1 Signaling protein abundance modulates the strength of the Spindle Assembly Checkpoint

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11 Summary

12 During mitosis, unattached kinetochores in a dividing cell signal to the Spindle Assembly Checkpoint to delay anaphase onset and prevent chromosome missegregation ¹⁻⁴. The signaling 13 14 activity of these kinetochores and the likelihood of chromosome missegregation both depend on 15 the amount of SAC signaling proteins that each kinetochore recruits ⁵⁻⁸. Therefore, factors that 16 control SAC protein recruitment to signaling kinetochores must be thoroughly understood. 17 Phosphoregulation of kinetochore and SAC signaling proteins emerging from the concerted action 18 of many kinases and phosphatases is a major determinant of SAC protein recruitment to signaling 19 kinetochores⁹. Whether the abundance of SAC proteins also influences their recruitment and signaling activity at human kinetochores has not been studied ^{8,10}. Here, we reveal that the low 20 21 cellular abundance of the SAC signaling protein Bub1 limits kinetochore recruitment of Bub1 and 22 BubR1 and reduces the SAC signaling activity of the kinetochore. Conversely, Bub1 23 overexpression results in higher protein recruitment and SAC activity producing longer delays in 24 anaphase onset. We also find that the number of SAC proteins recruited by a signaling 25 kinetochore is inversely correlated with the total number of signaling kinetochores in the cell. This 26 correlation likely arises from the competition among the signaling kinetochores to recruit from a 27 limited pool of signaling proteins. The inverse correlation between the number of signaling 28 kinetochores in the cell and the signaling activity of individual kinetochores may allow the dividing 29 cell to prevent the large number of signaling kinetochores in prophase from generating an 30 unnecessarily large signal, while enabling the last unaligned kinetochore to signal at the maximum 31 possible strength.

32 Results and Discussion

The stoichiometry of SAC proteins at signaling kinetochores depends on the number of signaling kinetochores in the cell

35 We set two interrelated goals for this study. First, we wanted quantify the steady-state 36 stoichiometry of key SAC proteins in human kinetochores to directly view the signaling activity of individual kinetochores ⁶. This has been accomplished in fungi ^{8,10}, but the data in vertebrates 37 remains indirect and fragmentary ^{5,11}. Therefore, we quantified the recruitment of three SAC 38 39 proteins representing the three layers of the well-defined signaling cascade that generates the 'Mitotic Checkpoint Complex' (MCC, Figure 1A)^{5,12-21}. To accomplish this, we used three genome-40 edited HeLa-A12 cell lines wherein either Bub1, BubR1, or Mad1 was fused with mNeonGreen 41 (mNG)^{22,23}. These HeLa cell lines are only partially genome-edited; fully edited cell lines could 42 not be obtained because of low efficiency of CRISPR-Cas9-mediated knock-ins and the pseudo-43 44 tetraploid nature of HeLa cells. Therefore, we also quantified the relative amounts of labeled and unlabeled proteins in the three cell lines using quantitative immunoblot analysis on whole-cell 45 46 lysates of mitotic cells (Figure S1A-B). This analysis defined the labeled-to-unlabeled protein ratio 47 for each of the three cell lines: ~40% for Bub1 and ~70% for both BubR1 and Mad1 (assuming similar transfer efficiencies for the labeled and unlabeled protein bands, Figure S1C). We used 48 49 these ratios to estimate the total protein recruitment per kinetochore from the average signal for 50 the mNG-tagged protein per kinetochore. It should be noted that these fusions do not detectably 51 affect SAC signaling (shown later in Figure 4).

52 For our second goal, we studied whether the number of signaling kinetochores in the cell affects 53 the signaling activity of individual kinetochores. In budding yeast, the recruitment of SAC proteins 54 per kinetochore is higher in cells that contain fewer signaling kinetochores, likely because of the 55 limited abundance of SAC proteins ¹⁰. Whether this factor influences the signaling activity of 56 human kinetochores is unknown. Therefore, we observed SAC protein recruitment per 57 kinetochore in mitotically arrested cells containing two distinctly different numbers of signaling 58 kinetochores. To obtain cells with nearly all kinetochores signaling, we released G1/S arrested 59 HeLa cells into the cell cycle and then arrested them in mitosis using 330 nM nocodazole to depolymerize microtubules (Figure 1B, right). These cells serve as a model for late prophase. To 60 61 obtain mitotic cells with a much smaller number of signaling kinetochores, we released G1/S 62 arrested HeLa cells into the cell cycle and treated them with 236 nM GSK923295, a small molecule inhibitor of the mitotic kinesin CENP-E, 1.5 hours prior to imaging ²⁴. These cells model 63 64 late prometaphase, because most of the chromosomes align at the metaphase plate, but a

smaller, variable number of chromosomes become stranded at the spindle poles. The unattached
or laterally attached kinetochores of these chromosomes activate the SAC (Figure 1B, left) ^{25,26}.
Relying on visual inspection, we selected only those cells containing ~ 10 or less polar
chromosomes for analysis.

69 The guantitative comparison of mNG-labeled protein recruitment revealed that the recruitment of 70 all three proteins per kinetochore was significantly higher in GSK923295-treated compared to 71 nocodazole-treated cells (Figure 1C). We combined these measurements with the quantitative 72 immunoblotting data to estimate the total (labeled + unlabeled) protein amount to understand how 73 BubR1 and Mad1 recruitment correlates with Bub1 recruitment after the two treatments (Figure 74 1D). We normalized total protein amounts with the amount of total Bub1 per kinetochore in 75 GSK923295-treated cells ^{13,27-32}. In GSK923295-treated cells, BubR1 and Mad1 per kinetochore 76 was ~ 60 and 45% respectively of total Bub1 per kinetochore. In nocodazole-treated cells, the 77 BubR1 and Mad1 amount per kinetochore was significantly lower ~ 40% and 25% respectively of 78 total Bub1. Thus, the overall protein recruitment per kinetochore was higher in GSK923295-79 treated cells compared to nocodazole-treated cells: 1.45-fold in case of Bub1, 2.3-fold and 2.6-80 fold for BubR1 and Mad1 respectively (Figure 1D). Thus, the number of SAC proteins recruited per kinetochore in HeLa cells is inversely correlated with the number of signaling kinetochores in 81 the cell, consistent with observations in budding yeast ¹⁰. 82

We used a fluorescence standard to estimate the copy number of each SAC protein recruited per 83 84 kinetochore. It is known that: (a) the human kinetochore contains ~ 250 molecules of the Ndc80 complex and (b) the stoichiometry between Ndc80 and Knl1 is 3:2³³. Therefore, under identical 85 imaging conditions we imaged HeLa cells exogenously overexpressing Spc25-mNG (Spc25 is an 86 87 Ndc80 complex subunit) and treated with siRNA targeting the endogenous Spc25 to guantify the 88 mNG signal per kinetochore in metaphase cells (Figure S1D). Using the above-mentioned parameters, we converted the estimated total amount of each protein per kinetochore into the 89 90 number of molecules of the protein recruited per Knl1 molecule in HeLa cells. These calculations 91 indicate that each Knl1 molecule recruits 5 ± 0.9 Bub1 molecules, 3 ± 0.7 BubR1 molecules, and 92 2 ± 0.7 Mad1 molecules in GSK923295-treated cells. The numbers are lower in nocodazole-93 treated cells: 3 ± 0.7 Bub1 molecules, and 1 ± 0.4 molecules of BubR1 and Mad1 each per Knl1. 94 The number of Bub1 molecules per kinetochore in nocodazole-treated cells is slightly lower than the previously reported number of 4-6 molecules ^{5,12,13}. A more recent study found that Bub1 95 96 recruitment per kinetochore was not significantly different in cells treated with nocodazole- and

97 GSK923295 ³⁴. However, this difference could arise from the lack of filtering based on the number
98 of signaling chromosomes per cell.

99 Mad1 is recruited to metazoan kinetochores by Bub1 and the RZZ complex ³⁵⁻³⁷. To quantify the contribution of the RZZ complex, we depleted ZW10 using RNA interference (RNAi) and then 100 101 quantified SAC protein recruitment as before. ZW10 siRNA treatment lowered Bub1 and BubR1 recruitment by ~ 25%, as noted by others 38 . As expected, Mad1 recruitment was significantly 102 103 lower in both GSK923295 and nocodazole-treated cells (Figure S1). The number of Mad1 104 molecules recruited per Bub1 molecule was ~ 50% lower revealing the contribution of the RZZ 105 under these conditions. Although the number of Bub1 and BubR1 molecules per kinetochore was 106 also reduced, the two proteins were recruited in approximately equal amounts.

