GTSE1 tunes microtubule dynamics for chromosome alignment and segregation through MCAK inhibition

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Abstract

The dynamic regulation of microtubules during mitosis is critical for accurate chromosome segregation and genome stability. Cancer cell lines with hyperstabilized kinetochore microtubules have increased segregation errors and elevated chromosomal instability (CIN), but the genetic defects responsible remain largely unknown. The microtubule depolymerase MCAK can influence CIN through its impact on microtubule stability, but how its potent activity is controlled in cells remains unclear. Here we show that GTSE1, a protein found overexpressed in aneuploid cancer cell lines and tumours, regulates microtubule stability during mitosis by inhibiting MCAK microtubule depolymerase activity. Cells lacking GTSE1 have defects in chromosome alignment and spindle positioning due to microtubule instability caused by excess MCAK activity. Reducing GTSE1 levels in CIN cancer cell lines reduces chromosome missegregation defects, while artificially inducing GTSE1 levels in chromosomally stable cells elevates chromosome missegregation and CIN. Thus, GTSE1 overexpression leading to hyperstabilization of kinetochore microtubules via MCAK inhibition defines a new potential mechanism driving CIN.
Introduction

The precise regulation of microtubule (MT) dynamics is essential to the accurate execution of mitosis and faithful segregation of chromosomes. Defects in the regulation of MT stability and dynamics can result in errors in spindle positioning and chromosome segregation, two processes found defective in cancers (Gordon et al., 2012; Noatynska et al., 2012). Persistent errors in chromosome segregation lead to chromosomal instability (CIN), the increased rate of gain or loss of chromosomes within a cell population. CIN is present in most solid tumours, and recent evidence suggests CIN plays a causal role in tumorigenesis (Schvartzman et al., 2010). The genetic and molecular defects that lead to CIN in tumours, however, remain largely unknown.

In several cancer cell lines with CIN, kinetochore-MT attachments are hyperstabilized (Bakhoum et al., 2009a). This leads to an increased frequency of chromosome missegregation, and ultimately to CIN, due to a reduced ability of cells to correct erroneous kinetochore-MT attachments, in particular merotelic attachments, where one kinetochore is connected to microtubules from both spindle poles (Bakhoum et al., 2009a; b). Cells must therefore be able to precisely regulate MT dynamics so that kinetochore MTs are dynamic enough to correct erroneous attachments, yet stable enough to efficiently capture and align chromosomes (Bakhoum et al., 2009a; b).
A major direct regulator of MT stability is the kinesin-13 MT depolymerase MCAK/Kif2C. In vitro, MCAK has extremely potent depolymerase activity (Desai et al., 1999; Hunter et al., 2003; Helenius et al., 2006). In cells, reduction of MCAK activity leads to an increase in microtubule polymer (Rankin and Wordeman, 2010; Rizk et al., 2009). Microtubule-kinetochore attachments are also hyperstabilized, leading to defects in correcting merotelic attachments and in chromosome segregation (Bakhoum et al., 2009a; Maney et al., 1998; Kline-Smith et al., 2004). Excessive MCAK activity induced by overexpression of MCAK leads to loss of MT stability throughout the cell and defects in the capture and alignment of chromosomes (Maney et al., 1998; Moore and Wordeman, 2004; Zhang et al., 2011). MCAK MT depolymerase activity must therefore be precisely controlled in time and cellular space to ensure both chromosome alignment and segregation and to avoid CIN. While interest in MCAK regulation has led to the identification of proteins that enhance or counteract MCAK activity in cells (Cross and Powers, 2011; Ohi et al., 2003; Jiang et al., 2009; Meunier and Vernos, 2011), only NuSAP has been recently reported attenuate MCAK activity via direct interaction (Li et al., 2015). In vitro studies of MCAK have uncovered potential mechanisms by which intramolecular rearrangements of MCAK can determine MT depolymerase activity (Ems-McClung et al., 2013; Talapatra et al., 2015; Burns et al., 2014). Based on this knowledge, proposed mechanisms for direct regulation of MCAK activity in cells have thus largely relied on intramolecular rearrangements induced from interaction with microtubules, nucleotide exchange, and phosphorylation by
mitotic kinases (Ems-McClung et al., 2013; Talapatra et al., 2015; Burns et al., 2014; Cooper et al., 2009).

Because MCAK activity affects kinetochore MT stability, its deregulation may impact CIN. Indeed, artificially destabilizing kinetochore MTs in CIN lines by overexpressing MCAK reduces chromosome missegregation and CIN (Bakhoum et al., 2009b). While these key experiments point to hyperstability of kinetochore MTs in cancer cell lines as a direct cause of CIN, they do not resolve the genetic origin of this defect, as MCAK protein levels are not generally downregulated in cancer cell lines or tumours (Bakhoum et al., 2009a; Sanhaji et al., 2011). Therefore, investigation into cellular regulation of MCAK activity, as well as the molecular basis of kinetochore MT hyperstabilization in cancer cells, is highly desirable.

GTSE1 is a microtubule-associated and EB1-dependent plus-end tracking protein (Monte et al., 2000; Scolz et al., 2012). In interphase, recruitment of GTSE1 to growing microtubule plus-ends by EB1 is required for cell migration. During mitosis, GTSE1 is heavily phosphorylated and no longer tracks microtubule plus-ends nor interacts directly with the microtubule lattice, but localizes to the centrosomes and mitotic spindle in a TACC3-dependent manner (Scolz et al., 2012; Hubner et al., 2010). GTSE1 levels are upregulated in several tumour types and cancer cell lines, and GTSE1 expression correlates with tumour grade and poor clinical outcome in breast cancers (Scolz et al., 2012). It is yet unclear, however, how GTSE1 overexpression may facilitate tumour development.
Here we show that GTSE1 tunes MT stability in mitosis to ensure both chromosome alignment and accurate segregation by suppressing MCAK MT depolymerase activity. GTSE1 interacts directly with MCAK and inhibits its activity in vitro, thus extending our understanding of the mechanisms controlling MCAK activity. Cellular GTSE1 levels influence CIN, as increasing GTSE1 expression in diploid cell lines to mimic the tumorigenic state leads to chromosome segregation defects and induces CIN. Depletion or knockout of the high GTSE1 levels in CIN cancer cell lines reduces chromosome segregation defects in an MCAK-dependent manner. Thus increased GTSE1 levels may represent a genetic defect by which CIN is elevated in cancer cells, through deregulation of MCAK activity and stabilization of kinetochore fibers.
**Results**

GTSE1 stabilizes microtubules in mitosis and promotes correct spindle orientation.

To investigate the role of GTSE1 during mitosis, we first reduced GTSE1 protein levels by RNAi in human U2OS cells and analyzed them by immunofluorescence (Fig. 1 A and S1 A). GTSE1-depleted mitotic cells displayed a loss of microtubule stability as observed by immunofluorescence labeling of tubulin. Astral microtubules were often absent from both prometaphase and metaphase cells (Fig. 1 A), and microtubule density within the inner spindle was significantly reduced (Fig. 1 B). To quantify the defect in astral microtubule stability, we measured the length in three dimensions of the 10 longest astral MTs visible per cell. The length of the few astral MTs remaining after GTSE1 RNAi was less than half that of control cells (Fig 1 C). To verify that the loss of MT stability following RNAi was specific to depletion of GTSE1, we depleted GTSE1 in two independent and clonal cell lines containing stably integrated RNAi-resistant, GFP-tagged GTSE1 genes harbored on bacterial artificial chromosomes (BACs) (Scolz et al., 2012; Bird et al., 2011). Following depletion of endogenous GTSE1, GTSE1-GFP localized to the mitotic spindle and centrosomes, consistent with the localization of endogenously expressed GTSE1 tagged with GFP via Cas9/CRISPR-mediated homologous recombination(Cong et al., 2013; Ran et al., 2013) (Figs. 1D, S1 B, and S2 A). RNAi-resistant GTSE1-GFP expressing cells maintained astral MTs after GTSE1 RNAi, confirming specificity (Figs. 1 E). To further confirm the role for GTSE1 in microtubule stability, we knocked out the genomic copies of the GTSE1 gene in
U2OS cells via Cas9/CRISPR nuclease targeting (Cong et al., 2013; Ran et al., 2013). Interestingly, we were able to isolate viable clonal GTSE1 knockout cell lines (Fig S2, B-D). Two independent knockout clones were analyzed and also displayed a significant loss of astral microtubules in mitosis (Fig. 1A,E).

We previously found that the interaction between GTSE1 and EB1 is required for GTSE1’s function in cell migration and microtubule-dependent focal adhesion disassembly (Scolz et al., 2012). To determine whether this interaction was important for the role of GTSE1 in mitotic MT stability, we analyzed cells expressing RNAi-resistant, BAC-based GTSE1 that does not interact with EB1 due to mutation of conserved EB-binding (SxIP) motifs (Scolz et al., 2012). After depletion of endogenous GTSE1, SxIP-mutated GTSE1 was able to maintain astral stability similar to the wild-type GTSE1 gene (Fig. 1 E). Thus, the interaction between GTSE1 and EB1 is not important for MT stability in prometaphase and metaphase.

