

Kinetochores–microtubule error correction is driven by differentially regulated interaction modes

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Abstract

For proper chromosome segregation, sister kinetochores must interact with microtubules from opposite spindle poles (bi-orientation). To establish bi-orientation, aberrant kinetochores–microtubule attachments are disrupted (error correction) by Aurora B kinase (Ipl1 in budding yeast). Paradoxically, during this disruption, new attachments are still formed efficiently to allow fresh attempts at bi-orientation. How this is possible remains an enigma. Here we show that kinetochores attachment to the microtubule lattice (lateral attachment) is impervious to Aurora B regulation, but attachment to the microtubule plus-end (end-on attachment) is disrupted by this kinase. Thus, a new lateral attachment is formed without interference, then converted to end-on attachment and released if incorrect. This process continues until bi-orientation is established and stabilized by tension across sister kinetochores. We reveal how Aurora B specifically promotes disruption of the end-on attachment through phospho-regulation of kinetochores components Dam1 and Ndc80. Our results reveal fundamental mechanisms for promoting error correction for bi-orientation.

Introduction

Proper chromosome segregation during mitosis relies on correct kinetochore–microtubule (KT–MT) interaction¹. The KT initially interacts with the lateral surface of a single MT (lateral attachment) and is then tethered at the MT plus end (end-on attachment)²⁻⁴. Subsequently sister KTs attach to MTs extending from the opposite spindle poles, establishing chromosome bi-orientation. If an aberrant attachment is formed (Fig 1a, left), it must be removed by Aurora B kinase (Ipl1 in budding yeast), which phosphorylates KT components and disrupts the KT–MT interaction (red arrow)⁵⁻⁷. However, during this disruptive process, new KT–MT interactions can still be formed efficiently (Fig 1a, black arrows), promoting the KT–MT turnover. If bi-orientation is established and tension is applied, the KT–MT attachment is stabilized (Fig 1a, right), completing error correction. However it remains a mystery how, despite KT-MT attachments being weakened and disrupted by Aurora B (Fig 1a, red arrow), new attachments can be formed efficiently (black arrows), ensuring the KT–MT turnover. Here we address this question, using *Saccharomyces cerevisiae* as a model organism.

The Ndc80 and Dam1 complexes (Ndc80c and Dam1c) are outer KT components comprising the KT–MT interface in budding yeast⁸⁻¹¹. The Ndc80c is an integral KT component since before the initial KT capture by spindle MTs, and is subsequently required for both lateral and end-on MT attachment³. On the other hand, the Dam1c is not present on unattached KTs and shows a high accumulation at the MT end⁴. When the lateral attachment is converted to an end-on attachment, the Dam1c at the MT end interacts physically with Ndc80c, forming the KT–MT interface of the end-on attachment¹²⁻¹⁵. Nevertheless the Dam1c also localizes along the MT lattice^{4,16,17} and it is unknown whether this Dam1c fraction plays any role during lateral KT–MT attachment.

The Dam1c and Ndc80c components have been identified as Aurora B substrates whose phosphorylation is important for bi-orientation¹⁸⁻²⁰. Aurora B phosphorylates Dam1 at several residues, and clustered phosphorylation at the Dam1 C-terminus is crucial for bi-orientation¹⁸, and disrupts the Dam1c–Ndc80c and Dam1c–Dam1c interactions *in vitro*^{12,13,17,21}. The N-terminal ~100 residues (N-tail) of Ndc80, protruding from the calponin-homology domain (the essential MT-binding domain)^{22,23}, is also heavily phosphorylated by Aurora B, and this phosphorylation facilitates bi-orientation in both yeast and metazoan cells^{19,24,25}. The Ndc80 N-tail strengthens

the Ndc80c–MT association, and its phosphorylation diminishes this association *in vitro*^{15,26-28}. Here we study how the Dam1 C-terminus and the Ndc80 N-tail regulate KT–MT interaction *in vivo* and how their phosphorylation by Aurora B promotes error correction for bi-orientation.

Results

KTs repeatedly detach from spindle MTs and are quickly recaptured by them, when the Dam1 C-terminus and Ndc80 N-tail are deleted.

To investigate the roles of the Dam1 C-terminus and Ndc80 N-tail in KT–MT interaction *in vivo*, we generated their deletions (*dam1Δ206-343* [*dam1ΔC*] and *ndc80Δ1-112* [*ndc80ΔN*]). Although *ndc80ΔN* showed normal growth, *dam1ΔC* showed very slow growth, and their combination was lethal. To address the reason for this lethality, we created a conditional Dam1 C-terminus deletion by introducing TEV protease cleavage sites in the middle of Dam1 (Fig 1b). When the TEV protease was expressed, Dam1 was cleaved within two hours (Fig 1c). As expected, the Dam1 C-terminus cleavage (Dam1ΔCclv) showed very slow growth, and was lethal when combined with *ndc80ΔN* (Fig 1d). When a chosen *CEN* (*CEN5*) was visualized in the Dam1ΔCclv Ndc80ΔN double mutant, sister *CEN5*s often failed to separate on the metaphase spindle, in contrast to wild-type, implying a bi-orientation defect (Fig 1e,f). This defect was more frequent than in Ndc80ΔN and Dam1ΔCclv single mutants. Intriguingly, in the double mutant, non-separated *CEN5* sisters often detached from the metaphase spindle and were quickly recaptured by spindle MTs. This cycle was repeated in many cells (Fig 1e,f). When all KTs were visualized instead of *CEN5*, their detachment and recapture on the spindle were observed in most of the cells having both Dam1ΔCclv and Ndc80ΔN (Supplementary Fig 1). Thus, when both the Dam1 C-terminus and Ndc80 N-tail are deleted, bi-orientation becomes defective, and centromeres show repeated detachment from, and quick reattachment to, the metaphase spindle.

KTs interact normally with the MT lateral surface, when the Dam1 C-terminus and Ndc80 N-tail are deleted.

To characterize the above defects, we analysed individual KT–MT interactions, using an engineered assay system³. In this assay, KT assembly on *CEN3* was delayed by transcription from an adjacently inserted promoter²⁹. This increased the distance between *CEN3* and the mitotic spindle³ (Fig 2a). In wild-type control (*NDC80*⁺ *DAM1*⁺) cells, after *CEN3* was reactivated for KT assembly, *CEN3* was rapidly captured by the MT lateral surface (Fig 2c, decline of blue line). Soon after *CEN3* reached the spindle, its sisters separated, i.e. bi-orientation was established (Fig 2b, left; c, red line). In Dam1ΔCclv Ndc80ΔN cells, the initial *CEN3* capture by the MT

lateral surface occurred with normal kinetics (Fig 2f, decline of blue line; g), but subsequent bi-orientation establishment was substantially delayed (Fig 2b, right; f, red line; h). Instead, *CEN3* often detached from spindle MTs (Fig 2f, purple line showing cumulative percentage of detachment), followed by recapture by other MTs (not counted again as a fresh MT interaction in Fig 2f). *Dam1ΔCclv* cells also showed similar, but slightly milder, defects (Fig 2e,g,h). *Ndc80ΔN* cells showed only a slight delay in sister *CEN3* separation on the spindle (Fig 2d,g,h).

Thus, the lateral KT–MT attachment does not require the Dam1 C-terminus or the Ndc80 N-tail, although it relies on the calponin-homology domain of Ndc80^{15,23} (Supplementary Fig 2). Nonetheless, when both the Dam1 C-terminus and the Ndc80 N-tail are deleted, bi-orientation becomes defective and KTs detach from spindle MTs. Subsequently, KTs can interact efficiently with the MT lateral surface, which explains the rapid recapture of KTs by MTs after their detachment from the spindle (see Fig 1e).

The Ndc80 N-tail and the Dam1 C-terminus promote distinct steps in the end-on conversion, and together prevent KT detachment from the MT end.

To address how *Dam1ΔCclv Ndc80ΔN* causes KT detachment from spindle MTs, we investigated the change in *CEN3*–MT interaction following lateral attachment. During lateral attachment, the plus ends of depolymerizing MTs often reached *CEN3*. Then, in wild-type cells, *CEN3* became tethered at the MT end and was pulled towards a spindle pole as the MT depolymerized (end-on pulling), i.e. end-on attachment was established⁴ (Fig 3a,b,c). However, even in wild-type cells, the conversion from the lateral to end-on attachment (end-on conversion) often failed; then, the MT grew again (MT rescue) (Fig 3a,b,d), as previously reported³⁰. By contrast, in *Dam1ΔCclv Ndc80ΔN*, after the MT plus end reached *CEN3* on the MT lateral surface, the *CEN3* detached from the MT end (end-on drop-off) in majority (Fig 3a,b,f). *Ndc80ΔN* cells showed MT rescue more frequently than wild-type (Fig 3b). In *Dam1ΔCclv* cells, *CEN3* often localized continuously (> 5 min) at the MT end, but without subsequent MT depolymerisation (end-on standstill) (Fig 3a,b,e).

The different phenotypes of the *Ndc80ΔN* and *Dam1ΔCclv* single mutants can be explained if the Ndc80 N-tail and Dam1 C-terminus promote distinct steps in the end-on conversion. The Ndc80 N-tail could be involved in an early step of the end-on conversion; i.e. tethering the KT (e.g. on *CEN3*) at the end of a MT (see Fig 4e). This

would explain frequent MT rescue with Ndc80 Δ N, as MT rescue takes place when the end-on conversion is defective ³⁰. Meanwhile, the Dam1 C-terminus could facilitate completion of this conversion; i.e. allowing KTs tethered at the MT end to be pulled as the MT shrinks (end-on pulling; see Fig 4e). This would explain frequent end-on standstill with Dam1 Δ Cclv. The Dam1 C-terminus supports physical interaction with Ndc80 ¹³ (Fig 3g); this interaction likely contributes to complete the end-on conversion ¹⁴. On the other hand, the Dam1 C-terminus was dispensable for Dam1c assembly ¹⁷ (Fig 3g) and for Dam1c accumulation at the MT end (Fig 3h).

During the process towards lateral to end-on conversion, Dam1 Δ Cclv Ndc80 Δ N cells showed frequent *CEN3* detachment from the MT end. This suggests that these cells are defective in both the end-on conversion, and the failsafe mechanism that would prevent *CEN* detachment ³⁰. Otherwise, when the end-on conversion fails, it should lead to MT rescue, which would maintain lateral attachment and prevent KT detachment ³⁰ (see Fig 3d). Why is this failsafe mechanism defective in Dam1 Δ Cclv Ndc80 Δ N cells? The following evidence suggests that the Dam1 C-terminus is required for the MT rescue upon failure in the end-on conversion (see Fig 4e).

This MT rescue is promoted by Stu2 (the MT polymerase whose vertebrate orthologues are XMAP215 and ch-TOG ³¹), which is transferred from the KT to the MT end ³⁰ (Fig 4a, MT rescue at KT). We addressed whether the Dam1-C terminus could assist in this Stu2 function. Stu2 at the KT promotes MT rescue, not only upon the failure in the end-on conversion, but also during the lateral KT–MT attachment, during which Stu2 is occasionally transported from the KT along the MT, promoting MT rescue upon its arrival at the MT end ³⁰ (Fig 4a, MT rescue distal to KT; see Fig 4c). The latter provides a convenient test for possible Stu2 assistance by Dam1, because the MT rescue happens without the KT at the MT end (which would exclude involvement of other KT components) but Dam1c is still at the MT end ⁴ (see Fig 3h). In Dam1 wild-type cells, Stu2 was occasionally transported from a KT along the MT and, on reaching the MT plus end, it promoted MT rescue (Fig 4b,c) ³⁰. With Dam1 Δ Cclv, Stu2 was still normally transported from a KT to the MT end, but often failed to promote MT rescue on reaching the MT plus end (Fig 4b,d). Moreover, in a two-hybrid assay, Dam1 and Stu2 showed a physical interaction, relying on the Dam1 C-terminus (Supplementary Fig 3a,b). This suggests Dam1 C-terminus physically interacts with Stu2 and helps rescue a MT at its plus end. This would be the case not only for MT rescue distal to KT, but also for MT rescue at KT (see Fig

4a), as both are dependent on Stu2 accumulation at the MT end where Dam1c also accumulates⁴.

In summary, frequent KT detachment from the MT end in Ndc80ΔN Dam1ΔCclv is explained by distinct roles of the two domains in the end-on conversion (Fig 4e). In this mutant, the end-on conversion is not initiated efficiently but MT rescue often fails; i.e. both steps are often blocked, leading to frequent KT detachment from the MT end (Fig 4e).

Phospho-mimicking mutants of the Ndc80 N-tail and Dam1 C-terminus at Aurora B sites show similar phenotypes to their deletions

To address how Aurora B phosphorylation of the Dam1 C-terminus and the Ndc80 N-tail affects KT–MT interaction, we generated phospho-mimicking mutations at these domains (*dam1C-4D[AurB]* and *ndc80N-7D[AurB]*; see Fig 5a,b legends). *dam1C-4D[AurB]* showed very slow growth and *dam1C-4D[AurB] ndc80N-7D[AurB]* was lethal, although Dam1C-4D[AurB] was expressed similarly to Dam1 wild-type (Fig 5a). To maintain growth of these phospho-mimicking mutants, we integrated wild-type *DAM1* containing an auxin-inducible degron tag (*dam1-aid*). With auxin, Dam1-aid was degraded (Fig 5a), *dam1C-4D[AurB]* cells showed very retarded growth, and *dam1C-4D[AurB] ndc80N-7D[AurB]* cells were lethal (Fig 5b).

