all 16 chromosomes are clustered throughout the cell cycle in budding yeast. Assuming a Cse4 dimer at each nucleosome, an anaphase cluster is presumed to contain 32 Cse4 molecules (Joglekar et al., 2006; Camahort et al., 2009), although the nucleosome structure at CENs is still disputed (Dalal et al., 2007; Mizuguchi et al., 2007; Furuyama and Henikoff, 2009; Black and Cleveland, 2011; Xiao et al., 2011). This value was used as a standard to count the kinetochore proteins and to build an architectural model of kinetochores (Joglekar et al., 2006, 2009). However, whether anaphase clusters contain only CEN-associated Cse4 has not been evaluated (Westermann et al., 2007).

In contrast to budding yeast, fission yeast and most other eukaryotes have regional CENs that span several kilobases to megabases and contain many CENP-A nucleosomes (Partridge et al., 2000; Takahashi et al., 2000; Irvine et al., 2004). A major question is whether the kinetochore assembled on point CENs represents a conserved subunit of each kinetochore-MT attachment site in regional CENs. To address this

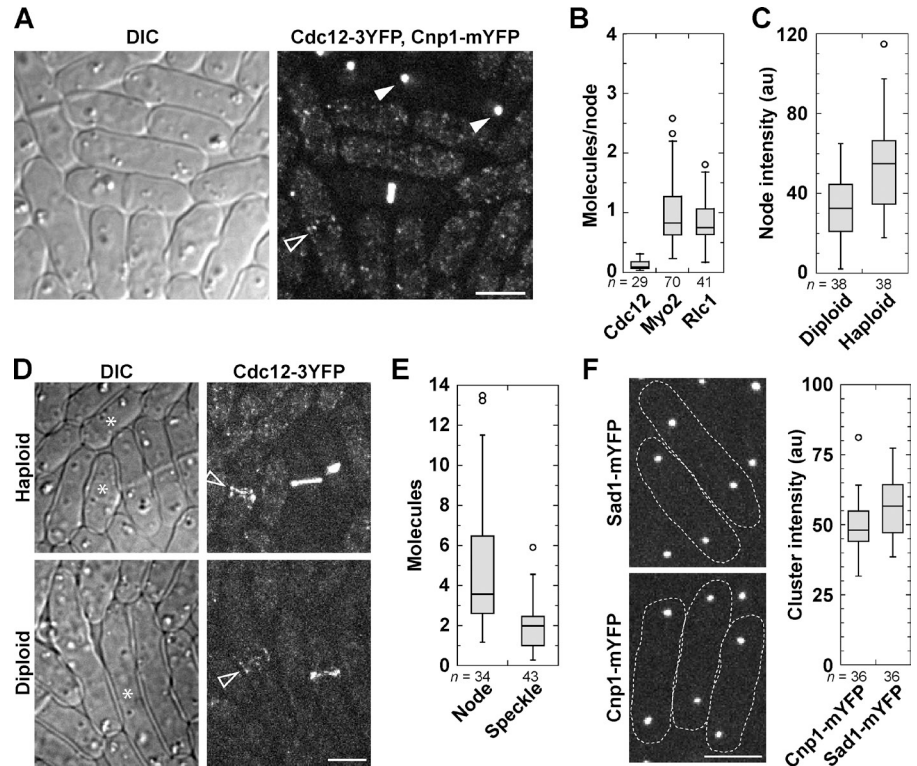
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Abbreviations used in this paper: CEN, centromere; ChIP, chromatin immunoprecipitation; DIC, differential interference contrast; FWHM, full width half maximum; mEGFP, monomeric EGFP; MNase, micrococcal nuclease; MT, microtubule; mYFP, monomeric YFP; ROI, region of interest; YPD, yeast peptone dextrose.

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Figure 1. Relative quantification of Cnp1 and *S. pombe* cytokinesis proteins. (A) Differential interference contrast (DIC) and fluorescence image of *S. pombe* cells showing Cdc12-3YFP nodes (strain KV346; open arrowhead) or Cnp1-mYFP anaphase clusters (strain JW1469; closed arrowheads). (B) Box plots (see Materials and methods) of the number of molecules in individual 3YFP- or mYFP-tagged cytokinesis nodes using Cnp1 as a standard (set at 15 Cnp1 molecules/anaphase cluster, $n = 60$ clusters; strain JW1469). From left to right, strains KV346, JW1110, and JW949 are shown. (C) Relative intensity of individual Cdc12-3YFP nodes in a heterozygous diploid strain and a haploid strain (JW1404-1). au, arbitrary unit. (D) DIC and fluorescence images of cells quantified in C. Arrowheads mark cells with nodes. Asterisks on the DIC images mark interphase cells with speckles (haploid) or without speckles (diploid). (E) Quantification of Cdc12-3YFP in nodes and speckles in haploid cells. (F) Micrographs and quantification of spindle pole body protein Sad1 (JW1141) and Cnp1 (JW1469) during anaphase. Dashed lines on micrographs show cell boundaries. Bars, 5 μ m.



question, Joglekar et al. (2008) compared kinetochore protein intensities in *Schizosaccharomyces pombe* with Cse4-GFP. They reported that the *S. pombe* CENP-A/Cnp1 is present in approximately two to three nucleosomes per CEN and concluded that numbers of CENP-A nucleosomes do not scale with CEN lengths. They also found that the ratio of kinetochore proteins is conserved on each MT attachment site (two to four per CEN in *S. pombe*) except for too few Dam1–DASH complexes to form a ring (Joglekar et al., 2008). Numerous other studies have also used a standard based on Cse4 to determine stoichiometry of cellular structures (Gardner et al., 2008; Moore et al., 2008; Yeh et al., 2008; Anderson et al., 2009; Markus et al., 2009; Ribeiro et al., 2009; Shimogawa et al., 2009; Tang et al., 2009; Thorpe et al., 2009; Gao et al., 2010; Johnston et al., 2010; Lin et al., 2010).

