Mitotic checkpoint gene expression is tuned by codon usage bias

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Abstract

The mitotic checkpoint (also called spindle assembly checkpoint, SAC) is a signaling pathway that safeguards proper chromosome segregation. Correct functioning of the SAC depends on adequate protein concentrations and appropriate stoichiometries between SAC proteins. Yet very little is known about the regulation of SAC gene expression. Here, we show in the fission yeast *Schizosaccharomyces pombe* that a combination of short mRNA half-lives and long protein half-lives supports stable SAC protein levels. For the SAC genes *mad2* and *mad3*, their short mRNA half-lives are caused, in part, by a high frequency of nonoptimal codons. In contrast, *mad2* mRNA has a short half-life despite a higher frequency of optimal codons, and despite the lack of known RNA-stabilizing motifs. Hence, different SAC genes employ different strategies of expression. We further show that Mad1 homodimers form co-translationally, which may necessitate a certain codon usage pattern. Taken together, we propose that the codon usage of SAC genes is fine-tuned to ensure proper SAC function. Our work shines light on gene expression features that promote spindle assembly checkpoint function and suggests that synonymous mutations may weaken the checkpoint.

Keywords co-translational assembly; gene expression noise; mitosis; mRNA decay; spindle assembly checkpoint

Subject Categories Cell Cycle; RNA Biology; Translation & Protein Quality

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Introduction

The spindle assembly checkpoint (SAC; also called mitotic checkpoint) is a eukaryotic signaling pathway that delays cell cycle progression when chromosomes have not yet become properly attached to microtubules during mitosis (Lara-Gonzalez et al., 2012; Musacchio, 2015; Kops et al., 2020). Proper function of the SAC needs appropriate SAC protein concentrations (both too low and too high expression can be detrimental) and needs adequate stoichiometries between proteins in the pathway (Chung & Chen, 2002; Ryan et al., 2012; Schuyler et al., 2012; Heinrich et al., 2013; Gross et al., 2018). This makes it important to quantitatively understand SAC gene expression. Yet, the expression of these genes has not been studied in any detail.

The protein network of the SAC, on the other hand, is well understood. While the SAC is active, it forms the mitotic checkpoint complex (MCC), which prevents the anaphase-promoting complex (APC/C) from initiating anaphase (Pines, 2011). A key effector of the SAC is the Mad1/Mad2 complex, a tetramer of two Mad1 and two Mad2 molecules (Chen et al., 1999; Sironi et al., 2002; Fig 1A). Mad1 homodimerizes through a long, parallel intermolecular coiled-coil at its N-terminus, which is followed by the Mad2-binding motif and a C-terminal RWD (RING finger-, WD-repeat-, and DEAD-like proteins) domain (Chen et al., 1999; Sironi et al., 2002; Kim et al., 2012; Piano et al., 2021; preprint: Fischer et al., 2022). The Mad1-binding partner Mad2 is a HORMA domain protein (named after Hop1, Rev7, and Mad2) that can change its conformation between open (O) and closed (C) (Aravind & Koonin, 1998; Luo et al., 2002, 2004). To bind Mad1, the C-terminus of Mad2 wraps around the Mad1 polypeptide similar to a seat belt and Mad2 adopts the closed conformation (Luo et al., 2002; Sironi et al., 2002). This results in a tight complex with no measurable dissociation rate in vitro (Chen et al., 1999; Sironi et al., 2001; Vink et al., 2006). If and to what extent the formation of the intricate Mad1/Mad2 complex is aided by other factors is unknown.

Through a different surface, Mad2 can form heterodimers between its two conformations (O–C) (Mapelli et al., 2007). Dimerization of Mad1/C-Mad2 with O-Mad2 facilitates binding of this O-Mad2 molecule to the APC/C activator Cdc20 (Slp1 in *Schizosaccharomyces pombe*) (De Antoni et al., 2005; Piano et al., 2021; preprint: Fischer et al., 2022). O-Mad2 changes its conformation in the process, forming C-Mad2/Cdc20 through the same seat belt type of binding (Luo et al., 2002). Subsequent binding of BubR1 (Mad3 in yeast) to C-Mad2/Cdc20 results in the mitotic checkpoint complex (MCC) (Sudakin et al., 2001; Chao et al., 2012). The MCC then inhibits the APC/C to block anaphase (Pines, 2011; Allerfi et al., 2016).

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Because the SAC plays a central role in preventing chromosome mis-segregation and because persistent chromosome mis-segregation is a driver of tumor evolution, SAC malfunction is suspected to contribute to carcinogenesis (Gordon et al., 2012; Funk et al., 2016). Mouse models have shown that impairing the SAC promotes chromosome mis-segregation and tumor formation (Baker...
et al., 2005; Holland & Cleveland, 2009; Schwartzman et al., 2010). Completely abolishing the SAC, however, is detrimental to human cells (Dobles et al., 2000; Kops et al., 2004; Michel et al., 2004; Schukken et al., 2021), and suppression of the SAC may in fact be a successful therapeutic strategy against some cancer types (Cohen-Sharir et al., 2021; Quinton et al., 2021). Together, these results indicate that tuning SAC function can make the difference between normal growth, cancerous growth, and cell death.

Although the SAC network has been studied in much detail from a protein-centric view, little is known about SAC gene expression. Understanding this regulatory layer is important, because the changes in SAC protein concentrations can cause SAC malfunction—at least partly because proper stoichiometries, such as between Mad1 and Mad2, are important for function (Chung & Chen, 2002; Ryan et al., 2012; Schuyler et al., 2012; Heinrich et al., 2013; Gross et al., 2018). Here, using fission yeast (Schizosaccharomyces pombe), we study the mRNA layer of SAC gene expression and provide evidence that a combination of short mRNA and long protein half-lives ensures a stable concentration of SAC proteins over time and between cells. Our findings indicate that codon usage bias in mad2<sup>+</sup> and mad3<sup>+</sup>, but not mad1<sup>+</sup>, contributes to their short mRNA half-lives, and that the coding sequence of mad1<sup>+</sup> carries other features that influence expression of this gene. We provide evidence that Mad1 homodimers form co-translationally, which may necessitate a certain codon usage pattern. Overall, our findings shine light on gene expression features that promote SAC function and raise the possibility that synonymous mutations may impair the SAC.

**Results**

**SAC mRNA numbers are approximately Poisson-distributed with means of two to four per cell**

We previously quantified the concentration of SAC proteins fused to green fluorescent protein (GFP) in S. pombe and determined protein concentrations in a range between 30 and 150 nM with strikingly little intercell variability (i.e., low gene expression “noise”) (Heinrich et al., 2013). In these strains, GFP had been fused by traditional tagging, changing the endogenous 3′ UTR to that of the Saccharomyces cerevisiae ADH1 gene and appending an antibiotic-resistance gene, which both may alter gene expression. To avoid such effects, we now employed CRISPR/Cas9-mediated scarless genome editing (Jacobs et al., 2014). We fused ymEGFP (yeast codon-optimized, monomeric enhanced GFP; in the following just “GFP”) to the SAC genes mad1<sup>+</sup>, mad2<sup>+</sup>, and mad3<sup>+</sup> without any change to the surrounding sequences (Fig 1B). Immunoblots showed concentrations broadly similar to the previous strains (Fig EV1A), and strains were not sensitive to the microtubule drug benomyl, suggesting that SAC functionality was maintained (Fig EV1B).

The mean SAC mRNA numbers per cell, determined by single-molecule mRNA fluorescence in situ hybridization (FISH) with probes targeting GFP, were in the range of 3 to 4, even lower than the means of 4.5 to 6 that we had previously observed (Figs 1C and D, and EV1C; Heinrich et al., 2013). This indicates that the traditional tagging strategy indeed influenced gene expression. To test whether the expression in the new strains resembles endogenous expression, we used FISH probes against endogenous mad1<sup>+</sup> and mad2<sup>+</sup> and compared strains expressing the endogenous untagged gene with strains expressing the GFP-tagged gene. For mad2<sup>+</sup>, the mean mRNA number for untagged and tagged mad2<sup>+</sup> was comparable (Figs 1E and EV1E). However, untagged mad1<sup>+</sup> showed even fewer mRNA molecules than mad1<sup>+</sup>-GFP (Fig 1E and EV1E), suggesting that the mere addition of GFP, without any changes in the UTRs or surrounding sequences can change expression of mad1<sup>+</sup>.

[Note that for mad2<sup>+</sup>, the efficiency of the gene-specific probe was slightly lower than the GFP probe (Fig EV1D, both probes measured on mad2<sup>+</sup>-GFP), but this is not expected to influence the conclusion in an experiment that only uses the gene-specific probe (Fig 1E).]

While the mean mRNA numbers per cell for the GFP-tagged genes were in the range of 3 to 4, the numbers in single cells ranged from 0 to around 9 (Fig 1D and E). As expected (Zhurinsky et al., 2010; Padovan-Merhar et al., 2015; Sun et al., 2020), smaller cells had on average lower numbers than larger cells (Fig EV1E). However, even cells of the same size could differ in mRNA number by 8 or more (Fig EV1E). The spread of mRNA numbers in the cell population was well approximated by a Poisson distribution (Fig 1D and E). A Poisson distribution is expected from constitutive expression, where mRNA is synthesized and degraded in uncorrelated events but with a uniform probability over time. In contrast, “bursty” expression (characterized by alterations of promoter activity and inactivity) would result in an even wider distribution (Zenklausen et al., 2008). These results therefore indicate that SAC mRNA numbers vary considerably, but that this variation is within the expected range for constitutive expression.

**mad1<sup>+</sup> and mad2<sup>+</sup> mRNAs do not co-localize in the cytoplasm**

The mRNA FISH data also provide the location of mRNAs. Recent work has suggested that co-translational assembly of protein complexes is more prevalent than previously thought (Schwarz & Beck, 2019). How the stable Mad1/Mad2 complex assembles is unknown. When heterodimeric complexes assemble while both subunits are being translated, their mRNAs will co-localize (Panasenko et al., 2019). We asked whether this is the case for Mad1 and Mad2. We stained mad1<sup>+</sup> mRNA (using a mad1<sup>+</sup> probe) and mad2<sup>+</sup>-GFP mRNA (using a GFP probe) in the same cells, where both were expressed from their respective endogenous loci. While a mad1<sup>+</sup>-GFP strain, used as positive control, showed strong co-localization of the mad1<sup>+</sup> and GFP probes, there was no evidence for co-localization of mad1<sup>+</sup> and mad2<sup>+</sup>-GFP mRNA (Fig 1F). This absence of mRNA co-localization excludes that the Mad1/Mad2 complex forms by synchronous co-translational assembly. We will discuss other possibilities below.

**Low protein noise can be explained through long protein and short mRNA half-lives**

To analyze if and to what extent the strong mRNA variation propagates to the protein level, we quantified GFP-tagged Mad1, Mad2, and Mad3 in single cells using our “Pomegranate” image analysis pipeline, which allows for 3D segmentation (Appendix Fig S1 and S2A; Baybay et al., 2020). To subtract autofluorescence, we mixed the GFP-expressing cells with cells not expressing GFP (Appendix Fig S1). Unlike for the mRNA, we observed little cell-to-cell
variability in the SAC protein concentrations (Fig 2A). As a comparison, we imaged a “noisy” S. pombe protein, Nmt1 (Saint et al., 2019), which indeed showed pronounced cell-to-cell variability (Fig 2A; Appendix Fig S1C). A measure of variability is the coefficient of variation (CV; standard deviation divided by mean). The CVs for Mad1-, Mad2-, or Mad3-GFP were in the range of 0.2, whereas that for Nmt1-GFP was around 0.5 (Fig 2A).

This raised the question how the protein concentrations of Mad1, Mad2, and Mad3 can be homogeneous across the population when the mRNA numbers are highly variable. We considered a simple gene expression model with a constitutively active promoter, and different mRNA and protein synthesis and degradation rates (see Methods for details) that would all yield mean mRNA and protein numbers similar to those that we observe for mad1<sup>+</sup>, mad2<sup>+</sup>, and mad3<sup>+</sup>. The longer the mRNA half-life, the longer a state of low or high mRNA numbers persists; and the shorter the protein half-life, the more closely protein concentrations follow the mRNA numbers (Fig 2B). Hence, long mRNA half-lives and short protein half-lives favor noise, whereas short mRNA half-lives and long protein half-lives suppress noise (Fig 2B and C; Appendix Fig S2B). In the latter case, the long persistence time of proteins buffers fast fluctuations at the mRNA level (Fig 2B).