The *dam1C-4D[AurB] ndc80N-7D[AurB]* cells showed bi-orientation defects and repeated detachment of *CEN3* from the spindle (followed by recapture) (Fig 5c,d), similarly to Dam1ΔCclv Ndc80ΔN cells (see Fig 1e). The *dam1C-4D[AurB]* alone also showed such phenotypes, but less frequently (Fig 5d). The Dam1 C-terminus (and perhaps the Ndc80 N-tail) is normally dephosphorylated when bi-orientation is established and tension is applied³². However such de-phosphorylation is not recapitulated with phospho-mimicking mutations, which explains the bi-orientation defects and repeated detachment from the spindle with these mutations (Fig 5c,d).

In the assay shown in Fig 2a, *dam1C-4D[AurB] ndc80N-7D[AurB]* cells showed normal kinetics in lateral *CEN3*-MT interaction (Fig 6a,d; decline of blue line: e), but demonstrated both a delay in bi-orientation (Fig 6a,d; red line: f) and *CEN3* detachment from the spindle and the MT end (Fig 6d, purple line; g), again similarly to Dam1ΔCclv Ndc80ΔN cells. *dam1C-4D[AurB]* made greater contribution to these defects than did *ndc80N-7D[AurB]*, and the defects were enhanced in combination.

Moreover, Dam1C-4D[AurB] showed reduced interaction with Ndc80 (Fig 6h) and with Stu2 (Supplementary Fig 3c), as did Dam1 Δ C. Dam1C-4D[AurB] often failed to assist Stu2-dependent MT rescue (Supplementary Fig 3d), as did Dam1 Δ C. Thus, Aurora B phospho-mimicking mutants of the Dam1 C-terminus and the Ndc80 N-tail caused *CEN* detachment from the MT end, without affecting *CEN* interaction with the MT lateral surface, probably due to suppression of functions of these Dam1 and Ndc80 domains.

In addition to Aurora B, Mps1 kinase is also required for the error correction to achieve chromosome bi-orientation^{33,34}. The substrates of Mps1 for this process have not yet been identified in yeast. Nevertheless Mps1 phosphorylates the Dam1 C-terminus at different sites from Aurora B phosphorylation³⁵. We made and tested phospho-mimicking Dam1 mutants at Mps1 sites, but they did not show the above defects found with *dam1C-4D[AurB]* (Supplementary Fig 4).

Physiological Aurora B activity is sufficient to promote KT detachment from the MT plus end but does not affect the lateral attachment

In the above experiments, we used phospho-mimicking mutations of Dam1 and Ndc80 at Aurora B phosphorylation sites. These mutations may represent their hyper-phosphorylation states. Next we aimed to study whether the physiological Aurora B activity is sufficient to cause *CEN* detachment from the MT end. However, *CEN* didn't detach from the MT end during end-on pulling in Aurora B wild-type cells (see Fig 3c). We speculated KT detachment may occur from the MT end in Aurora B wild-type cells, but its detection may be difficult for two reasons: First, during end-on pulling, *CEN* is pulled towards a spindle pole with telomeres (on the same chromosome) trailing³. Here the viscosity of the nucleoplasm may impose a weak force onto the trailing chromosome arms, generating a weak tension on the KT–MT interface. In addition, once the *CEN* is on the spindle, chromosome arms may collide with other chromosome arms, making the *CEN* less mobile after detaching from the spindle, and thus *CEN* detachment difficult to detect. In either case, use of a minichromosome (MC) may change the outcome, as it has minimal chromosome arms. Second, when a KT detaches from the MT end, its sister KT may still maintain MT attachment; for example, its sister KT may attach to the lateral surface of the same MT, close to the MT end. If so, we may detect detachment of unreplicated *CEN* from the MT end.

To test these two possibilities, we investigated the behaviour of a circular MC (~18 kb) containing *CEN3*, after inhibiting DNA replication initiation by Cdc6 depletion. Intriguingly, the unreplicated MC showed a higher rate of detachment from the metaphase spindle, followed by quick recapture on it, than two controls; a replicated MC, and an unreplicated *CEN3* on chromosome *III* (Fig 7a,b). This suggests that, to detect the KT detachment with wild-type Aurora B, the KT should be on an unreplicated *CEN* that is on a MC. Next, to investigate MC interaction with individual MTs, we inactivated *CEN3* (under the *GAL* promoter) on the unreplicated MC, and then reactivated it during metaphase arrest (similarly to Fig 2a). The MC was caught efficiently on the MT lateral side (see Fig 7f, Aurora B wild-type), but showed frequent detachment from the MT plus end (Fig 7c,d). To address whether Aurora B causes this detachment, we depleted Aurora B using an auxin-dependent degron tag. Indeed, Aurora B depletion led to reduction of the MC detachment from the spindle when *CEN3* on the MC was always active (Fig 7e). When *CEN3* on the MC was inactivated and then reactivated, Aurora B depletion did not change the kinetics in the lateral KT–MT attachment (Fig 7f, decline of blue line), but suppressed MC detachment from the MT end (Fig 7f, purple line; g). Thus, physiological Aurora B activity is sufficient to promote KT detachment from the MT end, but does not affect the lateral attachment.

Physiological phosphorylation of the Ndc80 N-tail and Dam1 C-terminus by Aurora B promotes KT detachment from the MT plus end

To address the effect of physiological phosphorylation of the Dam1 C-terminus and Ndc80 N-tail by Aurora B, we generated non-phosphorylatable mutations of Dam1 and Ndc80 (*dam1C-4A[AurB]* and *ndc80N-7A[AurB]*; see Fig 8a legend). The *dam1C-4A[AurB] ndc80N-7A[AurB]* double mutant showed a severe growth defect, when Dam1-aid was depleted in the presence of auxin (Fig 8a; *dam1-aid* was integrated to maintain cell viability, as in Fig 5a,b). When a chosen *CEN* (*CEN3*) was visualized, in 45 % of *dam1C-4A[AurB] ndc80N-7A[AurB]* cells sister *CEN3*s stayed together without separation in the vicinity of one spindle pole in metaphase, suggesting a bi-orientation defect (Fig 8b). Thus, Aurora B phosphorylation of the Dam1 C-terminus and Ndc80 N-tail is essential for bi-orientation.

Next, using these mutants, we studied *CEN3* interaction with individual MTs in two experiments: First, we inactivated *CEN3* (using *GAL* promoter) on chromosome *III* and then reactivated it (as in Fig 2a). In *dam1C-4A[AurB] ndc80N-7A[AurB]* cells,

CEN3 was caught on the MT lateral side with normal kinetics but subsequent establishment of bi-orientation was slower ([Supplementary Fig 5](#)). Second, as in [Fig 7c](#), we depleted Cdc6 to inhibit DNA replication; and then inactivated *CEN3* on the MC, and subsequently reactivated it, to observe interaction of an unreplicated MC with MTs. *dam1C-4A[AurB] ndc80N-7A[AurB]* cells showed normal kinetics of lateral KT-MT interaction ([Fig 8c](#), decline of blue line) and suppressed MC detachment from the MT end ([Fig 8c](#), purple line; [d](#)). *dam1C-4A[AurB]* cells also suppressed MC detachment, albeit modestly ([Fig 8d](#)). Thus, the physiological level of Aurora B phosphorylation of the Dam1 C-terminus and Ndc80 N-tail can cause KT detachment from the MT plus end without changing the kinetics of lateral KT–MT interaction.

Discussion

Error correction of the KT–MT attachment requires its turnover, i.e. the disruption of aberrant KT–MT attachments and formation of new ones. The disruption of KT–MT attachment is promoted by Aurora B phosphorylation of KT components that weakens KT–MT association ^{1,20}. However, it has long been unclear how new attachments can still be formed, and errors corrected, despite attachment being weakened and disrupted by Aurora B. KT–MT interaction is initiated efficiently by lateral attachment ^{2,3}, which presents a larger contact surface than the MT end, to the KT. It appears that this mode of attachment is impervious to Aurora B regulation (Fig 8e). Lateral attachment is converted to end-on attachment (but not vice versa) ⁴; end-on attachment is then weakened by Aurora B phosphorylation of KT components (unless bi-orientation is established) leading to KT detachment from the MT end. Thus, the differential regulation of the KT–MT attachment modes promotes the turnover of KT–MT interaction.

How does Aurora B specifically disrupt end-on attachment without affecting lateral attachment? The Dam1 C-terminus and the Ndc80 N-tail are known Aurora B targets whose phosphorylation is important for bi-orientation ¹⁸⁻²⁰. These domains promote distinct steps in the conversion from the lateral to end-on attachment (see Fig 4e). Aurora B phosphorylation of these domains suppresses their functions, causing KT detachment from the MT plus end. The Dam1c accumulates at the MT end and this Dam1c fraction plays an important role in the end-on KT–MT interaction ^{4,12-15,36}. The Dam1c also localizes along the MT lattice ^{4,16,17} but this fraction seems important neither for lateral attachment nor for Aurora B-dependent KT–MT regulation. Indeed, neither the Dam1 C-terminus nor the Ndc80 N-tail is required for the lateral KT–MT attachment. Presumably, the calponin-homology domains of Ndc80c are sufficient for lateral attachment, which is not altered by Aurora B activity.

KT detachment from the MT end was demonstrated using Dam1 and Ndc80 phospho-mimicking mutants. Such detachment was also promoted by physiological phosphorylation of Dam1 and Ndc80, but it could only be detected when the KT was on an unreplicated MC (see Fig 7 a–d). Why do we need an unreplicated MC to observe KT detachment in physiological conditions? Presumably, in physiological conditions, the level of phosphorylation by Aurora B does not reach 100%, in contrast to a phospho-mimicking state; thus, to observe KT detachment, we need conditions, such as an unreplicated MC, where a lower phosphorylation level suffices KT

detachment. The level of physiological phosphorylation may be adjusted to prevent sister KTs from frequently undergoing simultaneous dissociation from the MT (a single MT in the above context and different MTs in syntelic attachment ¹). If so, this would explain how cells avoid generating many unattached KTs (i.e. both sister KTs unattached to MTs) by Aurora B activity, but rather use its activity for error correction by maintaining MT attachment at least to one sister KT.

During error correction of KT–MT interaction, the lateral attachment is formed without interference. But, when it is converted to end-on attachment, the KT is released from the MT end by the action of Aurora B if tension is not applied (low-tension state). This process may repeat but, if bi-orientation is established and tension is applied across sister KTs, the end-on attachment is stabilized (high-tension state). However, a transition from low- to high-tension state still remains a mystery. For example, stable KT–MT end-on attachment and tension are mutually dependent on each other. Thus, it seems difficult to acquire tension and stable attachment when neither of them is present (initiation problem of bi-orientation) ³⁷. Intriguingly, we find that conversion from lateral to end-on attachment may provide a solution to this problem (Zhang, et al, manuscript in preparation).

In metazoan cells, KT–MT interaction is also initiated by lateral attachment, which is then converted to end-on attachment ². Our conclusion that differential regulation of lateral and end-on attachment promotes error correction in budding yeast may be essentially applied in metazoan cells, although the Aurora B targets are not exactly the same ²⁰. In metazoan cells, the Ndc80 N-tail plays a more important role in facilitating the end-on attachment, and its phosphorylation seems more important for bi-orientation ^{23,24,27,38,39}, than it is in budding yeast ^{19,25,40}. The metazoan Ska1 complex (Ska1c), a functional equivalent to yeast Dam1c, also assists the end-on attachment, and its assembly is under negative regulation by Aurora B ⁴¹⁻⁴³. Given that Ndc80c and Ska1c do not seem to be involved in lateral KT–MT attachment ^{42,44,45}, we speculate that end-on, but not lateral, attachment is regulated by Aurora B in metazoan cells, similarly to budding yeast.

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Author contributions

M.K., E.K. and T.U.T. designed experiments and interpreted results. M.K. and E.K. performed experiments and analysed data. T.U.T, M.K., E.K. T.Z. and B.N. wrote the manuscript. A.M. gave technical support.

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Figure legends

Figure 1. The KT–MT attachments are turned over repeatedly when Dam1 C-terminus and Ndc80 N-tail are deleted.

(a) Diagram shows the process of error correction. To resolve aberrant KT–MT attachment (left), it must be weakened and disrupted by Aurora B phosphorylation of KT components (red arrow). This disruption is followed by formation of new KT–MT attachment (black arrows). Such turnover (i.e. disruption followed by new formation) of KT–MT attachment continues (green arrow) until bi-orientation is established and stabilized by tension across sister KTs (right).

(b) Schematic representation of Dam1 protein showing the position of integrated TEV cleavage sites.

(c) Western blotting (with a Dam1 antibody) showing Dam1 C-terminus cleavage by TEV protease. Cells were incubated for 2 h with or without galactose. A full scan of the western blot is shown in Supplementary Fig 6.

(d) Serially diluted cells were incubated with or without TEV protease expression. Glc, glucose. Gal, galactose.

(e, f) *NDC80⁺ DAM1⁺* (T7549), *ndc80ΔN* (T8053), *dam1-TEVsites* (producing Dam1ΔCclv; T8723) and *ndc80ΔN dam1-TEVsites* (T8725) cells with *P_{GAL}-TEV* (except for T8053) *CEN5-tetOs TetR-3×CFP Venus-TUB1 P_{MET3}-CDC20* were treated with α factor for 3 h and released to fresh media with methionine (for Cdc20 depletion), in the presence of galactose (for TEV protease expression). From 70 min after the release, images were acquired every minute. **e** shows a representative cell with Ndc80ΔN plus Dam1ΔCclv (0 min: start of image acquisition). **f** shows percentage of cells that showed sister *CEN5* non-separation (left) and *CEN5* detachment from the spindle (usually followed by reattachment; right); n= 45, 35, 87 and 38 cells were analyzed (from left to right). The rate of *CEN5* detachment was probably underestimated since it would be overlooked if *CEN5* were immediately recaptured by MTs after detachment. Experiments were performed three times (statistics source data are shown in Supplementary Table 2) and a representative experiment is shown here. *p*-values (two tailed) were obtained by Fisher's exact test.

Figure 2. The KT attaches normally to the MT lateral surface but detaches afterwards, when the Dam1 C-terminus and the Ndc80 N-tail are deleted.