107 To test whether the inverse correlation is specific to HeLa cells, we examined Bub1 and BubR1 recruitment per kinetochore in hTERT-RPE1 cells. We released RPE-1 cells synchronized in G1/S 108 109 into media containing either nocodazole or GSK923295, and measured the total amounts of Bub1, 110 BubR1 recruited per kinetochore by immunofluorescence (Figure 1E). These measurements 111 show that the Bub1 amount per kinetochore in GSK923295-treated RPE-1 cells as ~1.6-fold 112 higher the Bub1 amount in nocodazole-treated cells; BubR1 amount was ~2.5-fold higher (Figure 1F). We did not quantify the amounts of Mad1 per kinetochore because a ring-like pool of Mad1 113 114 proximal to the spindle poles, especially in GSK923295-treated cells (see inset in Figure 1E), 115 prevented accurate quantification. These data demonstrate that the inverse correlation between 116 the amount of Bub1 and BubR1 per kinetochore and the number of signaling kinetochores is not 117 specific to HeLa or budding yeast cells. In fact, Bub1 and BubR1 show a similar degree of 118 enrichment at RPE-1 and HeLa kinetochores in GSK923295-treated cells.

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120 Mps1-mediated phosphorylation within signaling kinetochores is the same in nocodazole-121 and GSK923295-treated cells

122 Changes in the Mps1-mediated phosphoregulation of the kinetochore in GSK923295- and 123 nocodazole-treated cells may change SAC protein recruitment. Indeed, a prior study found that 124 the net Mps1-mediated phosphorylation decreased rather than increasing in GSK923295-treated 125 cells when compared to nocodazole-treated cells suggesting that SAC protein recruitment should 126 decrease in GSK923295-treated cells ³⁴. Therefore, we quantified the net Mps1 kinase activity 127 within the kinetochore using a recently developed FRET-based sensor (MPS1sen-KT) ^{34,39}. We 128 modified this sensor to use mNeonGreen/mScarlet-I as the acceptor/donor combination (Figure S2A). Since the acceptor and the donor has a fixed 1:1 stoichiometry in the sensor, we used the ratio between the green channel readout and the FRET channel readout (without correcting for the cross-excitation of the acceptor fluorescence and the bleed-through of donor fluorescence) as the normalized measurement of the FRET efficiency, which is positively correlated with the net phosphorylation (Figure S2B-D).

134 We confirmed that MPS1sen-KT recruited to unaligned kinetochores in both nocodazole- and 135 GSK923295- treated cells had lower FRET efficiencies (higher activities of Mps1) compared to 136 kinetochores aligned at metaphase plates in GSK923295-treated cells (Figure S2D). Importantly, 137 we did not detect any difference in the net Mps1-mediated phosphorylation within unaligned 138 kinetochores in nocodazole- and GSK923295-treated cells irrespective of the drug concentration 139 used (Figure S2D). This divergence which may arise from different properties of the two cell lines 140 as was found in a panel of colorectal cancer cell lines ³⁴. In either case, the increased kinetochore 141 recruitment of SAC proteins in GSK929395-treated cells does not stem from increased Mps1-142 mediated phosphorylation within the kinetochore.

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144 The number of MELT motifs per Knl1 controls Bub1, BubR1, and Mad1 recruitment to 145 signaling kinetochores

146 We next studied how the number of MELT motifs per Knl1 affects the SAC protein recruitment. We knocked down endogenous Knl1 using RNAi and replaced it with Knl1^Δ-M3 or Knl1^Δ-M3-M3. 147 which are mScarlet-I-tagged recombinant Knl1 versions containing three or six MELT motifs: 148 149 (Figure 2A)⁵. As before, we quantified Bub1, BubR1, and Mad1 signals at individual kinetochores 150 in cells treated with either GSK923295 or nocodazole (Figure 2B). For comparing the amounts of 151 proteins recruited under these conditions, we normalized all localized fluorescent intensities by 152 the average fluorescence signal per kinetochore of each protein in nocodazole-treated cells 153 expressing Knl1^Δ-M3. As before, we used the labeled-to-unlabeled protein ratios to estimate the 154 total amount of protein per kinetochore (Figure 2C).

In the GSK923295-treated cells, $Knl1^{\Delta}$ -M3 recruited ~ 1.4-times higher Bub1 per kinetochore compared to nocodazole-treated cells. $Knl1^{\Delta}$ -M3-M3 recruited more Bub1 than $Knl1^{\Delta}$ -M3 under both conditions. Importantly, $Knl1^{\Delta}$ -M3-M3 recruited 2-fold higher Bub1 than $Knl1^{\Delta}$ -M3 in GSK923295-treated cells consistent with its 2-fold higher number of MELT motifs. These data again highlight that the recruitment of SAC proteins per kinetochore is higher in cells containing small numbers of signaling kinetochores. BubR1 and Mad1 recruitment showed similar trends. Importantly, the total BubR1 recruitment per Knl1^Δ-M3-M3 was 2-fold than Knl1^Δ-M3 under both conditions. BubR1 amount per Knl1^Δ-M3 was 2-fold higher in GSK923295-treated cells compared to nocodazole-treated cells, which is consistent with the increase in Bub1 recruitment by Knl1^Δ-M3 under the two conditions. Interestingly, total Mad1 recruitment by Knl1^Δ-M3-M3 and Knl1^Δ-M3 was similar in nocodazole-treated cells but increased by 5-fold in GSK923295-treated cells. The kinetochore recruitment of the two recombinant Knl1's was very similar and cannot explain the observed trends in SAC protein recruitment (Figure S3A).

- 168 As before, we combined these data with the immunoblotting quantitation to estimate the number of molecules per Knl1 (Figure S3B). Knl1^Δ-M3-M3 cells recruits ~ 5 ± 0.3 Bub1 molecules, but 169 170 only 3 ± 0.3 molecules in nocodazole-treated cells. These numbers are very similar to the 171 numbers of Bub1 molecules recruited by wild-type Knl1 under the two conditions (5 \pm 0.9 and 3 \pm 172 0.7 molecules respectively, see Figure S1), indicating that our estimates are internally consistent. 173 A prior study reported that HeLa kinetochores recruit 4-6 Bub1 molecules per kinetochore in nocodazole-treated cells ⁵, whereas we find that there are 3 Bub1 molecules per Knl1 in 174 175 nocodazole-treated cells and 5 per kinetochore in GSK923295-treated cells. The reasons for this 176 discrepancy remain unclear.
- 177 These data provide several insights. First, they confirm that the number of MELT motifs per Knl1 178 strongly influences Bub1, BubR1, and Mad1 recruitment. Second, in the case of Bub1 and Mad1 179 this influence is detected only in GSK923295-treated cells containing small numbers of signaling kinetochores. Third, Knl1^Δ-M3-M3 recruits ~2-fold more BubR1 than Knl1^Δ-M3 under both 180 181 conditions. Increased BubR1 recruitment to the kinetochore is likely to be functionally significant 182 because it promotes SAC signaling ^{14,23}. Mad1 recruitment is disproportionately high GSK923295-183 treated cells, possibly because of changes in the contribution of the RZZ pathway to Mad1 184 recruitment ⁴⁰. Given the catalytic role of Mad1 in the rate-limiting step in the SAC signaling 185 cascade, the higher Mad1 recruitment may also increase the signaling activity of these 186 kinetochores. Therefore, we next tested whether the higher SAC protein recruitment by Knl1^Δ-187 M3-M3 translates into a stronger SAC and longer mitotic arrest durations.

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189 The number of MELT motifs per Knl1 affects the SAC signaling strength

Previous studies concluded that the number of MELT motifs per Knl1 does not significantly influence the duration of the SAC-mediated mitotic arrest unless the MELT motifs are weak in their activity ^{5,13}. However, these studies characterized SAC signaling only in cells treated with high doses of nocodazole. Our quantification shows that in the nocodazole-treated cells, kinetochores containing Knl1^{Δ}-M3-M3 recruit only modestly larger amounts of Bub1 and Mad1 compared to kinetochores containing Knl1^{Δ}-M3. Therefore, the SAC signaling activity of both types of kinetochores may have been similar in nocodazole-treated cells. A further complicating factor is that one of the studies partially inhibited Mps1 during the experiment, which may have masked differences in SAC signaling due to the two Knl1 versions ⁵.