To ascertain whether this prediction is met by SAC genes, we measured mRNA and protein half-lives. We determined mRNA half-life by metabolic labeling followed by depletion of the labeled pool and quantification of the remaining pool by quantitative PCR. The

**Figure 2.** The checkpoint genes mad1<sup>+</sup>, mad2<sup>+</sup>, and mad3<sup>+</sup> combine short mRNA and long protein half-lives, explaining low noise.

A. Cellular protein noise (coefficient of variation, CV = std / mean) in live-cell microscopy images of S. pombe; n = 7 images (Nmt1-GFP), 11 (Mad1-GFP), 19 (Mad2-GFP), 10 (Mad3-GFP); single images had 16–79 GFP-positive and 6–94 GFP-negative (control) cells. Boxplots show median and interquartile range (IQR); whiskers extend to values no further than 1.5 times the IQR from the first and third quartile, respectively. Mad1, Mad2, and Mad3 all showed significantly lower noise than Nmt1 (Wilcoxon rank sum test, all P < 0.001).

B. Simulations of stochastic gene expression noise from selected mRNA/protein half-life combinations assuming a constantly active promoter (see Methods). Synthesis rates were set to obtain a mean mRNA number of 4 per cell, and a mean protein number of 6,000 per cell. The x-axis of each graph shows time, the y-axis shows mRNA number per cell (blue) or protein number per cell (black).

C. Theoretical prediction for the coefficient of variation (CV = std/mean) of the protein number per cell, assuming different mRNA and protein half-lives, using the same underlying model as in B. Synthesis rates were adjusted to maintain a mean mRNA number per cell of 3, and a mean protein number per cell of 6,000 (approx. 100 nM).

D. mRNA abundances by qPCR following metabolic labeling and removal of the labeled pool (two independent experiments). Lines are regression curves from generalized linear mixed model fits, excluding the measurements at t = 0 in order to accommodate for noninstantaneous labeling by 4<sup>UT</sup>-Act1<sup>+</sup> and ecm33<sup>+</sup> were used as long and short half-life controls, respectively; qPCR was performed for the endogenous mRNAs. Half-lives (95% confidence interval): mad1<sup>+</sup> 5.6 min (4.3–8.4), mad2<sup>+</sup> 7.7 min (6.2–10.4), mad3<sup>+</sup> 5.2 min (4.3–6.9), act1<sup>+</sup> 61.8 min (37.2–1723), ecm33<sup>+</sup> 50.0 min (45.5–57).

E. Protein abundances after translation shut-off with cycloheximide (CHX); n = 3 experiments, error bars = std. Lines indicate fit to a one-phase exponential decay. Cdc2 and Cdc13 were used as long and short half-life controls, respectively. Immunoblots for the endogenous proteins (no tag). A representative experiment shown in Appendix Fig S2E.

Source data are available online for this figure.
mRNA half-lives for mad1\( ^+ \), mad2\( ^+ \), and mad3\( ^+ \) were all in the range of a few minutes (mad1\( ^+ \): 5.6 min, mad2\( ^+ \): 7.7 min, and mad3\( ^+ \): 5.2 min) (Fig 2D). This was consistent with the half-lives determined for these genes in a large-scale study using metabolic labeling (Appendix Fig S2D) (Eser et al., 2016). RNA half-lives have been notoriously difficult to measure, with much variability between studies (Carneiro et al., 2019; preprint: Agarwal & Kelly, 2022). An earlier S. pombe study (Hasan et al., 2014) found longer half-lives across the entire transcriptome, but even in this study, SAC genes were at the lower end of mRNA half-lives (Appendix Fig S2D). As controls, we measured two unrelated genes with reportedly long and short half-life (Eser et al., 2016), act1\( ^+ \) and ecm33\( ^+ \), which behaved as expected (Fig 2D). We determined protein half-lives by translation shut-off using cycloheximide, followed by immunoblotting. The half-lives of Mad1, Mad2, and Mad3 were in the range of many hours, considerably longer than the typical S. pombe cell cycle of 2.5 h (Fig 2E; Appendix Fig S2E) and broadly consistent with previous data (Sczaniecka et al., 2008; Horikoshi et al., 2013; Christiano et al., 2014). This large difference in mRNA and protein half-lives explains the low cell-to-cell variability in protein concentration despite the considerable variation in mRNA numbers (Fig 2C). The short mRNA half-life is therefore important to mitigate the effect of the large variation in mRNA numbers.

**mad2\( ^+ \) and mad3\( ^+ \) have low codon stabilization coefficients**

One of the determining factors for mRNA half-life is codon optimality, which positively correlates with mRNA stability in several eukaryotes (Presnyak et al., 2015; Hanson & Collier, 2018; Narula et al., 2019; Wu et al., 2019; Forrest et al., 2020). The codon stabilization coefficient (CSC) describes the correlation between the occurrence of a codon in mRNA transcripts and experimentally determined mRNA stability (Presnyak et al., 2015). The CSC for a codon is positive if this codon is overrepresented in stable mRNAs and negative if overrepresented in unstable mRNAs. Similar to Hargayaga & Parker (2016), we determined CSC values for S. pombe based on large-scale mRNA half-life measurements (Hasan et al., 2014; Eser et al., 2016). The CSC value for each gene (CSC\( _g \)) is the arithmetic mean of the CSC values of all codons in that gene. As had been seen before (Presnyak et al., 2015; Hargayaga & Parker, 2016), the CSC\( _g \) correlated with other measures of codon optimality such as the percentage of optimal codons or the tRNA adaptation index (tAI) (Appendix Fig S3A). As the SAC genes had short mRNA half-lives, we expected them to have low CSC\( _g \) values. Indeed, mad2\( ^+ \) and mad3\( ^+ \) were among the 20% of protein-coding genes with the lowest CSC\( _g \) values (Fig 3A and B). This result was independent of which large-scale mRNA half-life data or which correlation parameter was used (Appendix Fig S3C and D). These results raise the interesting possibility that codon usage in mad2\( ^+ \) and mad3\( ^+ \) contributes to their short mRNA half-life. The mad1\( ^+ \) gene showed different characteristics, which we will discuss below.

**Codon-optimization increases the mRNA concentration of mad2\( ^+ \) and mad3\( ^+ \)**

To test if codon usage contributes to the short mRNA half-lives, we codon-optimized mad2\( ^+ \) and mad3\( ^+ \) and inserted the codon-optimized sequence at the respective endogenous locus (Fig 3C; Appendix Fig S3B and F). The GFP tag, which remained unchanged, mitigated but did not abolish the effect of the codon-optimization on the CSC\( _g \) value of the fusion genes (Appendix Fig S3B). An increase in mRNA half-life should result in an increased steady-state mRNA number if synthesis was unchanged. Indeed, we found an increased mRNA number for codon-optimized mad2\( ^+ \) and mad3\( ^+ \) compared with the wild-type gene (Fig 3D). Cytoplasmic mRNAs showed a 27% increase (Fig EV3). For mad2\( ^+ \), the increase was restricted to the cytoplasm and not observed in the nucleus, strongly suggesting stabilization of the mRNA (Fig EV3).

In S. cerevisiae, the RNA helicase Dhh1 (S. pombe Ste13) is involved in specifically lowering the mRNA half-life of genes with a high fraction of nonoptimal codons (Radhakrishnan et al., 2016; Cheng et al., 2017; Webster et al., 2018; Buschauer et al., 2020). Consistently, we observed that the deletion of ste13\( ^+ \) significantly increased mad2\( ^+ \) and mad3\( ^+ \) mRNA half-lives—from about 8 to 14 min for mad2\( ^+ \), and 5 to 10 min for mad3\( ^+ \) (Figs 3E and EV2). This indicates that mad2\( ^+ \) and mad3\( ^+ \) mRNA are subject to Ste13-mediated degradation. The steady-state mRNA numbers were not greatly affected by ste13\( ^+ \) deletion (Figs 3D and EV2B, EV3). This is consistent with a global “buffering” of mRNA concentrations that has been observed in budding yeast when mRNA degradation rates or synthesis rates are globally reduced (Haimovich et al., 2013; Sun et al., 2013; Timmers & Tora, 2018; Fischer et al., 2020). Buffering has been found to be a global phenomenon, not observed when the mRNA of single genes is stabilized (Garcia-Martinez et al., 2021). This may explain why mRNA numbers increased after codon-optimization, but not after ste13\( ^+ \) deletion. Overall, our results support the hypothesis that nonoptimal codons in mad2\( ^+ \) and mad3\( ^+ \) contribute to the short mRNA half-life of these genes.

**Codon-optimization, but not ste13\( ^+ \) deletion, increases the protein concentration of Mad2 and Mad3**

To ask whether the consequences of codon-optimization propagate to the protein level, we quantified Mad2- and Mad3-GFP protein expressed from the wild-type or codon-optimized genes. Both immunoblotting (Fig 4A and C) and fluorescence microscopy (Fig 4D and E) showed an increase in protein concentration after codon-optimization, which can partly be explained by the increase in mRNA (Fig 3) and might be enhanced by an increased translation efficiency. In contrast, the Mad2 and Mad3 protein concentrations in ste13\( ^+ \) cells remained largely stable when analyzed by immunoblotting (Fig 4B and C), consistent with the mRNA results (Fig 3D). Altogether, these data support that codon usage bias toward nonoptimal codons in mad2\( ^+ \) and mad3\( ^+ \) lowers their protein concentration but supports a short mRNA half-life, thereby establishing a gene expression pattern that lowers cell-to-cell variability.

**mad1\( ^+ \) expression regulation differs from that of mad2\( ^+ \) and mad3\( ^+ \)**

The mad1\( ^+ \) gene shares a short mRNA half-life with mad2\( ^+ \) and mad3\( ^+ \) (Fig 2D). Different from mad2\( ^+ \) and mad3\( ^+ \), though, mad1\( ^+ \) has a higher fraction of optimal codons and a CSC\( _g \) value above the median of all protein-coding S. pombe genes (Fig 3A and B;
Figure 3.
Figure 3. Codon-optimization increases the steady-state mRNA numbers of mad2 and mad3.
A The mean CSC value for each S. pombe gene (CSCg) relative to protein number per cell by mass spectrometry (Carpy et al, 2014). CSC was determined using the mRNA half-life data by Eser et al (2016) as described in Methods. Colored dots highlight proteins of interest. For Mad2 and Bub1, no protein abundance data was available. 
B Cumulative frequency distribution of the CSCg values for protein-coding S. pombe genes. The position of spindle assembly checkpoint genes is highlighted. 
C Schematic of the mad2 and mad3 genes. Regions coding for important structural features are highlighted. Black lines in the bottom graph indicate synonymous codon changes in the codon-optimized version. 
D Scatter plots of whole-cell RNA counts versus cell length. Solid lines are regression curves from generalized linear mixed model fits (gray: wild type, black: codon-optimized or ste13Δ). Dashed lines: 95% bootstrap confidence bands for the regression curves. Model estimates of the ratio relative to wild-type mRNA are included with bootstrap 95% confidence interval in brackets. Two to five replicates per genotype. 
E Time course of RNA abundances by qPCR following metabolic labeling and removal of the labeled pool (two independent experiments). Solid lines: regression curves from generalized linear mixed model fits (dark blue: ste13Δ, light blue: ste13Δ), excluding t = 0 to accommodate for non-instantaneous labeling by 4tU. Shaded area: 95% bootstrap confidence band for ste13Δ, dashed lines: 95% bootstrap confidence band for ste13Δ. Half-life estimates are included with 95% bootstrap confidence intervals in brackets. See Fig EVOC for additional statistics. The ste13Δ data are the same as in Fig 2. 
Source data are available online for this figure.