GTSE1-depleted cells lacking astral microtubules often displayed a defect in the positioning of the spindle within the cell, consistent with known roles of astral MT-cortex interactions in establishing spindle orientation (Pearson and Bloom, 2004). We quantified this defect in spindle orientation by measuring the angle of the spindle relative to the coverslip surface, and found that cells depleted of GTSE1 more frequently failed to align the spindle axis within 20 degrees relative to the
surface (Fig. 1 F). Thus, GTSE1 is necessary for the stabilization of microtubules in mitosis and proper spindle orientation.

**GTSE1 stabilizes kinetochore MTs to ensure timely chromosome alignment**

To determine whether GTSE1 plays a role in mitotic progression and chromosome segregation, we imaged live U2OS cells expressing histone mH2A.Z-mCherry progressing through mitosis following RNAi (Figs 2 A and S1 C, Videos 1 and 2). GTSE1-depleted cells took significantly longer than control-depleted cells to align all of their chromosomes and enter anaphase (control RNAi: 28.4 ± 20.5 min; GTSE1 RNAi: 42.5 ± 36.3 min) (Fig 2 B). To quantify the chromosome alignment defect, we fixed cells after RNAi and determined the percentage of cells containing misaligned chromosomes but otherwise displaying a metaphase-like morphology (Fig. 2, C and D). Approximately 40% of GTSE1-depleted cells contained misaligned chromosomes, and this defect was rescued by expression of a wildtype RNAi-resistant GTSE1-GFP transgene, as well as the SxIP-mutated transgene, confirming the specificity of GTSE1 RNAi (Fig 2D). Misaligned chromosomes were also significantly increased in GTSE1 knockout clones, although to a lesser extent than after RNAi (Fig. 2D).

The defect in chromosome alignment and delay in anaphase onset in GTSE1-depleted cells indicated that the spindle assembly checkpoint was active in these cells, and suggested that they may be compromised for their ability to establish proper and stable kinetochore-MT attachments. To more specifically analyze this,
we first quantified the abundance of MAD1, which is recruited to kinetochores that do not have proper kinetochore-MT attachments (Howell et al., 2004), in metaphase cells with aligned chromosomes. Indeed, GTSE1-depleted metaphase cells displayed increased kinetochore localization of MAD1 (Fig. 2 E). The delays in chromosome alignment and MAD1 persistence at kinetochores following GTSE1 depletion could arise from destabilization of kinetochore MTs and kinetochore-MT attachments. To determine whether GTSE1 was required specifically for the stability of kinetochore MTs, we first tested whether there was a reduction in the cold-stable kinetochore MT population of metaphase cells following GTSE1 depletion. Indeed, GTSE1-depleted cells had a significant reduction of cold-stable kinetochore MTs as compared to control-treated cells (Fig. 2 F). To quantify kinetochore MT stability, we then assayed kinetochore MT turnover by measuring loss of fluorescence after photoactivation in metaphase U2OS cells expressing photoactivatable (PA) GFP-tubulin (Zhai et al., 1995; Bakhour et al., 2009b). Analysis of fluorescence dissipation after photoactivation indicated that cells depleted of GTSE1 showed a decrease in the half-life of kinetochore MTs as compared to control RNAi (Figs. 2 G, S1 D, and S3), indicating a reduction in the stability of kinetochore-MT attachments. Together, these results implicate GTSE1 as an important regulator of microtubule stability in mitosis, necessary to stabilize kinetochore MTs and promote efficient alignment of chromosomes.
Mitotic defects following GTSE1-depletion are dependent on the activity of MCAK

We next asked the mechanism by which GTSE1 promotes microtubule stability in mitosis. GTSE1 is highly phosphorylated when cells enter mitosis, and does not associate directly with growing microtubule tips nor the microtubule lattice (Scolz et al., 2012) (and unpublished results), indicating that the stabilizing effect of GTSE1 on microtubules in mitosis is mediated through other protein interactors. We previously identified mitotic interactors of GTSE1 by immunoprecipitation and mass spectrometry analysis (Hubner et al., 2010). Of these interactors, the microtubule depolymerase MCAK (Kif2C) was consistently enriched across repeated experiments. We confirmed this interaction by immunoprecipitating either endogenous or stably BAC-expressed GFP-tagged GTSE1 from mitotic cells and probing with antibodies against MCAK (Fig. 3, A and B).

The perturbed mitotic phenotypes following GTSE1 depletion (loss of MT stability and chromosome alignment defects) are reminiscent of the reported mitotic phenotypes upon increasing MCAK activity via overexpression (Maney et al., 1998; Moore and Wordeman, 2004; Zhang et al., 2011), suggesting that GTSE1 may normally attenuate MCAK activity, which becomes unregulated and hyperactive upon loss of GTSE1. We reasoned that if the loss of microtubule stability in mitosis following GTSE1 depletion resulted from excessive MCAK activity, then reducing MCAK levels/activity in these cells should restore MT stability. To test this, we depleted GTSE1 and MCAK by RNAi either individually or together and imaged
mitotic cells by immunofluorescence (Figs. 3 C and S1, E and F). RNAi of MCAK alone in U2OS cells strongly decreased MCAK protein levels and led to an increase in mitotic cells containing longer, curved, and more dense astral MTs, consistent with previous reports (Rankin and Wordeman, 2010; Rizk et al., 2009). When MCAK was codepleted from cells with GTSE1, in contrast to GTSE1 depletion alone, most cells displayed abundant, long astral MTs, indicating that the loss of microtubule stability observed after GTSE1 depletion is dependent on MCAK activity (Fig. 3 C, D, and E). Microtubule intensity within the inner spindle regions was also increased after codepletion of GTSE1 and MCAK as compared to GTSE1 alone, indicating this was not specific to astral MTs (Fig. 3F). Because astral MTs are known to mediate spindle orientation (Pearson and Bloom, 2004), we asked whether the spindle orientation defect after GTSE1 RNAi was restored after MCAK codepletion as well. Indeed, the spindle orientation defect following GTSE1 RNAi was also dependent on MCAK (Fig. 3 G).

If the defects in chromosome alignment seen upon depletion of GTSE1 resulted from MCAK-mediated defects in microtubule stability, we expected they would also be ameliorated by co-depletion of MCAK. Remarkably, upon co-depletion of GTSE1 and MCAK, there was a dramatic reduction in misaligned chromosomes as compared to GTSE1 RNAi alone (Fig. 3H). Together, the above results show that all observed mitotic phenotypes associated with depletion of GTSE1 are alleviated by co-depletion of MCAK. This is consistent with GTSE1 functioning to negatively regulate MCAK microtubule depolymerase activity to ensure stability of...
microtubules throughout the spindle required for spindle orientation and chromosome alignment.

**GTSE1 interacts directly with MCAK and inhibits its microtubule depolymerase activity in vitro**

To determine the mechanism by which GTSE1 antagonizes MCAK activity, we first asked if we could detect a direct interaction of MCAK with a defined domain of GTSE1. We performed *in vitro* pull-down assays with purified MCAK and either a purified GST-tagged GTSE1 N-terminal fragment containing residues 1-460, or a C-terminal fragment containing residues 463-739, the latter of which contains the regions of GTSE1 previously identified to interact with EB1 (Scolz et al., 2012) and p53 (Monte et al., 2003). The N-terminal fragment abundantly pulled down purified MCAK protein, while the C-terminal fragment did not (Figs. 4 A and S4 A), indicating that GTSE1 binds to MCAK via the N-terminal half of the protein.

To test whether GTSE1 could inhibit MCAK activity *in vitro*, we assayed the MT depolymerase activity of purified MCAK protein on labeled MTs using total internal reflection fluorescence (TIRF) microscopy (Helenius et al., 2006) (Figure 4B). Addition of MCAK alone to GMPCPP-stabilized MTs resulted in rapid MT depolymerization (Video 3). Addition of either equimolar or 5-fold excess amounts of GTSE1 protein to this reaction completely eliminated MCAK-induced depolymerization (Fig. 4, C and D; Video 4). This was not due to any inherent ability of GTSE1 to stabilize MTs, because GTSE1 does not have the same impact on
MCAK-independent MT depolymerization. We could only detect a minimal reduction of MT shrinkage rate following catastrophe when the higher concentration of GTSE1 alone was added to dynamic MTs, and catastrophe frequency was not reduced (Fig. S4 C-E, data not shown). Consistently, we also observed a concentration-dependent inhibition of MCAK depolymerase activity using taxol-stabilized MTs and decreasing amounts of GTSE1 protein in bulk MT sedimentation assays (Desai et al., 1999) (Fig. S4 F). Thus, GTSE1 inhibits MCAK depolymerization activity in vitro.

Elevated GTSE1 levels induce missegregation of chromosomes and chromosomal instability

Overexpression of MCAK in CIN cancer cell lines reduces MT-kinetochore attachment hyperstability, thereby allowing correction of erroneous merotelic MT-kinetochore attachments and decreasing the rate of chromosome missegregation and CIN (Bakhoum et al., 2009b). Because GTSE1 is a negative regulator of MCAK activity, and has upregulated protein levels in several cancer cell lines and tumours relative to non-transformed cells (Scolz et al., 2012), we wondered whether the increased levels of GTSE1 found in cancer cells may be inducing CIN. First, we analyzed anaphase chromosome segregation defects, including lagging chromosomes, which correlate with CIN (Thompson and Compton, 2008; Bakhoum et al., 2014). In HeLa and U2OS cells, highly CIN cancer cell lines with high levels of GTSE1 protein, we found that reduction of GTSE1 levels by either RNAi or Cas9-mediated gene knockout significantly reduced the frequency of anaphase
chromosome segregation defects (Fig. 5 A, B, and C). Closer analysis revealed that the reduction in anaphase chromosome segregation defects could be attributed specifically to lagging chromosomes, which generally arise from merotelic attachments, as opposed to chromosome bridges or acentric fragments (Fig. S 5). Importantly, this impact of GTSE1 on chromosome missegregation is also mediated through MCAK, as reducing GTSE1 levels in cells also depleted of MCAK did not have an affect on the frequency of anaphase chromosome segregation defects (Fig. 5 C).