- 199 Because of these considerations, we re-evaluated the strength of SAC signaling mediated by 200 Knl1^Δ-M3 and Knl1^Δ-M3-M3 specifically in GSK923295-treated cells. We used time-lapse imaging 201 to monitor mitotic progression in cells treated with different concentrations of GSK923295 to 202 achieve different degrees of CENP-E inhibition resulting in different durations of mitotic arrests. 203 In all cases, cells expressing Knl1^Δ-M3-M3 arrested for significantly longer durations compared 204 to cells expressing KnI1^Δ-M3 (Figure 2D-E). The same trend was observed when these cells were 205 treated with Taxol (Figure 2F). Thus, the number of MELT motifs in Knl1 can determine the 206 duration of checkpoint-induced mitotic delays in cells wherein a few signaling kinetochores sustain 207 the SAC. When combined with the results described in the previous section, this finding implies 208 that the higher recruitment of Bub1, BubR1, and Mad1 per kinetochore is correlated with a higher 209 SAC activity, and this correlation is readily detected only in when the mitotic cells contain small numbers of signaling kinetochores ⁷. 210
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212 Bub1 over-expression increases recruitment of SAC proteins to the kinetochore

213 A key question then what limits SAC protein recruitment to the kinetochore especially in 214 nocodazole-treated cells. Changes in phosphoregulation cannot explain the differences in SAC 215 protein recruitment under the two conditions studied here. One responsible factor may be the low 216 abundance of SAC proteins, especially Bub1, which is responsible for recruiting BubR1 and 217 Mad1. Bub1 overexpression results in higher Bub1 and Mad1 recruitment to the budding yeast 218 kinetochore ¹⁰. Mad1 recruitment may also be limited by its own low abundance compared to 219 Bub1 and BubR1²³. Therefore, to assess SAC protein abundance, we used Fluorescence 220 Correlation Spectroscopy (FCS). We performed FCS measurements on mitotically cells arrested 221 either using MG132 or nocodazole. These measurements identified Mad1 is the protein with the 222 lowest abundance amongst the three SAC proteins. They also suggest that cytosolic 223 concentration of Bub1 is modestly lower in nocodazole-treated compared to MG132-treated cells 224 (Figure S3C).

225 To test whether higher Bub1 expression can lead to higher kinetochore recruitment of SAC 226 proteins, we exogenously expressed mNG-Bub1 in HeLa cells using a doxycycline-inducible 227 promoter (Figure S4A) and guantified Bub1, BubR1, and Mad1 recruitment per kinetochore in 228 cells treated with GSK923295 using immunofluorescence. As expected, mNG-Bub1 localization 229 at signaling kinetochores was detected only after doxycycline treatment (Figure 3A). To measure 230 the total amount of Bub1 per kinetochore, we fixed and stained these cells with anti-Bub1 231 antibodies. Fluorescence quantification confirmed that signaling kinetochores in the doxycycline-232 treated cells recruited ~35% more Bub1 than signaling kinetochores in untreated cells (Figure 3A, 233 plot on the left). The kinetochores in the doxycycline-treated cells also recruited 26% more BubR1 234 (Figure 3A, middle plot), however, Mad1 amount per kinetochore did not change (Figure 3A, right 235 plot). Thus, signaling kinetochores in HeLa cells can recruit more Bub1 and BubR1 if more Bub1 236 is made available. However, Mad1 recruitment does not increase likely because of the low Mad1 237 abundance, although behavior of the RZZ pathway will also strongly influence Mad1 recruitment.

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Bub1 overexpression strengthens the SAC in the presence of a small numbers of signalingkinetochores

241 To test whether Bub1 overexpression results in longer delays in anaphase onset, we released 242 either untreated or doxycycline treated, G1/S synchronized cells into media containing 18-35 nM 243 GSK923295 and observed their mitotic progression (Figure 3B). We found that the doxycycline-244 treated cells arrested for ~ 145 min on average, whereas the untreated cells arrested for ~ 94 min 245 under the same condition (Figure 3B, left, also see Supplementary Videos S1 and S2). To 246 understand the correlation between the expression level of exogenous Bub1 and the mitotic 247 duration, we plotted the mitotic duration against the cellular mNG signal in both cases (the 248 fluorescence values were normalized to the average mNG fluorescence per cell from samples not 249 treated with doxycycline, 3B right). As expected, there was no correlation between the cellular 250 mNG signal and mitotic duration in untreated cells (gray circles, Pearson's correlation coefficient 251 displayed at the top right of the figure). The mitotic duration of doxycycline-treated cells was 252 positively correlated with the mNG-Bub1 signal (orange circles in Figure 5B). It should be noted 253 that the mNG-Bub1 over-expression did not affect the mitotic progression of normally dividing 254 cells, and the degree of overexpression was also uncorrelated with the duration of mitosis (Figure 255 S4B-C). Thus, the longer mitotic duration seen in GSK923295-treated cells arises from higher 256 SAC signaling activity of the unaligned kinetochores.

257 We also examined whether Bub1 overexpression produces longer delays in anaphase onset in 258 nocodazole-treated cells (Figure 3C, also see Supplementary Videos S3 and S4). Interestingly, 259 the cellular mNG signal and the duration of mitosis did not statistically significant correlation in 260 this case (Figure 3C). The likely explanation for this observation is that even with increased Bub1 261 recruitment, the large number of signaling kinetochores cannot recruit higher amounts of 262 BubR1/MAD1 because of their depletion (implied by the reduced BubR1/MAD1 recruitment in the 263 nocodazole-treated cells, Figure 1). Therefore, the Bub1 overexpression does not translate into 264 more SAC signaling activity. In conclusion, increased Bub1 levels in HeLa cells increases Bub1 265 and BubR1 recruitment to the kinetochore and strengthens the SAC. Importantly, the effect on 266 SAC strength is detected only in GSK923295-treated cells, which contain small numbers of 267 signaling kinetochores.

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Partial depletion of Bub1 and BubR1 weakens the SAC in cells containing small numbers of signaling kinetochores

Mad1 abundance is lower than Bub1 abundance, which suggests that Mad1 availability should also limit the signaling activity of individual kinetochores (Figure S3C). Because Mad1 overexpression can weaken the SAC by depleting the Mad2 pool available for MCC formation, we used a different approach ^{8,41-43}. Rather than over-expressing Mad1, we partially knocked down Mad1-mNG using an siRNA against mNeonGreen in the genome-edited cell line heterozygous for Mad1-mNG, and then quantified the duration of SAC-induced mitotic delays. We also lowered mNG-Bub1 and mNG-BubR1 protein using the same strategy.

278 As expected, the mNG-tagged protein was significantly depleted by the mNeonGreen siRNA after 279 two days, whereas the unlabeled protein was unaffected (Figure 4A). To study the effect of a mild 280 depletion of an SAC protein on the signaling activity of the kinetochore, we treated the three cell 281 lines with mNG siRNA for one day and observed the effects of protein depletion on mitotic duration 282 in GSK923295- and nocodazole-treated cells. Quantification of mNG fluorescence signal from 283 mitotic cells revealed that the mNG RNAi reduce the level of the mNG-tagged protein by ~ 50% 284 (Figure 4B). In GSK923295-treated cells, it also significantly affected the mitotic progression 285 (Figure 4C, left). In the case of mNG-Bub1 and mNG-BubR1 depletion, the average duration of 286 mitosis decreased from ~ 1000 minutes to 300-500 minutes (Figure 4C). We also confirmed that 287 the partial depletion of BubR1 in GSK923295-treated cells resulted in chromosome 288 missegregation (Supplementary Videos S5 and S6). Mad1 depletion had a relatively minor effect 289 on mitotic duration: the average time in mitosis decreased from ~ 1000 minutes to 868 minutes

(Figure 4C). Interestingly, in nocodazole-treated cells, the partial depletion of all three proteinsdid not significantly change the duration of mitosis (Figure 4C, right).

Many prior studies have shown that the depletion of Bub1 in HeLa cells has only a minor effect 292 on the duration of mitosis in nocodazole-treated cells ^{35,44-46}. Our results in nocodazole-treated 293 cells are consistent with these findings. Strikingly, we find that in GSK923295-treated cells even 294 295 a small (~ 20-40% on average) reduction in Bub1 or BubR1 level results in a significant decrease 296 in the duration of mitosis. This observation is consistent with our results involving Knl1^Δ-M3 and 297 Knl1^Δ-M3-M3 (Figure 2) and suggests that higher SAC protein recruitment to the kinetochore 298 results in higher SAC signaling activity. The likely reason why this is not seen in nocodazole-299 treated cells is that the combined activity of a large number of signaling kinetochores masks 300 defects in the signaling activity of individual kinetochores ⁷.

301 In conclusion, HeLa and RPE-1 cells containing small numbers of signaling kinetochores recruit 302 significantly higher amounts of Bub1, BubR1, and Mad1 per signaling kinetochore compared to 303 cells containing many signaling kinetochores. The recruitment of all three proteins strongly 304 correlates with the number of MELT motifs in Knl1. This correlation is readily detected in cells 305 containing small numbers of signaling kinetochores; it cannot be detected in cells containing many 306 signaling kinetochores likely because of a shortage of unbound SAC proteins. Importantly, the 307 increased SAC protein recruitment per kinetochore, either due to the larger number of MELT 308 motifs or Bub1 over-expression, results in longer delays in anaphase onset. Conversely, partial 309 depletion of Bub1 and BubR1, but not Mad1, leads to shorter delays in anaphase onset in the 310 presence of a small number of signaling kinetochores.