Figure 4. Codon-optimization increases the protein concentrations of Mad2 and Mad3. 
A Immunoblot of S. pombe protein extracts from cells expressing wild-type (WT) or codon-optimized (co) Mad2-GFP or Mad3-GFP probed with antibodies against GFP and Cdc2 (loading control). Lanes 3–5 are a 1:1 dilution series of the extract from cells expressing the codon-optimized version. 
B Immunoblot of protein extracts from wild-type (WT) or ste13Δ strains probed with antibodies against Mad2, Mad3, and tubulin (loading control). A 1:1 dilution series was loaded for quantification. 
C Estimates of the protein concentration relative to wild-type conditions from experiments such as in (A) and (B). Bars are experimental replicates, dots are technical replicates. Two-sided t-tests: P = 0.03 (Mad2-co), 0.004 (Mad3-co), 0.82 (Mad2 ste13Δ), 0.15 (Mad3 ste13Δ). 
D Whole-cell GFP concentration from individual live-cell fluorescence microscopy experiments (a.u., arbitrary units). Boxes show median and interquartile range (IQR); whiskers extend to values no further than 1.5 times the IQR from the first and third quartile, respectively. Codon-optimized concentration significantly higher than wild type for both genes (generalized linear mixed model). Mad2-GFP: n = 468 and 413; Mad2-co-GFP: n = 206 and 366; Mad3-GFP: n = 224 and 127; Mad3-co-GFP: n = 160, 450 and 212 cells. 
E Representative images from one of the experiments in (D). A single Z-slice is shown. Cells are outlined in gray. 
Source data are available online for this figure.
Appendix Fig S3A and B). This was surprising because we expected similar features within the SAC network. Unlike for mad2 and mad3, the mad1 mRNA number did not increase after codon-optimization, but rather decreased slightly (Figs 5A and B, and EV5). A second codon-optimized mad1 whose sequence was considerably different from the first (77% nucleotide identity; Appendix Fig S3F and Appendix Table S3) showed the same trend (Figs EV4A and EV5). Similar to mad2+ and mad3+, mad1+ mRNA half-life was still prolonged in ste13Δ cells (from 6 to 10 min; Fig 5C), but unlike for mad2+ and mad3+ not reaching statistical significance (Fig EV4E). Thus, the short mad1+ mRNA half-life is less dependent on codon usage bias and Ste13, and hence, different modes of regulation bring about the short mRNA half-life of these SAC genes.

The ecm33+ control mRNA was strongly stabilized in ste13+-deleted cells (Figs 5C and EV4E), despite a high fraction of optimal codons in ecm33+ mRNA (Fig 5D). This highlights that—despite some overall correlation—the relationships between codon optimality, mRNA half-life, and susceptibility to ste13+ deletion are far from predictable (Fig EV4F) (He et al., 2018). It is worth noting that Ecm33 is a plasma membrane-binding protein. The budding yeast and human orthologs of Ste13 (Dhh1 and DDX6, respectively) influence translation and mRNA degradation of membrane-binding proteins, and budding yeast Dhh1 has been shown to bind Ecm33 and its paralog Pst1 (Jungfleisch et al., 2017; Weber et al., 2020). If conserved in S. pombe, this could explain the strong destabilizing effect of Ste13 on ecm33+ mRNA.

Codon-optimization of mad1+ decreases its protein concentration

Unlike Mad2- and Mad3-GFP, whose protein concentration increased after codon-optimization, that of Mad1-GFP decreased, both by immunoblotting (Fig 6A and C) and by fluorescence microscopy (Fig 6D and E). Mad1 protein formed from the codon-optimized mRNA had a similar stability to that formed from wild-type mRNA (Appendix Fig S4A and B) and still bound Mad2 (Appendix Fig S4C). The reduction, rather than increase, in protein concentration after codon-optimization of mad1+ corroborates that the codon usage pattern of mad1+ serves a different purpose than that of mad2+ and mad3+. Deletion of ste13+ had hardly any influence on the Mad1 protein concentration (Fig 6B and C), consistent with the largely unchanged mRNA concentration (Fig 5B).

We previously found that SAC function was well preserved when Mad1 levels were lowered to 30% (Heinrich et al., 2013). Consistently, we did not observe an obvious growth defect when cells expressing codon-optimized mad1 were grown in the presence of the microtubule drug benomyl (Fig EV1B), and we did not observe a SAC defect in a live-cell imaging assay where microtubules were depolymerized (Appendix Fig S4D and E). To test SAC function in a more sensitive assay, we deleted the gene for the microtubule-interacting protein Alp7 (Sato et al., 2003). This also activates the SAC, but less robustly than microtubule-depolymerization. Using this assay, cells expressing codon-optimized mad1 tended to exit mitosis more quickly than cells expressing wild-type mad1+ (Fig 6F; Appendix Fig S4F). The difference did not reach the level of statistical significance but was reproducible with independent strains. This suggests that synonymous codon changes, without any change in the protein sequence, can impair SAC function.

Upstream and downstream sequences of mad1+ are insufficient for proper expression

The lower mRNA concentration after mad1 codon-optimization (Figs 5B and EV4A) suggested that the concentration of mad1+ mRNA is not purely determined by regulatory sequences upstream and downstream of the coding sequence. This is supported by our observation that merely fusing GFP to mad1+, without altering surrounding sequences, significantly increases its mRNA number (Figs 1E and EV1E). Further supporting this notion, but rather surprisingly, we found that replacing the mad1+ coding sequence with GFP produced neither significant amounts of mRNA nor protein (Appendix Fig S5A and B). This again contrasted with the mad2+ and mad3+ genes, which produced comparable amounts of mRNA and protein when the original coding sequence was replaced with GFP (Appendix Fig S5C and D). Hence, the sequences surrounding the mad1+ coding sequence are insufficient to establish mad1+like expression, and contributions from the coding sequence are required. Preserving the first 66 or 108 base pairs of mad1+ partly rescued both mRNA and protein levels but not completely (Appendix Fig S5A and B). This while suggests that the 5′ region of the mad1+ coding sequence carries signals that are important for mRNA synthesis or stabilization, some other genes contain sequences that can compensate. Introducing an nmt1+-GFP fusion gene or fusions between S. cerevisiae GCN4 and N-terminally truncated versions of S. pombe mad1+ (Heinrich et al., 2014) allowed for expression from the mad1+ locus (Appendix Fig S5A and B). What these genes share, that GFP does not, remains unclear.

Altogether, these results indicate that mad1+ expression has some unique aspects: mad1+ uses a different mode for reducing mRNA half-life than mad2+ or mad3+, and its coding sequence carries elements that help transcribe, stabilize, or translate RNA.

Mad1 homodimers assemble co-translationally

We considered whether mad1+ may have a certain codon usage pattern to facilitate protein production or complex formation (Liu et al., 2021). Mad1 forms a homodimer through a long N-terminal coiled-coil (Sironi et al., 2002; Piano et al., 2021), but—except in a very recent genome-wide study (Bertolini et al., 2021)—how this homodimer forms has not been examined. If formation was cotranslational rather than post-translational, this may require a certain pattern of codon usage for proper complex formation. To assess dimer formation, we examined cells expressing both tagged and untagged Mad1. If Mad1 dimer formation was post-translational, it should be possible to observe interactions between tagged and untagged Mad1. However, in haploid strains expressing a C-terminally GFP-tagged and an untagged mad1+ gene, a GFP immunoprecipitation almost exclusively precipitated Mad1-GFP, but not untagged Mad1 (Fig 7A). In contrast, a Mad1 immunoprecipitation precipitated Mad1-GFP and Mad1 in approximately the same ratio in which they were present in the extract. These experiments used a monomeric version of GFP. Thus, it is unlikely that this pattern is driven by dimerization of GFP. With two versions of Mad1 being expressed, a slight bias toward the form that is being pulled down would be expected even when heterodimers between these forms were generated with equal likelihood as homodimers (Fig EV6A). At a 1:1 ratio of the isoforms
in the extract, a 2:1 ratio would be expected in an immunoprecipitation or pull-down. However, the bias that we observed always exceeded the expected bias, usually vastly (Figs 7 and EV6). Hence, we propose that Mad1 forms homodimers between isoforms more efficiently than heterodimers. This is most easily explained by co-translational assembly of Mad1 dimers from the nascent chains of two ribosomes translating mad1+ from the same mRNA molecule (Fig 7B).

Figure 5. Codon-optimization and ste13+ deletion do not significantly affect the steady-state mRNA number of mad1+.

A Schematic of the mad1+ gene. Regions coding for important structural features are highlighted. Black lines in the bottom graph indicate synonymous codon changes in the codon-optimized version.

B Scatter plots of whole-cell mRNA counts versus cell length. Solid lines are regression curves from generalized linear mixed model fits (gray: wild type, black: codon-optimized or ste13Δ). Dashed lines: 95% bootstrap confidence bands for the regression curves. Model estimates of the ratio relative to wild-type mRNA are included with bootstrap 95% confidence interval in brackets. Two to three replicates per genotype.

C Time course of RNA abundances by qPCR following metabolic labeling and removal of the labeled pool (two independent experiments). Solid lines: regression curves from generalized linear mixed model fits (dark = ste13+, light = ste13Δ), excluding t = 0 to accommodate for non-instantaneous labeling by 4tU. Shaded area: 95% bootstrap confidence band for ste13+. Half-life estimates are included with 95% bootstrap confidence intervals in brackets. See Fig EV4E for additional statistics. The ste13+ data are the same as in Fig 2.

D Comparison between mean CSC values for selected genes (CSCg) and mRNA half-life measured with or without deletion of ste13+. mRNA half-life estimates from Figs 3E and 5C, and EV4D.

Source data are available online for this figure.
We further corroborated this finding using diploid strains expressing Mad1-GFP and Mad1-Strep from the two endogenous loci. Again, a GFP-immunoprecipitation isolated Mad1-GFP but very little Mad1-Strep, whereas a Strep pull-down isolated Mad1-Strep but very little Mad1-GFP (Figs 7C and EV6B). We obtained similar results after in vitro translation of Mad1 (Fig EV6C): when Mad1-GFP and Mad1-FLAG-His were co-translated in a rabbit reticulocyte lysate, a subsequent GFP immunoprecipitation isolated very little Mad1-FLAG-His, and a His pull-down isolated very little Mad1-GFP. Heterodimerization between C-terminal Mad1 fragments has previously been reported in an in vitro translation (Kim et al, 2012). However, in our experiments, even C-terminal fragments showed a strong bias toward the form that was being precipitated, both in yeast extracts and after in vitro translation (Appendix Fig S6). To exclude that heterodimer formation between Mad1-GFP and untagged Mad1 was nonphysiologically prevented by the large GFP tag, we tested a combination of Mad1-FLAG-His and untagged Mad1 in an in vitro translation. Again, his pull-down almost exclusively isolated Mad1-FLAG-His, whereas a Mad1 immunoprecipitation isolated both forms in approximately the same ratio in which they were present in the extract (Fig 7D).

To further test the idea that Mad1 dimer assembly occurs on a single mRNA molecule (Fig 7B), we examined mad1* mRNA. Consistent with few heterodimers on the protein level, we did not observe co-localization between two different mad1* isoform mRNAs present in the same cell (Fig 7E). Intensity measurements of mRNA FISH spots suggested the presence of single mRNAs, not mRNA doublets, when both untagged mad1* and mad1*-GFP were expressed and mRNA spots were detected with a mad1* probe (Fig 7E, left; EV6D). Further supporting this finding, the number of mad1* mRNA spots that were co-localizing with GFP spots (indicating mad1*-GFP) or not (indicating untagged mad1*) was identical in strains expressing one or both isoforms (Fig 7E, right), indicating that the isoforms do not co-localize. We additionally tested the possibility that mRNAs of the same isoform may co-localize by comparing FISH spot intensities with probes against GFP between mad1+-GFP mRNA and mad3*-GFP mRNA (the latter coding for Mad3 monomers). We did not find any difference in spot intensity (Fig 7F). Hence, we conclude that mad1* mRNAs rarely, if ever, co-localize, and we favor the idea that Mad1 homodimers emerge from two ribosomes co-translating a single mRNA (Fig 7B).

The fact that Mad1 homodimers form co-translationally is consistent with the idea that synonymous codon changes may subtly
Figure 7. Mad1 homodimers assemble co-translationally.

A Top: Immunoprecipitation (IP) with anti-GFP or anti-Mad1 from extracts of haploid S. pombe cells expressing both untagged and GFP-tagged Mad1, probed with antibodies against Mad1 and tubulin; in = input (2.5% of extract for IP), sup = supernatant after IP. Bottom: Comparison between the observed (obs.) and the expected (expect.) ratio between Mad1-GFP and untagged Mad1 in the IP given their ratio in the input (see Fig EV6A), two and one experiment(s), respectively. One more GFP-IP from the same strain was unquantifiable, because no second band was visible in the IP.