We next asked if overexpression of GTSE1 induced anaphase chromosome segregation defects in HCT116 cells, which are near-diploid, relatively chromosomally stable, and have relatively low levels of GTSE1 protein (Fig. 5 D) (Thompson and Compton, 2008). We stably transfected HCT116 and HCT116 p53-/- cells with a GTSE1-GFP cDNA construct that allowed us to maintain cells overexpressing GTSE1 over many generations through antibiotic selection (Fig. 5 D). We isolated clonal transformants overexpressing GTSE1-GFP to levels comparable to that observed in HeLa and U2OS cells (~5 to 20-fold) and grew them for approximately 60 generations before analyzing mitotic cells. All HCT116 cell clones overexpressing GTSE1 showed a significant increase in the frequency of anaphase chromosome segregation defects (Fig. 5, E and F).

Finally, we asked whether increased levels of GTSE1 in HCT116 cells would induce CIN in these cells. Clonal cell lines overexpressing GTSE1 were analyzed for the
frequency with which cells contained deviations from the disomic state after approximately 60 generations by fluorescence in situ hybridization (FISH) analysis of chromosomes 7 and 11 (Fig. 6A), as compared to mock-transfected clones. Independent HCT116 clones overexpressing GTSE1-GFP displayed a significant increase in CIN (Figure 6B-D). As p53 has been reported to induce apoptosis following chromosome missegregation (Thompson and Compton, 2008), we also performed these experiments in HCT116 p53-/- cells, which also showed elevated frequencies of anaphase segregation defects after GTSE1 overexpression (Fig. 5F). HCT116 p53-/- clones overexpressing GTSE1-GFP displayed a yet greater increase as compared to mock-transfected clones (Fig. 6B,E,F). Thus, reduction of the high GTSE1 levels in a highly CIN cancer cell line reduces the frequency of lagging chromosomes, while overexpression of GTSE1 in chromosomally stable cell lines induces lagging chromosomes and whole chromosome CIN.
Discussion

Here we have demonstrated that GTSE1 inhibits MCAK MT depolymerase activity to determine the stability of microtubules. Depletion of GTSE1 enhances MCAK activity in mitotic cells to levels that are incompatible with the stabilization of astral microtubules and MT-kinetochore interactions, leading to defects in spindle positioning and chromosome capture and alignment. These findings, corroborated by experiments in vitro, support a model in which elevated GTSE1 levels in cancer cells cause excessive inhibition of MCAK activity, leading to hyperstabilization of kinetochore-MT attachments, increased lagging chromosomes, and chromosomal instability.

The precise regulation of MCAK activity is essential for a number of cellular processes that rely on dynamic microtubules (Maney et al., 1998; Rankin and Wordeman, 2010) (Kline-Smith et al., 2004; Lan et al., 2004; Braun et al., 2014; Walczak et al., 2002; Domnitz et al., 2012). The potent MT depolymerase activity of MCAK observed in vitro suggests that its activity must be generally repressed in cells. This may allow for precise spatial and temporal activation of its activity for discrete functions. Here we have elucidated a novel regulatory mechanism responsible for inhibition of MCAK activity, through interaction with GTSE1. In vitro, GTSE1 protein present at equimolar amounts to MCAK is sufficient to completely inhibit MCAK depolymerase activity, suggesting GTSE1 may be a target for differential regulation of MCAK activity in cells.
Most proposed mechanisms to date for direct inhibition of MCAK activity involve phosphorylation of MCAK and/or intramolecular rearrangements (Ems-McClung et al., 2013; Talapatra et al., 2015; Burns et al., 2014). Several phosphorylation sites dependent on multiple mitotic kinases have been identified on MCAK, and been shown to be required for both positive and negative modulation of its activity (Lan et al., 2004; Zhang et al., 2011; Tanenbaum et al., 2011; Ems-McClung et al., 2013; Andrews et al., 2004; Sanhaji et al., 2010; Zhang et al., 2007), but how these various phosphorylation events control MCAK activity in the cellular context remain largely unknown. It was recently shown that NuSAP modulation of MCAK activity is dependent on Aurora B phosphorylation (Li et al., 2015). It will be interesting in the future to determine if GTSE1 association with MCAK is regulated by, or influences, MCAK’s phosphorylation state. A conformation of MCAK where the C-terminal tail interacts with its motor/neck domain is emerging as an important transition state during its catalytic cycle, and this transition could be a target for controlling its cellular regulation (Ems-McClung et al., 2013; Talapatra et al., 2015; Zong et al., 2016). Indeed, phosphorylation by Aurora B has been shown to affect this conformation in vitro (Ems-McClung et al., 2013). Determining the impact of GTSE1 interaction with MCAK on its ability to transition between distinct conformational states will be informative.

Our finding that GTSE1 regulates MT stability and induces CIN when overexpressed is interesting to consider in the context of Aurora A kinase control of MT stability and CIN. Aurora A and its activator TPX2 are commonly found
overexpressed in tumours, highly associated with CIN (Carter et al., 2006), and proposed to comprise a oncogenic holoenzyme (Asteriti et al., 2010). Aurora A kinase activity controls GTSE1 localization to the spindle (Hubner et al., 2010). During mitosis, Aurora A phosphorylates the TACC3 protein, which allows TACC3 to interact with clathrin heavy chain (CLTC), forming a microtubule-interaction interface that brings the pTACC3-clathrin complex to the spindle (Hubner et al., 2010; Booth et al., 2011; Hood et al., 2013; Lin et al., 2010; Fu et al., 2010; Kinoshita, 2005). GTSE1 spindle localization is dependent on Aurora A, TACC3, and clathrin, and GTSE1 is the most downstream of these components recruited to the spindle (Hubner et al., 2010). Loss of Aurora A, TACC3, or clathrin, like GTSE1, results in defects in MT stability and chromosome alignment (Lin et al., 2010; Fu et al., 2010; Booth et al., 2011; Marumoto et al., 2003; Bird and Hyman, 2008; Giet, 2002). The TACC3-clathrin complex has been shown to be required for kinetochore MT stability and the presence of a “mesh” of inter-MT connectors within kinetochore MT bundles, which have been proposed to physically stabilize and maintain the integrity of these bundles (Booth et al., 2011; Hood et al., 2013; Nixon et al., 2015). Our results suggest that Aurora A-dependent recruitment of the TACC3-clathrin complex also facilitates MT stability by recruiting GTSE1 to inhibit MCAK.

Overexpression of Aurora A has recently been shown to induce kinetochore MT hyperstability and CIN in colorectal cancer cells, and these affects were attributed to increased MT polymerization rates (Ertynch et al., 2014). Because GTSE1 spindle
recruitment is dependent on Aurora A activity, it is possible that overexpression of Aurora A also results in enhanced recruitment of GTSE1 to the spindle and increased inhibition of MCAK, which in turn alters MT dynamics and stabilizes kinetochore MTs, leading to CIN.

Here we have uncovered a novel role of GTSE1 in regulating chromosome stability through microtubule stability. Inhibition of MCAK activity by GTSE1 provides a new mechanism by which cells tune MT dynamics to ensure the precise balance of MT stability required for chromosome alignment and segregation. Control of cellular GTSE1 levels is thus essential to ensure genome stability, and deregulation of GTSE1 expression levels may represent a molecular defect contributing to the stabilization of MT dynamics and induction of CIN in tumours.
Materials and methods

Cloning and Plasmids

The hGTSE1-GFP-T2A-BSD plasmid was generated by in-frame cloning of GTSE1 cDNA, eGFP and T2A-BSD in a EGFP-N2 vector (Clontech). The GST-hGTSE1 construct used for protein purification from insect cells was generated by cloning GTSE1 cDNA using EcoRI and BamHI restriction sites into pFLMultiBac vector modified by adding GST (Fitzgerald et al., 2006). The GST-hGTSE1 fragment expression constructs were generated by sub-cloning the sequences encoding amino acids 1-460 (N-terminus) and 463-739 (C-terminus) in a pGEX-6p-1rbs vector (GE Healthcare) using BamHI and SalI restriction sites. An N-terminal FLAG-LAP “NFLAP” cassette was recombined onto the N-terminus of GTSE1 encoded on the BAC RP11-1152E11 via Red E/T-based recombination (Poser et al., 2008; Zhang et al., 1998). An mCherry-BSD cassette was recombined onto the C-terminus of mH2A.Z encoded on the BAC RP24-363J17 via Red E/T-based recombination (Poser et al., 2008; Zhang et al., 1998).