Here, we used independent standards to count molecules in CEN/kinetochore clusters. We found that each anaphase cluster contains 680 Cnp1 and 122 Cse4 molecules, much higher than the earlier studies. Thus, CENP-A copy number scales to CEN lengths in fission yeast and exceeds MT attachments in both yeasts. Our data present several challenges to the current CEN/kinetochore models.

Results and discussion

Visible cytokinesis nodes contain less than one Cdc12 molecule?

Cytokinesis nodes, discrete protein clusters at the equatorial plasma membrane during the G2/M transition (Fig. 1, A and D), are precursors of the contractile ring in fission yeast cytokinesis (Wu et al., 2006; Vavylonis et al., 2008; Coffman

et al., 2009; Laporte et al., 2011). We used the Wu and Pollard (2005) sum intensity method (see Materials and methods) to compare fluorescence intensity of cytokinesis node proteins with Cnp1 anaphase clusters. Based on the prediction of ~ 15 molecules of Cnp1 per cluster, each visible cytokinesis node contained <0.2 molecules of Cdc12 and ~ 1 molecule of myosin-II heavy chain Myo2 or its light chain, Rlc1 (Fig. 1, A and B). These results contrast the reported levels of ~ 4 molecules of Cdc12 and ~ 40 molecules of Myo2 and Rlc1 per cytokinesis node (Wu and Pollard, 2005) and suggest either severe quenching of tagged node proteins or more Cnp1 at anaphase clusters.

We hypothesized that if quenching could make several Cdc12 molecules appear to be <0.2 molecules, nodes in diploid cells expressing one tagged and one untagged copy of Cdc12 should display an intensity similar to nodes in haploids. Instead, nodes in diploids were about half as intense as in haploids (Fig. 1, C and D), suggesting that quenching was not significant. Cytoplasmic speckles of Cdc12 are the dimmest visible structure and are assumed to contain at least a dimer because formins dimerize with strong affinity (Moseley et al., 2004; Xu et al., 2004; Coffman et al., 2009). Consistent with this assumption, we observed fewer cytoplasmic speckles in the diploid cells (Fig. 1 D). Cdc12 nodes appeared two to three times as bright as Cdc12 speckles (Fig. 1 E), indicating at least four to six Cdc12 molecules per node (Laporte et al., 2011). Moreover, the intensities of Cnp1 clusters and Sad1 at spindle pole bodies were similar (Fig. 1 F). These data suggest >450 Cnp1 per cluster (Wu and Pollard, 2005) and indicate that another standard is needed to resolve the discrepancy.