B Schematic illustrating that Mad1-Mad1 complex assembly likely takes place co-translationally with only proteins synthesized from the same mRNA being combined.

C Top: Anti-CFP immunoprecipitation (IP) and Strep pull-down (PD) from extracts of diploid cells expressing Mad1-GFP and Mad1-Strep from the two endogenous loci, membrane probed with anti-Mad1; in, input (7% of extract for IP/PD), sup, supernatant after IP/PD. Bottom: as in (A), 2 experiments each. See Fig EV6A for a quantified experiment. The experiment shown at the top and two more GFP-IPs from the same strain were unquantifiable, because no second band was visible in the IP.

D In vitro translation (IVT) of Mad1-flag-His and untagged Mad1 in the presence of [35S]labeled Methionine and Cysteine, followed by Mad1 immunoprecipitation (IP) or His pull-down (PD); in, input (95% of extract for IP/PD), sup, supernatant after IP/PD. An IVT with only untagged Mad1 was used to check for specificity of the His PD (right side). Shown is the autoradiograph after SDS-PAGE with quantification of the Mad1-flag-His to untagged Mad1 ratio in select lanes.

E Test for mRNA dimerization by single-molecule mRNA FISH; probes against mad1+ and GFP. Top: Schematic of genotypes. Example pictures in Fig EV6. Bottom left: Intensity of cytoplasmic mad1+ mRNA spots in the different strains. For the 2 copy strain, a mad1+ spot was classified as mad1+ if it was co-localizing with a GFP spot, and as mad1+ otherwise. Colors as indicated in the schematic. Vertical solid line: peak of each density plot; dashed line: theoretical position of a double-intensity peak. Number of spots analyzed: mad1+=923, mad1+ (2 copy strain) = 637, mad1+ (2 copy strain) = 992, mad1+ (2 copy strain) = 1,699. Bottom right: Counts of cytoplasmic mad1+ or mad1+GFP mRNA from the same experiment with generalized linear mixed model fits as lines. Number of cells: 1 copy strain mad1+= 478, 2 copy strain = 327, 1 copy strain mad1+GFP = 1,299.

F Experiment similar to (E), except that cells expressing both mad1+GFP and mad3+GFP from the respective endogenous locus were probed with FISH probes against mad1+ and GFP mRNA. A GFP spot was classified as mad1+GFP if it was co-localizing with a mad1+ spot (arrowheads), and as mad3+GFP otherwise. The intensity of GFP spots was quantified. Vertical solid line: peak of each density plot; dashed line: theoretical position of a double-intensity peak. Number of spots analyzed: mad1+GFP = 987, mad3+GFP = 1,299.

Source data are available online for this figure.
improve complex formation and therefore translation efficiency and mRNA stability. Overall, these results suggest that codon usage bias within mad1* contributes to maintaining proper mRNA and protein levels, possibly by supporting Mad1 folding and dimerization.

Discussion

Proteins are the workhorses of cells. The deployment of this workhorse army is controlled by regulatory elements encoded in DNA that are still incompletely understood. The spindle assembly checkpoint is sensitive to expression changes, and we therefore asked which features of gene expression may be important for its proper function. Our results suggest that a combination of short mRNA half-lives and long protein half-lives is important to keep protein variability low. We also find that—despite their closely shared function—mad1* differs in its expression features from mad2* and mad3*. The coding sequences of mad2* and mad3* contribute to the short mRNA half-life of these genes, whereas that of mad1* contributes to maintaining mRNA (Appendix Fig S5) and protein levels (Fig 6). We propose that the choice of synonymous codons in mad1* is optimized for the formation of the Mad1 homodimer, and ultimately the Mad1/Mad2 complex.

Short mRNA half-life of constitutively expressed SAC genes favors low noise

The short mRNA half-lives of mad1*, mad2*, and mad3*, along with their long protein half-lives, can explain the low protein noise of SAC genes despite low and variable mRNA numbers (Figs 1 and 2) (Thaats-tai & van Oudenaarden, 2001). In human cells, a long protein half-life has also been shown to affect the expression rates of different mRNA numbers (Raj et al, 2006). Human Mad1, Mad2, and BubR1 (Mad3 ortholog) are also highly stable proteins (Suijkerbuijk et al, 2010; Varetti et al, 2011; Schweizer et al, 2013; Rodriguez-Bravo et al, 2014), which will support stable protein concentrations over time and between cells. SAC genes are certainly not unique in combining a short mRNA and long protein half-life to achieve low noise. Other constitutively expressed genes that produce low or modest amounts of protein will likely show a similar behavior. Keeping noise low in this manner requires a high turnover of mRNA that consumes some energy cost. An alternative way to keep protein noise low would be to produce the same amount of protein from a larger number of more stable mRNA molecules (Appendix Fig S2C). Several side-effects likely prohibit this solution as a general strategy. For example, the cytoplasm would be much more crowded with mRNAs, and stable mRNAs may accumulate chemical damage. Indeed, genes using an expression strategy of high transcription and low translation rates are exceedingly rare among different eukaryotes (Hauser et al, 2019).

Different SAC genes employ different strategies for achieving short mRNA half-life

The half-life of an mRNA is influenced by sequence motifs, codon usage, and other factors that influence translation. Currently, known factors predict around 50–60% of mRNA half-life in budding yeast (Neymotin et al, 2016; Cheng et al, 2017). At least two elements seem to play a role for mad2* and mad3*: Our data suggest that the mRNA half-lives are shortened by a high fraction of nonoptimal codons (Fig 3); in addition, the mad2* and mad3* 3′ UTRs contain sequence motifs that are associated with a short mRNA half-life (Eser et al, 2016). We previously found higher mRNA numbers after traditional tagging, which changed the 3′ UTR to that of a highly expressed gene (Heinrich et al, 2013), suggesting that the predicted motifs in the 3′ UTR may indeed be functional. For mad1*, in contrast, overall codon usage bias seems to play a lesser role (Fig 5), and the mad1* 3′ UTR does not contain reported motifs implicated in half-life shortening (Eser et al, 2016). We suspect that other elements that influence translation efficiency may be important. Generally, less efficiently translated mRNAs are less stable (Hanson & Coller, 2018), and mad1* seems to be translated less efficiently than mad2* or mad3* (Rubio et al, 2020).

Formation of the Mad1/Mad2 complex involves co-translation assembly of the Mad1 dimer but not synchronous co-translational assembly of the tetramer

Mad1 and Mad2 form a tight tetrameric complex (Sironi et al, 2002; Kim et al, 2012), but how this complex assembles is unknown. Our experiments suggest that the Mad1 homodimer forms between two polypeptides translated from the same mRNA, and that Mad1 molecules translated from different mRNA molecules associate less efficiently with each other, if at all (Fig 7). This assembly mode is further supported by a recent study in human cells, which analyzed footprints of ribosome disomes on mRNA and found wide-spread evidence for co-translational assembly of protein homomers (Bertolini et al, 2021). Coiled-coils were the most prominent class driving co-translational assembly, and co-translational assembly was more likely when the dimerization domain was N-terminal. Mad1 meets both these criteria and was indeed identified in this study as probably assembling co-translationally.

At least two studies have expressed Mad1 N-terminal fragments and full-length Mad1 from two different loci and have interpreted the failure to see association between those two as an inability of the N-terminal fragment to dimerize (Jin et al, 1998; Ji et al, 2018). Based on the evidence for co-translational homodimer assembly, we suggest that the capacity of an N-terminal Mad1 fragment to dimerize would need to be based on assessing self-association rather than assessing association with Mad1 expressed from a different locus. Of note, C-terminal Mad1 fragments also dimerize, possibly post-translationally (Kim et al, 2012), although our own experiments still suggest a preference of homodimerization (Appendix Fig S6).

While we propose that assembly of the Mad1 homodimer occurs co-translationally, the assembly of the Mad1/Mad2 tetramer does not occur in synchronous co-translational fashion, as the mRNAs for mad1* and mad2* do not co-localize in the cytoplasm (Fig 1). This leaves open the possibility of post-translational assembly of the tetramer or of asynchronous co-translational assembly, where one protein is already fully formed and binds the other that is being translated (Duncan & Mata, 2011; Shiber et al, 2018). Formation of the C-Mad2/Cdc20 complex necessitates catalysis (Kulikian et al, 2009; Lad et al, 2009; Simonetta et al, 2009; Faesen et al, 2017; Piano et al, 2021), making it likely that C-Mad2/Mad1 formation also needs to be facilitated. We favor the idea that the tetramer assembles while one of the proteins is being translated, and it will be interesting to test whether the mad1* mRNA binds Mad2 protein or vice versa to facilitate such an assembly. It will also be interesting
to examine whether different eukaryotes use the same assembly pathway for the highly conserved Mad1/Mad2 complex.

**Potential SAC malfunction from synonymous mutations**

Overall, our data suggest that the coding sequences of mad1+, mad2+, and mad3+ modulate gene expression. Hence, even synonymous mutations carry some risk of impairing the SAC. We suspect that mad1+ is most susceptible to single synonymous substitutions, given the need for co-translational homodimer assembly (Fig 7), which may be facilitated by controlling the speed of ribosome movement (Liu et al., 2021). In *S. pombe*, a cluster of nonoptimal codons follows the coiled-coil region of mad1+ (Appendix Figs S3E and S7), which may ensure that the N-terminal coiled-coil is fully formed before the remainder of Mad1 is translated.

It will be interesting to test whether synonymous mutations found in cancer samples can modulate SAC gene expression or function. Within MAD2L1 (H.s. mad2+), synonymous mutations detected in cancer samples seem to cluster in a conserved region with high CSC values preceding the “seat belt,” (Appendix Fig S7) suggesting that codon usage bias in this region may be functionally important. Although most synonymous mutations will only have small effects, they may fuel carcinogenesis. This is particularly true in the context of the SAC, because drastic impairment is more likely to be detrimental for cancer cells, whereas subtle impairment may promote carcinogenesis (Kops et al., 2004; Funk et al., 2016; Cohen-Sharir et al., 2021; Quinton et al., 2021). Synonymous mutations and changes in tRNA expression have been implicated in carcinogenesis (Sauna & Kimchi-Sarfaty, 2011; Supek et al., 2014). Our data suggest that this may partly occur by impairing the SAC.