(a) Experimental system to analyse the individual KT–MT interaction ³.

(b–h) *NDC80⁺ DAM1⁺* (T9162), *ndc80ΔN* (T8049), *dam1-TEVsites* (producing Dam1ΔCclv; T8921) and *ndc80ΔN dam1-TEVsites* (T8965) cells with *P_{GAL}-TEV* (except for T8049) *P_{GAL}-CEN3-tetOs TetR-3×CFP GFP-TUB1 P_{MET3}-CDC20* were treated with α factor for 3 h and released to fresh media with methionine (for Cdc20 depletion), in the presence of galactose (for TEV expression and *CEN3* inactivation). After 3 h, cells were suspended in medium with glucose (for *CEN3* reactivation) and images were acquired every 30 sec. **b** shows representative images acquired at 20 min after *CEN3* reactivation. Graphs show percentage of cells at each step of the KT–MT interaction (**c–f**); Purple lines (step 5) show cumulative percentage of *CEN3* detachment; *CEN3* detachment was followed by recapture by a MT, but such recapture was not included in counting for other steps. Percentage of cells at steps 1 and 4 are compared in four strains (**g** and **h**). n= 30 cells were analysed for each strain. Data represent one out of two independent experiments.

Figure 3. The KT often detaches from the MT end, when the Dam1 C-terminus and the Ndc80 N-tail are deleted.

(a) After the plus end of a shrinking MT had caught up with *CEN3*, four kinds of events were observed.

(b) Percentage of the events shown in **a**, in each strain. The four strains in [Fig 2b–h](#) were treated as in [Fig 2b–h](#), except that images were acquired every 10 sec. n= 27, 22, 27 and 28 events were analysed (from top to bottom). *p*-values (two tailed) were obtained by a chi-square test for trends. Data represent one out of two independent experiments.

(c–f) Examples of the end-on pulling (**c**, T9162), MT rescue (**d**, T8049), end-on standstill (**e**, T8921) and end-on drop-off (**f**, T8965), observed in **b**.

(g) Dam1 and Ndc80 interact physically in the two-hybrid assay, and this interaction requires the Dam1 C-terminus. Duo1 is a component of the Dam1 complex and serves as a control. Ras and Raf were also used as controls. AD, BD: fused with the transcription activation domain and the DNA-binding domain, respectively.

(h) The Dam1 C-terminus is dispensable for Dam1c accumulation at the MT end. *NDC80⁺ DAM1⁺* (T8875), *dam1-TEVsites* (producing Dam1ΔCclv; T8915) and *ndc80ΔN dam1-TEVsites* (T8916) cells with *P_{GAL}-TEV CEN5-tetOs TetR-3×CFP ASK1-4×mCherry Venus-TUB1* were treated with α factor for 3 h and released to fresh media in the presence of galactose (for TEV protease expression). Images were acquired 70 min after release. Ask1 is a component of Dam1c. Representative cells in metaphase (left). *CEN5* was on the spindle, not at the indicated MT end. Quantification of Ask1 signals at the MT ends (right). The maximum Ask1 signals at

the MT ends during image acquisition were quantified in $n = 10$ cells of each strain. Bars represent mean $\pm$ SE. Data represent one out of two independent experiments.

Figure 4. The Dam1 C-terminus helps Stu2 to rescue a MT.

(a) Diagrams illustrating three events following lateral KT–MT attachment; end-on attachment, MT rescue at the KT, and MT rescue distal to the KT. Relevant mechanisms are shown in pale blue ^{4, 30}.

(b–d) Dam1 C-terminus assists Stu2 in rescuing a MT after Stu2 is transported from a KT along the MT to the MT plus end, during the lateral KT–MT interaction. *DAM1*⁺ (T9229) and *dam1-TEVsites* *P*_{GAL}-TEV (producing Dam1 Δ Cclv; T9166) cells with *P*_{GAL}-CEN3-*tetOs TetR-3xCFP STU2-4xmCherry GFP-TUB1* *P*_{MET3}-CDC20 were treated as in Fig 2b–h. Images were acquired every 7 sec. Percentage of Stu2 transport events along a MT, leading, or not leading, to MT rescue; $n = 25$ and 37 Stu2 transport events were analysed for T9229 and T9166, respectively (b).

Representative examples of Stu2 transport along a MT, leading (c), or not leading (d), to MT rescue. In b, experiments were performed twice (statistic source data are shown in Supplementary Table 2), a representative experiment is shown here, and *p*-value (two tailed) was obtained by Fisher's exact test.

(e) Diagram illustrating the roles of the Ndc80 N-tail and the Dam1 C-terminus, in the conversion to end-on attachment and in MT rescue. The diagram explains the results in Fig 3 and 4, as follows: 1) with Ndc80 Δ N, the end-on conversion is not initiated efficiently, explaining frequent MT rescue; 2) with Dam1 Δ Cclv, the end-on conversion is initiated but not completed efficiently, i.e. end-on pulling often fails, accounting for the end-on standstill; 3) with Ndc80 Δ N plus Dam1 Δ Cclv, the end-on conversion is not initiated efficiently and MT rescue often fails; i.e. all subsequent steps are often blocked, leading to frequent KT detachment from the MT end.

Figure 5. Aurora B phospho-mimicking Dam1 and Ndc80 mutants show repeated turnover of KT–MT attachments.

(a) Western blotting (with Dam1 antibody) showing Dam1-aid depletion and Dam1 protein (wild-type or Dam1C-4D[AurB]) expression from an additional *DAM1* construct. Cells were incubated with or without auxin NAA for 4 h in methionine drop-out medium. In Dam1C-4D[AurB], four Aurora B target sites (S257, S265, S292 and S327 ¹⁸) were replaced with aspartates at the Dam1 C-terminus. A full scan of the western blot is shown in Supplementary Fig 6.

(b) Serially diluted cells were incubated with or without auxin NAA. In Ndc80 N-7D[AurB], seven Aurora B target sites (T21, S37, T54, T71, T74, S95 and S100¹⁹) were replaced with aspartates at the Ndc80 N-tail.

(c, d) *NDC80⁺ DAM1⁺* (T9530), *ndc80N-7D[AurB]* (T11102), *dam1C-4D[AurB] dam1-aid* (T11326) and *ndc80N-7D[AurB] dam1C-4D[AurB] dam1-aid* (T11452) cells with *TIR P_{GAL}-CEN3-tetOs TetR-3xCFP GFP-TUB1 P_{MET3}-CDC20* were treated with α factor for 3 h and released to fresh media with methionine (for Cdc20 depletion) in the presence of NAA (to deplete Dam1-aid) and glucose (*CEN3* under the *GAL* promoter was always active). From 90 min after release, images were acquired every 30 sec. **c** shows a representative cell with *ndc80N-7D[AurB] dam1C-4D[AurB]* (0 sec: start of image acquisition). **d** displays percentage of cells showing sister *CEN3* non-separation (left) and *CEN3* detachment from the spindle (usually followed by reattachment; right); n= 62, 48, 75 and 56 cells were analyzed (from left to right). Experiments were performed twice (statistics source data are shown in Supplementary Table 2) and a representative experiment is shown here. *p*-values (two tailed) were obtained by Fisher's exact test.

Figure 6. Aurora B phospho-mimicking Dam1 and Ndc80 mutants show normal lateral KT–MT attachment, but subsequent detachment from the MT end.

(a–g) The cells shown in Fig 5c, d were treated as in Figs 2b–h and 3b (*CEN3* under the *GAL* promoter was inactivated and then reactivated), except that NAA was added 30 min before image acquisition. **a–f** shows percentage of cells at each step of KT–MT interaction (as in Fig 2c–h). n= 25, 25, 30 and 30 cells were analysed in **a–d**, respectively. Data represent one out of two independent experiments. **g** shows an example of *CEN3* detachment from the MT end in an *ndc80N-7D[AurB] dam1C-4D[AurB]* cell.

(h) The Dam1 and Ndc80 interaction, detected by the two-hybrid assay, is disrupted by Aurora B phospho-mimicking mutations at the Dam1 C-terminus (Dam1C-4D[AurB]). Duo1 is a component of the Dam1c.

Figure 7. Aurora B facilitates detachment of an unreplicated minichromosome from the MT end without changing kinetics of its lateral MT attachment.

(a, b) An unreplicated minichromosome (MC) frequently shows detachment from, and recapture on, the spindle. *cdc6-anchor-away TetR-3xCFP GFP-TUB1 P_{MET3}-CDC20* cells with pT111 (MC with *P_{GAL}-CEN3-tetOs*) (T11561) or *P_{GAL}-CEN3-tetOs* on chromosome III (T11629) were arrested in metaphase after Cdc6 depletion.

Incubation with glucose kept *CEN3* always active. Cells, like T11561, but with wild-type *CDC6* (T11562) were also analyzed. Images were acquired every 10 sec. **a** shows a representative T11561 cell. In **b**, *n* = 51, 48 and 50 cells were analysed (left to right).

(c, d) An unreplicated MC frequently shows detachment from the MT plus end. The cells in **b** were treated as above, but *CEN3* under the *GAL* promoter was inactivated and subsequently reactivated. **c** shows a representative T11561 cell. In **d**, *n* = 11, 21 and 26 end-on attachment events were analysed (left to right).

(e) Aurora B (*Ipl1*) depletion reduces detachment of an unreplicated MC from the spindle. *IPL1* wild-type (T11695) and *ipl1-321-aid* (T11694) cells with *TIR* (otherwise like T11651; see **a**), were treated as in **a** (*CEN3* was always active), except that NAA was added to deplete *Ipl1*; *n* = 68 and 48 cells were analysed for T11695 and T11694, respectively.

(f, g) Aurora B depletion reduces detachment of an unreplicated MC from the MT end without changing the kinetics of lateral MT attachment. T11695 and T11694 cells were treated as in **c** (*CEN3* was inactivated and reactivated), except that NAA was added as in **e**. **f** shows percentage of cells at each step of KT–MT interaction. Purple lines show cumulative percentages of detachment from the spindle or the individual MTs (subsequent MT re-attachment was not counted). **g** shows percentage of MC detachment from the MT ends during end-on attachment. *n* = 31 and 31 cells (**f**) and *n* = 29 and 11 end-on pulling events (**g**) were analysed for T11695 and T11694, respectively. In **b**, **d**, **e** and **g**, experiments were performed twice (statistics source data are shown in Supplementary Table 2) and a representative experiment is shown here. *p*-values (two tailed) were obtained by Fisher's exact test.

Figure 8. Non-phosphorylatable Dam1 and Ndc80 mutants at Aurora B sites reduce the detachment rate of a KT from the MT plus end.

(a) Serially diluted cells were incubated with or without auxin NAA. In *Dam1C-4A[AurB]*, four Aurora B target sites (S257, S265, S292 and S327¹⁸) were replaced with alanines in the *Dam1* C-terminus. In *Ndc80N-7A[AurB]*, seven Aurora B target sites (T21, S37, T54, T71, T74, S95 and S100¹⁹) were replaced with alanines at the *Ndc80* N-tail.

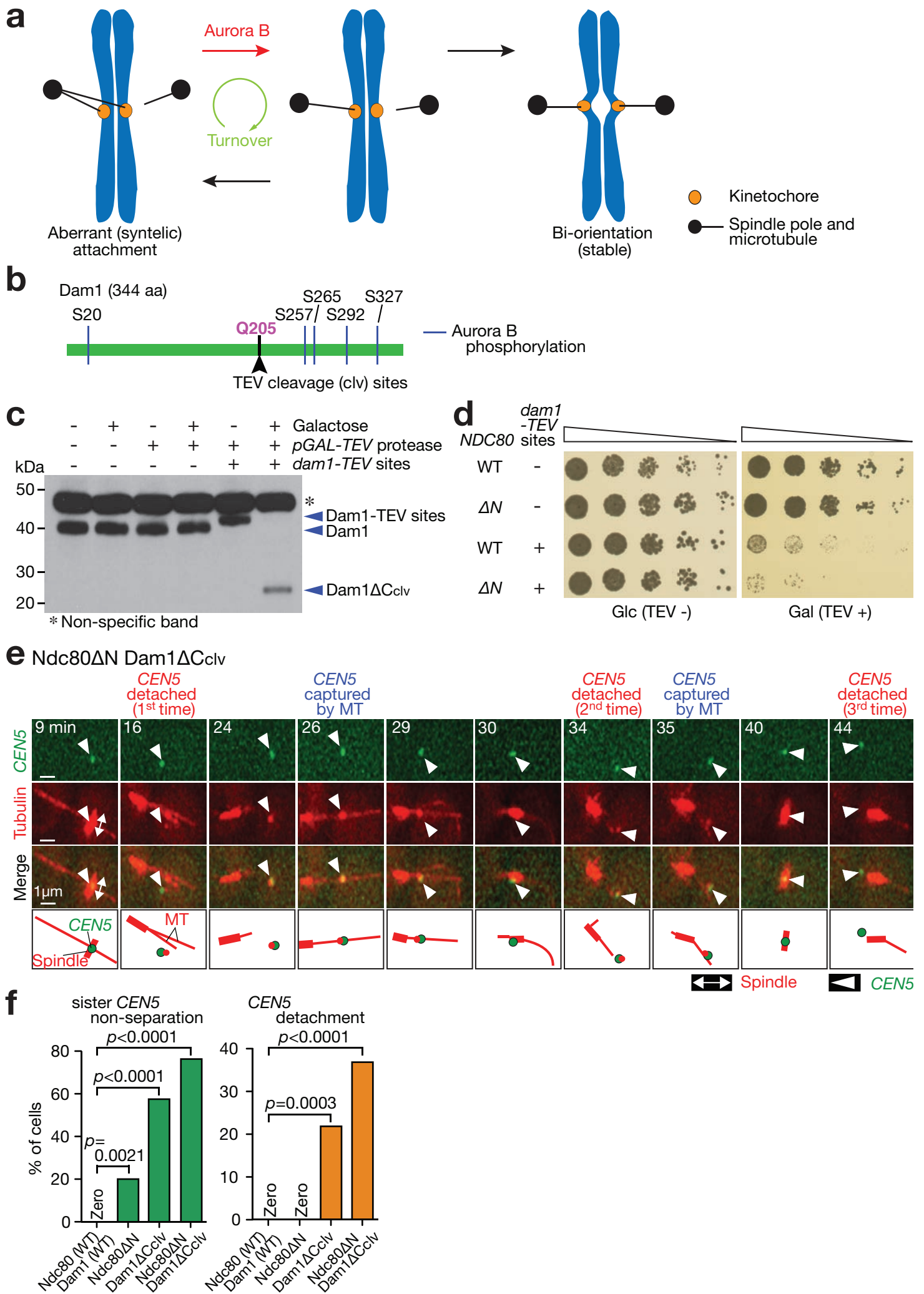
(b) *NDC80⁺ DAM1⁺* (T10638), *ndc80N-7A[AurB]* (T11813), *dam1C-4A[AurB] dam1-aid* (T11795) and *ndc80N-7A[AurB] dam1C-4A[AurB] dam1-aid* (T11794) cells with *TIR* (except for T11813) *P_{GAL}-CEN3-tetOs TetR-3×CFP GFP-TUB1 P_{MET3}-CDC20* were treated and analysed as in Fig 5c; i.e. arrested in metaphase with glucose (*CEN3* was always active). Representative images (of T10638 and T11794) and

percentages of sister *CEN3* non-separation (n= 39, 61, 45, 31 cells were analysed, respectively) are shown.