Cell culture and cell lines

All cell lines were grown in DMEM (PAN Biotech) containing 10% fetal bovine serum (Gibco), 2 mM L-Glutamine (PAN Biotech), 100U/mL Penicillin and 0.1mg/ml streptomycin (PAN Biotech) at 37 °C in 5% CO₂. U2OS cells stably expressing RNAi-resistant wild-type and SxIP-mutated GTSE1-LAP were described previously(Scolz et al., 2012). HCT116 p53−/− and U2OS-PA-GFP-tubulin cells were kind gifts from Duane Compton. U2OS cells expressing NFLAP-GTSE1 and mH2A.Z-
mCherry were generated by transfecting the respective BACs described above using Effectene according to manufacturers protocol, and selecting for stable transfectants. HCT116 and HCT116 p53−/− cells grown in 6 cm dishes were transfected with 1.5 μg of hGTSE1-GFP-T2A-BSD cDNA using 5 μL of Lipofectamine 2000 (Invitrogen) according to manufacturer’s protocol. Stable line populations were selected on BSD (4 μg/mL), and individual clones isolated. HCT116 and HCT116 p53−/− control clones were isolated after treating the cells with Lipofectamine 2000 in absence of DNA.

**Generation of GTSE1 knockouts using CRISPR-Cas9 system**

To generate GTSE1 knockout cells using the CRISPR-Cas9 system, two different target sites were chosen in GTSE1 downstream of the ATG in exon 2 (GGCAGGCTGAAGGCTCATCG) and exon 9 (AAGGCCAGACGCAGCCCGGC). Plasmids expressing the Cas9 nuclease and the corresponding guide RNAs were obtained by cloning the following primers pairs: (5’-CACCGAAGGCCAGACGCAGCCCGGC-3’ and 5’-AAACGCCGGCGCTGCTCTGGCCTTC-3’) and (5’-CACCGGCAGGCTGAAGGCTCATCG-3’ and 5’-AAACCGATGAGCCTTCAGCCTGCC-3’)) into the pX330 background (from Addgene), as described in(Ran et al., 2013). U2OS cells were transfected with individual plasmids targeting a single site, or with two plasmids targeting both sites simultaneously, and clones were isolated by serial dilution 48 h after transfection. Genomic DNA was prepared using the DNeasy Blood and Tissue kit (Qiagen). The regions surrounding the CRISPR-Cas9 sites were amplified using the primers 5’- CGGCAATGAGTCTCCCTCAG-3’ and 5’-
AATCGCTTGAACCGAAAGG-3’ (for the site in exon 2) or 5’-CAGTCCACAGCAAATGCCAG-3’ and 5’-CACGACTGAGGTGTGACTTC-3’ (for the site in exon 9) and the corresponding PCR products were sequenced using primers 5’-CCCTGGGATGCGATCATTTC-3’ or 5’-CCCTCAGCACTGCATTAGCAC-3’, respectively. Heterozygous insertions and deletions were resolved using the TIDE software (http://tide.nki.nl) as described in (Brinkman et al., 2014).

**Generation of GFP knock in at endogenous GTSE1 via Cas9/CRISPR**

To generate the U2OS GTSE1-GFP-knock in, a BAC containing a GTSE1-GFP-T2A-BSD<sup>R</sup> fusion was first generated via Red E/T-based recombination of a GFP-T2A-BSD<sup>R</sup> cassette (gift of Tony Hyman) amplified with the primers TGAGCCCTGAGGCTGACAAGGAGAACGTGGATTCCCCACTCCTCAAGTTCGAGAATCTT TATTTTCAGGGCG and CAAGTGTAAGCCACTGCGACCAGCCTAAAGCTTGTTT CTGAGGTTGAAAAATTAACCCTCCCACACGTAGCC into a BAC containing GTSE1. The template for Cas9 mediated HR was obtained by amplifying from the GTSE1-GFP-T2A-BSD<sup>R</sup> BAC a region encompassing the GFP-T2A-BSD<sup>R</sup> cassette and 1kbp homology arms to the C-terminus of GTSE1 using primers CATCAGCCAAATGAGCGAC and GAATCAGCAGTAACCCGAG. The plasmid expressing the Cas9 nuclease and the guide RNA targeting the C-terminus of GTSE1 was obtained by cloning the following primers pairs: (CACCGTCTGAAAGACCCCTAAGAAG and AAACCTTTAGGGCTTCTTCTTCAAC) into the pX330 background.
RNAi

siRNA against hGTSE1 (5′-GAUCAUACAGGUCAGAA-3′), hMCAK (5′-GAUCCAACGCAGUAAU-3′) and control siRNA (Silencer negative control #2) were purchased from Ambion. Approximately 35,000 U2OS cells were added to prewarmed media in 24 well plates or 8-well imaging chambers (ibidi), and transfection complexes containing 2.5 μl oligofectamine and siRNA were added immediately afterwards. Media was changed after 6–8 h. A final concentration of 80 nM and 12 nM RNAi were used for GTSE1 and MCAK, respectively. All experiments were performed 48 hours after siRNA transfection.

Antibodies

Rabbit antibodies against hGTSE1 were previously described (Scolz et al., 2012). Goat anti-GFP antibodies used in immunoprecipitation assays were previously described (Poser et al., 2008). Rabbit antibody against CEP135 was previously described (Bird and Hyman, 2008). Mouse monoclonal antibody conjugated with 488 against Mad1 was a kind gift from Andrea Musacchio. The following antibodies were obtained from commercial sources: mouse anti-alpha-tubulin (DM1alpha, Sigma Aldrich), mouse anti-MCAK (Abnova Corporation, Clone 1G2), human nuclear antibodies to nuclear antigens–centromere autoantibody (CREST; CS1058; Europa Bioproducts Ltd.), mouse anti-c-myc (Oncogene/Calibiochem). The following Secondary antibodies were used: donkey anti-mouse, –rabbit, or –rat conjugated to Alexa 488, 594, or 647 (Bethyl laboratories) and donkey anti-human conjugated to Texas Red or Cy5 (Jackson ImmunoResearch Laboratories).
**Immunofluorescence**

Cells on coverslips were fixed using −20 °C methanol for 10 minutes. Cells were blocked with 0.2% fish skin gelatin (Sigma-Aldrich) in PBS. Cells were incubated with primary antibodies in 0.2% fish skin gelatin in PBS for 1 hour at 37 °C in a humidified chamber, washed, and the same repeated with secondary antibodies. Coverslips were mounted with ProLong gold with DAPI (Molecular Probes, Life technology).

**Microscopy and live cell imaging**

Fluorescence dissipation after photoactivation (FDAPA) analysis and images used for quantifying astral length, inner spindle intensity, and spindle orientation were acquired using a Marianas (3i) spinning disk confocal system based on an Axio Observer Z1 microscope (Zeiss) equipped with a Hamamatsu ORCA-Flash 4.0 Camera. Images were taken using 63x 1.4 NA Apochromat objective (Zeiss,). The images were Z-projected using Slidebook software 5.5. All other images were acquired using a DeltaVision imaging system (GE Healthcare) equipped with an sCMOS camera (PCO edge 5.5). Images were taken using a 60x 1.42 NA PlanApo-N objective (Olympus) at room temperature. Serial Z-stacks of 0.2 μm thickness were obtained and deconvolved using SoftWoRx 6.1.1 software. For live cell imaging, media was changed to CO₂ independent media (Gibco) 12 hours prior to imaging. Live cell image sequences were acquired at 1 min intervals for 12 hours in 2 μm serial Z sections using a 40x 1.42 NA UPlanFL-N objective (Olympus) at 37 °C.
**Image quantification and data analysis**

Inner spindle intensity was quantified in three dimensions using the surface module in IMARIS software (Bitplane). Astral microtubule lengths were measured in three dimensions using IMARIS software. To measure spindle tilt, both spindle poles were located in the Z-series and then using the angle tool in Slidebook software 5.5, the angle made by the spindle to the substratum was measured. Images were processed with ImageJ or Photoshop (Adobe).

**Determining kinetochore-MT half life by FDAPA**

FDAPA in U2OS cells expressing Photoactivatable (PA)-GFP-tubulin was performed as described in (Bakhoum et al., 2009b). U2OS-PA-GFP-Tubulin cells grown on poly-L-Lysine (Sigma-Aldrich) coated 3.5 cm glass bottom chamber (ibidi) were treated with either control or GTSE1 RNAi for 48 h. The medium was changed to CO₂-independent media containing DNA staining dye-Celllight histone 2B-RFP (Life technology) 12 h prior to the experiment. PA-GFP-tubulin in a small area around the kinetochore-MT attachment region on the metaphase spindle was activated with a 405 nm laser and images were taken every 10 sec for 5 minutes. Fluorescence intensity after activation was measured at each time point using ImageJ. Fluorescence intensity was measured within a similar region on the other non-activated half spindle was used for background subtraction. Values were corrected for photobleaching by normalizing to values obtained from taxol-treated stabilized spindles. Following background subtraction and correction for
photobleaching, values were normalized to the first time-point and were fitted to a double exponential decay curve \( F = A_1 \times \exp(-k_1 t) + A_2 \times \exp(-k_2 t) \), using Prism, where \( A_1 \) and \( A_2 \) are the percent total fluorescence contribution of the non-kinetochore and kinetochore microtubules, \( k_1 \) and \( k_2 \) are their respective decay rate constants and \( t \) is the time after photoactivation. The half life of kinetochore microtubules was calculated using \( T_{1/2} = \ln 2 / k_2 \).