311 The inverse correlation between the recruitment level of SAC proteins at a signaling kinetochore 312 and the total number of signaling kinetochores in a cell is likely to be biologically significant (Figure 313 4D). In human cells, the number of signaling kinetochores drops by almost two orders of 314 magnitude: from ~ 92 following nuclear envelop break-down to a few or just one in prometaphase. 315 If the amount of MCC in the cell scales linearly with the number of signaling kinetochores, the 316 nearly 100-fold drop in the number of kinetochores will result in a similar drop in the MCC creating 317 the possibility of a premature anaphase onset. Pre-mitotic MCC generation at the nuclear envelop also buffers SAC signaling activity of the kinetochores ⁴⁷. However, the strong decrease in the 318 duration of mitotic arrest in cells expressing Knl1^Δ-M3 or cells with partial Bub1 depletion suggests 319 320 that the buffering mechanism cannot compensate for lower signaling activity of kinetochores. 321 Instead, our data suggest that as the number of signaling kinetochores drops, the availability of 322 SAC signaling proteins, especially Bub1, increases, and enables the remaining signaling

kinetochores to recruit more SAC proteins. Consequently, the signaling strength of each
 kinetochore will become inversely correlated to the number of signaling kinetochores in the cell.
 Additionally, synergistic signaling by the Knl1 phosphodomain under this condition may further
 strengthen the signaling activity of these remaining kinetochores ⁴⁸.

327 Our results also raise additional questions requiring further investigation. First, it remains unclear 328 why kinetochores in HeLa cells and other eukaryotes use only a small fraction of their signaling 329 strength. One possibility is that this modulation of the signaling strength of the kinetochore 330 represents a compromise for balancing SAC signaling strength with its responsiveness to 331 silencing mechanisms ⁷. Higher Bub1 expression may also have drawbacks in other aspects of mitosis and cell biology ⁴⁹. Finally, these results bring into focus the contribution of perturbations 332 333 or imbalance in the expression levels of SAC proteins on chromosome missegregation commonly 334 seen in cancer cells ⁵⁰.

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343 Materials and Methods

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345 Cell culture

346 HeLa A12 cells (a kind gift from the Makeyev lab) were grown in DMEM media supplemented with 347 10% fetal bovine serum (FBS), Pen-Strep, 25 mM HEPES, and 1x GlutaMAX at 37 °C and 5% 348 CO₂, 0.5-2 µg/ml Puromycin or 10 µg/ml Blasticidin was used as needed. To express recombinant 349 proteins, we modified a bicistronic vector designed by the Makeyev lab as necessary ⁵¹. The 350 modified vectors were transfected into HeLa A12 cells that have the LoxP-Blasticidin-Lox2272 351 cassette engineered into their genome. Transformed cells were pooled and used for further 352 experimentation. The methodology used to construct the genome-edited HeLa A12 cell lines has 353 been described in detail in ⁵².

354 Imaging experiments were performed in Fluorobrite media supplemented with FBS, Pen-Strep,

355 HEPES as noted above. G1/S synchronization was achieved using a double thymidine block

procedure (2.5 mM thymidine). On the day of imaging, cells were released from the second block

- and treated with either nocodazole or GSK923295 ~ 1.5 hours prior to imaging.
- 358

359 Immunoblotting

For the quantitative immunoblot analysis shown in Figure S1, cells were synchronized using a single thymidine block (2.5 mM thymidine) for 24 hours. The cells were released into media containing 660 nM Nocodazole and harvested 15 hours post release. Anti-Bub1 (A300-373A, Bethyl Laboratories), anti-BubR1 (A300-386A, Bethyl Laboratories), and anti-Mad1 (ab184560, Abcam) primary antibodies were used to probe membranes at a dilution of 1:1000. Alexa Fluor 633-conjugated secondary antibodies (Goat anti-Mouse, Invitrogen A21052 or Goat anti-Rabbit, Invitrogen A21071 as appropriate) were used at a 1:15000 dilution.

367 For immunoblots in Figures 4A, S2A, and S4A: To acquire unsynchronized HeLa-A12 cells 368 lysates, cells were either scraped off the dish surface or trypsinized. To acquired mitotic HeLa-369 A12 cells, cells were first synchronized in G1/S with 2.5 mM thymidine and then arrested in mitosis 370 with 330 nM of nocodazole for 16 h before mitotic shake-off. Mitotic cells were then washed once 371 by phosphate-buffered saline (Gibco), pelleted down, and chilled on ice. Lysis was performed by 372 directly adding 2x Laemmli sample buffer (supplemented by 2-mercaptoethanol. Bio-Rad 373 Laboratories) at a ratio of 1 µL per 0.1 mg of cell pellets and pipetting up and down. Lysates were 374 boiled immediately afterward for 10 min and then chilled on ice. 8 µL of supernatant was loaded 375 onto each lane of a 15-well, 0.75-mm SDS-PAGE mini gel.

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376 Primary antibodies used included anti-BUBR1 (Bethyl Laboratories A300-995A, 1 : 1000), anti-

377 MAD1 (GeneTex GTX109519, 1 : 2000), anti-CDC20 (Santa Cruz Biotechnology sc-13162, 1 :

- 378 200), anti-BUB3 (Sigma-Aldrich B7811, 1 : 500), anti-GAPDH (Proteintech 60004-1-lg, 1 : 5000).
- 379

380 RNA interference

Cells were transfected with siRNA in the morning. 2.5 mM of thymidine was added 8 h later and cells were incubated overnight. The next morning, cells were released from the thymidine block into fresh media. This sequence was then repeated once and cells were released into FluoroBrite[™] DMEM (Gibco) supplemented with 9% (by volume) of FBS and 1× GlutaMAX and incubated for 6 h before the addition of mitotic drugs. Drugs were left to take effect for 1 h before imaging.

387 Sense-strand sequences and working concentrations of small interfering RNA duplexes (siRNAs) 388 used in this study include the SPC25 siRNA (5'- GCCUGCGAAGCAUUGUCCUACAUAA-3', 40 389 nM ⁵³), BUBR1 siRNA (5'-GAUGGUGAAUUGUGGAAUA-3', 40 nM ⁵⁴), BUB1 siRNA (5'-390 CGAAGAGUGAUCACGAUUU-3', 40 nM ⁵⁵), mNeonGreen siRNA (5'-391 ZW10 GAGCUGAAGCACUCCAAGACA-3'. 40 nM) and siRNA (5'the UGAUCAAUGUGCUGUUCAA-3', 100 nM ³⁸). The siRNAs against all five B56 isoforms were 392 taken from the second pool in ⁵⁶. Desalted siRNA duplexes modified by double-deoxythymidine 393 394 overhangs at 3'-ends of both strands were synthesized by Sigma. The AllStar siRNA (QIAGEN) 395 was used for the negative control. All siRNAs were transfected into the cells via Lipofectamine 396 RNAiMAX (Invitrogen).

397

398 Live-cell imaging

Cells were plated in a Nunc Lab-Tek II chambered cover glass (Thermo Scientific) or a 35-mm cover glass-bottomed dish (MatTek) and treated with drugs and/or siRNAs as described above. For imaging, the chambered cover glass or the cover glass-bottomed dish was loaded into a CU-501 temperature and gas control system (Live Cell Instrument). The sample holder was maintained at 37°C and ventilated by humidified 5% of CO₂ and the objective was maintained at 37°C by a heating band.

405 MPS1sen-KT FRET and immunofluorescence experiments were performed on a Nikon Eclipse
406 Ti-E/B inverted microscope, with a CFI Plan Apochromat Lambda 100×, 1.45 NA oil objective
407 (Nikon). The microscope was equipped with an H117E1 motorized stage (Prior Scientific) and a

408 NanoScanZ 100 piezo stage (Prior Scientific). A SPECTRA 5-LCR-XA Light Engine (Lumencor) 409 served as the excitation light source. The 475 nm-centered band of excitation light was used for 410 the green channel and the 575/30nm-filtered band of excitation light was used for the red channel. 411 An ET-EGFP/mCherry filter cube (Chroma Technology) was used as the dichroic mirror, where 412 the built-in emission filter on the cube has been removed. Emission light in the red channel was 413 filtered by an ET632/60m (Chroma Technology). Emission light in the green channel was filtered 414 by an ET525/50m (Chroma Technology). Emission filters were mounted on a high-speed filter 415 wheel (Prior Scientific) positioned in the light path. Images were acquired with an iXon3 EMCCD 416 camera (Andor Technology) operating in the conventional CCD mode. Signaling kinetochores in 417 GSK923295-treated cells were identified by their polar positioning and the enrichment of localized 418 SAC proteins on them.