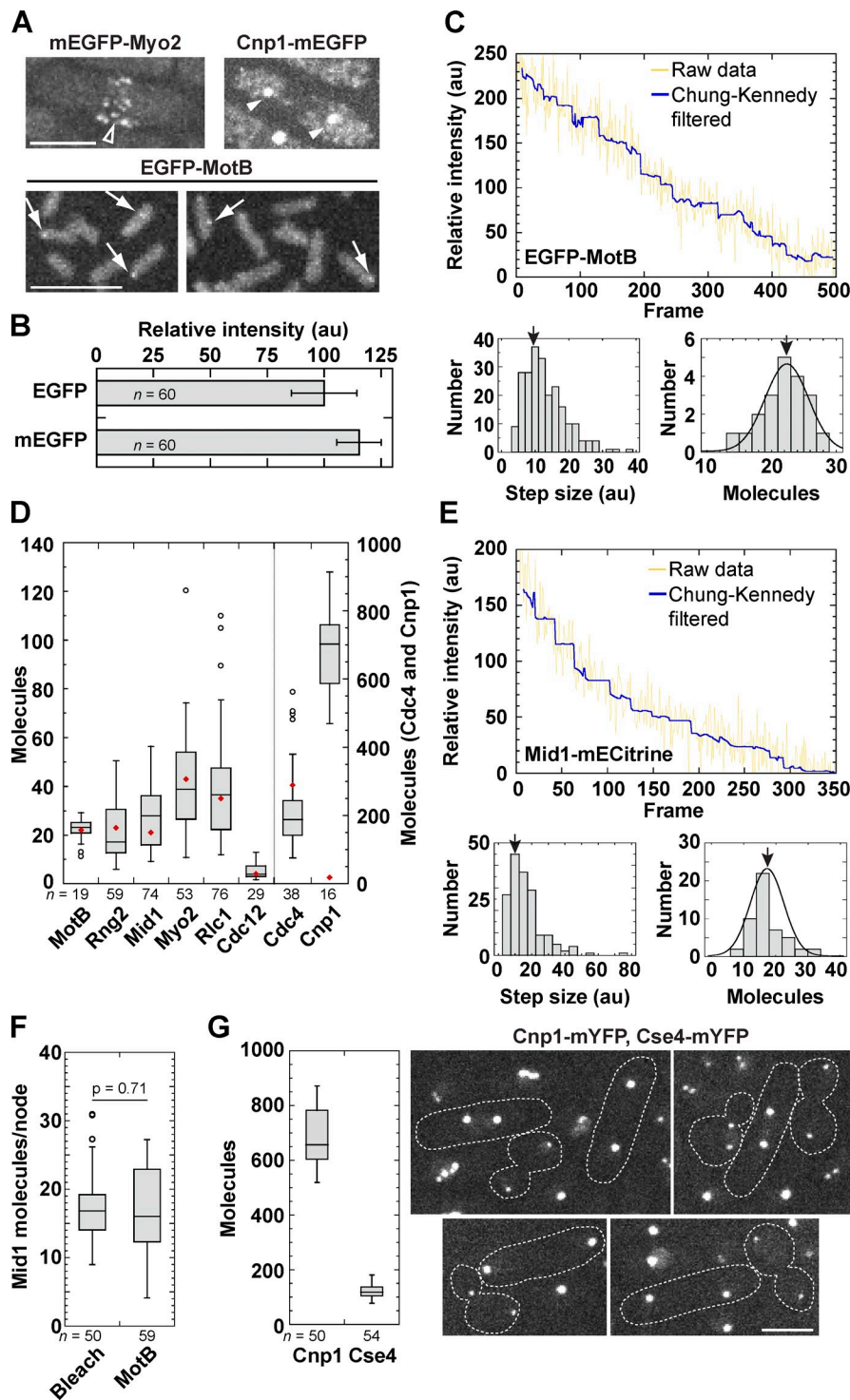


Figure 2. Quantification of node proteins and CENP-As using MotB as a reference. (A) Micrographs of Myo2 nodes (JW1109; open arrowhead), Cnp1 anaphase clusters (JW1470; closed arrowheads), and MotB motor of *E. coli* (JPA750; arrows) with the same imaging settings and contrast adjustment. (B) Comparison of EGFP (MLP198) and mEGFP (JW948) intensity (mean \pm SD). au, arbitrary unit. (C) Quantification of the intensity of a single EGFP molecule and MotB molecules in each motor using stepwise bleaching experiments. (top) A representative bleaching trace of EGFP-MotB. Each plateau ranges from 10 to 50 frames. The yellow line shows raw data after background subtraction, whereas the blue line shows Chung-Kennedy filtered data. (bottom left) The histogram shows the step sizes from 20 bleaching traces. (bottom right) The number of EGFP-MotB molecules per motor calculated from the measured step size. The arrows indicate the peak (9.52 units [left] and 22.5 ± 3.4 molecules [right]). (D) Molecules measured in individual cytokinesis nodes and Cnp1 anaphase clusters (JW1470) compared with the MotB motor. Red diamonds indicate published mean values (Wu and Pollard, 2005; Leake et al., 2006; Joglekar et al., 2008). (E) Quantification of Mid1-mECitrine (JW1790) in interphase nodes using stepwise bleaching as in C from eight bleaching traces. The arrows indicate the peak (left) or intensity ratio to the MotB motor (right). (F) Number of Mid1 molecules in interphase nodes using the bleaching method (left) or intensity ratio to the MotB motor (right). (G, left) Comparison of Cnp1 (JW1469) and Cse4 (JW2686-2) in anaphase clusters using MotB as a standard. (right) Cells expressing mYFP-tagged CENP-As in the same field for comparison. Dashed lines on micrographs show cell boundaries. Bars, 5 μ m.

Anaphase CEN clusters harbor additional CENP-A

The intensity of EGFP-MotB in *Escherichia coli* flagellar motors is 22 times the intensity of a single EGFP molecule, determined by stepwise bleaching (Leake et al., 2006). Cytokinesis nodes of monomeric EGFP (mEGFP)-Myo2 were similar to EGFP-MotB in intensity, whereas Cnp1-mEGFP was much brighter (Fig. 2 A). The mEGFP was 14% brighter than EGFP, and we used this number to correct our data (see Materials and

methods; Fig. 2 B). We repeated the bleaching experiments of Leake et al. (2006) to obtain the fluorescence intensity of each EGFP molecule. We calculated the average step size of intensity loss during bleaching and divided the initial intensity of the MotB motors by that value to find 22.5 ± 3.4 molecules per MotB motor (Fig. 2 C). Quantification of molecules in individual nodes of six cytokinesis proteins using MotB as a standard (Fig. 2 D) reproduced the previously estimated values (Wu and Pollard, 2005). However, each anaphase cluster had 680 ± 119 Cnp1-mEGFP

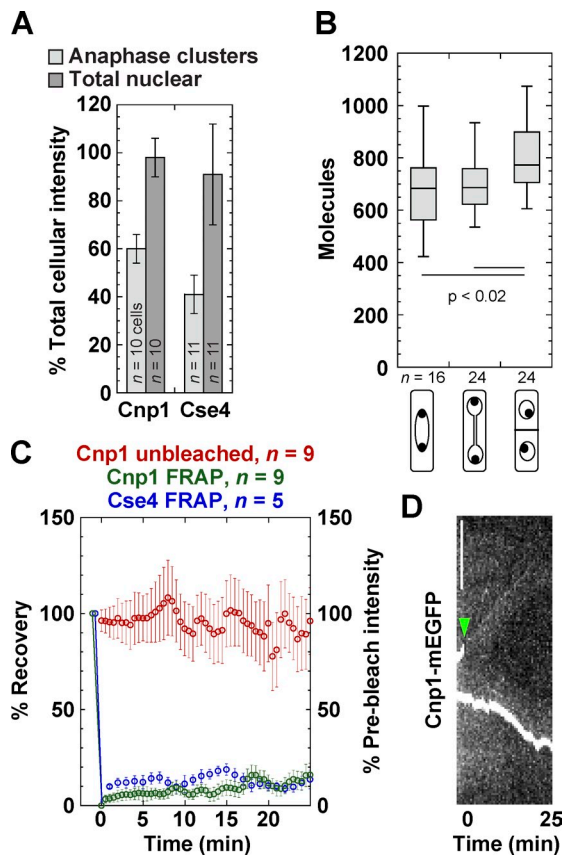


Figure 3. Cnp1 displays similar dynamics to Cse4 at anaphase clusters. (A) Percentage of total cellular CENP-A intensity (mean \pm SD) contained in anaphase clusters or the nucleus of *S. pombe* (JW1469) and *S. cerevisiae* (JW2687) cells. (B) Quantification of Cnp1 at early anaphase B, late anaphase, and G1/S phase (JW1469) with cellular features diagrammed below the graph (see Materials and methods). (C) FRAP is plotted as the percentage of recovery (left axis), whereas the unbleached cluster's intensity is plotted as the percentage of prebleach intensity (right axis; mean \pm SEM). One CEN cluster in each cell was bleached at early anaphase B, and the recovery was monitored with a 30-s delay for Cnp1 (JW1470) and 1 min for Cse4 (KBY7006). The change in intensity of the unbleached cluster was also monitored. (D) A kymograph of a representative bleached cell expressing Cnp1-mEGFP. The arrowhead marks the bleach point. Bar, 5 μ m.