**Kinetochore MT stability assays and determining Mad1 positive kinetochores**

U2OS cells treated with either control or GTSE1 RNAi were arrested using Cdk1 inhibitor (RO-3306, Calbiochem) at 31 hours after RNAi transfection. Cells were arrested for 17 hours and then released in normal media for 1 h. After 1 h of release the cells were immediately transferred to ice cold media for 17 minutes and fixed in -20 °C methanol. To determine the ratio of Mad1 positive kinetochores, U2OS cells treated with either control or GTSE1 RNAi were arrested using Cdk1 inhibitor (RO-3306, Calbiochem) as mentioned for kinetochore MT stability assay. After 1 hr of release the cells were fixed with 4% PFA and permeabilized with 0.1% triton X-100. The ratio of Mad1 positive kinetochores was determined by quantifying the total number of kinetochores in aligned metaphase plates and then the number of Mad1 positive kinetochores were quantified using Coloc tool of IMARIS software.

**Immunoprecipitation**
Cells with ~ 70% confluency were arrested in mitosis by adding 200 ng/mL nocodazole for 18 h. Mitotic cells were harvested by shakeoff and lysed using cell lysis buffer (50 mM HEPES pH 7.2, 50 mM Na<sub>2</sub>HPO<sub>4</sub>, 150 mM NaCl, 10% glycerol, 1% Triton X-100, 1 mM EGTA, 1.5 mM MgCl<sub>2</sub>, protease inhibitors) followed by centrifugation at 13,000 rpm for 10 minutes at 4 °C to clear the lysate. A part of the supernatant was taken as “input” and 1-2 µg of the indicated antibody was added to the remaining supernatant and incubated for 2 h at 4 °C with rotation. Dynabeads couple to protein G (Novex Life Technology) were added to the extracts and incubated for 4 hours at 4 °C. The beads were washed three times with cell lysis buffer and once with 1x PBS. The beads were resuspended in hot Lamelli buffer and analyzed by Western blotting.

**Western Blot**

For western blotting after RNAi, cells were harvested by directly adding hot Lamelli buffer in 24 well plates. The proteins were separated on SDS-page gels and transferred onto nitrocellulose membranes. The membrane was incubated with the indicted primary antibodies. The secondary antibodies were coupled to horseradish peroxidase and protein bands were detected using enhanced chemiluminescence (ECL, Amersham, GE Healthcare).

**Purification of GTSE1**

Full-length human *GTSE1* cDNA sequence was cloned into pFLMultiBac vectors and baculovirus generated. hGTSE1-FL protein was expressed in Tnao38 insect cells at
27 °C for 48 hours. The cells were harvested by centrifugation at 1800 rpm for 15 minutes in a Sorvall RC 3BP+ (Thermo scientific) centrifuge. The cell pellet was either stored at -80 °C or processed immediately. Cell pellet from 1 L culture was resuspended in 100 mL ice cold Buffer A (50 mM HEPES pH 8.0, 300 mM NaCl, 5% Glycerol, 2 mM TCEP and Protease inhibitors [Serva]) lysed by sonication and clarified by centrifuging at 29,000 rpm for 50 minutes at 4 °C. The cell lysate was incubated with 1 mL Glutathione resin (Amintra) for 1 hour at 4 °C. The resin beads were loaded on gravity flow columns and washed with 150 mL Buffer A at 4 °C. hGTSE1 was cleaved from the beads using GST Precision overnight at 4 °C. The protein was eluted and concentrated using Amicon concentrators. The protein was further purified by size exclusion in a Superdex 200 10/300 column (GE Healthcare) using gel filtration buffer (30 mM HEPES pH8, 300 mM NaCl, 5% glycerol, 2 mM TCEP). The peak fractions were collected and concentrated in Amicon concentrators to give a final concentration of 5–10 µM. The hGTSE1 1-460 and 463-739 fragments and GST were cloned into pGEX-6p-1rbs vector and expressed in bacteria. Bacteria were grown to O.D. 600 of 0.8 and were induced using 1 mM IPTG at 20 °C overnight. The bacterial cells were pelleted at 4000 rpm for 20 minutes at room temperature. Cells were resuspended in GST binding buffer (25 mM HEPES pH 7.5, 300 mM NaCl, 1 mM EDTA, 5% glycerol, 1% Triton X-100, DNase), lysed by sonication and cleared by centrifugation at 29,000 rpm for 30 minutes at 4 °C. The cleared lysate was incubated with Glutathione resin (Amintra) overnight at 4 °C. The beads were washed with GST binding buffer and the protein
was used for experiments.

**Pulldowns**

*In vitro* GST pull downs were performed in GST binding buffer by incubating equal amount of purified FL-MCAK with GST alone or GST-hGTSE1 1-460 and 463-739 fragments (immobilized on Glutathione resin) for 1 h at 4 °C. The reactions were washed with GST binding buffer and resuspended in hot Lamelli buffer and analyzed by Western blotting.

**Microtubule Pelleting Assays**

Taxol stabilized microtubules were prepared by incubating microtubules with 1 mM GTP at 37 °C for 30 minutes followed by incubation at 37 °C for 5 minutes after addition of 50 µM taxol. 1.66 µM tubulin was added to a reaction mixture of MCAK (200 nM) and increasing amounts of GTSE1. All reactions were performed in BRB80 buffer (80 mM Pipes pH 6.8, 1 mM EGTA, 1 mM MgCl₂) supplemented with 70 mM KCl, 1.5 mM ATP, 10 µM taxol for 1 h at room temperature. The reaction was then layered onto a cushion buffer (80 mM PIPES, 1 mM EGTA, 1 mM MgCl₂, 50% glycerol) in a microcentrifuge tube and was centrifuged at 90,000 rpm in a TLA-120.1 rotor (Beckman-Coulter) for 10 min at 25 °C. Supernatant and pellet fractions were separated by SDS–polyacrylamide gel electrophoresis and stained with Coomassie blue for analysis.
**Total internal reflection fluorescence microscopy assays for microtubule dynamics and MCAK activity**

**Tubulin and microtubule preparation:**

Tubulin was purified from juvenile bovine brains using a modified version of the high-PIPES method (Castoldi and Popov, 2003), wherein the first polymerization cycle was performed in 100 mM PIPES instead of 1 M PIPES. Labelling of tubulin with tetramethylrhodamine (TAMRA) and Alexa Fluor 546 succinimidyl esters (Life Technologies) was performed as described previously (Wieczorek et al., 2013). GMPCPP stabilized microtubules were prepared as follows: A polymerization mixture was prepared with BRB80 (80 mM PIPES-KOH, pH 6.9, 1 mM EGTA, 1 mM MgCl₂)+ 2 mM tubulin, 1 mM GMPCPP (Jena Biosciences), 1 mM MgCl₂ and a 1:4 molar ratio of TAMRA-labelled/unlabelled tubulin. The mixture was incubated on ice for 5 min, followed by incubation at 37 °C for 2 hr. The polymerized GMPCPP microtubules were centrifuged at maximum speed in a Beckman Airfuge and resuspended in BRB80. GMPCPP seeds were prepared by polymerizing a 1:4 molar ratio of TAMRA-labelled/unlabelled tubulin in the presence of guanosine-5’-[(α, β)-methylene]triphosphate (GMPCPP, Jena Biosciences) in two cycles, as described previously (Gell et al., 2010). GMPCPP seeds prepared in this way were stable for several months at −80 °C.

**Total internal reflection fluorescence microscopy and preparation of microscope chambers:**
The microscope set-up uses a Zeiss Axiovert Z1 microscope chassis, a ×100 1.45 NA Plan-apochromat objective lens, and the Zeiss TIRF III slider. A λ = 491 nm diode-pumped solid-state laser (Cobolt) was coupled to a fiber optic cable in free space and introduced into the Zeiss slider. Epifluorescence was achieved using a PhotofluorII excitation source (89 North) with wavelength-specific filter cubes (Chroma). Images were recorded using an Andor iXon + DV-897 EMCCD cameras. Microscope chambers were constructed using custom-machined mounts (Gell et al., 2010). In brief, cover glass was cleaned and silanized as described previously (Helenius et al., 2006). Cover glasses (22 × 22 mm and 18 × 18 mm) were separated by two layers of double-sided tape creating a channel for the exchange of solution. Image acquisition was controlled using MetaMorph (Molecular Devices).

Seeds or GMPCPP stabilized microtubules depending on the experiment were adhered to silanized glass slides as described previously (Bechstedt and Brouhard, 2012). On the day of each experiment for the dynamic assay, aliquots of unlabelled and Alexa Fluor 546-labelled tubulin were thawed, mixed to a 1:3 molar labelling ratio, aliquoted again, and stored in liquid nitrogen.

Microtubule growth from GMPCPP seeds was achieved by incubating flow channels with tubulin in standard polymerization buffer: BRB80, 1 mM GTP, 0.1 mg ml⁻¹ BSA, 1% 2-mercaptoethanol, 250 nM glucose oxidase, 64 nM catalase, 40 mM D-glucose. Assays were performed with an objective heater set to 35 °C. Time-lapse image sequences were acquired at 5 s intervals. MCAK depolymerization of
GMPCPP stabilized seeds was achieved by incubating flow channels with MCAK and standard polymerization buffer plus ATP: BRB80, 1 mM GTP, 0.1 mg ml⁻¹ BSA, 1% 2-mercaptoethanol, 250 nM glucose oxidase, 64 nM catalase, 40 mM D-glucose and 1mM ATP. Assays were performed with an objective heater set to 35 °C. Time-lapse image sequences were acquired at 5 s intervals.