419 The FRET measurement with MPS1sen-KT was performed on a Nikon Eclipse TiE/B inverted 420 microscope, with a CFI Plan Apochromat VC 100×, 1.40 NA oil objective (Nikon). The microscope 421 was equipped with an H117E1 motorized stage (Prior Scientific), a NanoScanZ 100 piezo stage 422 (Prior Scientific), and an X-Light V2 L-FOV confocal unit with 60-µm pinholes (CrestOptics). A 423 CELESTA Light Engine (Lumencor) served as the excitation laser source. The 477-nm line (at 424 25% power with an exposure time of 400ms for each frame) was used for both the green and the 425 FRET channels and the 546-nm line (at 50% power with an exposure time of 400ms for each 426 frame) was used for the red channel. A ZT488/543rpc (Chroma Technology) was used as the 427 dichroic mirror. Emission through both the red and the FRET channels was filtered by an 428 ET605/52m (Chroma Technology) while emission through the green channel was filtered by an 429 ET525/36m (Chroma Technology). Images were acquired by a Prime 95B 25mm sCMOS camera 430 (Teledyne Photometrics).

431 Time-lapse live-cell imaging related to the knockdown-rescue experiments was performed on an 432 ImageXpress Nano Automated Imaging System (Molecular Devices). A SOLA Light Engine 433 (Lumencor) served as the excitation light source. Cells were plated on 24-well cell imaging plates 434 (black plate with treated glass bottom, Eppendorf) and treated with drugs and/or siRNAs 435 accordingly. Input humidified 5% CO₂ flow was maintained at around 19 psi and the environment 436 chamber was maintained at 37°C. All SAC proteins tagged by mNeonGreen (BUBR1, BUB1, and 437 MAD1) feature inhomogeneous distributions between the cytosol and the nucleus/nuclear 438 envelope in the prophase (data not shown). In principle, they can indicate the accurate timing of 439 the nuclear envelope breakdown (NEBD) in each cell during the time-lapse imaging. However, 440 due to the resolution limit and for consistency, we determined mitotic duration mainly based on

441 cell morphology (from rounding-up at the NEBD to elongation at the anaphase onset) from
442 transmitted-light images. This was facilitated by the semi-automatic image analysis pipeline
443 available at https://github.com/CreLox/IXNAnalysis

444

445 Immunofluorescence

446 Hela cells that exogenously expressing mNG-Bub1 were plated on a 12-mm coverslip and 447 synchronized by thymidine (2.5 mM for 8 hours) followed by RO3306 treatment (15 nM for 20 448 hours). G2/M arrested cells were washed three times with PBS and released into media 449 containing GSK923295 (236 nM) for 4 hours) to get arrested in Mitosis. For immunolabeling, the 450 cells were pre-extracted with 0.5% Triton X-100 in PHEM (240 mM Pipes, 100 mM HEPES, 8 mM 451 MqCl₂ and 40 mM EGTA) and then fixed with 4% PFA for 10 min. The coverslips were washed 452 three times with PHEM buffer and blocked for 30 min at room-temperature with 5% Donkey serum. 453 Next, the coverslips were incubated in anti-Mad1 (1:1000, Sigma M80691), anti-Bub1 (1:500, 454 Thermo A300-373A), or anti-BubR1 (1:1000, Thermo Scientific A300-373A) over night at 4°C. 455 The next day, the coverslips were washed 4 times with PHEM containing 0.05% Tween-20 and 456 incubated with appropriate Alexa Fluor 633-conjugated secondary antibodies (Goat anti-Mouse, 457 Invitrogen A21052 or Goat anti-Rabbit, Invitrogen A21071) at a 1:10000 dilution for 45 min at 458 room temperature in the dark. Following secondary antibody labeling, the coverslips were washed four times and mounted in an antifade solution (ProLong; Molecular Probes). 459

460 Immunofluorescence images were acquired on the spinning disk confocal microscope described
461 above. For each field of view, 31 *z*-sections were acquired at 0.2-µm steps.

462

463 **Quantification of kinetochore-localized fluorescence signal**

464 Quantification of the kinetochore-localized fluorescence signal was performed using a custom 465 graphical user interface written in MATLAB. The methodology has been described previously ^{11,53}. 466 Briefly, individual kinetochores were identified in the mCherry channel based on the Spc25-467 mCherry fluorescence. A 6×6-pixel box was drawn centered on the maximum intensity pixel in 468 the in-focus mCherry plane, and the mNG intensity of all the pixels within this box was summed 469 to obtain the mNG fluorescence signal for that kinetochore. The local background was defined as 470 the median intensity value of the perimeter pixel intensities in a 10×10 pixel box concentric with 471 the 6×6 pixel kinetochore box.

472 For immunofluorescence quantification involving both RPE-1 and HeLa cells, anti-ACA antibodies

473 probed with Alexa Fluor 488-conjugated secondary antibodies were used to identify the

474 centromeres. Alexa Fluor 633-conjugated secondary antibodies were used to visualize primary

- 475 antibodies against the respective SAC proteins.
- 476

477 Fluorescence correlation spectroscopy (FCS)

The total number of fluorophores in a homogeneous solution is $N_{\text{total}} := N_A c V_{\text{total}}$, where N_A is the Avogadro constant, *c* is the molar concentration of the fluorophore, and V_{total} is the total volume of the solution. The probability that a specific fluorophore molecule is within the excitation volume $V_0(\ll V_{\text{total}})$ at any given time is $p_0 := V_0/V_{\text{total}} \ll 1$. For freely diffusive fluorophores in a diluted solution, whether or not a specific fluorophore is within the excitation volume is independent of each other. Thus, the number of fluorophores inside the excitation volume at any given time N_0 has a binomial distribution $B(N_{\text{total}}, p_0)$. Therefore,

$$G_0 \coloneqq \frac{\sigma_{N_0}^2}{\langle N_0 \rangle^2} = \frac{1 - p_0}{N_{\text{total}} p_0} \approx \frac{1}{N_{\text{A}} c V_0}$$

485

486 Under a fixed live-cell imaging setup [which includes the microscope (its alignment and the 487 objective), the wavelength of the excitation light, the thickness of the coverslip (affecting the actual 488 working distance), and the refractive index of the cytosol], V_0 is fixed. Therefore, G_0 is inversely 489 proportional to the molar concentration of the fluorophore. The average number (or the variance 490 of the number) of fluorophores inside the excitation volume observed over a long period should 491 be close to the theoretical mean $\langle N_0 \rangle$ (or the theoretical variance σ^2_{N0}).

492 All FCS data were collected on an Alba v5 Laser Scanning Microscope (ISS), connected to an 493 Olympus IX81 inverted microscope main body [equipped with a UPLSAPO60XW objective (1.2 494 NA, Olympus)]. A Fianium WL-SC-400-8 laser (NKT Photonics) with an acoustooptic tunable filter 495 was used to generate excitation pulses at a wavelength of 488 nm and a frequency of around 20 496 MHz. Excitation light was further filtered by a Z405/488/561/635rpc quadband dichroic mirror 497 (Chroma). Emission went through a 655DCSPXR short-pass dichroic mirror (Chroma) and an 498 FF01-531/40-25 filter (Semrock) and was finally detected by an SPCM-AQRH-15 avalanche 499 photodiode (Perkin Elmer). The time-correlated single photon counting module to register 500 detected photon events to excitation pulses was SPC830 (Becker & Hickl). Data acquisition was 501 facilitated by VistaVision (ISS). The excitation volume (V_0) was calibrated by taking FCS data from 502 TetraSpeck[™] microspheres (0.1 µm, Invitrogen) of known concentrations.

503 Statistical analysis

504 Statistical analysis was conducted using GraphPad Prism version 9.

505

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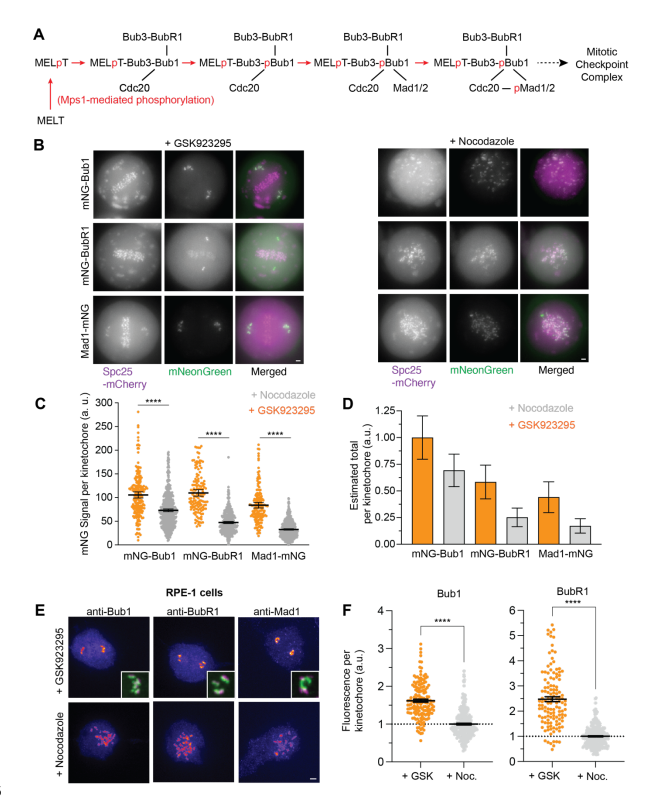
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Figure 1 The SAC protein recruitment at signaling kinetochores depends on the number of
 signaling kinetochores in the cell.