molecules (Fig. 2 D), which is much higher than the reported \sim 15 Cnp1 molecules (Joglekar et al., 2008). Counting Mid1 by stepwise bleaching in *S. pombe* yielded a similar result to the ratio measurements (Fig. 2, E and F). When *S. pombe* and *Saccharomyces cerevisiae* were observed together, each anaphase cluster contained 680 ± 100 Cnp1 and 122 ± 24 Cse4 molecules, respectively (Fig. 2 G).

We explored the discrepancy with published ratios using molecular tools. PCR and sequencing revealed that the Cnp1-GFP(S65T) strain (YWY277) used in Joglekar et al. (2008) has *cnp1-GFP(S65T)* inserted at the Cnp1 locus upstream of the native gene that lacks the start codon (Fig. S1 A). We hypothesized that the native gene is translated from the ATG for amino acid 7. This untagged Cnp1 likely inserts at CENs in strain YWY277 based on the N-terminal truncation data (Takayama et al., 2008). In a strain (JW3523) with *cnp1-GFP(S65T)* completely replacing the native gene, anaphase clusters were 4.4-fold as bright as in YWY277 cells (Fig. S1, B and C).

These data and the new Cse4 copy numbers explain the \sim 40-fold underestimation of Cnp1 and show that Cnp1 occupancy is proportional to CEN sizes as observed in *Candida albicans* and vertebrates (Irvine et al., 2004; Joglekar et al., 2008; Sullivan et al., 2011).

Cnp1 and Cse4 display similar dynamics at anaphase

Next, we investigated whether all CENP-A molecules at anaphase clusters are stably associated with chromosomes. Cse4 is incorporated into CEN nucleosomes during DNA replication and remains stably bound throughout most of the cell cycle in budding yeast (Pearson et al., 2004; Camahort et al., 2007). Cnp1 anaphase clusters (two per cell) and nuclei contained 61 ± 6 and $98 \pm 8\%$ of total Cnp1 intensity, respectively, whereas Cse4 anaphase clusters and nuclei contained 41 ± 8 and $91 \pm 21\%$ of total Cse4, respectively (Fig. 3 A). These data suggest that molecules could be available in the nucleus to exchange with those in the anaphase clusters. Cnp1 intensity increased during septum formation (Fig. 3 B; Takayama et al., 2008), which began \sim 25 min after the start of anaphase (Wu et al., 2003). Cse4 recovered little in anaphase clusters, as reported (Pearson et al., 2004) using FRAP assays. Similarly, Cnp1 clusters bleached at early anaphase B did not recover much until \sim 25 min later, nor did the unbleached clusters lose intensity (Fig. 3, C and D). This large immobile fraction ($>80\%$) suggests that Cnp1 stably associates with nucleosomes and kinetochores.

Some Cse4 bound to 2- μ m circles and other locations outside CENs in budding yeast

Most laboratory budding yeast (but not fission yeast) strains contain \sim 60–100 parasitic plasmids per cell, called 2- μ m circles (2- μ m), that partition using Cse4, cohesin, and other proteins (Velmurugan et al., 2000; Mehta et al., 2002, 2005; Hajra et al., 2006; Ghosh et al., 2007). The plasmids incorporate Cse4 nucleosomes and cluster near CENs (Velmurugan et al., 2000; Hajra et al., 2006), but it is unknown how many Cse4 molecules bind to 2- μ m (Yeh and Bloom, 2006; Huang et al., 2011). Compared with 122 Cse4 molecules in YEF473a, the strain used above, each anaphase cluster contained 105 ± 13 Cse4 molecules in a *cir⁰* strain (cured of 2- μ m) and 121 ± 19 molecules in the isogenic *cir⁺* strain (Fig. 4, A and B). Thus, 2- μ m plasmids can bind \sim 16 Cse4 molecules.

In budding yeast, MTs are attached to kinetochores throughout the cell cycle except briefly at S phase (Pearson et al., 2004; Westermann et al., 2007). Thus, Cse4 remains bound to CEN DNA during G1 phase. Cells in G1 phase from asynchronous cultures or after α -factor treatment had \sim 40–50% Cse4 intensity in each CEN cluster compared with anaphase cells. The intensity was further reduced in prolonged G1 arrest (Fig. 4, C and D), suggesting that most Cse4 molecules are lost from the cluster in G1 phase.

Collectively, Cse4 molecules in each anaphase cluster are almost fourfold higher than the assumed 32 molecules (Joglekar et al., 2006). Some of the additional Cse4 associates with 2- μ m plasmids (Fig. 4). However, the sizeable discrepancy

suggests that the original conclusion of one Cse4 nucleosome per CEN may need further verification. This value derives from elegant ChIP experiments performed on micrococcal nuclease (MNase)-digested chromatin (Furuyama and Biggins, 2007). A critical assumption in these experiments is that terminal centromeric chromatin fragments generated by MNase digestion are mononucleosomes. The MNase-resistant fragment containing CEN DNA was >200 bp (Furuyama and Biggins, 2007), but recent biochemical analyses of nucleosomes constructed in vitro indicate that Cse4-containing nucleosomes only protect ~115–130-bp DNA (Cole et al., 2011; Dechassa et al., 2011; Kingston et al., 2011). Thus, the size of the native MNase-resistant centromeric fragment is similar to a Cse4-containing dinucleosome. In addition, these experiments did not include a cross-linking step, so some less stably associated Cse4 at or near the CEN could have been lost. Alternatively, the additional Cse4 may bind to DNA surrounding the CENs and/or proteins at kinetochores (Akey and Luger, 2003; Riedel et al., 2006; Yong-Gonzalez et al., 2007), and these Cse4 may dissociate during G1 (Fig. 4, C and D).