Microtubule growth and shrinkage rates:

Microtubule growth rates were analyzed by manually fitting lines to kymographs of growing microtubules using the Kymograph and Linescan features in MetaMorph (Molecular Devices).

**Fluorescence in situ hybridization**

HCT116 and HCT116 p53⁻/⁻ cells grown on 22 mm coverslips were washed once with 1x PBS and fixed with (3:1) methanol-acetic acid solution at room temperature for 30 minutes. Cells on coverslips were dried at room temperature for 10 minutes and immersed in 2X SSC (0.3 M NaCl, 30 mM sodium citrate) for 5 minutes at room temperature. Cell were dehydrated in ethanol series (70%, 85%, 100%) each for 5 minutes at room temperature and then air-dried for 10 minutes. Cells were processed for FISH by using specific α-satellite probes against chromosome 7 and 11 (Cytocell) according to the manufacturers protocol. Coverslips were mounted using ProLong gold with DAPI overnight at room temperature and sealed. Chromosome signals from approximately 1000 cells were scored using the criteria outlined in (Van Stedum and King, 2002).
Statistical analysis

Statistical significance was determined by performing two-tailed Student’s t-test with unequal-variance unless otherwise stated. Statistical analysis of all karyotype studies, quantification of lagging chromosomes in GTSE1-overexpressing clonal cell lines, and phenotype quantification in GTSE1 knockout clonal cell lines were performed using chi-squared tests.

Supplemental Material

Figure S1. Western blots showing RNAi depletion efficiency for all siRNAs and cell lines analyzed. (A) Western blot of cell lysates following control (with dilutions indicated) or GTSE1 RNAi in U2OS cells, probed with anti-GTSE1 and anti-alpha-tubulin. GTSE1 levels are depleted to less than 10% (B) Western blot of cell lysates following control or GTSE1 RNAi in U2OS cells, two stable U2OS cell clones expressing RNAi-resistant GTSE1 (U2OS WT204 and U2OS WT212), and two stable U2OS cell clones expressing RNAi-resistant GTSE1 mutated to abolish interaction with EB1 (U2OS Sk202 and U2OS Sk208). Blots were probed with anti-GTSE1 and anti-alpha-tubulin. (C) Western blot of cell lysates following control (with dilutions indicated) or GTSE1 RNAi in U2OS H2A.Z-mCherry cells, probed with anti-GTSE1 and anti-alpha-tubulin. GTSE1 levels are depleted to less than 25%. (D) Western blot of cell lysates following control (with dilutions indicated) or GTSE1 RNAi in U2OS PA-GFP-tubulin cells, probed with anti-GTSE1 and anti-alpha-tubulin. GTSE1 levels are depleted to less than 20%. (E) Western blot of cell lysates
following control (with dilutions indicated) or MCAK RNAi in U2OS cells, probed with anti-MCAK and anti-alpha-tubulin. MCAK levels are depleted to less than 20%.

(F) Western blot of cell lysates following control (with dilutions indicated) or GTSE1 and MCAK RNAi in U2OS cells, probed with anti-GTSE1, anti-MCAK, and anti-alpha-tubulin. GTSE1 and MCAK levels are both depleted to less than 10%.

**Figure S2. Verification of Cas9 nuclease-mediated GTSE1-GFP and GTSE1 knockout clones.** (A) Western blots showing U2OS cells with endogenous GTSE1 protein tagged with GFP via Cas9-mediated homologous recombination. (B) Western blot of cell lysates from U2OS cells and two Cas9-mediated GTSE1 knockout U2OS clones run by SDS-PAGE and probed with antibodies against GTSE1 or alpha-tubulin. (C) Scheme representing the GTSE1 locus and the two chosen sites targeted by the Cas9 nuclease for knockouts. The sequence targeted by each Cas9/guide RNA pair is indicated. (D) Regions surrounding the CRISPR-Cas9 cutting sites were amplified by PCR from genomic DNA, sequenced, and heterozygous insertions and deletions were resolved using TIDE software. Histograms show the percentage of sequences showing no modification (n=0), an insertion (n>0) or a deletion of n nucleotides (n<0) at the CRISPR-Cas9 cutting site in exon 2 and exon 9 of the GTSE1 locus. Clones were generated by transfecting U2OS cells with a Cas9 nuclease targeting both exon 2 and 9 (Clone 1) or exon 9 alone (Clone 2), respectively.
Figure S3. FDAPA analysis to determine kinetochore microtubule half-life in control or GTSE1-depleted U2OS PA-GFP cells. Graph showing average normalized fluorescence intensity at each timepoint following photoactivation of PA-GFP-tubulin in U2OS metaphase spindles treated with control or GTSE1 RNAi (n=11 per condition over three independent experiments). Solid lines represent the double exponential fit for control and GTSE1 RNAi. Error bars represent standard error of the mean.

Figure S4. In vitro analysis of GTSE1 inhibition of MCAK. (A) Coomassie-stained SDS-PAGE gels of inputs from in vitro GST pull-down assays. (B) Gel showing full length MCAK and GTSE1 purified from baculovirus-infected insect cells. Molecular weights are indicated to the left. (C) Kymograph of a microtubule growing in the presence of 20 μM tubulin from a GMPCPP microtubule seed. (D) Kymograph of a microtubule growing in the presence of 20 μM tubulin and 250 nM GTSE1 from a GMPCPP microtubule seed. (E) Box plot of the shrinkage rate of dynamic microtubules in the presence of 20 μM tubulin and 20 μM tubulin with 250 nM GTSE1. (F) In vitro sedimentation assay to monitor inhibition of MCAK depolymerase activity by GTSE1. Taxol stabilized microtubules were added to 200 nM MCAK and varying concentrations of GTSE1 in presence of ATP. The reaction mixture was incubated for 1 h at room temperature followed by centrifugation at high speed. Supernatants and pellets were collected and equal amounts were run on SDS-PAGE gel and stained with Coomassie blue. The histogram shows the percentage of tubulin for each condition in the pellet fraction, quantified from the
density of the Coomassie gel.

**Figure S5. Quantification of anaphase chromosome segregation defects in U2OS and HeLa cells after GTSE1 and/or MCAK RNAi depletion.** (A) Quantification of anaphase lagging chromosomes. (C) Quantification of chromosome bridges. (D) Quantification of acentric fragments. Error bars represent standard error of the mean. n > 220 U2OS cells with defective anaphases over 3 independent experiments. n > 390 HeLa cells with defective anaphases over 3 independent experiments. p ≤ 0.05 *, p ≤ 0.01 **, p ≤ 0.001 ***

**Supplemental Video Legends**

**Movie 1. Chromosome dynamics and mitotic progression in control U2OS cells.** Mitotic cells efficiently align chromosomes and enter anaphase. U2OS histone H2A.Z-mCherry cells after control RNAi were imaged at 1 minute intervals, and displayed at 8 frames per second.

**Movie 2. Chromosome dynamics and mitotic progression in GTSE1-depleted U2OS cells.** Mitotic cells have difficulty aligning chromosomes and delayed mitotic timing, yet are able to eventually align chromosomes and enter anaphase. U2OS histone H2A.Z-mCherry cells after GTSE1 RNAi were imaged at 1 minute intervals, and displayed at 8 frames per second.

**Movie 3. GMPCPP stabilized microtubules in the presence of 50 nM MCAK.** Epifluorescence images of GMPCPP stabilized microtubules were captured at 5 s intervals for 5 minutes. The GMPCPP stabilized microtubules were observed to
depolymerize rapidly from both ends in the presence of 50nM MCAK. Video playback is 45x real-time (see time stamp).

**Movie 4. GMPCPP stabilized microtubules in the presence of 50 nM MCAK and 250 nM GTSE1.** Epifluorescence images of GMPCPP stabilized microtubules were captured at 5 s intervals for 5 minutes. The GMPCPP stabilized microtubules were observed to maintain a constant length in the presence of 50 nM MCAK and 250 nM GTSE1. Video playback is 45x real-time (see time stamp).
Acknowledgements

We thank D. Compton (Dartmouth Medical School) for providing the U2OS PA-GFP-tubulin and HCT116 p53/− cell lines. We thank A. Musacchio and G. Vader for comments on the manuscript. We thank S. Maffini for help with performing and analyzing photoactivation experiments, A. Faesen for advice and help with microtubule sedimentation assays, M. Mattiuzzo for help with FISH, and J. Beermann and K. Klare for help with protein purification. This work was supported by the Max Planck Institute of Molecular Physiology, a Worldwide Cancer Research project grant to A.B., and an IMPRS-CMB PhD fellowship to S.B.
### Abbreviation List

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<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>BAC</td>
<td>Bacterial artificial chromosome</td>
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<tr>
<td>CIN</td>
<td>Chromosomal instability</td>
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<td>FISH</td>
<td>Fluorescence in situ hybridization</td>
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<td>MT</td>
<td>Microtubule</td>
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<td>TIRF</td>
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Figure Legends:

Figure 1. GTSE1 stabilizes microtubules in mitosis and promotes correct spindle orientation. (A) Immunofluorescence images of U2OS cells after control RNAi, GTSE1 RNAi, and stable knockout of GTSE1 showing less astral microtubules following reduced GTSE1 levels. (B) Quantification of “inner-spindle” tubulin fluorescence intensity from fixed U2OS cells stained for alpha-tubulin following control or GTSE1 RNAi. The yellow dotted line in the sample image indicates the region analyzed. n ≥ 19 per experiment; 3 experiments per condition. (C) Quantification of the average length of astral microtubules in three dimensions from immunofluorescence analysis as shown in A. 10 astral MTs were measured per cell. n = 10 cells per experiment, 3 experiments per condition. (D) Live-cell fluorescence images of metaphase U2OS cells expressing either bacterial artificial chromosome-expressed GTSE1-GFP (U2OS WT212) or endogenously tagged GTSE1-GFP. Both constructs localize to the spindle and centrosomes. (E) Quantification of the percent cells lacking astral microtubules from immunofluorescence analysis as shown in A. The left histogram shows control or GTSE1 RNAi of U2OS cells, two stable U2OS cell clones expressing RNAi-resistant GTSE1 (U2OS WT204 and U2OS WT212), and two stable U2OS cell clones expressing RNAi-resistant GTSE1 mutated to abolish interaction with EB1 (U2OS Sk202 and U2OS Sk208). n > 150 cells over 3 experiments per condition. The right histogram shows two independent U2OS GTSE1 knockout clones. P-values were obtained from a chi-squared test comparing to the U2OS control condition (marked with •). n > 100 cells (F) Analysis of spindle orientation. Images of mitotic cells viewed from the side and stained for DNA (blue), kinetochores (red) and centrioles (green) depict a cell with...
normal spindle alignment parallel to the substrate (top image), and a cell with defective orientation (bottom image). The angle of spindle tilt was calculated by determining the angle between the substrate and a line connecting both centrosomes, as depicted. Quantification of the percent of metaphase cells with a spindle tilt angle greater than 20 degrees is shown. n > 140 cells over 3 experiments per condition. All scale bars represent 5 microns and all error bars represent standard error of the mean. p≤0.05 *, p≤0.01 **, p≤0.001 ***

**Figure 2.** GTSE1 stabilizes kinetochore microtubules to ensure timely chromosome alignment. (A) Still frames of mitoses from time-lapse movies (Supplemental Videos 1 and 2) of U2OS cells expressing histone H2AFZ-mCherry after control (top row) or GTSE1 RNAi (bottom two rows). GTSE1-depleted cells require a longer time to align all chromosomes and enter anaphase. (B) Mitotic duration (nuclear envelope breakdown (NEBD) to anaphase onset) of individual cells is plotted from the analysis of movies of control or GTSE1 RNAi-treated U2OS histone H2AFZ-mCherry cells as are shown in A. GTSE1-depleted cells have a longer average duration of mitosis (control RNAi: 28.4 ± 20.5 min, n=269; GTSE1 RNAi: 42.5 ± 36.3 min, n=295; black bars represent the average) and a higher percentage of cells with mitotic duration longer than 40 minutes (5.9% of control depleted cells versus 29.2% of GTSE1-depleted cells). (C) Immunofluorescence images of chromosomes (red) and kinetochores (white) in fixed U2OS cells after control or GTSE1 RNAi. (D) Quantification of the percent cells with misaligned chromosomes from immunofluorescence analysis as shown in C. The left histogram shows control or
GTSE1 RNAi of U2OS cells, two stable U2OS cell clones expressing RNAi-resistant GTSE1 (U2OS \textsuperscript{WT\textsubscript{204}} and U2OS \textsuperscript{WT\textsubscript{212}}), and two stable U2OS cell clones expressing RNAi-resistant GTSE1 mutated to abolish interaction with EB1 (U2OS \textsuperscript{Sk\textsubscript{202}} and U2OS \textsuperscript{Sk\textsubscript{206}}). n > 150 cells over 3 experiments per condition. The right histogram shows two independent U2OS GTSE1 knockout clones. P-values were obtained from a chi-squared test comparing to the U2OS control condition (marked with •). N > 100 cells (E) Representative immunofluorescence images of metaphase U2OS cells stained with DAPI (DNA), MAD1 and CREST showing MAD1 accumulation on kinetochores after GTSE1 RNAi but not after control RNAi. n > 90 cells from three independent experiments. (F) Fewer GTSE1-depleted cells contain cold-stable microtubules than control-depleted cells. Images show mitotic cells fixed following cold treatment and stained for tubulin and kinetochores. The histogram shows the proportion of cells containing a full complement of cold-stable microtubules following cold treatment, after control cells were normalized to 100%. N > 100 cells per experiment; 3 experiments per condition. (G) Graph showing kinetochore microtubule half-life in control or GTSE1-depleted U2OS cells expressing photoactivatable (PA) GFP-tubulin (see Fig. S 3). Each circle represents kinetochore microtubule half life of a cell from three independent experiments; bars represent the average kinetochore microtubule half life that was calculated by averaging fluorescence intensities at each timepoint from individual experiments and performing a double exponential decay fit (see Fig. S 3) n = 11 cells over 3 experiments per condition. All error bars represent standard error of the mean. All scale bars represent 5 microns. p ≤ 0.05 *, p ≤ 0.01 **, p ≤ 0.001 ***
Figure 3. Mitotic defects following GTSE1 depletion are dependent on the activity of the MT depolymerase MCAK. (A) MCAK coimmunoprecipitates with GTSE1. Western blot following immunoprecipitation from U2OS cell lysates using either anti-GTSE1 or anti-GFP antibodies, and probing with anti-GTSE1 or anti-MCAK antibodies. (B) Western blot following immunoprecipitation from U2OS NFLAP-GTSE1 cell lysates using either anti-GFP or anti-c-myc antibodies, and probing with anti-GFP or anti-MCAK antibodies. (C) Immunofluorescence images of mitotic U2OS cells following control, GTSE1, MCAK, or combined GTSE1 and MCAK RNAi, stained for DNA (DAPI), kinetochores (CREST), and microtubules (tubulin). Scale bar represents 5 microns. (D) Quantification of the average length of astral microtubules in three dimensions from immunofluorescence analysis as shown in c. 10 astral MTs were measured per cell. n = 10 cells per experiment, 3 experiments per condition. (E) Quantification of the percent cells lacking astral microtubules from immunofluorescence analysis as shown in c. n > 100 cells per experiment; 3 experiments per condition. (F) Quantification of “inner-spindle” tubulin fluorescence intensity from immunofluorescence analysis as shown in C. n ≥ 19 per experiment; 3 experiments per condition. (G) Analysis of spindle orientation. Quantification of the percent of metaphase cells with a spindle tilt angle greater than 20 degrees is shown for each RNAi condition. n > 140 cells over 3 experiments per condition. (H) Quantification of the percent cells with misaligned chromosomes from immunofluorescence analysis as shown in C. n > 100 cells per experiments; 3 experiments per condition. All error bars represent standard error of the mean. p≤0.05 *, p≤0.01 **, p≤0.001 ***
Figure 4. GTSE1 interacts directly with MCAK and inhibits its microtubule depolymerase activity in vitro. (A) Immunoblot showing MCAK pulled down in an in vitro GST pull down using GST alone and GST-GTSE1 1-460, GST-GTSE1 463-739 fragments. Input represents 2.5% of total MCAK protein used for GST pull down assay. (B) Kymograph depicting 50 nM MCAK depolymerizing a GMPCPP stabilized microtubule. The black dashed line represents the start of the experiment when MCAK was added. (C) Kymograph depicting a GMPCPP stabilized microtubule maintaining constant length in the presence of 50 nM MCAK plus 250 nM GTSE1. The black dashed line represents the start of the experiment when MCAK and GTSE1 were added. (D) Box plot of the depolymerization rate of GMPCPP stabilized microtubules in the presence of 50 nM MCAK alone, 50 nM MCAK with a 5 fold excess of GTSE1, 50 nM MCAK with an equimolar amount of GTSE1.

Figure 5. Reduction of GTSE1 levels reduces anaphase chromosome segregation defects in HeLa and U2OS cells, while overexpression induces segregation defects in HCT116 cells. (A) Immunofluorescence images of anaphase U2OS cells after control, GTSE1, MCAK, or GTSE1 and MCAK siRNA, stained for DNA (DAPI) and kinetochores (CREST). (B) Quantification of the percentage of anaphase HeLa cells with defective anaphase chromosome segregation events. n > 390 per condition over 3 experiments. (C) Quantification of the percentage of anaphase U2OS cells with defective anaphase chromosome segregation events from immunofluorescence analysis as shown in A. The left histogram shows U2OS cells after RNAi conditions as in A. n > 65 per
experiment; 3 experiments per condition. The right histogram shows two independent U2OS GTSE1 knockout clones. P-values were obtained from a chi-squared test comparing to the U2OS control condition (marked with •). N > 100 cells. (D) Western blots of cell lysates from U2OS, HeLa, and HCT116 and HCT116 p53−/− control and GFP-GTSE1 expressing clonal cell lines used for analysis in (E) and (F). Blots were probed with anti-GTSE1 and anti-alpha-tubulin. The relative abundance of total GTSE1 protein was quantified and normalized to tubulin levels. (E) Quantification of the percent of anaphase cells with lagging chromosomes for control or GTSE1-GFP expressing HCT116 clones. P-values were obtained from chi-squared tests comparing control clones designated with •. (F) Quantification of the percent of anaphase cells with lagging chromosomes for control or GTSE1-GFP expressing HCT116 p53−/− clones. P-values were obtained from chi-squared tests comparing control clones designated with •. p≤0.05 *, p≤0.01 **, p≤0.001 ***