(A) A simplified schematic of the MELT motif-mediated recruitment of SAC proteins to the
 kinetochore. Red arrows signify Mps1-mediated phosphorylation, black lines indicate protein protein interactions. The RZZ pathway, which is activated by Bub1 ^{29,35,40}, is not shown.

692 **(B)** Representative fluorescence micrographs of mitotic HeLa cells expressing the indicated 693 proteins treated with GSK923295 (left) and nocodazole (right; scale bar $\sim 1.6 \,\mu$ m)

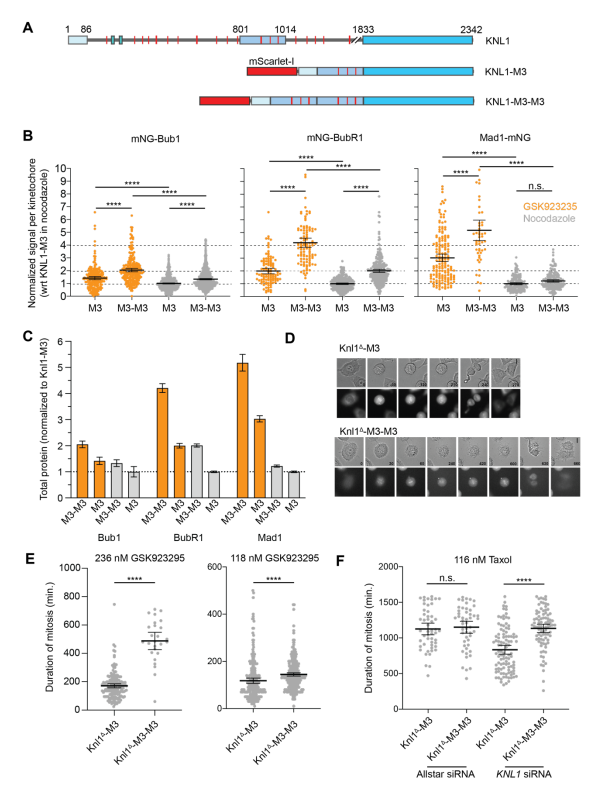
694 Quantification of the fluorescence signal per signaling kinetochore. Horizontal lines display (C) 695 mean \pm 95% confidence intervals (n = 200, 142, 160 in GSK923295-treated cells and 562, 446, 696 and 658 in nocodazole-treated cells for Bub1, BubR1, and Mad1 respectively, observations 697 pooled from \geq 2 technical repeats). Right: Spc25-mNG signal per kinetochore quantified from 698 images acquired under identical imaging conditions (n = 178). Welch's *t*-test was used to 699 determine whether the sample means are significantly different. In all the figures in this study, **** 700 indicates p < 0.0001, *** indicates 0.0001 , ** indicates <math>0.001 , * indicates701 0.01 , and n.s. (not significant) indicates <math>p > 0.05.

(D) Estimation of total protein recruited per kinetochore based on the fluorescence
 measurements shown in C and immunoblot quantification of labeled to unlabeled fraction for each
 protein shown in Figure S1.

705 **(E)** Representative immune-fluorescence micrographs (pseudo-colored as heatmaps) of 706 RPE-1 cells. Insets show magnified regions of interest around unaligned chromosomes with 707 antibodies labeling SAC proteins pseudo colored as green and antibodies labeling centromeres 708 pseudo colored as magenta. Notice in the case of Mad1 a ring-like structure in the polar region 709 presumably representing the Mad1 protein aggregated by its dynein-mediated transport from the 710 kinetochore. This structure prevented us from quantifying the amount of kinetochore localized 711 Mad1. Scale bar ~ $2.44 \mu m$.

712 **(F)** Quantitation of immunofluorescence per kinetochore for Bub1 and BubR1 (each dot 713 represents a single kinetochore; mean \pm SEM is overlaid for each group. For Bub1, n = 255 714 (GSK923235-treated) and 157 kinetochores (nocodazole-treated) from two technical repeats. For 715 BubR1, n = 125 (GSK923235-treated) and 288 kinetochores (nocodazole-treated) from two 716 technical repeats. Welch's *t*-test used to compare sample means.

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Figure 2 The number of MELT motifs in Knl1 affects SAC protein recruitment to the 723 kinetochore and SAC signaling strength.

724 **(A)** Schematics of the full-length, endogenous Knl1 and the recombinant Knl1^{Δ}-M3 and Knl1^{Δ}-725 M3-M3 versions. The expression of Knl1^{Δ}-M3 (or Knl1^{Δ}-M3-M3) is under the regulation of a 726 constitutive *EF1a* promoter.

727 Quantification of the fluorescence signal per signaling kinetochore for the indicated **(B)** 728 protein. In each case, the signals were normalized using the mean signal for each protein 729 measured in nocodazole-treated cells expressing Knl1^Δ-M3 (mNG-Bub1: n =844 and 602 for 730 Knl1^Δ-M3-M3 and Knl1^Δ-M3 respectively in nocodazole-treated cells and 222 and 120 for Knl1^Δ-M3-M3 and Knl1^{Δ}-M3 respectively in GSK923295-treated cells, pooled from \geq 2 technical repeats; 731 732 mNG-BubR1: n = 278 and 272 for Knl1^{Δ}-M3-M3 and Knl1^{Δ}-M3 respectively in nocodazole-treated cells and 110 and 107 for Knl1^Δ-M3-M3 and Knl1^Δ-M3 respectively in GSK923295-treated cells, 733 pooled from \geq 2 technical repeats; Mad1-mNG: n = 182 and 184 for Knl1^{\triangle}-M3-M3 and Knl1^{\triangle}-M3 734 respectively in nocodazole-treated cells and 52 and 166 for Knl1^Δ-M3-M3 and Knl1^Δ-M3 735 736 respectively in GSK923295-treated cells, pooled from \geq 2 technical repeats).

737 (C) Estimation of total protein amount per kinetochore relative to the total Bub1 amount per738 kinetochore in drug-treated cells (colors same as in as in B).

739 **(D)** Representative micrographs show mitotic progression of cells with either Knl1^{Δ}-M3 (top) 740 or Knl1^{Δ}-M3-M3 (bottom, scale bar: 10 µm). Time stamps are in minutes.

741 **(E-F)** Duration of mitosis for cells treated with the indicated concentration of GSK923295 and 742 Taxol (left: n = 162 and 26 for Knl1^{Δ}-M3 and Knl1^{Δ}-M3-M3 respectively from two technical repeats; 743 right: n = 235 and 339 for Knl1^{Δ}-M3 and Knl1^{Δ}-M3-M3 respectively, n = 51 and 52 for Allstar siRNA 744 + taxol treatment and n = 102 and 88 for KNL1 siRNA + taxol treatment for Knl1^{Δ}-M3 and Knl1^{Δ}-745 M3-M3 respectively). Welch's *t*-test was used to compare sample means.

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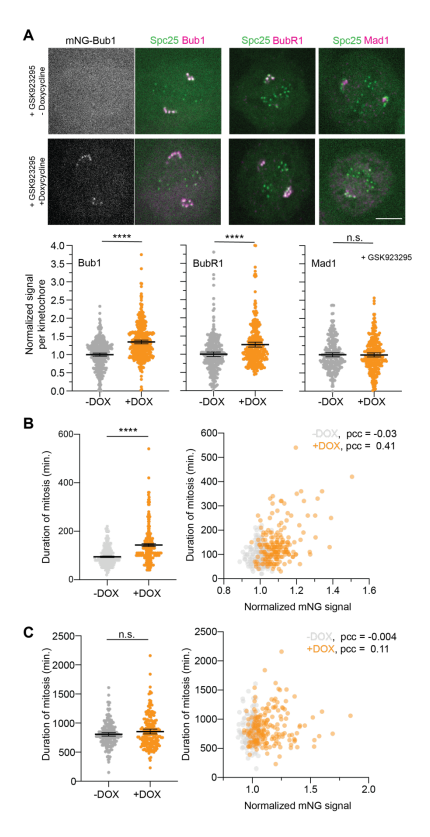


Figure 3 – Bub1 over-expression results in higher Bub1 and BubR1 recruitment per
 kinetochore and prolongs the delay in anaphase onset.

757 (A) Top panel: representative fluorescence micrographs of control (top) cells and cells treated 758 with doxycycline to exogenously express mNG-Bub1. Scale bar $\sim 4.92 \mu m$. Bottom panel: 759 guantification of immunofluorescence signal per kinetochore for the indicated proteins in control 760 cells and cells treated with doxycycline to induce exogenous mNG-Bub1 expression. Signal 761 values were normalized with the average signal for the respective protein in doxycycline-untreated 762 cells. (n = 344 and 390 for Bub1, 346 and 387 for BubR1, and 234 and 275 for Mad1 in untreated 763 and doxycycline-treated cells respectively, observations pooled from 3 technical repeats). 764 Welch's *t*-test used to compare sample means.