Fission yeast kinetochores have sufficient Dam1-DASH complex to form a ring

Because of the underestimation of Cse4, the copy numbers of kinetochore components need to be adjusted accordingly and the architecture revisited to account for the additional proteins. The Ndc80 and Dam1–DASH complexes cooperate to processively maintain the kinetochore-MT attachment and produce force (Lampert et al., 2010; Tien et al., 2010). The Ndc80 complex binds the inner kinetochore at one end and MTs at the other (Wei et al., 2005; McIntosh et al., 2008; Powers et al., 2009). In vitro, Dam1 complexes can assemble into MT-binding rings with 16-fold symmetry (Miranda et al., 2005; Westermann et al., 2005), which harness the forces produced from depolymerizing MT ends and function as processivity factors for kinetochore-MT binding (Grishchuk et al., 2005; Asbury et al., 2006; Efremov et al., 2007; Burrack et al., 2011).

Previous data indicated that the Dam1 complexes at *S. pombe* kinetochores are insufficient to form rings around the MTs (Joglekar et al., 2008). Calculations using our data yield enough Dam1 complexes at each MT attachment site to form one ring in fission yeast and several in budding yeast. Indeed, occupancy of Dam1 complex proteins was 120 ± 33 Dam1 and 141 ± 40 Ask1, whereas Ndc80 was 351 ± 49 at each anaphase cluster (Fig. 5, A and C). Dam1 and Ask1 also formed foci on the spindle with 14 ± 4 Dam1 molecules each (Fig. 5 B). As a control, the ratio between Cse4 and Cnp1 was unchanged (Fig. 5, A and C). *S. pombe* has three chromosomes with two to four MT attachment sites each. Thus, enough Dam1 complexes exist at kinetochore-MT attachment sites to form rings. Single Dam1 complexes and small oligomers can interact with MTs with load-bearing capacity, but the cooperative binding of Dam1 complexes suggests a tendency toward oligomerization (Gestaut et al., 2008; Grishchuk et al., 2008; Gao et al., 2010; Lampert et al., 2010; Tien et al., 2010). Dam1 complex rings have not been observed in vivo (McIntosh, 2005), so the structure of Dam1 complexes requires further studies.

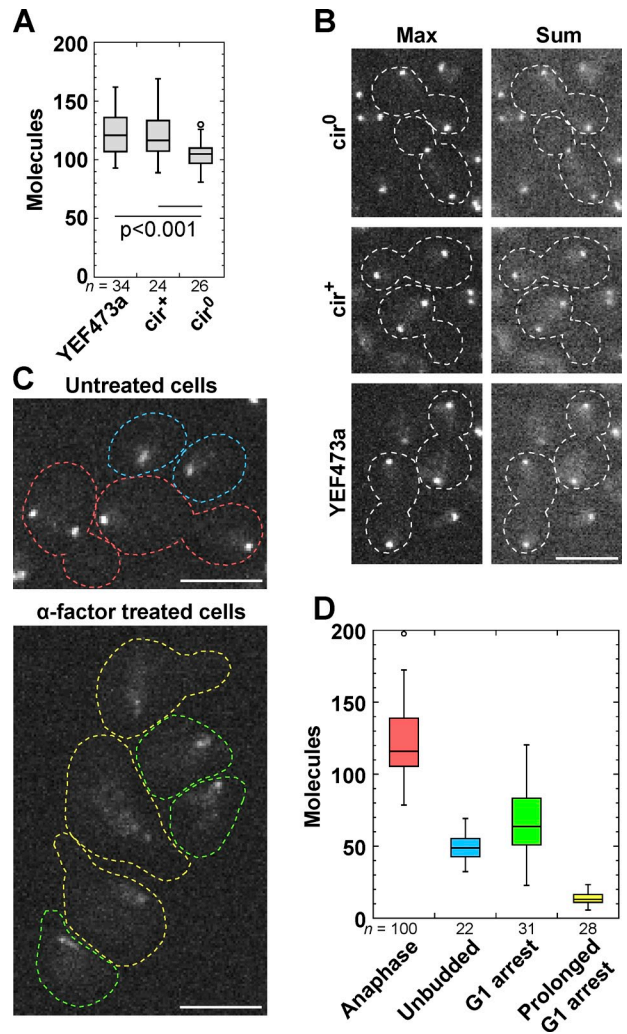


Figure 4. **Some Cse4 in *S. cerevisiae* binds to 2- μ m, and most is lost from the CEN cluster during G1 phase.** (A) Molecules in each anaphase cluster in Cse4-linker-mYFP strains with and without 2- μ m. From left to right, strains JW2687, JW2679, and JW2683 are shown. (B) Maximum and sum intensity projections of representative anaphase clusters of the strains in A. Dashed lines on micrographs show cell boundaries. (C) Cells expressing Cse4-linker-mYFP (JW2687) imaged with the same settings and contrast adjustments. (D) Molecules in each CEN cluster for cells, color coded as in C. Prolonged G1 arrest is defined as cells with a long shmoo. Bars, 5 μ m.

How to specify the sites for kinetochore assembly?

Our data raise several challenging questions. Where are the additional Cse4 and Cnp1 molecules located? As each chromosome has one to a few MT attachment sites, how do cells select kinetochore assembly sites among the CENP-A nucleosomes? Could kinetochores assemble on several CENP-A nucleosomes instead of only one, as proposed in previous models (Joglekar et al., 2006, 2008)? What are the architectures of kinetochores from yeast to vertebrate cells based on the revised numbers of molecules?