**Figure 6.** Overexpression of GTSE1 induces CIN in HCT116 cells. (A) Fluorescence images from fixed control or GTSE1-GFP overexpressing clonal HCT116 cells processed for fluorescence in situ hybridization (FISH). Probes for α-satellite regions on Chromosomes 7 (green) and 11 (red) are shown and were used to count the number of copies of each chromosome in cells. (B) Statistical significance values from comparing deviances from the modal number of chromosomes in control versus GTSE1-GFP overexpressing HCT116 cells from data presented in (C-F). P-values were determined from chi-squared analysis comparing the indicated clonal cell lines. (C) Percentage of HCT116 control or GFP-GTSE1 expressing clonal cell lines containing
numbers of chromosome 7 deviating from the mode. n > 980 cells for each condition. 
(D) Percentage of HCT116 control or GFP-GTSE1 expressing clonal cell lines containing numbers of chromosome 11 deviating from the mode. n > 980 cells for each condition. (E) Percentage of HCT116 p53−/− control or GFP-GTSE1 expressing clonal cell lines containing numbers of chromosome 7 deviating from the mode. n > 1000 cells for each condition. (F) Percentage of HCT116 p53−/− control or GFP-GTSE1 expressing clonal cell lines containing numbers of chromosome 11 deviating from the mode. n > 1000 cells for each condition.
Figure 1. GTSE1 stabilizes microtubules in mitosis and promotes correct spindle orientation. (A) Immunofluorescence images of U2OS cells after control RNAi, GTSE1 RNAi, and stable knockout of GTSE1 showing less astral microtubules following reduced GTSE1 levels. (B) Quantification of “inner-spindle” tubulin fluorescence intensity from fixed U2OS cells stained for alpha-tubulin following control or GTSE1 RNAi. The yellow dotted line in the sample image indicates the region analyzed. n ≥ 19 per experiment; 3 experiments per condition. (C) Quantification of the average length of astral microtubules in three dimensions from immunofluorescence analysis as shown in A. 10 astral MTs were measured per cell. n = 10 cells per experiment, 3 experiments per condition. (D) Live-cell fluorescence images of metaphase U2OS cells expressing either bacterial artificial chromosome-expressed GTSE1-GFP (U2OS WT212) or endogenously tagged GTSE1-GFP. Both constructs localize to the spindle and centrosomes (E) Quantification of the percent cells lacking astral microtubules from immunofluorescence analysis as shown in A. The left histogram shows control or GTSE1 RNAi of U2OS cells, two stable U2OS cell clones expressing RNAi-resistant GTSE1 (U2OS WT204 and U2OS WT212), and two stable U2OS cell clones expressing RNAi-resistant GTSE1 mutated to abolish interaction with EB1 (U2OS Sk202 and U2OS Sk208). n > 150 cells over 3 experiments per condition. The right histogram shows two independent U2OS GTSE1 knockout clones. P-values were obtained from a chi-squared test comparing to the U2OS control condition (marked with •). n > 100 cells (F) Analysis of spindle orientation. Images of mitotic cells viewed from the side and stained for DNA (blue), kinetochores (red) and centrioles (green) depict a cell with normal spindle alignment parallel to the substrate (top image), and a cell with defective orientation (bottom image). The angle of spindle tilt was calculated by determining the angle between the substrate and a line connecting both centrosomes, as depicted. Quantification of the percent of metaphase cells with a spindle tilt angle greater than 20 degrees is shown. n > 140 cells over 3 experiments per condition. All scale bars represent 5 microns and all error bars represent standard error of the mean. p≤0.05 *, p≤0.01 **, p≤0.001 ***
Figure 2. GTSE1 stabilizes kinetochore microtubules to ensure timely chromosome alignment. (A) Still frames of mitoses from time-lapse movies (Supplemental Videos 1 and 2) of U2OS cells expressing histone H2AFZ-mCherry after control (top row) or GTSE1 RNAi (bottom two rows). GTSE1-depleted cells require a longer time to align all chromosomes and enter anaphase. (B) Mitotic duration (nuclear envelope breakdown (NEBD) to anaphase onset) of individual cells is plotted from the analysis of movies of control or GTSE1 RNAi-treated U2OS histone H2A:FZ-mCherry cells as are shown in A. GTSE1-depleted cells have a longer average duration of mitosis (control RNAi: 28.4 ± 20.5 min, n=269; GTSE1 RNAi: 42.5 ± 36.3 min, n=295; black bars represent the average) and a higher percentage of cells with mitotic duration longer than 40 minutes (5.9% of control depleted cells versus 29.2% of GTSE1-depleted cells). (C) Immunofluorescence images of chromosomes (red) and kinetochores (white) in fixed U2OS cells after control or GTSE1 RNAi. (D) Quantification of the percent cells with misaligned chromosomes from immunofluorescence analysis as shown in C. The left histogram shows control or GTSE1 RNAi of U2OS cells, two stable U2OS cell clones expressing RNAi-resistant GTSE1 (U2OS WT204 and U2OS WT212), and two stable U2OS cell clones expressing RNAi-resistant GTSE1 mutated to abolish interaction with EB1 (U2OS Sk202 and U2OS Sk208), n > 150 cells over 3 experiments per condition. The right histogram shows two independent U2OS GTSE1 knockout clones. P-values were obtained from a chi-squared test comparing to the U2OS control condition (marked with *). N > 100 cells (E) Representative immunofluorescence images of metaphase U2OS cells stained with DAPI (DNA), MAD1 and CREST showing MAD1 accumulation on kinetochores after GTSE1 RNAi but not after control RNAi. n > 90 cells from three independent experiments. (F) Fewer GTSE1-depleted cells contain cold-stable microtubules than control-depleted cells. Images show mitotic cells fixed following cold treatment and stained for tubulin and kinetochores. The histogram shows the proportion of cells containing a full complement of cold-stable microtubules following cold treatment, after control cells were normalized to 100%. N > 100 cells per experiment; 3 experiments per condition. (G) Graph showing kinetochore microtubule half-life in control or GTSE1-depleted U2OS cells expressing photoactivatable (PA) GFP-tubulin (see Fig. S3). Each circle represents kinetochore microtubule half life of a cell from three independent experiments; bars represent the average kinetochore microtubule half life that was calculated by averaging fluorescence intensities at each timepoint from individual experiments and performing a double exponential decay fit (see Fig. S3) n = 11 cells over 3 experiments per condition. All error bars represent standard error of the mean. All scale bars represent 5 microns. p<0.05 *, p<0.01 **, p<0.001 ***
Figure 3. Mitotic defects following GTSE1 depletion are dependent on the activity of the MT depolymerase MCAK. (A) MCAK coimmunoprecipitates with GTSE1. Western blot following immunoprecipitation from U2OS cell lysates using either anti-GTSE1 or anti-GFP antibodies, and probing with anti-GTSE1 or anti-MCAK antibodies. (B) Western blot following immunoprecipitation from U2OS NFLAP-GTSE1 cell lysates using either anti-GFP or anti-c-myc antibodies, and probing with anti-GFP or anti-MCAK antibodies. (C) Immunofluorescence images of mitotic U2OS cells following control, GTSE1, MCAK, or combined GTSE1 and MCAK RNAi, stained for DNA (DAPI), kinetochores (CREST), and microtubules (tubulin). Scale bar represents 5 microns. (D) Quantification of the average length of astral microtubules in three dimensions from immunofluorescence analysis as shown in c. 10 astral MTs were measured per cell. n = 10 cells per experiment, 3 experiments per condition. (E) Quantification of the percent cells lacking astral microtubules from immunofluorescence analysis as shown in c. n > 100 cells per experiment; 3 experiments per condition. (F) Quantification of “inner-spindle” tubulin fluorescence intensity from immunofluorescence analysis as shown in C. n ≥ 19 per experiment; 3 experiments per condition. (G) Analysis of spindle orientation. Quantification of the percent of metaphase cells with a spindle tilt angle greater than 20 degrees is shown for each RNAi condition. n > 140 cells over 3 experiments per condition. (H) Quantification of the percent cells with misaligned chromosomes from immunofluorescence analysis as shown in C. n > 100 cells per experiments; 3 experiments per condition. All error bars represent standard error of the mean. p≤0.05 *, p≤0.01 **, p≤0.001 ***
Figure 4. GTSE1 interacts directly with MCAK and inhibits its microtubule depolymerase activity in vitro. (A) Immunoblot showing MCAK pulled down in a in vitro GST pull down using GST alone and GST-GTSE1 1-460, GST-GTSE1 463-739 fragments. Input represents 2.5% of total MCAK protein used for GST pull down assay. (B) Kymograph depicting 50 nM MCAK depolymerizing a GMPCPP stabilized microtubule. The black dashed line represents the start of the experiment when MCAK was added. (C) Kymograph depicting a GMPCPP stabilized microtubule maintaining constant length in the presence of 50 nM MCAK plus 250 nM GTSE1. The black dashed line represents the start of the experiment when MCAK and GTSE1 were added. (D) Box plot of the depolymerization rate of GMPCPP stabilized microtubules in the presence of 50 nM MCAK alone, 50 nM MCAK with a 5 fold excess of GTSE1, 50 nM MCAK with an equimolar amount of GTSE1.
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