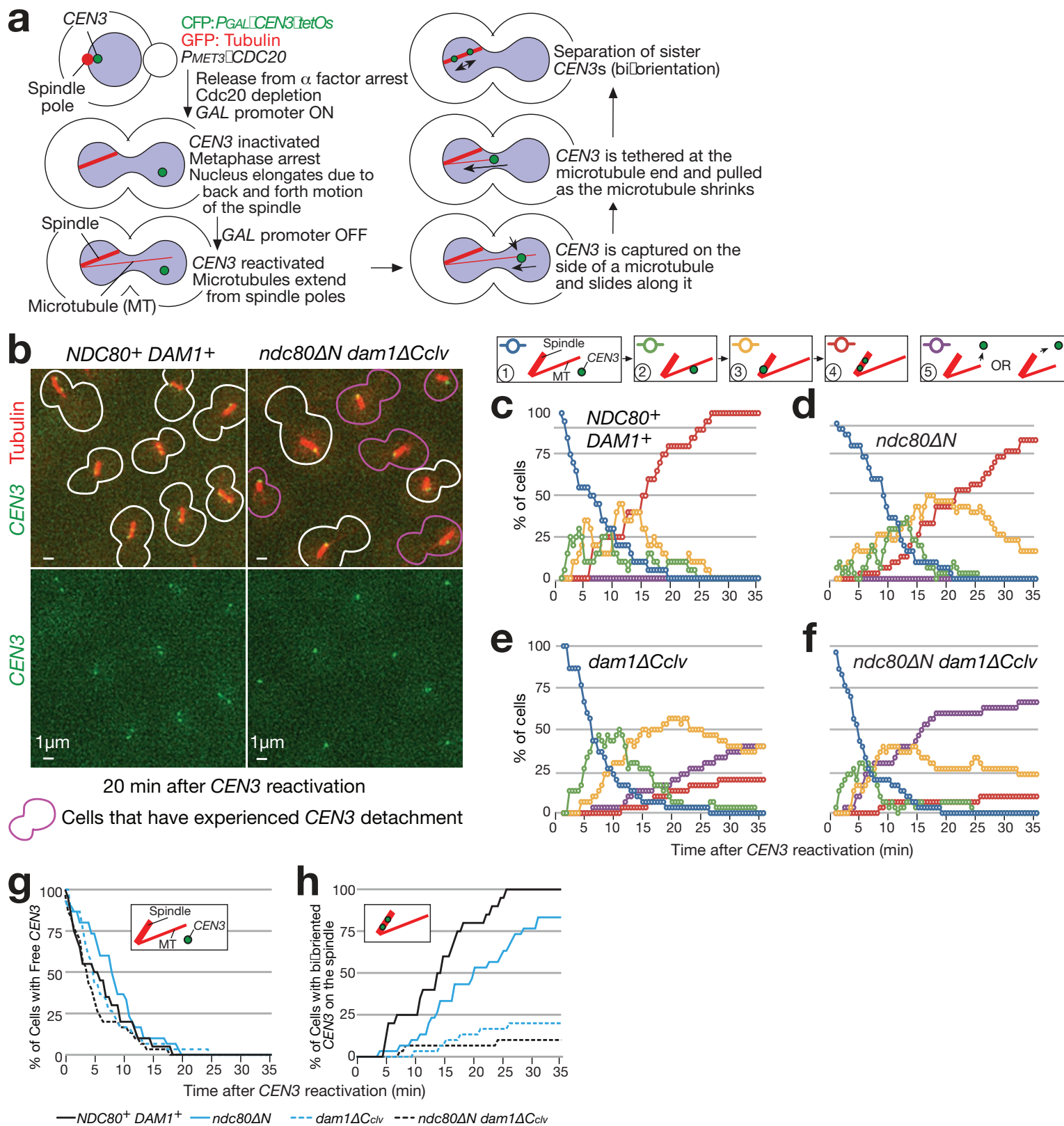
(c, d) *NDC80*⁺ *DAM1*⁺ (T11675), *ndc80N-7A[AurB]* (T11788), *dam1C-4A[AurB]* (T11790) and *ndc80N-7A[AurB] dam1C-4A[AurB]* (T11811) with *TIR dam1-aid* (except for T11788) *cdc6-anchor-away TetR-3xCFP GFP-TUB1 P_{MET3}-CDC20* and pT111 (MC with *P_{GAL}-CEN3-tetOs*) were treated as in Fig 7f,g; i.e. Cdc6 was depleted, *CEN3* under the *GAL* promoter was inactivated and subsequently reactivated, and Dam1-aid was depleted. c shows percentage of cells at each step of KT–MT interaction, as in Fig 7f. d shows percentages of MC detachment from the MT ends during end-on attachment. n= 24 and 23 cells (c) and n= 25, 24, 16 and 14 end-on pulling events (d) were analysed (left to right).

In b and d, experiments were performed three times or twice, respectively (statistics source data in Supplementary Table 2), and a representative experiment is shown here. *p*-values (two tailed) were obtained by Fisher's exact test.

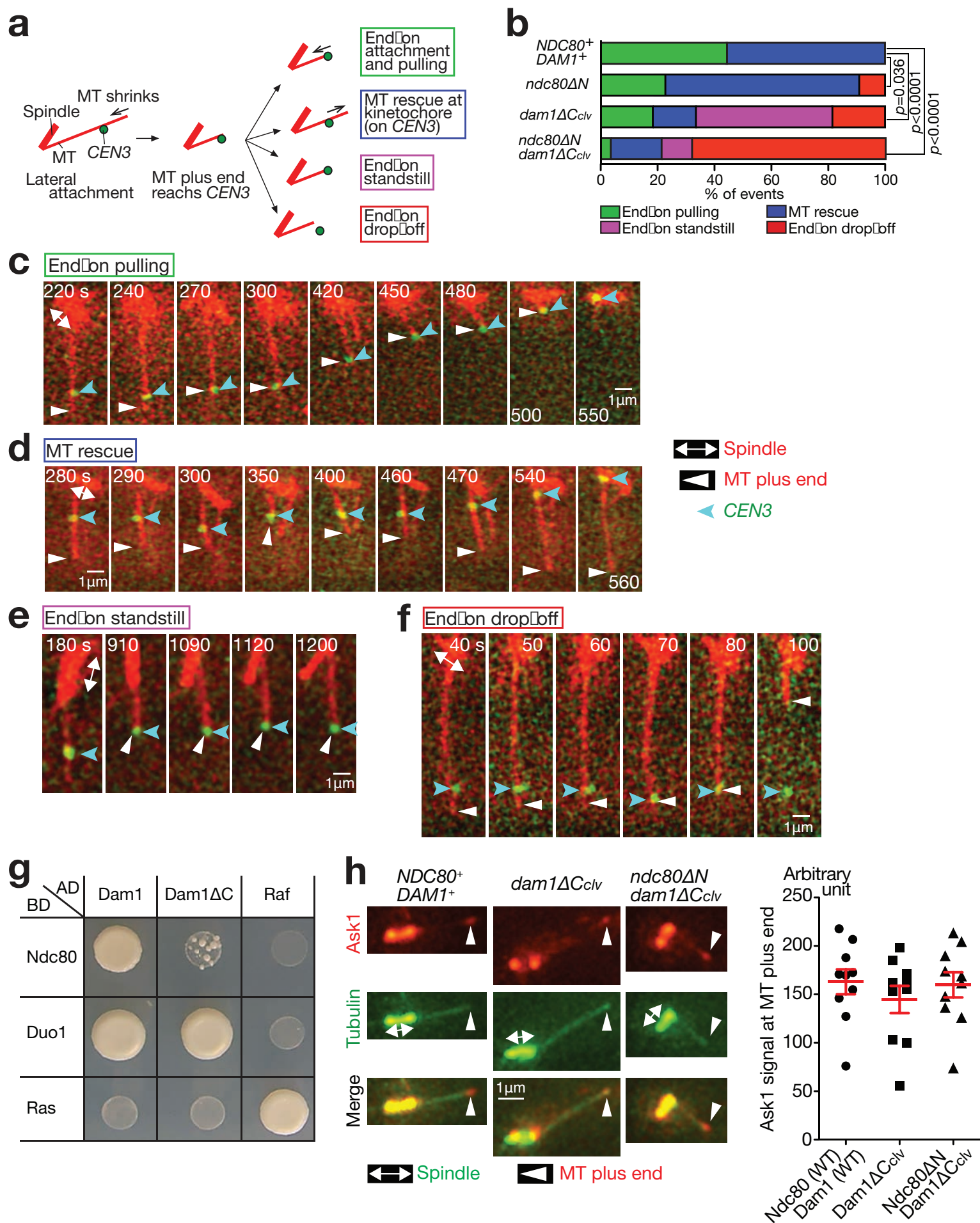
(e) Summary diagram. Aberrant KT–MT attachment (left) is dissolved through disruption of end-on attachment by Aurora B phosphorylation of KT components such as Dam1 and Ndc80. Subsequently lateral attachment is formed efficiently as this form of attachment is impervious to Aurora B regulation (insets). If this leads to establishment of bi-orientation (right), tension is applied across sister KTs and KT–MT attachment is stabilized. Alternatively, if that leads to aberrant attachment (left), error correction must continue to establish bi-orientation.