In fission yeast, a temperature-sensitive allele of Mis6, which is required to recruit Cnp1 to CENs, displays nearly 100% minichromosome loss (Takahashi et al., 1994, 2000). In contrast, Fft3 deletion causes a 2.7 \times reduction in Cnp1 binding and only 11% minichromosome loss (Strålfors et al., 2011),

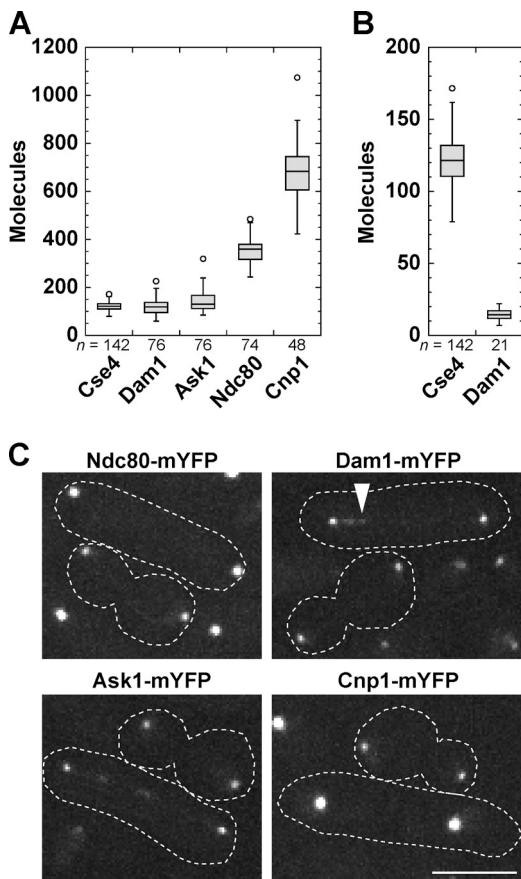


Figure 5. Fission yeast kinetochores have sufficient Dam1–DASH complex to form a ring. (A) Molecules of Dam1 (JW3667), Ask1 (JW3666), Ndc80 (JW3668) and Cnp1 (JW1469) in anaphase clusters using Cse4-mYFP in the same field as a standard. (B) Quantification of Dam1 foci along the spindle. (C) Representative images of budding and fission yeast anaphase cells used for quantification in A and B. The arrowhead marks a Dam1 structure as quantified in B. Dashed lines on micrographs show cell boundaries. Bar, 5 μ m.

suggesting that loss of some Cnp1 is tolerated. In human cells, 10% of the normal CENP-A level is sufficient to drive kinetochore assembly (Liu et al., 2006). Recent studies indicate that the constitutive CEN-associated network components CENP-C, -N, and -T are crucial for kinetochore assembly in fission yeast and other cells (Hori et al., 2008; Tanaka et al., 2009; Trazzi et al., 2009; Carroll et al., 2010; Gascoigne et al., 2011; Przewloka et al., 2011). In human and chicken cells, targeting CENP-C and -T to ectopic loci results in MT-attached kinetochores in the absence of CENP-A (Gascoigne et al., 2011). *Drosophila melanogaster* CENP-A and -C are interdependent for CEN localization (Erhardt et al., 2008). Based on these data, we hypothesize that a subset of CENP-A might recruit constitutive CEN-associated network components to assemble kinetochores in regional CENs. Distinguishing these assembly sites may be related to the mechanism of neo-CEN formation among ectopic CENP-A deposition sites (Heun et al., 2006; Zeitlin et al., 2009; Olszak et al., 2011). Thus, the CEN environment and interaction partners of CENP-As may both be necessary to direct kinetochore assembly at specific sites.

Collectively, our experiments challenge the idea that in yeast, each CENP-A nucleosome directs the assembly of one kinetochore-MT attachment. Instead, CENP-A occupancy exceeds the number of kinetochore-MT attachment sites in yeast as it does in higher eukaryotes. Our experiments also highlight the importance of understanding how kinetochore assembly sites are selected on CENP-A chromatin.

Materials and methods

Strains and growing conditions

Table S1 lists the *E. coli*, *S. pombe*, and *S. cerevisiae* strains used in this study. All tagged yeast genes were integrated at their native chromosomal loci under the control of endogenous promoters using pFA6a modules as previously described (Bähler et al., 1998; Longtine et al., 1998). The diploid strain expressing one copy of *cdc12-3YFP* was constructed using complementary *ade6* mutant alleles by crossing strain JW81 to JW1404-1 and maintained in YE5S– adenine using standard genetic methods (Moreno et al., 1991). We found that all the tags for Cnp1, Cse4, and kinetochore proteins were free of mutations by sequencing genomic DNA after gene targeting. The *cnp1* locus of strain YWY277 *cnp1-GFP(S65T)* was amplified and sequenced, which led to the discovery of the insertion and duplication at that locus (Fig. S1 A).

Because tagged histones are often less functional (Takayama et al., 2008), we tagged Cse4 in several strains using a flexible linker of 24 amino acids between the *cse4* ORF and the monomeric YFP (mYFP) tag (Sandblad et al., 2006). The linker made the tagged Cse4 strains healthier than strains without a linker from 25 to 32°C, although we observed no obvious difference in growth rates at 25°C (Fig. S1, D and E). The linker strains also had no more than a 10% difference in anaphase cluster intensity from those without it, suggesting that the functionality of Cse4 makes little difference in its incorporation at 25°C. The strains did not show increased sensitivity to 20 μ g/ml nocodazole (Fig. S1 E), suggesting that the tagged Cse4 is functional at 25°C but not fully functional at higher temperatures. Most of the tagged Cnp1 strains were functional, with the exceptions of the GFP(S65T)-tagged strains (Fig. S1 G), perhaps as a result of the thermosensitivity of GFP(S65T) itself. Cnp1 tagged at its N terminus under the endogenous promoter (*Pcnp1* from –717 to +6 bp with respect to the ATG) affects its expression and level at CENs either with or without the *kanMX6* marker (Fig. S1 F; Takayama et al., 2008).