**Materials and Methods**

### Reagents and Tools table

| Reagent/Resource | Reference or Source | Identifier or Catalog Number |
|------------------|---------------------|-----------------------------|
| **Experimental models** | | |
| *Schizosaccharomyces pombe* strains | This study | Appendix Table S1 |
| *Saccharomyces cerevisiae* strain | Nick Buchler, NC State University, USA | Appendix Table S1 |
| **Recombinant DNA** | | |
| sgRNA sequences | This study | Appendix Table S2 |
| Codon-optimized mad1, mad2, and mad3 | This study | Appendix Table S3 |
| PCR fragments for *in vitro* transcription | This study | Appendix Table S6 |
| **Antibodies** | | |
| Mouse anti-Cdc13 (monoclonal) | Novus | Cat # NB200-576; RRID: AB_10003103 |
| Rabbit anti-Cdc2 (polyclonal) | Santa-Cruz | Cat # sc-53; RRID: AB_2074908 |
| Mouse anti-GFP (mix of 2 monoclonals) | Roche | Cat # 11814460001; RRID: AB_390913 |
| Rabbit anti-Mad1 (polyclonal, against peptide ADSPRDPFQSRSQLC) | Heinrich et al (2013), PMID: 24161933 | N/A |
| Rabbit anti-Mad2 (polyclonal, against recombinant protein) | Sewart and Hauf (2017), PMID: 28366743 | N/A |
| Rabbit anti-Mad3 (polyclonal, against recombinant protein) | Sewart and Hauf (2017), PMID: 28366743 | N/A |
| Rabbit anti-Strep-tag II (monoclonal, recombinant) | Abcam | Cat # ab180957 |
| Mouse anti-tubulin | Sigma | Cat # T5168; RRID: AB_477579 |
| Goat anti-mouse HRP | Jackson ImmunoResearch Labs | Cat # 115–035-003; RRID: AB_10015289 |
| Goat anti-rabbit HRP | Jackson ImmunoResearch Labs | Cat # 111–035-003; RRID: AB_2313567 |
| **Oligonucleotides and other sequence-based reagents** | | |
| FISH probes | This study | Appendix Table S4 |
| qPCR primers | This study | Appendix Table S5 |
| **Chemicals, enzymes, and other reagents** | | |
| 4-thiouracil (4tU) | Chem Impex | Cat # 21484 |
| MTSEA biotin-XX | Biotium | Cat # 90066 |
| Cycloheximide (from Streptomyces griseus) | Chem Impex | Cat # 00083 |
| Wizard SV Gel and PCR Clean-Up System | Promega | Cat # A9285 |
| Reagent/Resource                                      | Reference or Source       | Identifier or Catalog Number |
|------------------------------------------------------|---------------------------|------------------------------|
| SuperScript IV First Strand Synthesis System         | ThermoFisher              | Cat # 18091050               |
| HiScribe T7 ARCA mRNA Kit (with tailing)             | New England Biolabs       | Cat # E20605                 |
| Monarch RNA Cleanup Kit                              | New England Biolabs       | Cat # T20405                 |
| Rabbit Reticulocyte Lysate, Nuclease-Treated         | Promega                   | Cat # L4960                  |
| EasyTag EXPRESS 35S Protein Labeling Mix             | Perkin Elmer               | Cat # NEG772007MC            |
| SUPERase•In RNase Inhibitor                          | ThermoFisher              | Cat # ACM2694                |
| SuperSignal West Pico PLUS Chemiluminescent Substrate| ThermoFisher              | Cat # 34580                  |
| cComplete, EDTA-free Protease Inhibitor Cocktail     | Roche                     | Cat # 04693132001            |
| Halt Protease Inhibitor Cocktail, EDTA-Free (100X)   | ThermoFisher              | Cat # 87785                  |
| PhosSTOP                                             | Roche                     | Cat # 04906837001            |
| Halt Phosphatase Inhibitor Cocktail                  | ThermoFisher              | Cat # 78420                  |
| Dynabeads Protein G                                  | ThermoFisher              | Cat # 10003D                 |
| Dynabeads His-Tag Isolation and Pull-down XT beads   | IBA Lifesciences           | Cat # 2-4090-002             |
| Dynabeads MyOne Streptavidin C1                     | Thermo Fisher              | Cat # 65001                  |
| Oligo d(T)25 Magnetic Beads                          | New England Biolabs       | Cat # 514195                 |
| Pierce BCA Protein Assay Kit                         | ThermoFisher              | Cat # 23225                  |
| EMM (Edinburgh’s Minimal Medium)                    | MP Biomedicals             | Cat # 114110022              |
| Lectin                                               | Sigma                     | Cat # L1395                  |

**Software**

| Software                                      | Reference                                      | RRID: SCR_002798 |
|-----------------------------------------------|------------------------------------------------|------------------|
| Fiji/ImageJ                                   | Schindelin et al (2012), PMID: 22743772        | https://imagej.net/software/fiji/|
| SoftWoRx                                      | Applied Precision, GE Healthcare               | https://download.cytivalifesciences.com/cellanalysis/download_data/softWoRx6.5.2/SoftWoRx.htm;|
| MetaMorph                                     | Molecular Devices                              | Version 7.10.1    |
| YeaZ                                          | Dietler et al (2020), PMID: 33184262           | N/A              |
| ImageLab                                      | Bio-Rad Laboratories                           | Version 6.0.1 build 34 |
| Matlab                                        | Mathworks                                      | https://www.mathworks.com; |
| FISH-Quant                                     | Mueller et al (2013), PMID: 23338861           | N/A              |
| Trainable Weka Segmentation                    | Arganda-Carreras et al (2017), PMID: 28369169  | N/A              |
| Prism 9                                       | GraphPad Software, Inc                         | https://www.graphpad.com; |
| R                                             | Cran.R                                         | https://cran.r-project.org; |
| R studio                                      | N/A                                            | https://www.rstudio.com; |
| tidyverse package                             | Cran.R                                         | https://tidyverse.tidyverse.org; |
| ggplot2 package                               | Cran.R                                         | https://ggplot2.tidyverse.org; |
| alphashape3d package                          | Cran.R                                         | https://CRAN.R-project.org/package=alphashape3d, Version 1.3.1 |
| boxcoxmix package                             | Cran.R                                         | https://cran.r-project.org/src/contrib/Archive/boxcoxmix/, Version 0.28 |
| broom package                                 | Cran.R                                         | https://CRAN.R-project.org/package=broom, Version 0.7.9 |
| broom.mixed package                            | Cran.R                                         | https://CRAN.R-project.org/package=broom.mixed, Version 0.2.7 |
| cairo package                                 | Cran.R                                         | https://CRAN.R-project.org/package=Cairo, Version 1.5–12.2 |
Reagents and Tools table (continued)

| Reagent/Resource | Reference or Source | Identifier or Catalog Number |
|------------------|---------------------|------------------------------|
| cowplot package  | Cran.R              | https://cran.r-project.org/package=cowplot; RRID: SCR_018081, Version 1.1.1 |
| descTools package| Cran.R              | https://cran.r-project.org/package=DescTools, Version 0.99.43 |
| egg package      | Cran.R              | https://CRAN.R-project.org/package=egg; Version 0.4.5 |
| geometry package | Cran.R              | https://CRAN.R-project.org/package=geometry; Version 0.4.5 |
| gridExtra package | Cran.R             | https://CRAN.R-project.org/package=gridExtra; Version 2.3 |
| lemon package    | Cran.R              | https://CRAN.R-project.org/package=lemon; Version 0.4.5 |
| lime4 package    | Cran.R              | https://cran.r-project.org/web/packages/lime4/index.html; RRID: SCR_015654 |
| mclust package   | Cran.R              | https://cran.r-project.org/package=mclust |
| nabor package    | Cran.R              | https://cran.r-project.org/package=nabor |
| pbkrtest package | Cran.R              | https://cran.r-project.org/package=pbkrtest |
| plotly package   | Cran.R              | https://plotly.com/r/; RRID: SCR_013991, Version 4.1.0 |
| plyr package     | Cran.R              | https://cran.r-project.org/package=plyr |
| readxl package   | Cran.R              | https://cran.r-project.org/web/packages/readxl/index.html; RRID: SCR_018083, Version 1.3.1 |
| rgl package      | Cran.R              | https://CRAN.R-project.org/package=rgl; Version 0.107.14 |
| sf package       | Cran.R              | https://CRAN.R-project.org/package=sf |
| shotGroups package| Cran.R             | https://CRAN.R-project.org/package=shotGroups; Version 0.8.1 |
| spatstat package | Cran.R              | https://cran.r-project.org/package=spatstat |
| Other            |                     |                              |
| Mixer mill MM400 | Retsch              | Cat # 20.745.0001 |
| Grinding jar 10 ml| Retsch            | Cat # 01.462.0236 |
| Grinding jar 25 ml| Retsch            | Cat # 01.462.0213 |
| Adapter for reaction vials | Retsch | Cat # 22.008.0008 |
| Glass beads, acid-washed | Sigma | Cat # G8772 |
| μ-Slide 8-well, glass bottom | Ibidi | Cat # 80827 |
| YO4C Microfluidic Plate for Haploid Yeast | Celliasic / Sigma | Cat # YO4C-02-SPK |
| Invitrogen NuPAGE 4 to 12%, Bis-Tris, 20-well | Invitrogen | Cat # WG1402BOX |
| Invitrogen NuPAGE 4 to 12%, Bis-Tris, 20-well | Invitrogen | Cat # NP0322BOX |
| Immobilon-P PVDF membrane | Millipore | Cat # IPVH00010 |

Methods and Protocols

Yeast strains

Yeast strains are listed in Appendix Table S1. Tagging of nmt1+ and deletion of ste13+ and alp7+ were performed by conventional PCR-based gene targeting (Bähler et al., 1998). Marker-less insertion at the endogenous locus was performed either by replacement of a counter-selectable rpl42-hphNT1 cassette in an rpl42::cyhR(sP56Q) background (Roguev et al., 2007) or by using CRISPR/Cas9 (Jacobs et al., 2014). Sequences used for targeting Cas9 are listed in Appendix Table S2. The mad2+-ymEGFP strain contains a single, silent (AGG to AGA) PAM site mutation at amino acid position 173 of Mad2. The mad3+-ymEGFP strain contains a single, silent (TTG to TTA) PAM site mutation at amino acid position 199 of Mad3. Yeast, monomeric-enhanced GFP (ymEGFP) was derived from yEGFP (yeast codon-optimized green fluorescent protein (Watson et al., 2008)) by mutation of Alanine 206 to Arginine (A206R), which is expected to reduce dimerization (Zacharias et al., 2002). Codon-optimization used proprietary algorithms by two different companies, and sequences are listed in Appendix Table S3. The haploid strain with two differently tagged versions of mad1+ has mad1+-ymEGFP along with 110 bp upstream and 164 bp downstream of the coding sequence integrated between the leu1+ and apc10+ gene.

Yeast cultures

Schizosaccharomyces pombe cultures were grown at 30°C either in rich medium (yeast extract supplemented with 0.15 g/l adenine; YEA) or in Edinburgh minimal medium (EMM, MP Biomedicals,
4110012) supplemented with 0.2 g/l leucine, 0.15 g/l adenine or 0.05 g/l uracil if required (Petersen & Russell, 2016). When cultures in minimal medium were started at low concentration, “preconditioned medium” was added to a maximum of 50%. Preconditioned medium was obtained by growing cells in EMM and then removing the cells by filtration. For growth assays, cells were grown in YEA to a concentration of around 1 × 10^6 cells/ml, diluted to 4 × 10^5 cells/ml in YEA and further diluted in a 1:5 dilution series. 10 μl were spotted on indicated plates. S. cerevisiae cultures were grown at 30°C in yeast extract supplemented with 20 mg/ml each of Bacto peptone and dextrose (YPD).

**Cycloheximide treatment for determination of protein half-lives**

Cells were grown in EMM (plus supplements required for auxotrophic mutations) to a final concentration of around 1 × 10^7 cells/ml. Cultures were diluted to 8 × 10^6 cells/ml, transferred to a 30°C water bath for 30 min and a sample was taken prior to the addition of cycloheximide (CHX) to a final concentration of 1 mg/ml. Cells were collected at specified time points, spun down at 980 rcf, and frozen in liquid nitrogen before processing.

**In vitro transcription and translation**

The T7 promoter was appended 5’ of the mad1+ transcription start site by PCR. Precise sequences are available in Appendix Table S6. Full-length mad1+ was amplified from cDNA generated using the SuperScript IV First Strand Synthesis System (ThermoFisher). Mad1 fragments 3’ of the intron were amplified from genomic DNA. PCR fragments were purified using the Wizard SV Gel and PCR Clean-Up System (Promega). In vitro transcription was carried out with the HiScribe T7 ARCA mRNA Kit (with tailing) (New England Biolabs) using between 25 and 70 ng/ml template DNA. Reactions were run at 32°C or 37°C for 2 h. RNA was purified using the Monarch RNA Cleanup Kit (New England Biolabs). RNAs were mixed and diluted as required before adding them to rabbit reticulocyte lysate (Promega). Translation reactions contained amino acid mix without Methionine, approx. 1 mM/ml ^35S-Methionine/Cysteine mix (Perkin Elmer, NEG772007MC), 0.2 μl/μl SUPERase^In RNase Inhibitor (ThermoFisher), and between 0.35 and 40 ng/μl RNA. Incubation was at 30°C for 1 h 30 min.

**Denatured whole-cell extracts**

Cells were grown to a final concentration of around 1 × 10^7 cells/ml and collected by centrifugation (1 × 10^6 cells per sample). Supernatant was removed, and cells were washed with 1 ml of 20% trichloroacetic acid (TCA). Supernatant was removed, and cells were resuspended in 500 μl of water. 75 μl of NaOH/beta-mercaptoethanol (final conc. = 0.22 M NaOH, 0.12 M β-ME) was added, and samples incubated on ice for 15 min. 75 μl of 55% TCA was added and samples incubated on ice for another 10 min. Samples were spun at 16,900 rcf for 10 min at 4°C, and supernatant was removed. Pellets were resuspended in 100 μl sample buffer (50 μl of 2x HU buffer [8 M urea, 5% SDS (w/v), 200 mM Tris–HCl pH 6.8 (v/v), 20% glycerol (v/v), 1 mM EDTA (v/v), 0.1% (w/v) bromophenol blue], 40 μl water, and 10 μl of 1 M DTT) to a final concentration corresponding to 1 × 10^9 cells/ml. Approximately 150 μl of acid-washed beads (Sigma) were added before agitation in a ball mill (Mixer Mill 400; Retsch) for 2 min at 30 Hz. Tubes were pierced at the bottom, cell extract was collected from the beads by centrifugation at 2,350 rcf for 1 min and heated at 75°C for 5 min. Typically, the extract equivalent of 2–3 × 10^6 cells was loaded for immunoblotting.