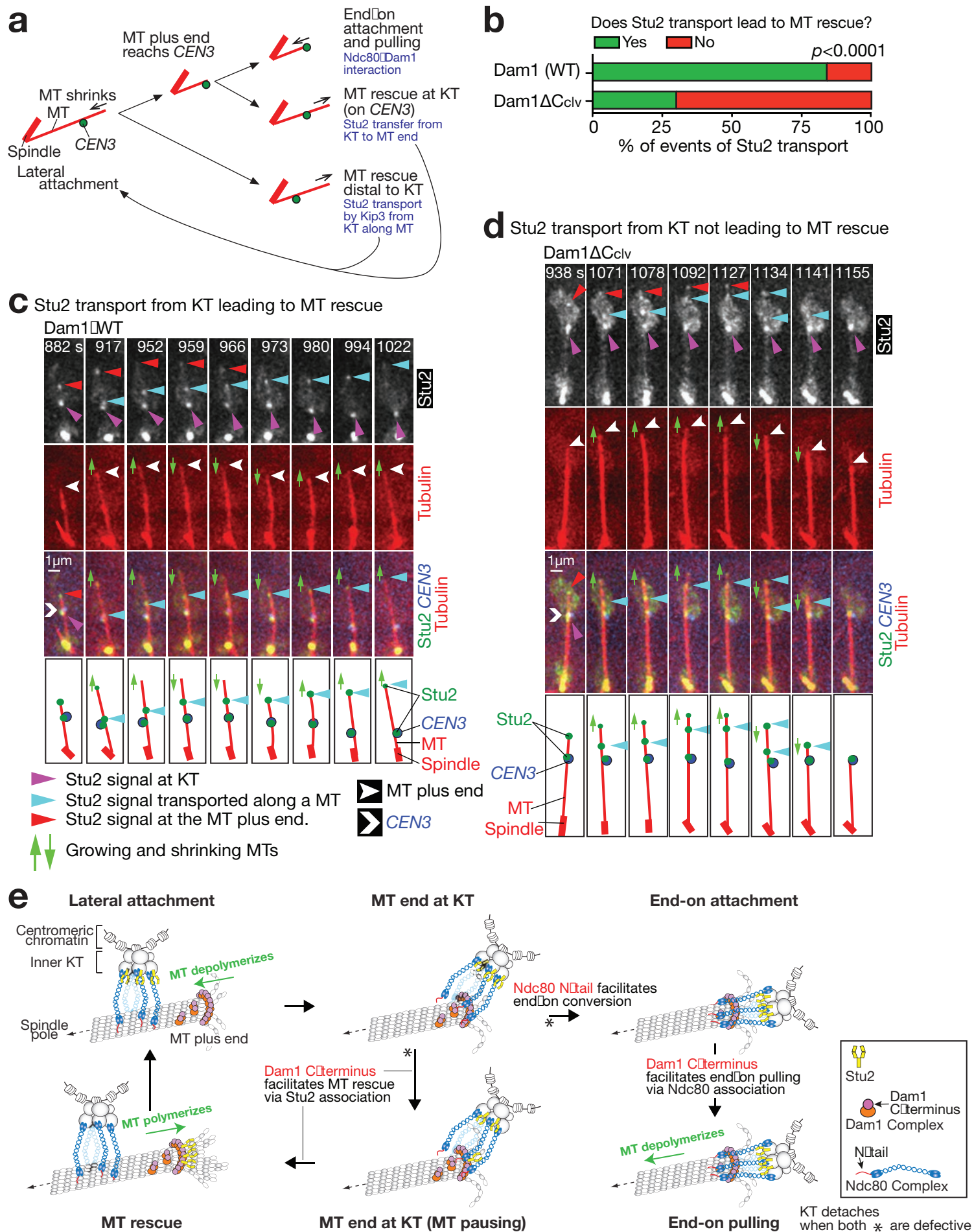
Kalantzaki et al, Fig.1

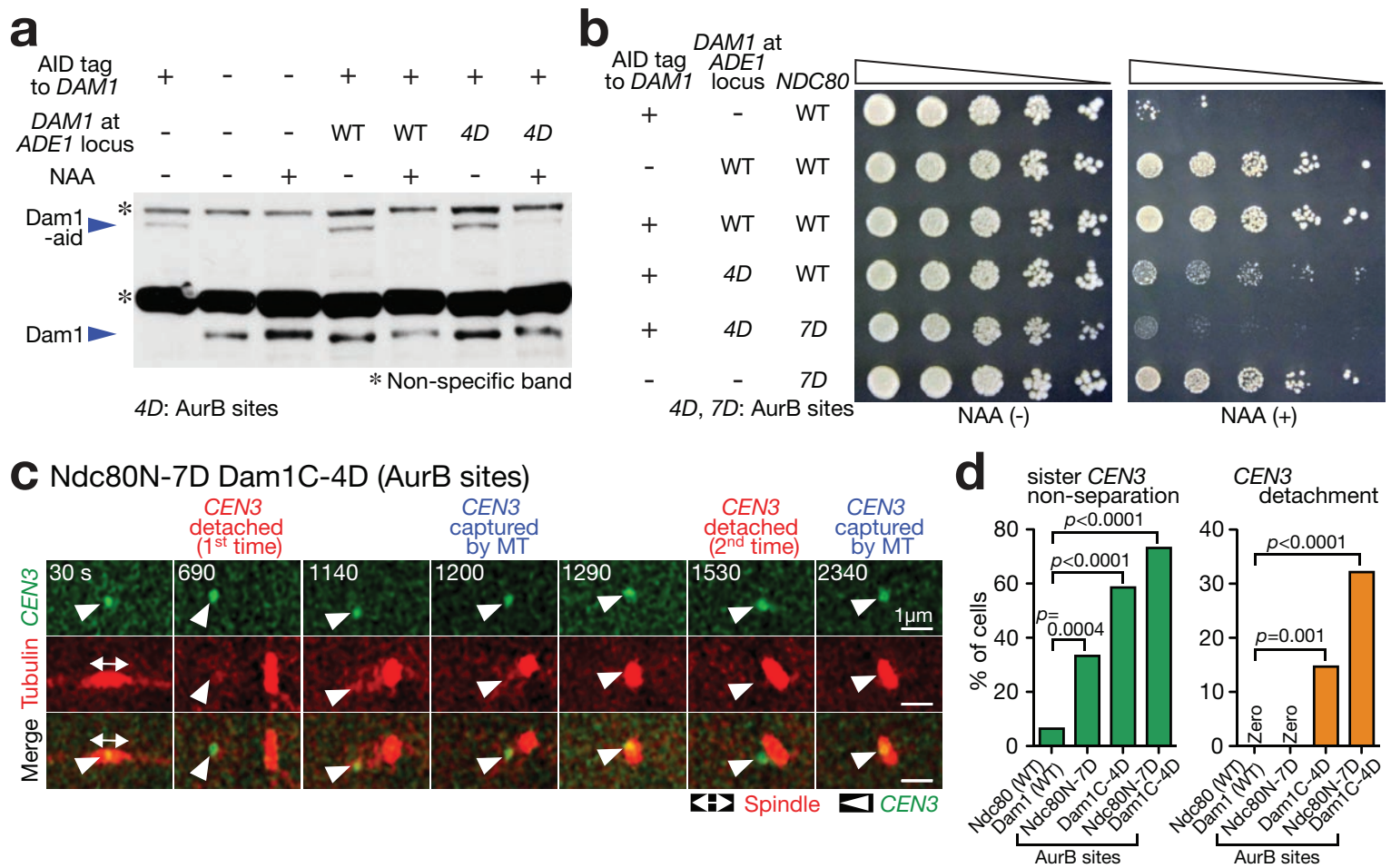


Kalantzaki et al, Fig. 2

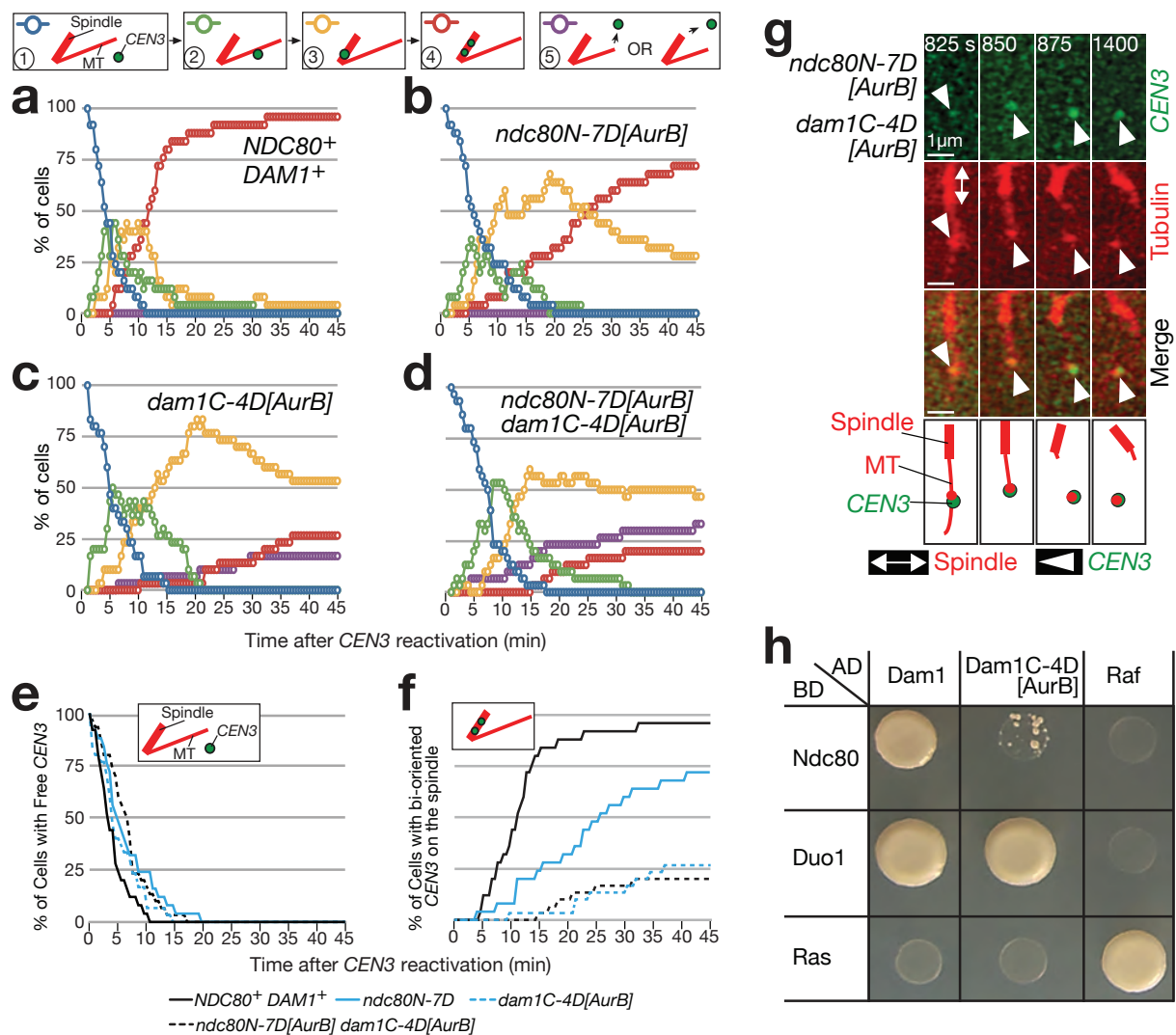


Kalantzaki et al, Fig. 3



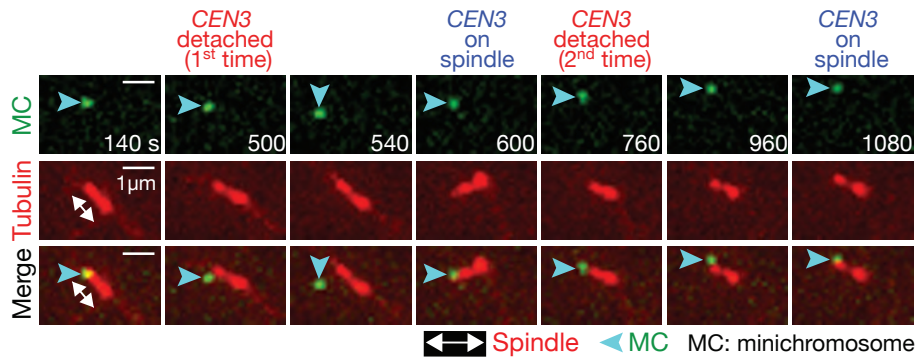


Kalantzaki et al, Fig. 5

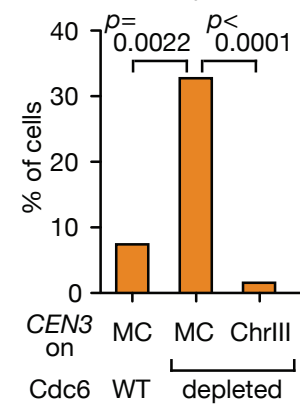


Kalantzaki et al, Fig. 6

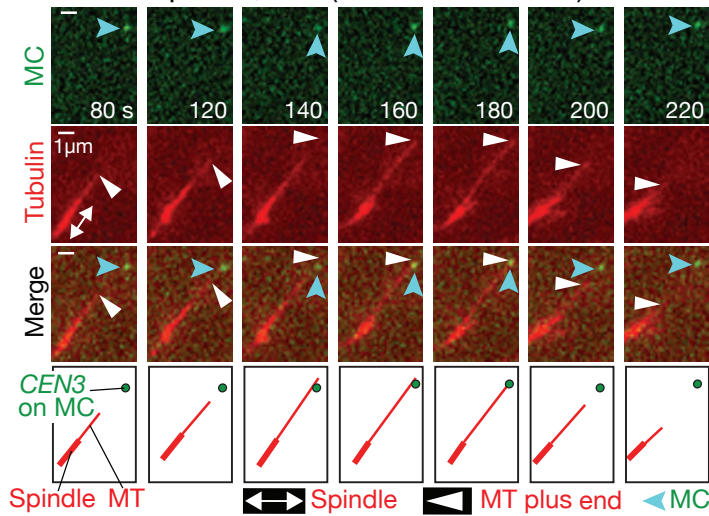
a Cdc6 depleted, MC (*CEN3* continuously active)



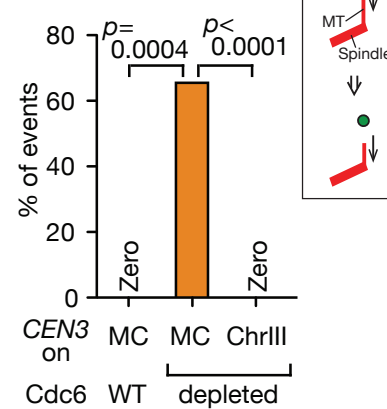
b *CEN3* detachment from the spindle



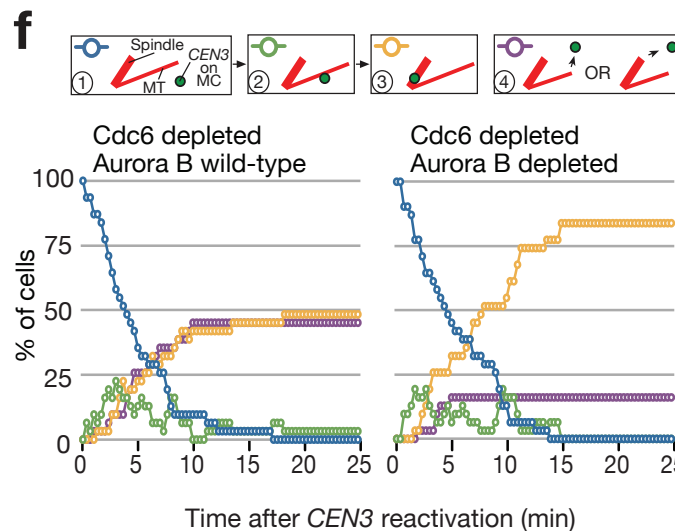
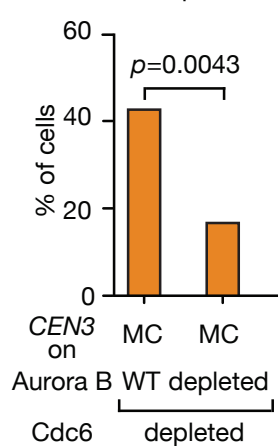
c Cdc6 depleted, MC (*CEN3* reactivated)