765 Left: Duration of mitosis in control and doxycycline-treated cells in media containing 25-**(B)** 766 37 nM GSK923295 (n = 195 and 187 for untreated and doxycycline-treated cells respectively, 767 from 2 technical repeats; p < 0.0001 for doxycycline-treated and GSK923295 treated cells). 768 Horizontal lines represent mean ± 95% confidence intervals on the mean. Right: Correlation 769 between the duration of mitosis and the average cellular level of the exogenously expressed 770 mNG-Bub1. mNG-Bub1 signal per cell was normalized with the average signal measured in 771 control cells. Pearson's correlation coefficient (pcc) for each data set is noted in the figure (p =772 0.6222 and < 0.0001 respectively for untreated and doxycycline-treated cells respectively)

773 Left: Duration of mitosis in control and doxycycline-treated cells in media containing 330 (C) 774 nM nocodazole (n = 189 and 185 for untreated and doxycycline-treated cells respectively, from 2 775 technical repeats). Horizontal lines represent mean ± 95% confidence intervals on the mean. 776 Right: Correlation between the duration of mitosis and the average cellular level of the 777 exogenously expressed mNG-Bub1. mNG-Bub1 signal per cell was normalized with the average 778 signal measured in control cells. Pearson's correlation coefficient (pcc) for each data set is noted 779 in the figure (p = 0.053 and 0.1309 respectively for untreated and doxycycline-treated cells 780 respectively).

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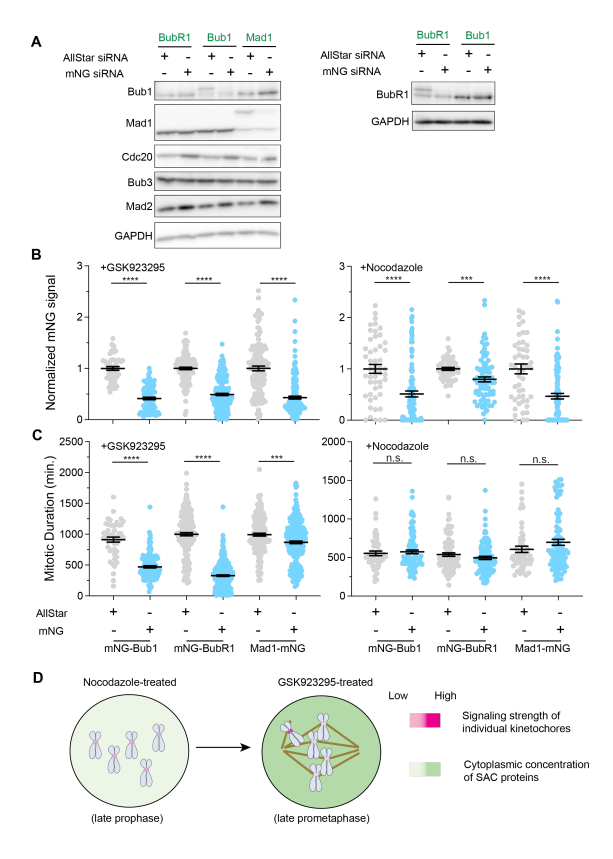


Figure 4 – Partial depletion of Bub1 and BubR1, but not Mad1, weakens the signaling potential of the SAC.

(A) Immunoblot analysis of total cell lysates shows that the mNG-tagged SAC protein is
 specifically targeted for RNAi after a 48-hour treatment with mNG siRNA.

787 **(B)** Quantification of the mNG-signal per cell under the indicated conditions (For the 788 experiment involving GSK923295 treatment, n = 50, 126, 130 for mNG-Bub1, mNG-BubR1, and 789 Mad1-mNG cells respectively treated with Allstar siRNA; n = 100, 219, and 177 for mNG-Bub1, 790 mNG-BubR1, and Mad1-mNG cells respectively treated with mNG siRNA; data collected in two 791 technical repeats). For the experiment involving nocodazole treatment, n = 50, 50, and 50 for 792 mNG-Bub1, mNG-BubR1, and Mad1-mNG cells respectively treated with Allstar siRNA; n = 100, 793 85, and 100 for mNG-Bub1, mNG-BubR1, and Mad1-mNG cells respectively treated with mNG 794 siRNA; data collected from 2 technical repeats). Each dot represents a measurement from one 795 cell, mean ± SEM overlaid. Welch's *t*-test used to compare sample means.

796 **(C)** Duration of mitosis for same cells as in B in media containing GSK923295 and nocodazole 797 (for GSK923295-treated cells: $p \le 0.0001$ based on ordinary one-way ANOVA; for nocodazole-798 treated cells, p > 0.253 based on ordinary one-way ANOVA). Each dot represents a measurement 799 from one cell, mean ± SEM overlaid. Pairwise comparisons of sample means performed using 800 Tukey's multiple comparison test.

(D) Model depicts the combined effect of the number of signaling kinetochores per cell and
 the availability of free, cytoplasmic SAC proteins on the signaling strength of individual
 kinetochores.

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807 Supplementary figures

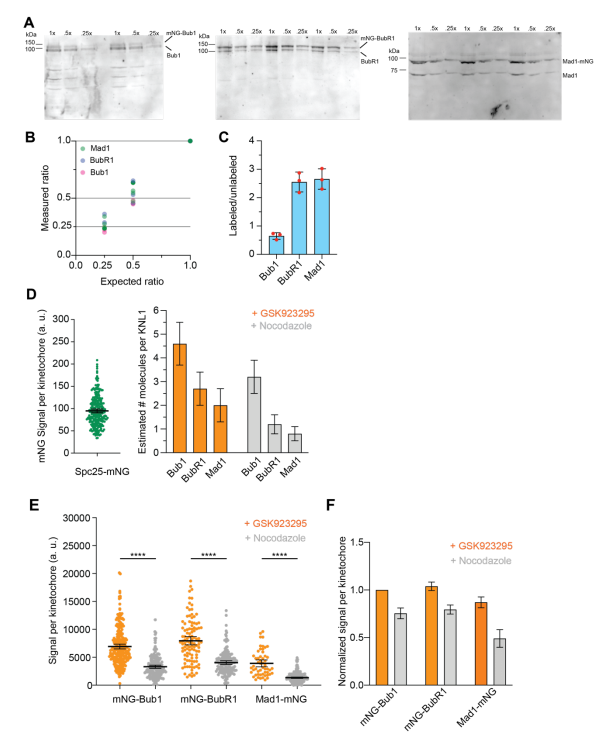


Figure S1 (related to Figure 1) Assessment of the relative amounts of mNG-labeled and
 unlabeled proteins in the three cell lines.

(A) Immunoblot analysis of whole mitotic cell lysates of the three cell lines probing for the indicated proteins. Two out of the three technical repeats for the mNG-Bub1 cell line and all three technical repeats for the mNG-BubR1 and Mad1-mNG cell lines are displayed. Numbers at the top denote the lysate volume loaded to ascertain the linearity of the fluorescent secondary antibody signal.

(B) Ratio metric analysis of serial dilutions confirms that band intensities are directly
proportional to the amount of protein loaded. The band intensities were quantified using the 'Gel
Analyzer' tool in Fiji. The ratio of labeled to unlabeled protein for each technical repeat was defined
as the average of the ratios for the three 'repeats'.

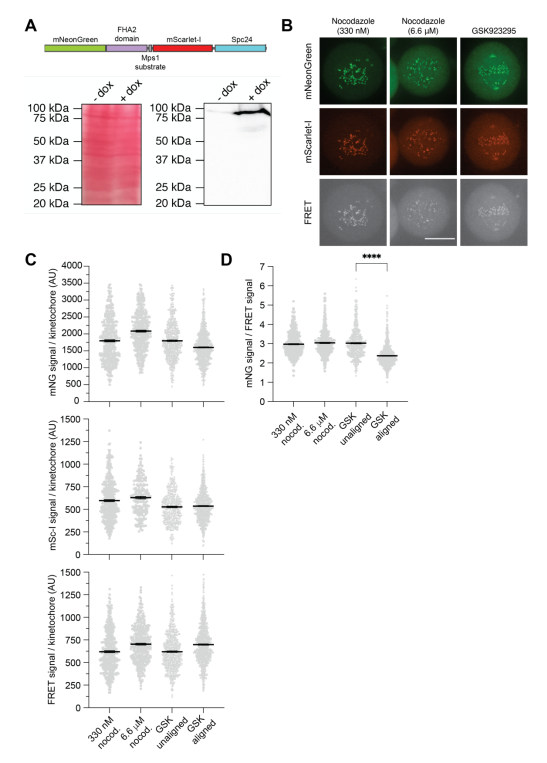
820 (C) The ratio of the mNG-tagged to unlabeled species for the three proteins. Error bars display821 the standard deviation.