**Immunoprecipitation or pull-down from yeast cell extract**

Asynchronously growing cultures were harvested, washed with deionized water, or with 20 mM Tris pH 7.5/150 mM NaCl, and frozen as droplets in liquid nitrogen. Cell powder was prepared from these droplets using a ball mill (Mixer Mill 400; Retsch) for 30 s at 30 Hz under cryogenic conditions. Cell powder was resuspended in lysis buffer (20 mM Tris pH 7.5, 150 mM NaCl, 5% glycerol, and 0.1% NP-40), and protein concentration was determined by BCA assay (ThermoFisher). For immunoprecipitations, powder was resuspended to a final concentration of 15–20 mg/ml in lysis buffer supplemented with a 5–10x final concentration of protease inhibitor cocktail and a 1x final concentration of phosphatase inhibitor cocktail. Extracts were spun down for 10 min at 4°C and 16,900 rcf. For the input sample, supernatant was mixed with an equal volume of sample buffer (2x HU buffer with 200 mM DTT, or 2x NuPAGE LDS sample buffer with 10% beta-mercaptoethanol) and heated for 3–5 min at 75°C. For immunoprecipitations, Protein G Dynabeads (ThermoFisher) were covalently coupled with anti-GFP antibodies (Roche, 160 μg antibody per 1 ml bead suspension) or anti-Mad1 antibodies (80 μg antibody per 1 ml bead suspension). Strep-tag pull-downs used MagStrep “type3” XT beads (IBA Lifesciences). Immunoprecipitations used around 30 μl bead suspension per 200 μl of extract and were performed for 10 min at 4°C on a rotating wheel. Strep pull-downs used around 200 μl bead suspension per 200 μl of extract and were performed for 45 min to 1 h at 4°C on a rotating wheel. Beads were washed with lysis buffer (IPs), or with a more stringent wash buffer (20 mM Tris pH 7.5, 300 mM NaCl, 5% glycerol, 1% NP-40) for some Strep pull-downs. Elution from anti-GFP or anti-Mad1 beads was performed by the addition of 7–25 μl 100 mM citric acid and gentle agitation for 5 min at 4°C. Samples were neutralized by the addition of 1.5 M Tris pH 9.2, mixed with an equal volume of sample buffer and heated at 75°C for 3 min. Elution from MagStrep beads was performed with sample buffer and incubation at 95°C for 2 min, or 85°C for 5 min.

**Immunoprecipitation or pull-down after in vitro translation**

In vitro translation reactions (IVTs) were diluted to 6- to 13-times the original volume with either Tris buffer for immunoprecipitations (final concentration: 20 mM Tris pH 7.5, 150 mM NaCl, 0.1% NP-40), or with sodium-phosphate buffer for Ni-NTA pull-downs (final concentration: 50 mM sodium-phosphate pH 8.0, 300 mM NaCl, 0.01% Tween-20). Immunoprecipitations used 10 μl Dynabeads suspension, Ni-NTA pull-downs used 40 μl Ni-NTA Dynabeads suspension per 15 μl original IVT (volume prior to dilution). Immunoprecipitations were processed as above, Ni-NTA beads were washed with sodium-phosphate buffer plus 10–20 mM imidazole and 0.1% NP-40 and eluted with sodium-phosphate buffer plus an additional 300 mM imidazole.

**Immunoblotting**

Proteins were separated by SDS-PAGE (NuPAGE, Bis-Tris, MOPS buffer, Thermo Fisher) and transferred onto a PVDF membrane (Immobilon-P, Millipore) in a semidry blotting assembly (Amer sham Biosciences TE-70 ECL) using transfer buffer (39 mM glycine,
48 mM Tris base) with 10% methanol, 0.01% SDS, and 1:1,000 NuPAGE Antioxidant. Membranes were probed with mouse anti-GFP (Roche, 1181446001), rabbit anti-Cdc2 (CDK1, Santa Cruz, SC-53), mouse anti-Cdc13 (cyclin B, Novus, NB200-576), rabbit anti-Mad1 (Heinrich et al., 2013), rabbit anti-Mad2 (Heinrich et al., 2013), rabbit anti-Mad3 (Heinrich et al., 2013), rabbit anti-Strep (Abcam, ab180957 and ab76949), or mouse anti-tubulin (Sigma, T5168). Secondary antibodies were either anti-mouse or anti-rabbit conjugated to HRP (Dianova) and quantified by chemiluminescence using SuperSignal West Dura ECL (ThermoFisher) and imaged on a Bio-Rad Gel Doc system. Chemiluminescence signals were quantified on nonsaturated images using Image Lab software (Bio-Rad). Measurements from a reference dilution series were used to create a standard curve, which was used to determine the concentration of sample relative to the reference. Membranes with radioactive proteins were dried and exposed to a phosphorscreen (GE Healthcare), which was read-out on a Typhoon phosphorimager (GE Healthcare/Cytiva).

Quantification of GFP fusion proteins in single cells (3D segmentation)

To quantify GFP fusion proteins in single cells, cells were grown in EMM (plus supplements that were required for auxotrophic mutations) at 30°C to a final concentration of 6–9 × 10^6 cells/ml. Cultures of GFP-positive and GFP-negative cells were mixed at a 1:1 ratio to a final concentration of 2.5–6.0 × 10^6 cells/ml and incubated for 30 min at 30°C. To ensure a uniform and flat imaging plane, cells were loaded into a Y04C microfluidics trapping plate (Millipore Sigma) and incubated inside a climate-controlled microscope chamber for 2 h at 30°C with constant flow of fresh media. Imaging was performed on a DeltaVision Elite system equipped with a PCO edge sCMOS camera and an Olympus 60x/1.42 Plan APO oil objective. Images were acquired for ymEGFP, tdTomato, and brightfield as 7.2 μm or 10 μm stacks with images separated by 0.1 μm. The acquired image area was 1,024 × 1,024 pixels with 1 x 1 binning. All images were deconvolved using SoftWoRx software. To correct for uneven illumination, deconvoluted fluorescence images were flat-fielded individually for each channel using a custom FIJI script (Baybay et al., 2020).

The Pomegranate image analysis pipeline (Baybay et al., 2020) was used to segment nuclei (using TetR-tdTomato-NLS) and whole cells (using brightfield signal and spherical extrusion of the mid-plane segmentation) (Appendix Fig S1A). We corrected for chromatic aberration and for stretching of distances in the Z direction (Baybay et al., 2020). Further analysis was conducted in R (R Core Team, 2020), and figures were produced using the package ggplot2 (Wickham, 2016).

Only information from mono-nucleated cells for which both the whole cell and the nucleus had been segmented was retained. Cells were excluded if one or more of the following conditions were met: the nuclear segmentation protruded beyond the three-dimensional bounds of the cell; whole-cell segmentation was cut-off by more than two slices because insufficient slices in Z had been recorded; cell was at the image edge and incompletely recorded; the nucleus had an aspect ratio (diameter in Z to diameter in XY) of less than 0.8 or more than 1.2; cell volume was lower than the 0.1st or higher than the 99.9th percentile. Cells with or without GFP signal were distinguished by k-means (k = 2) clustering (Appendix Fig S1D–F), except for Nmt1-GFP, where the threshold for each image was set manually. One image, where the autofluorescence of GFP-negative cells deviated by more than three standard deviations from that of other images, was excluded. One additional image, where the cells had visibly moved during acquisition, was also excluded.

To subtract autofluorescence and other background, we averaged the fluorescence intensity per cell or nuclear volume for GFP-negative cells in an image and subtracted that value from the fluorescence intensity per cell or nuclear volume of each GFP-positive cell in the image. For a rough estimate of absolute concentration in nanomolar, we used our previous estimate of about 70 nM Mad3-GFP in the cell nucleus (Heinrich et al., 2013) and normalized all background-subtracted data to this value. Even after background subtraction, we observed some variation of mean intensities between single images (Appendix Fig S1F), and we could not distinguish whether these differences were a consequence of sampling or came from conditions on the microscope stage while recording the image. We therefore opted to determine the coefficient of variation (CV = standard deviation/mean) for each protein not across all images, but instead for each image separately; Fig 2A shows the variation across images.

Generalized linear mixed models were used to test for differences in whole-cell GFP concentration between wild-type and codon-optimized Mad1, Mad2, and Mad3 from the single-cell measurements. A separate model was fit for each gene and included whole-cell GFP concentration as the response variable and genotype (wild-type versus codon-optimized) as a categorical fixed effect predictor variable. Two nested random effects variables, experimental replicate and image, were also included in the model (random intercepts only). To meet the model assumptions of normality and constant variance, GFP concentration was transformed with a Box Cox transformation using “optim.boxcox” from the boxcoxmix package. Wild-type and codon-optimized genotypes were determined to have significantly different GFP concentrations if the 95% bootstrap confidence interval for the genotype coefficient excluded 0.

Quantification of GFP in single cells (2D segmentation and projection)

For experiments evaluating fluorescence signals after replacing the coding sequences of mad1*, mad2*, and mad3* (Appendix Fig S5), quantification was performed on projections, using 2D segmentation of cells. Cells were grown in minimal medium, collected by centrifugation from liquid cultures, mounted in medium on a slide, and brightfield and fluorescence images were collected immediately at room temperature. At least two slides were prepared and imaged for each strain. Images were recorded on a Zeiss Axiolmager M1, using Xcite Fire LED illumination (Excitellis), a Zeiss Plan-Apochromat 63x/1.40 Oil DIC objective, and an ORCA-Flash4.0LT sCMOS camera (Hamamatsu) with Z sections spaced by 0.2 μm.

Cells were segmented based on an in-focus brightfield image using YeaZ (Dietler et al., 2020). Falsely segmented cells (e.g., background, or cells falsely combined into one) were manually excluded in Fiji. Only cells in the center of the image, where fluorescence illumination was homogeneous, were included. Flatfielding was not performed. The brightfield images were systematically shifted relative to the fluorescence images and we corrected for that error. Quantification of signals was performed on an average projection of the 23 most in-focus Z-slices (covering 4.6 μm, which is slightly...
larger than the width of a typical S. pombe cell). For each image, the median extracellular background in the same central area of the image was subtracted.