Yeast cells were restreaked from –80°C stocks, grown 2–3 d on plates at 25°C, and then inoculated into 5–15 ml of liquid media. *S. pombe* cells were grown in YE5S and *S. cerevisiae* in yeast peptone dextrose (YPD) at 25°C. Cells were grown in liquid media at exponential phase for ~48 h before microscopy (Wu et al., 2006). *E. coli* cells were streaked from –80°C stock, grown overnight on lysogeny broth plates, and then inoculated into 10 ml of liquid tryptone broth. Liquid cultures were grown overnight in tryptone broth at 30°C and washed into motility buffer at room temperature (10 mM potassium phosphate and 0.1 mM EDTA, pH 7.0) for imaging (Leake et al., 2006).

Microscopy

Yeast cells for microscopy were collected from liquid cultures. For some experiments, cells of different strains were mixed together just before collection to image under identical conditions. Cells were centrifuged at 5,000 rpm for 30 s and then washed into EMM5S or synthetic dextrose medium for imaging. Live-cell microscopy of *S. pombe* was performed using a thin layer of EMM5S liquid medium with 20% gelatin and 0.1 mM *n*-propyl-gallate and observed at 23–25°C as previously described (Wu et al., 2006). Live-cell microscopy of *S. cerevisiae* was performed either on synthetic dextrose medium plus 2% agar pads or on EMM5S with 20% gelatin with *S. pombe* cells to prevent clumping while immobilizing the cells. For synchronization, *S. cerevisiae* cells were treated with 8 μ g/ml α -factor for 3 h (approximately one generation) at 25°C and imaged on agar pads containing 8 μ g/ml α -factor. *S. pombe* cells for FRAP experiments were treated with 20 mM hydroxyurea for 4 h at 25°C to synchronize in S phase and were then released for 3 h before FRAP to ensure that all bleached cells and cells used for corrections were at similar cell cycle phases. *E. coli* cells were observed on bare slides under a coverslip sealed with Valap to observe rotational motility and to verify the functionality of EGFP-MotB motors (Leake et al., 2006) and were then imaged immediately on motility buffer plus 20% gelatin to immobilize the cells for quantification. To calculate the full width

half maximum (FWHM), we imaged 0.1- μm fluorescent beads (TetraSpeck microspheres; Invitrogen) at 0.05- μm spacing and calculated the FWHM of a Gaussian fit to the point spread function in the z axis.

We used a 100x/1.4 NA Plan-Apo objective lens (Nikon) on a spinning-disk confocal microscope (UltraView ERS [PerkinElmer] with a CSU22 confocal head on an Eclipse TE2000-U microscope [Nikon]) with 488- and 514-nm argon ion lasers and a cooled charge-coupled device camera (ORCA-AG; Hamamatsu Photonics). Image analyses were performed using ImageJ (National Institutes of Health). Images of yeast and *E. coli* cells in figures are maximum intensity projections of z sections spaced at 0.4 μm unless otherwise noted. We found no obvious signal variation as a result of focal plane depth, as previously reported (Joglekar et al., 2006), using our imaging system, perhaps because our imaging starts away from the coverslip (Fig. S2 A).

Comparison of the methods for measuring fluorescence intensity

A sum of intensity from multiple z sections spaced at the FWHM of the point spread function along the z axis was consistent with immunoblotting (Wu and Pollard, 2005), whereas a comparison of only the maximum intensity section was used to count molecules in kinetochores (Joglekar et al., 2006). We tested whether there is a difference in the quantification using these two methods.

We determined that the maximum intensity section does not represent the same fraction of the total intensity for structures with different apparent sizes by comparing *S. pombe* cells expressing fluorescent-tagged cytokinesis node proteins or Cnp1 and *S. cerevisiae* cells expressing Cse4-GFP (Fig. S2, B and C). Under our imaging conditions, 90% of the intensity in the maximum intensity plane of each CENP-A anaphase cluster was contained in a 0.5- μm^2 (25 pixel) region of interest (ROI), whereas each node was contained in a 0.2- μm^2 (9 pixel) ROI. The maximum z section of each node protein contained 51–65% of the sum intensity (Fig. S2 C), whereas anaphase clusters of Cnp1 and Cse4 had 46–48% in the maximum section. Comparing the maximum section of node and CEN cluster intensities yields a different ratio of fluorescence than the sum intensity ratio (Fig. S2 D). Because the maximum section contains a different fraction of the total intensity, the sum method is more accurate for comparing structures with different apparent sizes. Thus, throughout the manuscript, the data represented are the sum intensity of all 0.4- μm -spaced z sections that contain intensity above background for a given structure except where noted.