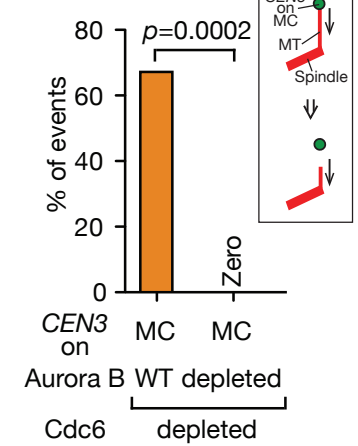
d *CEN3* detachment from MT end



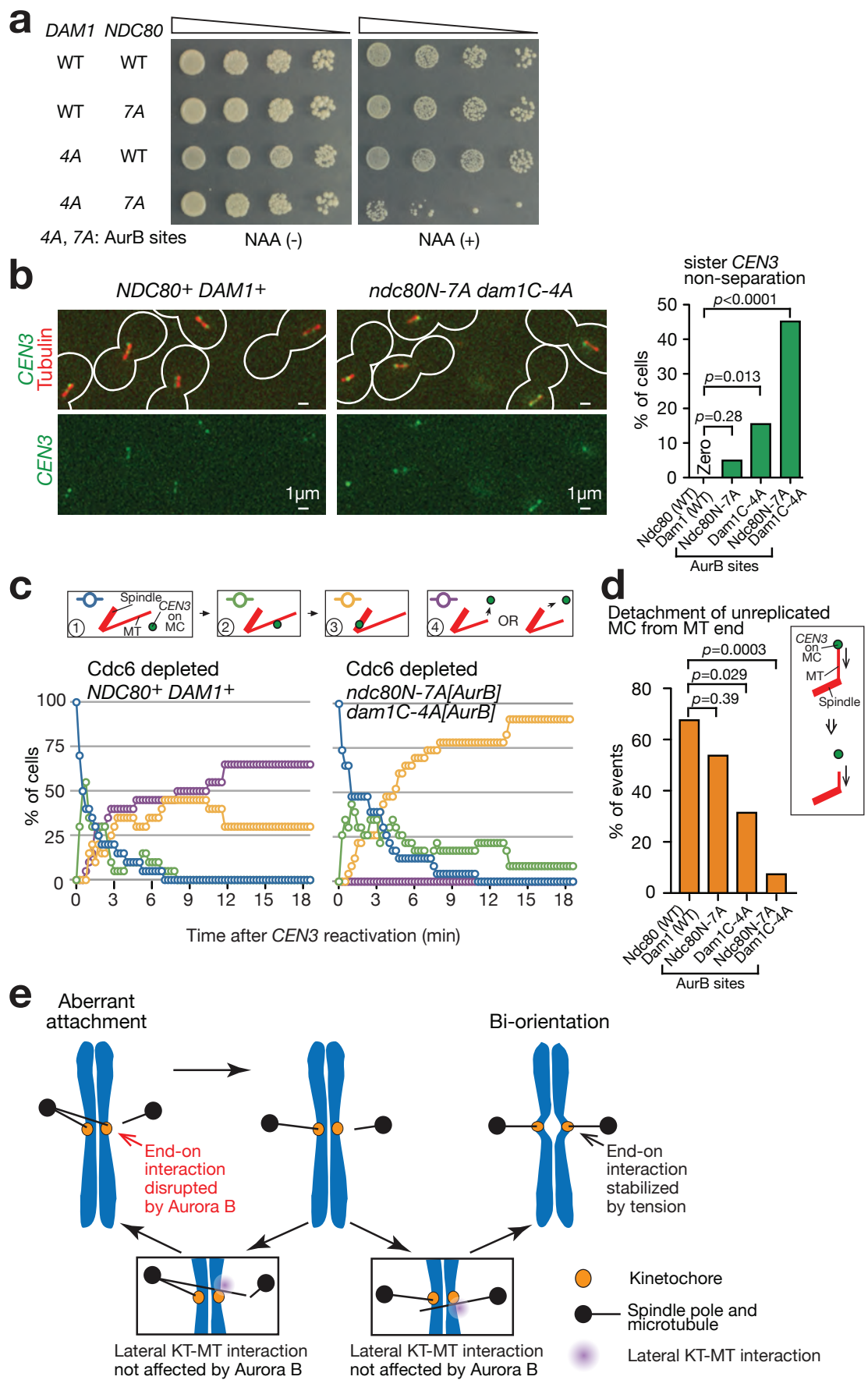
e *CEN3* detachment from the spindle



g *CEN3* detachment from MT end



Kalantzaki et al, Fig.7



Legends for supplementary figures

Supplementary Figure 1. KTs show detachment from the spindle, followed by recapture, when Dam1 C-terminus and Ndc80 N-tail are deleted.

NDC80⁺ DAM1⁺ (T9659), *ndc80 ΔN* (T9298), *dam1-TEVsites* (T9258), *ndc80 ΔN dam1-TEVsites* (T11454) cells with *P_{GAL}-TEV* (except for T9298) *MTW1-4×mCherry Venus-TUB1 P_{MET3}-CDC20* were treated as in Fig 1e, except that images were acquired every 30 sec. A representative cell with Ndc80ΔN plus Dam1ΔCclv (0 min: start of image acquisition) is shown here. The graphs show percentages of cells that showed KT detachment from the spindle (usually followed by reattachment); n= 33, 38, 34 and 35 cells were analysed (from left to right). Experiments were performed twice (statistics source data are shown in Supplementary Table 2) and a representative experiment is shown here. *p*-values (two tailed) were obtained by Fisher's exact test.

Supplementary Figure 2. Mutations at the Ndc80 calponin-homology domain lead to a defect in the lateral KT–MT attachment.

P_{CUP1}-ubi-DHFR-ndc80 (*ndc80-td*) *P_{GAL}-CEN3-tetOs TetR-GFP YFP-TUB1 P_{MET3}-CDC20* cells with *NDC80⁺* (T10069), *ndc80-CH-K6A* (T7427) [expressed from *NDC80* promoter] or no additional *NDC80* construct (T7428), inserted at *his3* locus, were treated with α factor at 25°C in methionine-dropout medium with raffinose, galactose and 2% CuSO₄. After 3 h, cells were released to YPA medium with raffinose, galactose (to inactivate *P_{GAL}-CEN3*) and methionine (to deplete Cdc20) at 35°C (to degrade Ndc80-td). After 20 min, cells were transferred to synthetic complete medium with glucose (to re-activate *P_{GAL}-CEN3*) and methionine, and images were acquired every 5 min (start of image acquisition; 0 min) at 35°C. In the *ndc80-CH-K6A* mutant, six lysines were replaced with alanines within the calponin-homology domain. Representative images of T10069 and T7427 cells at 20 min (left) and the percentage of cells (n= 20–30 cells were analysed at each time point) with *CEN3* that is unattached to MTs (right).

The above result and Fig 2 suggest the Ndc80 calponin-homology domain, but not the Dam1 C-terminus or the Ndc80 N-tail, is required for the initial lateral KT-MT attachment. Note that the KT on *CEN3* before being caught on the lateral surface of a spindle MT (a MT extending from a spindle pole) often generates short MTs that are thought to facilitate a subsequent KT encounter with a spindle MT⁵⁸; such short KT-derived MTs were found similarly in the wild-type control, Dam1ΔCclv, Ndc80ΔN

and double deletion (our unpublished result). Once *CEN3* is loaded on the lattice of a spindle MT, *CEN3* showed sliding along this MT towards a spindle pole ³; this sliding was also found similarly in wild-type control and the deletion mutants (our unpublished result).

Supplementary Figure 3. Dam1 and Stu2 interact physically in a two-hybrid assay and this interaction is abolished with Dam1 Δ C and with Dam1C-4D[AurB].

(a) Stu2 protein and its C-terminus and N-terminus deletions are shown in the diagram. These deletions were used in **b** and **c**.

(b, c) **b** shows that Dam1 and Stu2 interact physically in a two-hybrid assay and this interaction requires the Dam1 C-terminus. **c** shows that Dam1–Stu2 interaction in a two-hybrid assay is abolished with Dam1C-4D[AurB], i.e. with phospho-mimicking mutations of the Dam1 C-terminus at Aurora B sites. Duo1 is a component of the Dam1 complex and serves as a control. Ras and Raf were also used as controls. AD and BD were as in [Fig 3g](#).

(d) Phospho-mimicking mutants of the Dam1 C-terminus at Aurora B sites are defective in assisting Stu2 in rescuing a MT. Graph shows percentage of Stu2 transport events along a MT, leading, or not leading, to MT rescue, as in [Fig 4b](#).

DAM1⁺ (T11596) and *dam1C-4D[AurB] dam1-aid* (T11595) cells with *TIR P_{GAL}-CEN3-tetOs TetR-3xCFP STU2-4xmCherry GFP-TUB1 P_{MET3}-CDC20* were treated as in [Fig 6a–g](#). n= 13 Stu2 transport events were analysed in each of T11595 and T11596. *p*-values (two tailed) were obtained by Fisher's exact test. Data represent one out of two independent experiments.

Supplementary Figure 4. Phospho-mimicking mutants of the Dam1 C-terminus at Mps1 sites do not show KT detachment from the spindle.

We tested whether phospho-mimicking Dam1 mutants at Mps1 sites (*dam1C-4D[Mps1]*; T217, S218, S221 and S232 replaced with aspartates) show phenotypes similar to those of *dam1C-4D[AurB]*. *dam1C-4D[Mps1]* (T11680) and *dam1C-8D[AurB+Mps1]* (T11642) cells with *dam1-aid TIR P_{GAL}-CEN3-tetOs TetR-3xCFP GFP-TUB1 P_{MET3}-CDC20* were treated and analysed as in [Fig 5 c and d](#). The results of T9530 and T11326 in [Fig 5 c and d](#) are shown again for comparison. n= 65 and 59 cells were analysed for T11680 and T11642, respectively. Experiments were performed twice (statistics source data are shown in Supplementary Table 2) and a representative experiment is shown here. *p*-values (two tailed) were obtained by Fisher's exact test.

In contrast to *dam1C-4D[AurB]*, *dam1C-4D[Mps1]* did not significantly increase the level of bi-orientation defects or *CEN3* detachment from the spindle. Furthermore, the combination of phospho-mimicking mutations at both Aurora B and Mps1 sites (*dam1C-8D[AurB+Mps1]*) did not exacerbate the defects found in *dam1C-4D[AurB]*. Thus Mps1 phosphorylation of the Dam1 C-terminus may not suppress the function of this domain and may not contribute to error correction. Consistently, non-phosphorylatable mutants of Dam1 at these Mps1 sites did not show a defect in bi-orientation³⁵ (our unpublished result).

Supplementary Figure 5. Non-phosphorylatable mutants of the Ndc80 N-tail and Dam1 C-terminus at Aurora B sites show normal lateral KT–MT interaction and slower bi-orientation establishment.

Cells in Fig 8b were treated as in Fig 6 a–g; i.e. *CEN3* under the *GAL* promoter was inactivated upon release from α factor treatment and then reactivated during metaphase arrest. Percentage of cells at each step of KT–MT interaction is shown (as in Fig 2c–f). Note that, in these cells, we did not observe any *CEN3* detachment from a MT or from the spindle after *CEN3* was caught on the MT or on the spindle. n= 21 cells were analysed in each strain. Data represent one out of two independent experiments.

Supplementary Figure 6. Full scans of western blots (for Figs 1c and 5a).

Positions of protein size markers are shown at left of each blot.

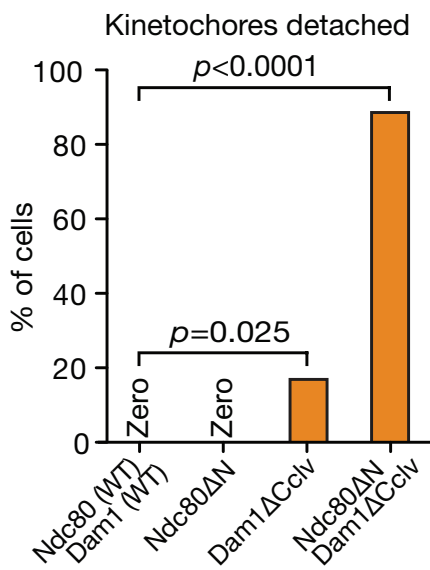
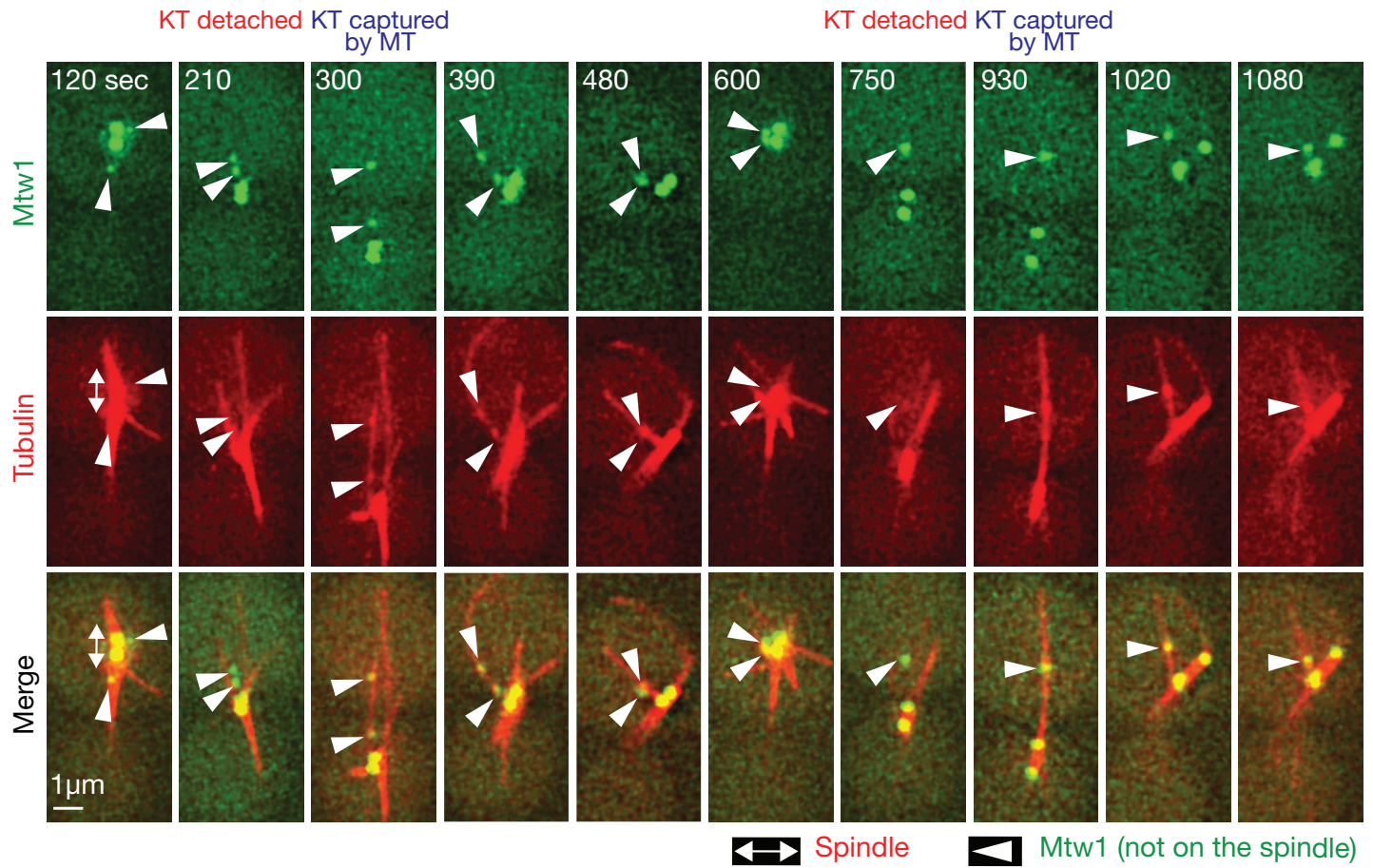
Supplementary Table 1. Genotypes of yeast strains used in this study.

