Kre28–Spc105 interaction is essential for Spc105 loading at the kinetochore

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1. Introduction

During eukaryotic cell division, kinetochores (KTs) facilitate faithful segregation of genetic material from mother to daughter cells. Each KT is a large protein machine that assembles on a specialized chromatin domain called the centromere and establishes end-on attachments between the sister chromatids and spindle MTs emanating from opposite spindle poles. The budding yeast Saccharomyces cerevisiae has one of the ‘simplest’ KT structures known to date, yet it harbours approximately 70 protein subunits. Components of the yeast KT can be divided into two main categories. The first category contains the centromeric DNA-binding components and their associated network known as CCAN (constitutive centromere-associated network). The second comprises the MT-binding protein network: the KMN super-complex, the fungi-specific Dam1 complex and the MT plus-end-binding protein Stu2 [1–5]. The budding yeast KT incorporates each of these proteins positioned at well-defined average locations along with the KT-MT attachment (figure 1a; electronic supplementary material, figure S1B) [7,8]. For molecular and cell biologists, the budding yeast KT serves as an excellent model to determine the mechanisms underlying KT functions.

Spc105 is an essential KT protein that gets co-purified with the COMA complex subunit Mcm21 and with the MIND complex (Mtw1-Nsl1-Nnf1-Dsn1) [9,10]. It forms a complex with another essential KT protein, Kre28, also known as Ydr532C. Kre28 is an orthologue of human Zwint1, C. elegans Kbp-5 and S. pombe Sos7 [10–13]. Previous studies, using in vitro and in vivo...
Figure 1. Defining the localization of the C-terminus of Kre28 in the metaphase kinetochore using FRET and high-resolution colocalization. (a) The organization of kinetochore proteins along with the microtubule axis in bioriented yeast kinetochores. Positions of C termini of Spc24, Ndc80 and the N terminus of Ndc80 are indicated. (b) Frequency distribution of the distance between the centroids of Kre28-mCherry (Kre28-C) and GFP-Ndc80 (N-Ndc80). The black curve line is the maximum-likelihood fit. (c) Proximity ratio for FRET between fluorophores fused to either Spc24-C or Ndc80-C and Kre28-C (mean ± s.e.m.) in bioriented kinetochore clusters. At least 29 bioriented kinetochores were analysed for this dataset. The p-values obtained from unpaired t-tests are displayed above the plot. (d) Proximity ratio for FRET between adjacent C termini of Kre28 in bioriented yeast kinetochores. Eighty-two kinetochores were analysed to obtain these data. (e) Top: line diagram of Spc105 molecule. The illustration was duplicated from our previous study [6]. Red bars represent PP1/Glc7 recruitment site (amino acid 75–79), and six MELT repeats. Amino acid locations of GFP fusion are shown at the top on amino acid residues 222, 455, 709 and C (917). Bottom: proximity ratio for FRET between Kre28-C and different amino acid positions of Spc105 molecules in bi-oriented kinetochores. At least 35 kinetochore foci were analysed for this graph. The p-values obtained from one-way ANOVA test performed on the data are mentioned above the plot. (f) Localization of C termini of Kre28 in KMN network of metaphase kinetochores of yeast cells.
experiments, provide some insights into how Spc105 and Kre28 are assembled at the KTs. Still, the specific function of Kre28 remains unclear. How Kre28 localizes within the KT-MT attachment site is also unknown [14–16]. Here we define the localization of Kre28 in KT-MT attachment sites of bioriented KTs and elucidate its functional role.

2. Results

2.1. Localization of Kre28 in the KMN network of bioriented kinetochores

We have previously shown that the precise organization and alignment of Spc105 can influence proper spindle assembly checkpoint (SAC) activation and silencing [17–20]. Kre28, being an essential component of the KT, may also contribute to the organization of Spc105 into the yeast KT. Zwint1, the human orthologue of Kre28, localizes very close to Cdc20 at the human KTs [21,22]. Therefore, Kre28 may play a role in determining the Spc105 organization.

To determine Kre28’s position with respect to Spc105, we first had to define the organization of the entire Spc105 protein within the yeast KT. Previous studies show that the C-terminal RWD domain of Spc105 binds directly to the Mtw1 complex and remains in the proximity of Spc24/Spc25 C-termini [15,23,24]. On the other hand, the N-terminus of Spc105 (abbreviated as N-Spc105) consists of a long, disordered phosphodomain that lies somewhere between the Dam1 complex and the C-termini of Ndc80 and Nuf2 within the Ndc80 complex [6–8]. To map out the overall organization of the Spc105 phosphodomain, we inserted a GFP at locations within Spc105 that demarcate domains predicted to possess secondary structure (see electronic supplementary material, figure S1). Additionally, we tagged three different KT subunits to position mCherry at different locations along with the KT-MT attachment (electronic supplementary material, figure S1). Quantification of FRET between the GFP inserted in Spc105 and one of the three mCherry acceptors shows that despite being discarded, the Spc105 phosphodomain localization is mainly limited to a span between the Dam1 complex and the C-terminus of the Ndc80 subunit (abbreviated at Ndc80-C) of the Ndc80 complex. The disordered nature of the phosphodomain also gave rise to FRET between different sections of adjacent Spc105 molecules (electronic supplementary material, figure S1D).

Having established Spc105, we examined the localization of Kre28 by centroid measurement and FRET assay. Previous literature suggested that the C-terminal structured domains of Spc105 harbour interaction sites for Kre28 [22,25].