(D) Left: the beeswarm plot displays the quantification of Spc25-mNG signal per kinetochore
in images obtained under identical imaging conditions as in Figure 1C. The bar plot on the right
displays the estimated number of molecules of the mNG-tagged SAC protein per Knl1 calculated
by dividing the mean signal per kinetochore for each protein with the mean Spc25-mNG signal
per kinetochore as the reference. Error bars represent mean ± SEM.

(E) Left: Quantification of the fluorescence signal per signaling kinetochore in cells treated with
siRNA against *ZW10* (n = 108, 242, 54 in GSK923295-treated cells and 166, 178, 219 in
nocodazole-treated cells for Bub1, BubR1, and Mad1, respectively, pooled from 2 technical
repeats).

(F) Average signals from E are normalized using the mNG-Bub1 signal per kinetochore.Horizontal lines display the standard error on the mean ratio.

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835 Supplementary Figure S2 (related to Figure 1) Net phosphorylation at signaling 836 kinetochores is the same in nocodazole- and GSK923295-treated cells.

(A) Top: Schematic of the modified MPS1sen-KT ³⁴ that uses the mNeonGreen and mScarlet-I
 as the FRET pair. Bottom: Using an antibody against DsRed2 (which detects mScarlet-I), we

839 confirmed that MPS1sen-KT (with a theoretical molecular weight of 97.2 kDa) is expressed as a

840 full-length protein with negligible partial degradation in HeLa A12 treated with doxycycline. A

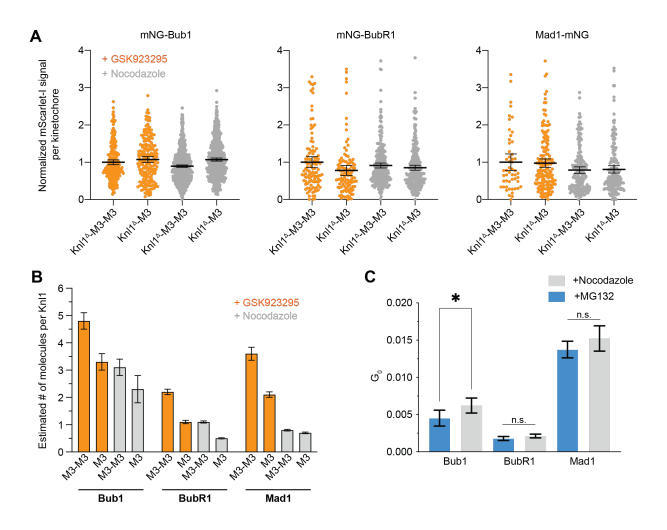
841 Ponceau S staining of the same blot before membrane blocking (left panel) serves as the loading

and membrane transferring control.

- (B) Representative images of cells expressing MPS1sen-KT. The look-up table for each channel
 (row) is universal for different groups (column). Scale bar, 10 μm.
- 845 (C) Raw fluorescence quantification at signaling kinetochores under various drug treatments in 846 the green, red, and FRET channels. Each gray dot represents a single kinetochore measurement. 847 Data from cells treated with 45 nM, 90 nM, and 200 nM GSK923295 were pooled together (the 848 mean values are not significantly different from one another; data not shown) to simplify the 849 presentation. Data were compiled from at least two independent experiments (more than 40 cells 850 in total for each group). Mean values ± 95% confidence intervals are overlaid. The variation across 851 different groups can be attributed to different FRET efficiencies (which we intend to quantify) and 852 day-to-day variations in the doxycycline-induced expression of MPS1sen-KT. To cancel the effect 853 of these variations, a normalized metric of the FRET efficiency (mNG/FRET) was used in this 854 study.
- 855 (D) A summary of a normalized FRET metric (mNeonGreen signal/FRET signal) in HeLa A12 856 cells treated with different drugs at various concentrations. Each gray dot represents a single 857 kinetochore measurement. Data were compiled from at least two independent experiments (more 858 than 40 cells in total for each group). As in C, Data from cells treated with 45 nM, 90 nM, and 200 859 nM GSK923295 were pooled together. Mean values ± 95% confidence intervals are overlaid. The 860 Welch's ANOVA test [W(DFn,DFd) = 1.339(2.000,917.0), P = 0.2626] was performed for the first 861 3 columns (unaligned, signaling kinetochores). The unpaired t-test with Welch's correction 862 (P<0.0001) was performed to compare aligned non-signaling kinetochores with unaligned 863 signaling kinetochores in GSK923295-treated cells.
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Figure S3 (related to Figure 2) Assessment of the kinetochore recruitment of mScarlet-I KnI1^Δ-M3-M3 and mScarlet-I-KnI1^Δ-M3.

(A) mScarlet-I fluorescence signal per kinetochore indicating the amount of recombinant Knl1
versions (noted by the X-axis labels). Minor differences in the incorporation of the recombinant
Knl1 cannot explain the observed differences in SAC protein recruitment in each case. Note that
the average value for the mScarlet-I-Knl1^Δ-M3-M3 signal in GSK939295 in each data set was
used as the normalization factor for each cell line. The numbers of data points are identical to
those in Figure 2.

(B) Estimation of the copy number of each SAC protein per Knl1 molecule using the Spc25-mNG
fluorescence signal per kinetochore as the fiduciary standard (shown in Figure S1). Error bars
indicate mean ± SEM.

(C) G_0 of the three proteins in metaphase-arrested (MG-132-treated) and nocodazole-treated cells as measured by Fluorescence Correlation Spectroscopy (FCS). This is the magnitude of the

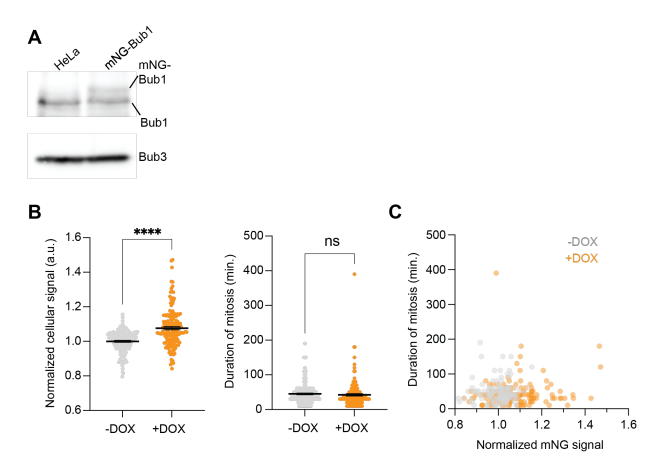
- 881 auto-correlation function for the fluorescence fluctuation data for 0-time delay. It is inversely
- proportional to the average number of fluorophores in the focal volume, i.e. the concentration of
- the fluorophore ⁵⁷. Mean values under the two conditions compared using Welch's *t*-test.

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Figure S4 (related to Figure 3) mNG-Bub1 over-expression does not affect normal mitotic progression of HeLa A12 cells.

(A) Immunoblot shows doxycycline-induced expression of exogenous mNG-Bub1.

(B) Left: The average mNG signal in mitotic HeLa cells that are either untreated or treated with doxycycline to induce exogenous mNG-Bub1 expression (normalized to the average value of mNG fluorescence in untreated cells, which also represents the background signal). Right: the duration of mitosis for the same cells. (n = 155 and 154 for -dox and +dox respectively pooled from 2 technical repeats). The horizontal lines indicate mean ± SEM. Welch's *t*-test was used to compare the sample means.

898 **(C)** Scatterplot displays the correlation between the cellular mNG signal (normalized as in B) and 899 the duration of mitosis (Pearson's correlation coefficient = 0.05 and 0.13 with p = 0.5 and 0.1 for 900 -Dox and +Dox respectively).

901 Supplementary Videos

- 902 Video S1 Time-lapse imaging of HeLa cells treated with 25 nM GSK923295. Time hr:min.
- 903 Video S2 Time-lapse imaging of HeLa cells treated with 25 nM GSK923295 and 1 μg/ml of
- 904 Doxycycline to exogenously over-express mNG-Bub1. Time hr:min.
- 905 Video S3 Time-lapse imaging of HeLa cells treated with 330 nM Nocodazole. Time hr:min.
- 906 Video S4 Time-lapse imaging of HeLa cells treated with 330 nM Nocodazole and 1 µg/ml of
- 907 Doxycycline to exogenously over-express mNG-Bub1. Time hr:min.
- 908 Video S5 Time-lapse imaging of genome-edited HeLa cells heterozygous for mNG-Bub1 and

909 exogenously expressing H2B-mCherry. Cells were released from a G1/S block into media

- 910 containing 236 nM GSK923295. Time hr:min.
- 911 Video S6 Time-lapse imaging of genome-edited HeLa cells heterozygous for mNG-Bub1 and
- 912 exogenously expressing H2B-mCherry. Cells were treated with siRNA targeting mNG to partially

913 deplete BubR1 and released from a G1/S block into media containing 236 nM GSK923295. Time

914 - hr:min.