The table shows genotypes of yeast strains used in this study. All strains used in this study are derivatives of *Saccharomyces cerevisiae* W303 (K699 and K700 from Kim Nasmyth lab).

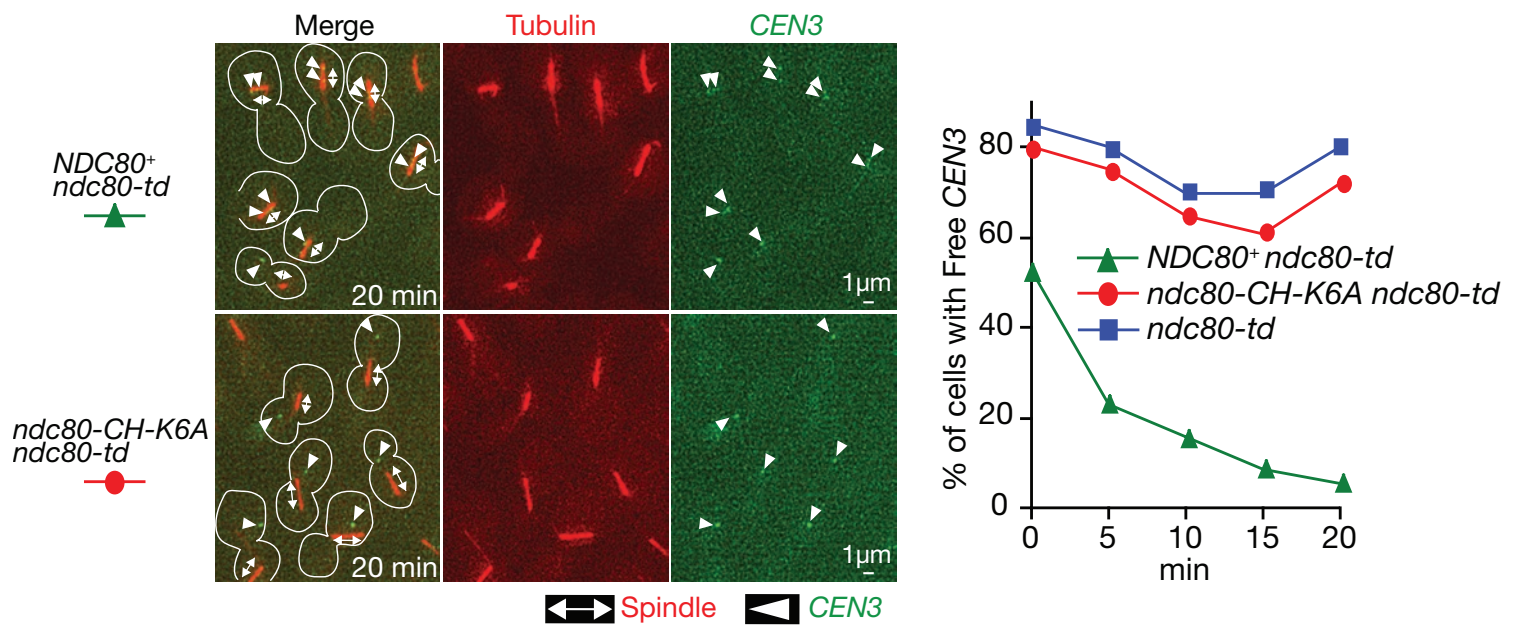
Supplementary Table 2. Statistics source data.

Tables show statistics source data. All *p* values were two-tailed, and calculated in comparison with wild-type controls (shown in the first row of each table) unless otherwise stated. Data from Experiment 1 (Exp1) are shown in each corresponding figure.

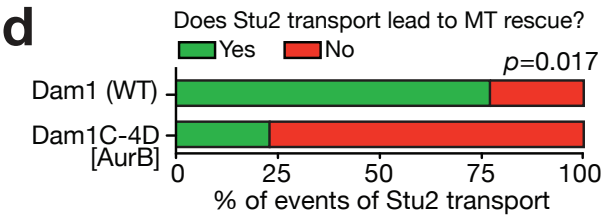
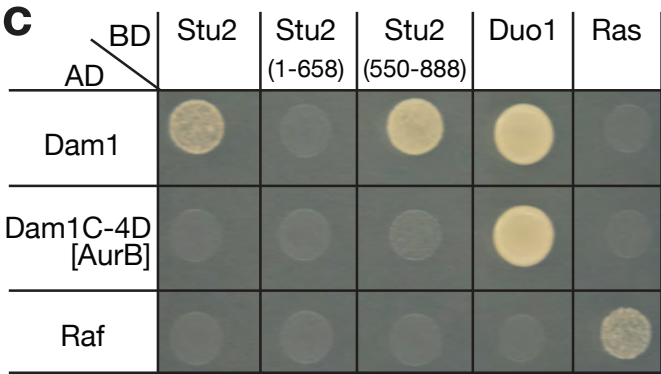
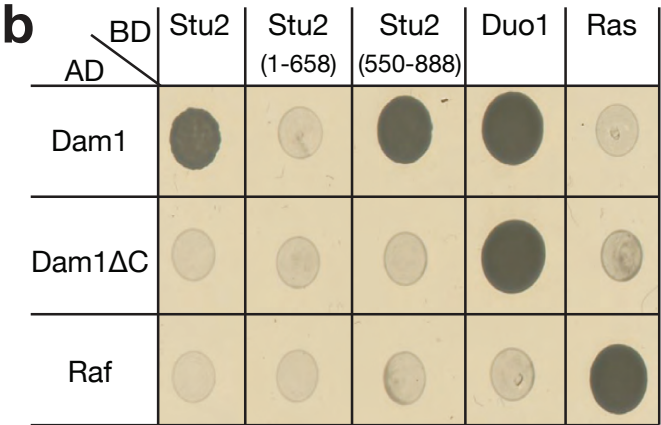
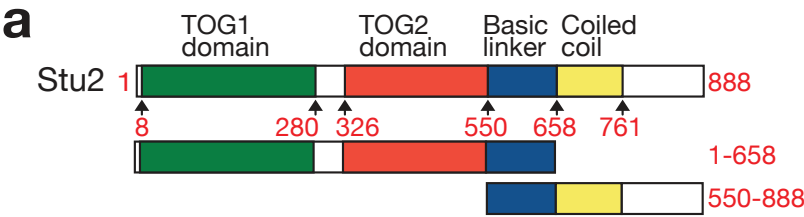
Kalantzaki et al, Supplementary Figure 1



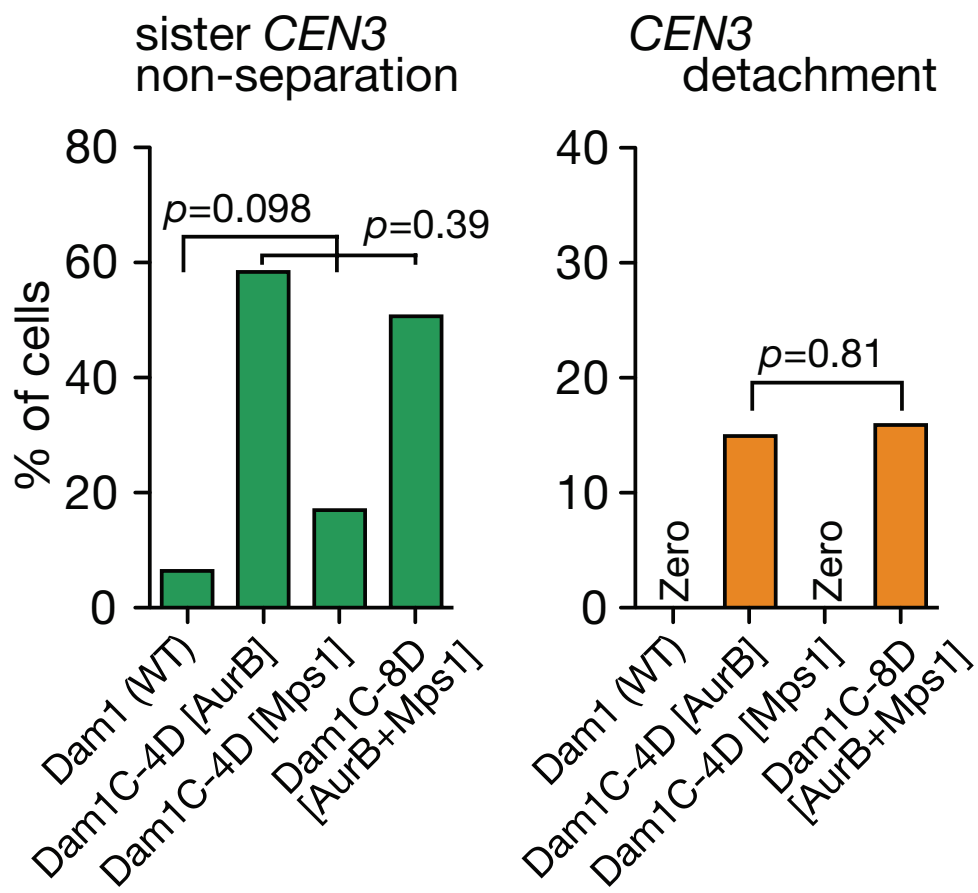
Kalantzaki et al, Supplementary Figure 2



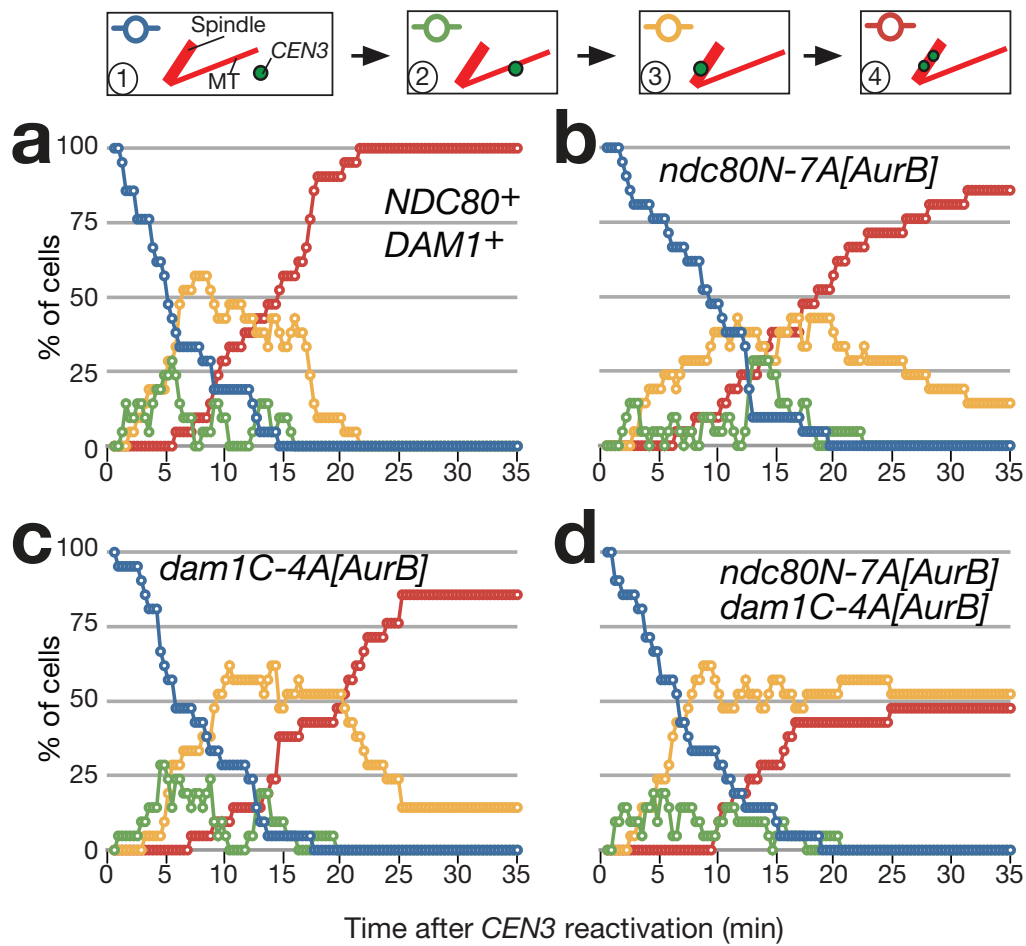
Kalantzaki et al, Supplementary Figure 3