**Single-molecule mRNA FISH**

For quantification of mRNA by single-molecule fluorescent in-situ hybridization, cultures of asynchronously dividing cells were grown to a concentration of about 1 x 10^7 cells/ml in EMM. Typically, 2 x 10^8 cells were fixed with 4% paraformaldehyde for 30 min before being washed three times with ice-cold Buffer B (1.2 M sorbitol, 100 mM potassium phosphate buffer pH 7.5) and stored at 4°C before digestion of the cell wall. Cells were resuspended in spheroplast buffer (1.2 M sorbitol, 0.1 M potassium phosphate, 20 mM vanadyl ribonuclease complex [NEB S1402S], 20 μM beta-mercaptoethanol) and digested with 0.002% 100 T zymolase (US Biological Z1000) for approximately 45–75 min. Zymolase reaction was quenched when the addition of water to the cells resulted in around 50% lysed cells. Reactions were quenched with 3 washes of Buffer B. Cell pellets were resuspended in 1 ml of 0.01% Triton X-100 in 1x PBS for 20 min and washed three times with Buffer B. For hybridization of probes, approximately 20–25 μl of CAL Fluor red 610 probes targeting ymEGFP or mad2*, or Quasar 570 probes targeting mad1+ were mixed with 2 μl each of yeast tRNA (Life Technologies) and Salmon sperm DNA (Life Technologies) per reaction. For two-color FISH experiments, 20–25 ng of each probe were combined, resulting in ~ 50 ng of total FISH probes per reaction. Sequences of probes are given in Appendix Table S4. Buffer F (20% formamide, 10 mM sodium-phosphate buffer pH 7.2; 45 μl per reaction) was mixed with the probe solution, heated at 95°C for 3 min, and allowed to cool to room temperature before mixing with Buffer H (4x saline-sodium citrate (SSC) buffer, 4 mg/ml acetylated BSA, 20 mM vanadyl ribonuclease complex; 50 μl per reaction). Each sample of digested cells was divided into two reactions, each of which was resuspended in 100 μl of this hybridization solution. Resuspended cells were incubated at 37°C overnight. Cells were washed with 10% formamide/2x SSC followed by 0.1% Triton X-100/2x SSC. For DAPI staining, cells were incubated in 1x PBS with 1 μg/ml DAPI for 10 min and washed once more with 1x PBS. Cell pellets were mixed with SlowFade Diamond Antifade Mountant (Thermo Scientific, S36972) and mounted on DEPC-cleaned slides using #1.5 glass coverslips. Imaging was performed on a Zeiss AxiosImager M1 equipped with Xcite Fire LED illumination (Excelsitas), a Zeiss αPlan FLUAR 100x/1.45 oil objective, and an ORCA-Flash4.0LT sCMOS camera (Hamamatsu). Images were acquired for 6 μm in Z separated by 0.2 μm steps for each channel. Images of labeled RNA were captured with either an mCherry filter or a “gold FISH” filter (Chroma, 49,304). Additional data on the cell and nucleus were captured with GFP, DAPI, and CFP filters. Images were dark noise-subtracted and flat-field-corrected. A custom FIJI macro, using trainable WEKA segmentation (Arganda-Carreras et al, 2017), was used to create two-dimensional outlines of cells by CFP autofluorescence and of corresponding nuclei by DAPI. For analyses with cytoplasmic or nuclear RNA counts (except the mad1/mad2 co-localization experiment; Fig 1F), nuclei were re-segmented in three dimensions using a FIJI macro adapted from https://github.com/haesleinhuepf/cca_benchmarking (Robert Haase, MPI-CBG, Dresden). Analysis was limited to cells whose nuclei were entirely contained within the image stack. RNA spot analysis was performed in FISHquant (Mueller et al, 2013). Spots were initially detected based on an automatic intensity threshold and filtered with an additional manual threshold following the suggestions of the FISHquant documentation. A subset of cells in each image was cross-checked manually for successful RNA spot detection.

To measure co-localization of mad1+ and mad2+ mRNA, a two-color FISH experiment was performed targeting mad1+ with gene-specific probes and mad2+-ymEGFP with ymEGFP probes. The three-dimensional coordinates of each spot were recorded and corrected for relative chromatic aberration in Z. Distances were then calculated from each mRNA to its nearest neighbor of the other species within the same cell. To determine a distance cut-off for classifying RNA molecules as either co-localized or unpaired, the same two probe sets were used in another two-color FISH experiment in which both probes targeted mad1+-ymEGFP. Nearest-neighbor distances were calculated in the same way, and the distribution of these distances was used to determine the co-localization distance cut-off value. This cut-off was applied to the distances in the original experiment to classify each mad1+ or mad2+ mRNA molecule as co-localized or unpaired.

To test if mad1+ mRNA forms dimers, we used RNA FISH experiments to measure spot intensities and counts of RNA in the cytoplasm in strains with the following genotypes: (1) untagged mad1+ expressed from the endogenous locus, (2) untagged mad1+ expressed from the endogenous locus and mad1+-ymEGFP expressed from the exogenous leu1+ locus, (3) endogenous mad1+ deleted and mad1+-ymEGFP expressed from the exogenous leu1+ locus, and (4) mad1+/-ymEGFP and mad3+/-ymEGFP expressed from the endogenous loci. All samples were hybridized with a combination of mad1- and ymEGFP-targeting probes in two-color FISH experiments. FISH probe spots were quantified separately for each imaging channel. Colocalized spots of different colors were then paired using the same co-localization method as described for mad1/mad2 co-localization above. Intensity analysis used the amplitude of the point spread function fit to each spot provided by FISHquant (Mueller et al, 2013). Using the intensity of each spot after background filtering, also provided by FISHquant, yielded the same result. All images quantified from the experiment for each probe set, regardless of genotype, gave consistent distributions of spot intensities, except for one image: while the distributions of spot intensities were qualitatively the same and the counts of spots were consistent with the other images, both its mad1+ and its GFP probe spot intensities were shifted substantially lower compared with the other images (including another image from the same slide and another slide prepared from the same FISH sample). Thus, we decided to remove it from the analysis.

Single-cell RNA counts from FISH experiments were fit with generalized linear mixed models. All models used a Poisson error distribution and natural log link function. The models included the fixed effect cell length and up to three nested random effects (when present): biological replicate (strain), experimental replicate, and microscopy image. Random effects included both random slopes and intercepts for all three variables. As the relationship between mean RNA count and cell length was approximately linear, cell length was natural log transformed. The transformed cell length was then centered so that a cell of average length had a value of zero.
To test for differences in mean RNA levels between genotypes, the categorical fixed effect variable genotype was added to the model. The interaction between cell length and genotype was also included if a likelihood ratio test comparing models with and without the interaction term showed that it improved the model's ability to explain the data significantly (P < 0.05). P-values for the likelihood ratio test were obtained both by comparing the test statistic to a chi-square distribution and generating a null distribution by bootstrapping (1,000 replicates) using the “PBmodcomp” function from the package pbkrtest. In all cases, the results were consistent between the two methods. Only a few models required the interaction term: comparison of wild-type with codon-optimized mad2 (whole cell, cytoplasmic, and nuclear RNA counts), comparison of wild-type with codon-optimized mad2 + ste13Δ (cytoplasmic RNA only), and comparison of untagged and GFP-tagged mad1Δ.

Genotype coefficients and corresponding 95% bootstrap confidence intervals presented in the paper were exponentiated (e^) and represent the ratio of expected RNA levels between the two genotypes in each comparison. RNA levels were considered significantly different between genotypes if the exponentiated confidence interval excluded 1.

**Assay for spindle assembly checkpoint function using nda3-KM311**

Strains expressing the tubulin mutant nda3-KM311 were grown in EMM (plus supplements required for auxotrophic mutations) at 30°C to a concentration of 0.5–1.0 × 10^7 cells/ml. Cells were diluted with EMM to a final concentration of 7.5 × 10^5 or 1.5 × 10^6 cells/ml. 300 μl of each strain were loaded into a lectin-coated Ibidi μ-Slide glass-bottom chamber and incubated about 1 h at 16°C on the microscope stage prior to imaging. Cells were imaged at 16°C on a DeltaVision Elite system with a PCO edge sCMOS (PCO), an Olympus 60x/1.42 Plan APO oil objective, and environmental chamber. Images were acquired every 5 min for GFP and mCherry over an 18-h period using an “optical axis integration” (sum projection) over a 3.2 μm Z-distance.

Plo1-mCherry localizes to spindle-pole bodies during mitosis and was used to identify cells in mitosis. Kinetochores cluster with spindle-pole bodies in *S. pombe* interphase (Funabiki et al., 1993) and dot-like GFP signals were therefore measured in the direct vicinity to Plo1-mCherry. An area of the same size for each cell was used to capture the kinetochore signal and was also used to measure the intensity in the nucleoplasm for background subtraction. GFP intensities from multiple cells were aligned to the time point of Plo1-mCherry appearance and averaged for each time point.

**Assay for spindle assembly checkpoint function using alp7Δ**

Cells were grown in EMM at 25°C to a concentration of 0.5–1.0 × 10^7 cells/ml, diluted to 1.5 × 10^6 cells/ml, and 300 μl of this dilution were loaded into a lectin-coated ibidi μ-Slide glass-bottom chamber. Cells were incubated on the microscope stage at 30°C for 35 min before imaging. Images were acquired at 30°C every 55 s to 1.5 min for 2–3.5 h using an “optical axis integration” (sum projection) over a 3.6 μm Z-distance. Cells were segmented based on the brightfield image using YeAZ (Dietler et al., 2020). All pixels within the cell were quantified and the 0.1st percentile value was subtracted from the 99.9th percentile value to obtain the “maximal intensity.” The localization of Plo1-TdTomato to spindle-pole bodies or Mad1-GFP to kinetochores (Appendix Fig S4F) is reflected in higher maximal intensities. Time in mitosis was determined from a custom Matlab script that detects strong increases and decreases in signal. Some cells could not be analyzed in an automated fashion (e.g., due to overlapping other cells) and were analyzed manually. The analysis mode is reported in the source data.

**RNA preparation**

Asynchronous *S. pombe* cultures were grown to a final concentration of approximately 0.7–1.5 × 10^7 cells/ml at 30°C in either EMM with 0.2 g/l leucine or YEA. 1 × 10^6 cells were collected by centrifugation, washed once with deionized water, and immediately flash-frozen in liquid nitrogen and stored at −80°C before processing. RNA was extracted by resuspending samples in 700 μl of TES buffer (10 mM Tris HCL pH 7.5, 10 mM EDTA, 0.5% SDS) and adding 700 μl of acidic phenol chloroform (Fisher Scientific). Samples were immediately vortexed for 20 s and incubated for 1 h at 65°C. Following incubation, samples were cooled on ice for 1 min, vortexed for an additional 20 s, and centrifuged for 15 min at 16,000 rcf at 4°C. The RNA was further purified by twice mixing the aqueous supernatant with 700 μl of acidic phenol chloroform and centrifuging the solution in a 5Prime Phase Lock Gel Heavy 2 ml tube (Andwin Scientific) at 16,000 rcf to separate the phases. Following overnight ethanol precipitation, samples were centrifuged at 16,000 rcf for 10 min at 4°C and washed with one equivalent of 70% ethanol before additional centrifugation. Samples were left to air-dry at room temperature and resuspended in nuclease free water before quantification. 50 μg of total RNA was subjected to DNase treatment (Roche, 10776785001) followed by ethanol precipitation.

**Quantitative PCR (qPCR)**

For quantitative PCR (qPCR), 1 μg of DNase-treated total RNA was subjected to Superscript IV cDNA synthesis using oligo d(T)20 primers. Transcript abundance was quantified on a QuantStudio 6 Real-Time PCR system using SYBR® Green PCR Master Mix (ThermoFisher) and gene-specific primers (Appendix Table S5). To estimate relative expression, raw Ct values (2–3 technical replicates per sample) were averaged and normalized according to the following formula (Pfaffl, 2001; Hellemans et al., 2007):

\[
\text{Relative Expression} = \frac{\left(\text{efficiency}_{\text{target}} + 1\right)^{\left(C_{\text{target sample}} - C_{\text{target control}}\right)}}{\left(\prod \left(\text{efficiency}_{\text{reference}} + 1\right)^{\left(C_{\text{reference sample}} - C_{\text{reference control}}\right)^{\frac{1}{n}}}\right)^{\frac{1}{n}}}
\]

where “target” is the mRNA of interest, “reference” is the reference gene, “sample” is the sample of interest, and “control” is the control sample being normalized to. The denominator is the geometric mean of the reference genes (*act1* and *cdc2*), and efficiencies were estimated from the slopes of four-step, serial 1:5 dilution standard curves.

**Determination of mRNA half-life**

The mRNA half-life measurement procedure was adapted from published protocols (Duffy et al., 2015; Chan et al., 2018). Asynchronous *S. pombe* cultures were grown to a final density of approximately 0.7–0.9 × 10^7 cells/ml at 30°C in EMM with 0.45 mM uracil and 1.5 mM leucine before collection. 4-thiouracil (4tU) in DMSO
(Chem-Impex International, CMX-21484) was added at 5 mM final concentration (Eser et al., 2016). Cells (5 x 10^7 cells per sample) were collected by centrifugation, immediately flash-frozen in liquid nitrogen, and stored at −80°C before processing. Samples were collected from the culture before addition of 4tU (time = 0) and at a series of time points after. For use as a spike-in control, *S. cerevisiae* cultures were grown to a final density of 1.4–3.7 x 10^7 cells/ml in YPD at 30°C, flash-frozen, and stored at −80°C.