Counting proteins by ratio fluorescence imaging and bleaching

Uneven illumination was corrected using images of purified 6His-mYFP solution as previously described (Wu and Pollard, 2005; Wu et al., 2008). In brief, at least 10 images of purified YFP were averaged after offset subtraction and then divided by the maximum value in the field using the ImageJ Math process to get an image with intensity values ranging from ~ 0.5 to 1. Offset-subtracted images of cells were divided by this correction image in ImageJ before measurements were taken. The offset is an image taken without laser lights. Background corrections were implemented as previously described (Wu and Pollard, 2005; Joglekar et al., 2006). In brief, an ROI size was chosen for each tagged protein that included at least 90% of the intensity (\pm two SDs of a Gaussian fit along two axes). The ROI sizes depended on the imaging conditions, but, in general, Cnp1 ROI was greater than Cse4 ROI, which was greater than cytokinesis node ROI. The same relationship exists between the numbers of z sections each protein occupied. Cnp1 and Cse4 were also compared with the same size ROI, and the ratio between their sum intensities was unchanged. Thus, as long as each ROI is chosen so that at least 90% of the intensity in the maximum plane is included, the ratios are consistent whether the ROIs are the same or different. The background intensity from a concentric ROI twice the size was calculated and subtracted from the measuring area in each imaging plane. All molecule number plots show the ratio of sum intensity except where noted. In box plots, the box contains the middle 50% of the data, the line in the box is the median, and circles above or below represent outliers. Outliers are defined by KaleidaGraph software based on the height of the box. Points $>1.5\times$ the box height above or below the box are defined as outliers.

The EGFP-tagged MotB structure in *E. coli* was verified to contain 22 molecules by imaging the cells in a single plane continuously with a 200-ms exposure for each frame and then quantifying the fluorescence intensity for each GFP molecule (Leake et al., 2006). Mid1-mECitrine nodes in interphase *S. pombe* cells were imaged continuously with a 100-ms exposure to verify the ratio comparison between fluorophores expressed in *E. coli* and *S. pombe*. The background-subtracted data were fit with a modified

Chung-Kennedy filter that preserves the edges of steps (Leake et al., 2006; Engel et al., 2009). In brief, starting at the tenth frame, the mean and SD of two consecutive sets of 10 points (up to and just after the tenth frame) were determined, and the mean of the set with the lower SD was used at that frame. Some of the beginning and ending data are lost in this filter. In essence, the filter is a rolling mean that does not cross step boundaries. Thus, a lower SD suggests that the dataset lies on a plateau, whereas a higher SD suggests that the dataset crosses a boundary and is not reported. On the plot of filtered data, the discrete changes in fluorescence are more obvious and thus easier to measure. The fluorescence of a single molecule is determined by fitting the step sizes to a γ distribution, and the starting intensity is divided by the step size to get the number of molecules.

Molecules in cytokinesis nodes and anaphase CEN/kinetochore clusters were counted based on fluorescence intensity. Z sections spaced at 0.4 μm (except where noted) were collected for strains expressing mYFP- or mEGFP-tagged proteins in *S. pombe* and *S. cerevisiae*. Intensities of cytokinesis node proteins were obtained from the sum of two to three consecutive sections and compared with the sum intensity of anaphase clusters in five to six consecutive sections to obtain the mean number of each protein in individual structures. Intensities of Cnp1-mEGFP in anaphase clusters and mEGFP-tagged node proteins (Mid1 [strain JW1088], Rng2 [JW1112], and Myo2 [JW1109]) were compared with the intensity of EGFP-MotB motors, with a slight correction for the difference in intensity between EGFP and mEGFP. Ratios between Cnp1 and the three node proteins were the same using both mEGFP and mYFP tags (Mid1 [strain JW1089], Rng2 [JW1761], and Myo2 [JW1763]), verifying the consistency of the method. Additional node proteins were measured using mYFP (Rlc1 [strain JW1757-2], Cdc12 [JW1404-1], and Cdc4 [JW1797-2]). The 3YFP intensity of Cdc12-3YFP is corrected to the intensity of mYFP for comparison (Wu and Pollard, 2005). Then, Cse4- and Cnp1-mEGFP strains were imaged under the same conditions to extrapolate the MotB standard for Cse4 using the Cnp1 value. The ratios between Cnp1 and Cse4 were the same for mYFP- and mEGFP-tagged proteins or by comparing only the best focal plane.

In Fig. 3 B, the cell cycle stages of the cells were estimated as follows from the left to right: early anaphase B, with two CEN clusters 3–6.5 μm apart; late anaphase, with clusters >6.5 μm apart; and G1/S phase, with a septum that has begun to form. The first two stages (without a septum) were used to quantify anaphase clusters and were not significantly different by the *t* test. The Student's two-tailed *t* test was used to calculate *p*-values.

FRAP analysis

FRAP was performed using the photokinesis unit on the (UltraView ERS) confocal system as previously described (Coffman et al., 2009; Laporte et al., 2011). In brief, the data were gathered using the Track-It function in the software (UltraView ERS) to find anaphase clusters in focus before bleaching. We collected at least three prebleach stacks (four z sections spaced at 0.3 μm) of anaphase clusters and 50 postbleach stacks at 30-s or 1-min intervals. For clusters that remained in focus during the movie, an ROI was selected at each site that was bleached $>50\%$ of the original signal. Unbleached cells at the same cell cycle stage as bleached cells were used for photobleaching corrections. Background corrections were performed using extracellular areas, and the data were normalized to the mean prebleach intensity set to 100% (Coffman et al., 2009). The intensity immediately after bleaching was set to 0% for the FRAP curves so that the graph shows the percentage of recovery. The plane with the maximum intensity from four sections was used for analysis at each time point, which is more accurate because the anaphase clusters do not always stay in the same focal plane. The unbleached Cnp1 cluster was also analyzed for loss of fluorescence because the *S. pombe* nuclei remained connected for most of the duration of our imaging. We attempted to fit the data to a single exponential curve equation given by $y = m_1 - m_2 \times \exp(-m_3 \times x)$, in which m_3 equals the off rate (Laporte et al., 2011), but the fit was not good as a result of the lack of recovery.

Online supplemental material

Fig. S1 shows that the abnormal *cnp1* locus affects the intensity of Cnp1-GFP anaphase CEN clusters in strain YWY277 and effects of different tags on CENP-A intensity and function. Fig. S2 shows a comparison of ratio measurement methods. Table S1 lists the *E. coli*, *S. pombe*, and *S. cerevisiae* strains used in this study. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201106078/DC1>.

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