Kalantzaki et al, Supplementary Figure 4



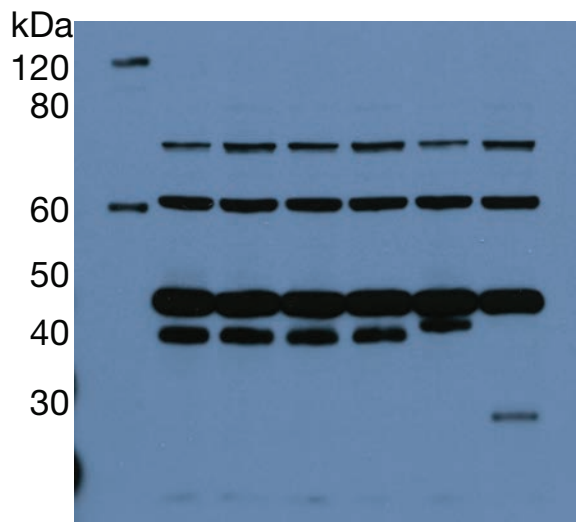
Kalantzaki et al, Supplementary Figure 5



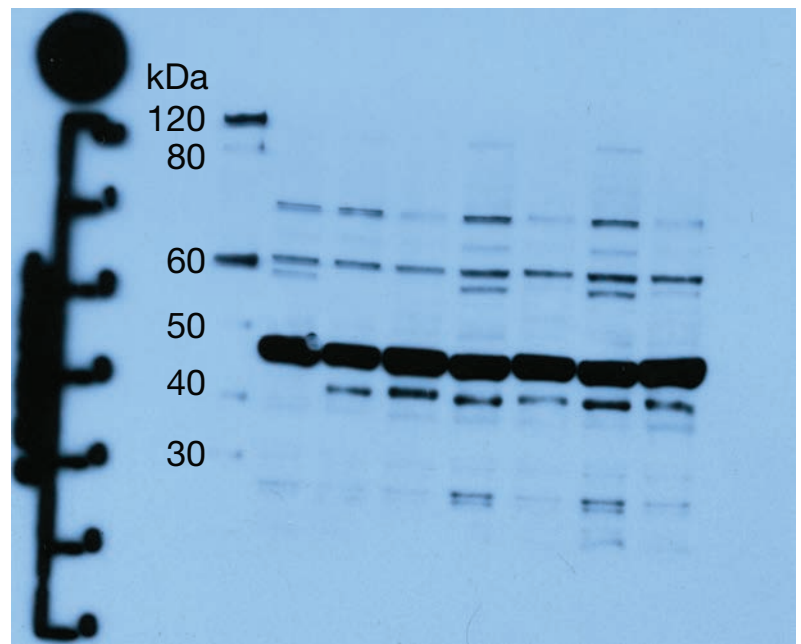
Kalantzaki et al, Supplementary Figure 6

Original uncropped western blots

For Figure 1c



For Figure 5a



Methods

Yeast strains and cell culture

The background of yeast strains (W303; K699 and K700 from Kim Nasmyth lab), the methods for yeast culture and α factor arrest have all been described previously^{4,46}. Unless otherwise stated, cells were cultured at 25°C in YPA medium containing 2% glucose (YPAD). To activate the *GAL* promoter, cells were pre-incubated in medium containing 2% raffinose (without glucose) at least for 3 h, and subsequently incubated in medium containing both 2% galactose and 2% raffinose. To suppress the *GAL* promoter (without subsequent activation), cells were incubated in medium containing 2% glucose. To activate the *MET3* promoter, cells were incubated in methionine drop-out media. To suppress it, 2mM methionine was added to the relevant media. Constructs of *CEN5-tetOs*⁴⁷, *P_{GAL}-CEN3-tetOs*^{3,29,48}, *TetR-3×CFP*^{48,49}, *P_{MET3}-CDC20*⁵⁰, *Venus-TUB1*⁵¹, *GFP-TUB1*⁵², *P_{GAL}-TEV*⁵⁰ were described previously. *STU2*, *MTW1* and *ASK1* genes were tagged with 4xmCherry at their C-terminus at their original gene loci using a one-step PCR method and the pT909 plasmid⁵¹ as a PCR template. The pT111 minichromosome was constructed by inserting *tetOs* (224 copies)⁴⁸ into the *pGAL-CEN3* plasmid²⁹. The *dam1* mutants and *ndc80* mutants were constructed as explained in the next section. Mutants of Aurora B (*ipl1* in budding yeast) and *cdc6* were constructed as explained in the section after next. Two-hybrid assays, and plating after serial dilution of cells, were as described previously⁵¹, and repeated at least twice to confirm reproducibility. Genotypes of yeast strains, used in this study, are shown in Supplementary Table 1.

Construction of *ndc80* and *dam1* mutants

ndc80ΔN: an *ndc80* gene with deletion of the N-terminus (1–112 amino acids) was generated by a reverse PCR. A single copy of the construct was inserted at the *his3* locus (while the original *NDC80* gene was deleted) or it replaced the original *NDC80* gene at its original locus. Ndc80ΔN protein was expressed at a similar level to that of wild-type protein.

P_{CUP1}-ubi-DHFR-ndc80 (*ndc80-td*): a heat-inducible degron tag⁵³ was added at the N-terminus of the *NDC80* gene at its original gene locus, using a one-step PCR method, as described previously⁵¹.

ndc80-CH-K6A: an *ndc80* gene, in which lysines at 122, 152, 160, 181, 192, 204 were replaced with alanines, was constructed by mutagenesis. These lysines correspond to those in the human Ndc80 calponin-homology domain, which interact with the MT surface²³. The *ndc80-CH-K6A* mutant was lethal. Therefore, to maintain

cell viability, a single copy of the *ndc80-CH-K6A* construct was inserted at *his3* locus while the original *NDC80* gene was replaced with *ndc80-td*.

ndc80N-7D: an *ndc80* gene, in which T21, S37, T54, T71, T74, S95 and S100 were replaced with aspartates, was constructed by gene synthesis (by DNA 2.0). These serines and threonines were the sites of phosphorylation by Aurora B ¹⁹. The mutant construct replaced the original *NDC80* gene at its original locus.

dam1ΔC: the C-terminus (206–343 amino acids) of *DAM1* gene was deleted at its original locus using the one-step PCR method.

dam1-aid: *DAM1* gene was fused with an auxin-inducible degron tag ⁵⁴ at their C-terminus at its original gene locus, using the one-step PCR method, as described previously ⁵⁴. To deplete Dam1-aid, cells with the *TIR* gene ⁵⁴ were incubated with auxin NAA (0.5 mM on plate, 1 mM in liquid media).

dam1-TEVsites: A *dam1* gene with two tandem copies of TEV protease cleavage sites (ENLYFQG×2) ⁵⁰ inserted between Q205 and V206 was generated by an overlap PCR. A single copy of the construct was inserted at *leu2* locus while the original *DAM1* was deleted.

dam1C-4D[AurB]: a *dam1* gene, in which serines at 257, 265, 292 and 327 were replaced with aspartates, was constructed by gene synthesis (by DNA 2.0). S257, S265 and S292 were identified as the sites of phosphorylation by Aurora B ¹⁸ while S327 matches a consensus site of Aurora B phosphorylation ¹⁸. A single copy of the mutant construct was inserted at an auxotroph locus while the original *DAM1* was replaced with *dam1-aid*.

dam1C-4D[Mps1], *dam1C-8D[AurB+Mps1]*: these *dam1* mutants were generated in the same way as above, but at different phosphorylation sites. In *dam1C-4D[Mps1]*, T217, S218, S221 and S232 (Mps1 phosphorylation sites ³⁵) were replaced with aspartates. In *dam1C-8D[AurB+Mps1]*, T217, S218, S221, S232, S257, S265, S292 and S327 were replaced with aspartates.

ndc80N-7A: an *ndc80* gene, in which T21, S37, T54, T71, T74, S95 and S100 were replaced with alanines, was constructed by gene synthesis (by DNA 2.0). The mutant construct replaced the original *NDC80* gene at its original locus.

dam1C-4A[AurB]: a *dam1* gene, in which serines at 257, 265, 292 and 327 were replaced with alanines, was constructed by gene synthesis (by DNA 2.0). A single copy of the mutant construct was inserted at an auxotroph locus or at *NDC80* locus (with *ndc80N-7A*) while the original *DAM1* was replaced with *dam1-aid*.

Western blotting was carried out, as described previously ³⁰, to evaluate protein expression in *dam1* mutants (Figs 1c and 5a; full scans of blots are shown in Supplementary Fig 6). Dam1 proteins were detected using an anti-Dam1 affinity-

purified polyclonal antibody ³². The experiment was repeated at least twice to confirm reproducibility.

Depletion of Cdc6 and Aurora B (Ipl1 in budding yeast)

cdc6 anchor away: To deplete Cdc6 protein, we used an anchor away system ⁵⁵, which consists of *cdc6-1-FRB*, *RPL13A-2×FKBP12*, *TOR1-1* and *fpr1Δ*. In the presence of rapamycin (10 μM), Cdc6 protein binds Rpl13A ribosomal protein due to the FRB-FKBP12 interaction, which leads to depletion of Cdc6 in the nucleus. We found that, by combining *cdc6-1* temperature sensitive mutant with the anchor away and by exposing cells to a high temperature, Cdc6 depletion or inactivation was more thorough (even if the temperature was shifted down afterwards). In *cdc6-anchor-away P_{MET3}-CDC20* cells, Cdc6 was depleted and inactivated as follows: Cells were arrested in metaphase, and then released from it, by suppressing and reactivating the *MET3* promoter. Upon release, rapamycin was added to deplete Cdc6. One hour later, the *MET3* promoter was suppressed again to arrest cells in metaphase. In this procedure, cells were incubated at a high temperature (35°C) for 1.5 h (for 30 min prior to the first release from metaphase arrest and for a further 1 h after the release) and subsequently incubated at 25°C before and during image acquisition (because image quality was better at 25°C than at 35°C). A lack of appreciable DNA replication was confirmed by FACS DNA content analysis after this procedure of Cdc6 depletion and inactivation. To inactivate and then reactivate *CEN3* under the *GAL* promoter during the procedure of Cdc6 depletion, cells were incubated with galactose after release from the first metaphase arrest, and then incubated with glucose, prior to image acquisition, after the second metaphase arrest.

ipl1-321-aid: *ipl1-321* mutant gene ⁵⁶ was fused with an auxin-inducible degron tag ⁵⁴ at the C-terminus, in its original gene locus, using the one-step PCR method, as described previously ⁵⁴. To deplete Ipl1-321-aid, cells with the *TIR* gene ⁵⁴ were incubated with auxin NAA (0.5 mM on plate, 1 mM in liquid media). In [Fig 7 e-g](#), NAA was added to deplete Ipl1-321-aid upon release from the first metaphase arrest (refer to the above procedure of Cdc6 depletion). Note that *ipl1-321-aid* showed higher sensitivity to NAA at 25°C than *ipl1-aid* (i.e. *IPL1* wild-type gene fused with an auxin-inducible degron tag).

Live-cell imaging and image analyses

The procedures for time-lapse fluorescence microscopy were described previously ⁵⁷. Time-lapse images were collected at 25 °C unless otherwise stated. For image acquisition, we used a DeltaVision RT or Core microscope (Applied Precision), an

UPlanSApo 100× objective lens (Olympus; NA 1.40), SoftWoRx software (Applied Precision), and either a CoolSnap HQ (Photometrics) or Cascade II 512B (Roper Scientific) CCD camera. We acquired 7–9 (0.5–0.7 µm apart) z-sections, which were subsequently deconvoluted, projected to two-dimensional images and analyzed with SoftWoRx and Volocity (Improvision) software. CFP, GFP, and mCherry signals were discriminated using the 89006 multi-band filter set (Chroma). In [Fig 2c–f](#), [Fig 6a–d](#), [Fig 7c–d](#) and [Supplementary Fig 5](#), 1) cells with free *CEN3* (not on MTs or the spindle) were 80–85% ([Fig 2c–f](#)), 60–70% ([Fig 6a–d](#)) and 80–90% ([Fig 7c–d](#), [Supplementary Fig 5](#)) of the whole cell population at 0 min and we analysed what percentage of these cells showed which steps of KT–MT interaction along the time course, and 2) purple lines show cumulative percentage of *CEN3* detachment and, although this *CEN3* detachment was followed by recapture by a MT, such recapture was not included in counts of other steps. In [Fig 8b](#) and [Supplementary Fig 5](#), up to 30% of T11794 cells showed two *CEN3* signals, presumably due to chromosome mis-segregation in a prior cell cycle; such cells were excluded from counting. Meanwhile, the presence and absence of *P_{GAL}-TEV* and *TIR* did not make a significant difference in *NDC80⁺ DAM1⁺* and *ndc80ΔN DAM1⁺* cells in the relevant experiments. Statistical analyses were carried out using Prism (Graph Pad) software. In [Fig 7d](#), [Fig 7g](#) and [Fig 8d](#), end-on attachment was analysed only when it started more than 1 µm away from a spindle pole. We used Fisher’s exact test in [Figs 1f](#), [4b](#), [5d](#), [7b](#), [7d](#), [7e](#), [7g](#), [8b](#) and [8d](#), and [Supplementary Figs 1](#), [3d](#) and [4](#); and a chi-square test for trends in [Fig 3b](#). Statistical analyses were repeated at least twice to confirm reproducibility of the results ([Supplementary Table 2](#)).

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