RNA extraction was performed as above except flash-frozen samples were initially resuspended in 600 μl of 96% ethanol and 100 μl of resuspended *S. cerevisiae* cells (approximately 5 x 10^7 cells) were added as a spike-in control. Care was taken to add the same amount of *S. cerevisiae* cells to all samples of a time series. After extraction, 200 μg of total RNA was subjected to DNase treatment. Following DNase treatment, 70 μg of RNA was biotinylated with MTSAP biotin-XX (Biotium, 90066) as previously described (Duffy et al., 2015; Chan et al., 2018). 50 μg of biotinylated RNA was subjected to oligo d(T) selection using oligo d(T)25 magnetic beads (NEB, S1419S), substituting SDS and NaCl for the recommended LiDS and LiCl. For streptavidin selection of biotinylated RNA, 500 ng of the oligo d(T) selected mRNA was used. 25 μl of MyOne Streptavidin C1 Dynabeads (ThermoFisher, 65001) were washed with 75 μl of 0.1 M NaOH two times, followed by a single 0.1 M NaCl wash, and two additional washes with Buffer 3 (10 mM Tris HCl pH 7.4, 10 mM EDTA, 1 M NaCl). Streptavidin beads were blocked by resuspension in 50 μl of Buffer 3 and 5 μl of 50X Denhardt’s reagent. Beads were incubated for 20 min with gentle agitation. Following blocking, beads were washed with 75 μl of Buffer 3 four times and resuspended in 75 μl of Buffer 3 with 4 μl of 5 M NaCl. 500 ng of mRNA was added to the beads and gently agitated for 15 min. Following incubation, beads were washed with 75 μl of Buffer 3 prewarmed to 65°C, once with Buffer 4 (10 mM Tris–HCl pH 7.4, 1 mM EDTA, 1% SDS) and twice with 10% Buffer 3. All flow-through was pooled before addition of 20 μg of linear acrylamide (ThermoFisher) followed by ethanol precipitation.

Expression relative to the time = 0 sample was quantified using qPCR as described above except 50 ng of recovered unlabeled mRNA was used in the Superscript IV cDNA synthesis reaction and a single reference gene (*S. cerevisiae* ACT1 from the spiked in *S. cerevisiae* cells) was used to normalize expression.

To estimate mRNA half-lives (HL), several different exponential decay models (adapted from Chan et al., 2018)) were initially fit to each time series using nonlinear least squares regression:

\[
\text{fit 1} = 2^{-\text{slope}} \\
\text{fit 2} = \text{efficiency} \times 2^{-\text{slope}} + (1 - \text{efficiency}) \\
\text{fit 3} = \text{mRNA}_0 \times 2^{-\text{slope}}.
\]

Fit 1 is a simple one-phase exponential decay model. Fit 2 incorporates an efficiency parameter to accommodate that mRNA levels may not decay to zero (Chan et al., 2018). Finally, to accommodate the effects of noninstantaneous labeling by 4tU on the decay curve, the fit 3 model was fit without the time = 0 measurements and instead allowed the mRNA level at time = 0 to be estimated as a separate free parameter (mRNA_0). Qualitatively, fit 3 consistently fit the time series best, but all models yielded similar results. To test for statistically significant differences in mRNA half-lives between ste13^+ and ste13Δ genotypes, we therefore proceeded with the fit 3 model, which was linearized, resulting in the equation

\[
\ln(\text{mRNA}_t) = \ln(\text{mRNA}_0) + \frac{-\ln(2)}{\text{HL}} t
\]

and was fit to the combined ste13^+ and ste13Δ time series data for each gene (excluding time = 0) using a general linear mixed model. In the model, natural log transformed relative mRNA expression was modeled as a function of the continuous fixed effects variable time (minutes since 4tU addition), the categorical fixed effect genotype (ste13^+ vs. ste13Δ), the interaction of time and genotype, and the random effect experimental replicate. The random effect included both random slopes and intercepts to allow the decay rate to vary across experimental replicates.

In the model, half-life is related to the change in expression with respect to time by the formula

\[
\text{HL} = \ln(2)/\text{slope}
\]

where slope is the time coefficient for the genotype coded 0 and the sum of the time and interaction coefficients for the genotype coded 1. To simplify the extraction of half-life estimates, models were fit with the genotype coded both ways: first ste13^+ = 0 and then ste13Δ = 0. Half-life estimates for ste13^+ and ste13Δ were derived from the version of the model in which the genotype of interest was coded zero using the formula

\[
\ln(2)/[\text{time coefficient}]
\]

Similarly, the difference in half-life between the two genotypes was estimated with the formula:

\[
\ln(2) \times [\text{interaction coefficient}]/(\text{time coefficient} \times [\text{time coefficient} + \text{interaction coefficient}])
\]

Significance of the change in half-life due to the deletion of ste13^+ was measured two ways. If the 95% bootstrap confidence interval for the interaction coefficient excluded 0, the slopes of expression with respect to time in the model, and thus half-lives, were considered to be different. Similarly, if the 95% bootstrap confidence interval for the half-life difference excluded 0, the change was considered significant.

**Codon usage bias calculations**

The “Codon occurrence to mRNA Stability Correlation coefficient” (CSC) for each codon was determined as in Presnyak et al. (2015) by calculating a Pearson’s correlation coefficient between the frequency of occurrence of individual codons in *S. pombe* mRNAs and the half-lives of these mRNAs. Coding sequences for *S. pombe* (protein-coding genes, excluding dubious and transposons) were downloaded from Pombase (ASM294v2.62, Release date 2017-01-30) (Lock et al., 2019). From this list, we excluded “Genome location: mitochondrial,” “Genome location: mating_type_region,” and “sequence error in Pombase (ASM294v2.62, Release date 2017-01-30)” (Lock et al., 2019).

A previous study (Harigaya & Parker, 2016) used the Spearman’s correlation coefficient to determine CSC values for *S. pombe*, because of outliers in the half-life data. We instead removed outliers...
from the half-life data and used the Pearson’s correlation coefficient. A comparison between the different strategies is shown in Appendix Fig S4C and D. Our criteria for removing outliers were as follows: (i) a value that was more than 10 interquartile ranges above the upper quartile (which removed three genes, based on their value in the Eser et al., 2016 data) and (ii) a deviation in rank position of > 2,500 between the two datasets (which removed 13 genes). After the removal of outliers, the Pearson’s correlation coefficient between the two mRNA half-life datasets was 0.80, the Spearman’s rank correlation 0.82. Using either the Hasan et al., 2014 or the Eser et al., 2016 half-life data for CSC calculation yielded highly similar results (Appendix Fig S4C and D). When not otherwise indicated, CSC values obtained from Eser et al., 2016 (the more recent study) were used. The CSC values for each gene was determined as the arithmetic mean of all codons, excluding the stop codons.

The percentage of optimal codons based on the “classical translational efficiency” (cTE) used the optimality table for S. pombe from Pechmann & Frydman (2013). For tAI (tRNA adaptation index), we used the values determined by Tuller et al. (2010) and also reported in Pechmann & Frydman (2013). The TAI value for each gene was determined as the geometric mean of all codons, excluding the stop codons.

CSC values for budding yeast were taken from Carneiro et al. (2019) and only values derived from mRNA half-life measurements within the last 10 years were included (Becskei: Baudrimont et al., 2017; Coller: Presnyakov et al., 2015; Cramer: Sun et al., 2012; Gresham: Neymotin et al., 2014; Struhl: Geisberg et al., 2014; Weis: Munchel et al., 2011). CSC values derived from the Struhl study are shown in gray in Appendix Fig S7, since they differed from the ones based on the other mRNA half-life datasets (Carneiro et al., 2019). CSC values for human cells were taken from Wu et al. (2019), Narula et al. (2019), and Forrest et al. (2020). CSC values from the Forrest study are shown in gray in Appendix Fig S7, as they were based on fewer mRNA half-life data and differed from the other two studies in their trend. The multiple CSC values from Wu et al. (2019) and Narula et al. (2019), respectively, were averaged, and the mean value obtained in each study was used. Mad1 and Mad2 sequences from opisthokonts were taken from Vleugel et al. (2012). The human Mad1 sequence was swapped for the canonical isoform (UniProt, Q9Y6D9, MDI11_HUMAN), the S. pombe Mad1 sequence was shortened N-terminally by 13 amino acids to start with what is now considered the correct start codon (pombase.org). All sequences shorter than 600 amino acids were omitted for Mad1. Protein sequences were aligned using MAFFT (G-INS-I, using DASH and Mafft-homologs [100 homologs, E = 1e-30]) (Katoh et al., 2019). Sequences from all species other than S.c., S.p., and H.s. were deleted for the display of conservation shown in Appendix Fig S7. The moving average of CSC values across nine codons was plotted along the length of the aligned sequence. The null distribution of the moving average was obtained by randomizing the codon order 10,000 times. Observed values that deviated by more than two standard deviations from the null mean were marked with filled circles.

**Gene expression model—Simulations and theoretical predictions**

Protein noise predictions (Fig 2C; Appendix Fig S2B and C) were made by assuming a constitutively active promoter, and only considering stochastic mRNA and protein synthesis and degradation and ignoring cell growth and division. The coefficient of variation (CV = standard deviation/mean) for protein is calculated as:

\[
CV_p = \sqrt{\frac{1}{P} + \frac{1}{M} \left( \frac{k_{degP}}{k_{degM} + k_{degP}} \right)}
\]

where \(P\) is the mean protein number per cell, \(M\) the mean mRNA number per cell, \(k_{degM}\) the mRNA degradation rate, and \(k_{degP}\) the protein degradation rate (Swain, 2004). For the predictions in Appendix Fig S2C, we assumed a mean protein number of 6,000 per cell, mean mRNA numbers of 1 to 1,000, and we varied RNA degradation rate in a range corresponding to half-lives of 1 to 60 min, and protein degradation rate in a range corresponding to half-lives of 15 to 600 min, which we consider a physiologically plausible range. Predictions were excluded when mRNA synthesis or protein synthesis rates became unrealistically high. We assumed this to be the case when mRNA synthesis rate was higher than 25 min\(^{-1}\) or protein synthesis rate higher than 20 mRNA\(^{-1}\) min\(^{-1}\). Assuming a gene with characteristics similar to a SAC gene (mean protein number = 6,000, mean mRNA number = 3.5, protein half-life = 360 min, mRNA half-life = 4 min) yields a CV prediction of 0.0575. In the figure, we labeled CV predictions less than 0.06 in light gray (low noise) and those equal or higher than 0.06 in dark gray (high noise).

The stochastic simulation of mRNA and protein numbers (Fig 2B) used the same simple underlying model and the Gillespie algorithm in a Matlab script written by Daniel Charlebois and available on Math-Works (“Gillespie’s Direct Method Stochastic Simulation Algorithm”).

**Statistical tests**

Data processing performed in R used the packages tidyverse (Wickham et al., 2019), alphashape3d, boxcoxmix, broom, broom.mixed, DescTools, geometry, Irescale, mclust, nabor, plyr, readxl, rgl, sg, shotGroups, and spatstat. Statistical tests were performed in Prism (GraphPad), or in R using the packages lim4 (Bates et al., 2015), MASS, pbkrtest, and stats. Figure plots were generated in Prism (GraphPad) or in R using the packages Cairo, cowplot, ggplot2, gridExtra, plotly, and lemon.

General linear mixed models and generalized linear mixed models were fit using the functions “lmer” and “glmer,” respectively, from the lme4 package. Default function settings were used except for the optimizer in “glmer,” which was set to “bobyqa.” Bootstrapping using the function “bootMer” (10,000 replicates, lme4 package) was used to obtain 95% confidence intervals for fixed effects model coefficients and mRNA half-life estimates, and 95% confidence bands for predicted regression curves. Wilcoxon rank sum tests, and \(t\)-tests were performed using “wilcox.test” and “t.test,” respectively, from the package stats. Poisson distributions were fit to data frequency distributions using “fitdistr” from the package MASS.

Sample sizes were not predetermined. Blinding was not performed, as most analyses were run in an automated fashion.

**Data availability**

This study includes no data deposited in external repositories.

**Expanded View** for this article is available online.

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**Author contributions**

**Eric Esposito:** Conceptualization; formal analysis; investigation; visualization; writing – original draft. **Douglas E Weidemann:** Conceptualization; formal analysis; investigation; visualization; writing – review and editing. **Claire M Morton:** Formal analysis; investigation. **Erod Keaton Baybay:** Investigation; methodology; writing – review and editing. **Jessie M Rogers:** Formal analysis; investigation; visualization; writing – review and editing. **Silke Hauf:** Conceptualization; formal analysis; supervision; funding acquisition; investigation; visualization; writing – original draft.

**Disclosure and competing interests statement**

The authors declare that they have no conflict of interest.

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