To define the localization of Kre28 within the KMN network of bioriented KTs, we performed high-resolution colocalization to measure the mean separation between Kre28-C and the N termini of the Ndc80 subunit (N-Ndc80) in the bioriented KTs of yeast (figure 1a) [8]. We observed that the C-terminus of Kre28 is positioned between 45 and 50 nm from N-Ndc80, which is consistent with previously published work with Zwint1 [22].

To determine Kre28 localization with higher resolution, we quantified FRET between Kre28-C with either Spc24-C or Ndc80-C in metaphase cells. We obtained a low to moderate proximity ratio in both cases, indicating that Kre28-C may localize somewhere between C-termini of Spc24 and Ndc80 (figure 1b). The absence of FRET between adjacent Kre28-C termini (Kre28-GFP/Kre28-mCherry) indicated that the C termini of Kre28 molecules are farther apart than 10 nm in metaphase (figure 1c). We also measured FRET between Kre28-mCherry and GFP inserted at different positions of Spc105 (222nd, 455th, 709th, or the C terminus) to find that the C terminus of Kre28 are proximal of the KT-binding RWD domain (RING finger, WD repeat, DEAD-like helicases) of Spc105 (figure 1d,e). A previous study suggested that the stoichiometry of Kre28 and Spc105 is 2:1 [12]. However, a comparison of the intensity of Spc105-GFP or Kre28-mCherry signal per KT revealed that there is one molecule of Kre28 per Spc105 molecule in bioriented KTs of yeast (electronic supplementary material, figure S2A–D).

2.2. Interaction of Kre28 with coiled coil domain of Spc105

Studies of Zwint1 (orthologue of Kre28 in humans) found that it interacts with a domain within amino acid 1980–2109 of human Spc105 [25]. Protein cross-linking experiments also revealed that coiled-coiled domains (CCs) Kre28212–169 and Kre28229–259 interact with Spc105201–210 in the predicted CC of Spc105 [14]. We wanted to uncover the domains within both Kre28 and Spc105 that are necessary for their mutual interactions.

To study these domains in the ex-vivo condition, we first used the yeast two-hybrid assay. We chose Kre28 fragments (amino acid 1–201 and 202–385, based on predicted secondary structure of Spc105, http://www.compbio.dundee.ac.uk/jpred4/index_up.html) and Spc105 CC (455–708) and the C-terminal RWD domain of Spc105 (amino acid 709–917 [15]; see also electronic supplementary material, figure S3A). Both Kre28FL and Kre28201 showed interactions with CC as indicated by the growth of colonies co-expressing GBD+spc105CC, GAD+Kre28FL, and GBD+spc105CC,GAD+kre28201 in synthetic dextrose plates lacking histidine. Interestingly, we did not see any interactions between Kre28FL and spc105CC+RWD (figure 2a). This may be because of the misfolding of spc105CC+RWD fusion with the GAL4-binding domain (GDB_C1). It is also possible that the RWD domain interferes with the interaction of CC and Kre28, pointing to a regulatory mechanism. To dissect the interaction between spc105CC and kre28201 more thoroughly, we used smaller fragments (1–126 and 1–80) for our yeast two-hybrid assay with spc105CC. We did not notice any interaction using these combinations (figure 2a). Furthermore, we saw a significant contrast in colony growth between the combinations of spc105CC+Kre28FL and that of spc105CC+kre28201, which denotes a change in the strength of interaction with spc105CC. In conclusion, our yeast two-hybrid assay data indicated that Spc105CC binding domain lies within Kre28212–201.

Next, we mapped the Kre28 interacting domain of Spc105 in vivo. We performed domain mapping experiments where we truncated the mid strand domain of Spc105 (amino acid 313–708) at different residues based on predicted secondary structure (figure 2b). We constructed versions of GFP-labelled Spc105 with different truncations in the mid strand domain (Δ313–455 harbouring only the unstructured region, Δ313–507 containing unstructured region and a small helical domain, Δ313–638 that contains unstructured region and an alpha helix-rich domain of CC and Δ313–709 that
encompasses the entire mid strand domain, figure 2b top right) and transformed them in a heterozygous diploid strain (AJY3278, SPC105/Δ::NAT). We examined the localization of these mutants by microscopy. First, KT localization of Δ313–455 and Δ313–507 displayed no discernable difference in localization compared to wild-type (figure 2b, right). Our nuclear localization signal (NLS) analyses indicated residues of 337–345 (SSNKRRKLD, score 9.0) and/or that of 599N-625L (score 6.9) contain NLSs. However, previously we have shown that the mutation of 340-KRRK-343 to alanine residues
does not affect the KT localization of the mutant [26]. Therefore, this region of 313–507 is not essential for the KT localization of Spc105. Truncation of 313–638 or 313–709 completely abrogated KT localization of Spc105. According to our analysis, spc105Δ313–625 may harbour an NLS, and deletion of this signal may have abrogated KT localization of the mutants expressing spc105Δ313–638 or spc105Δ313–709. We introduced SV40-NLS (NLSV40) at the N-terminus of spc105Δ313–638::GFP to check if nuclear localization of this mutant rescues its loading at the KTs (figure 2b, bottom left). However, we did not observe any KT-specific localization.

Subsequently, we wanted to check whether these truncation mutants can support cell viability. To address this, we induced sporulation/meiosis in heterozygous diploid strains expressing these truncated molecules of spc105 (Δ313–455 or Δ313–507). We observed that they were able to complement the deletion of endogenous SPC105 (spc105Δ). We can conclude that the domain of 313–507 is not essential for any activity of Spc105 that contributes to cell viability. On the contrary, Spc105 mutants with either 313–638 or 313–709 truncated could not rescue the viability of spc105Δ. Even fusion of SV40-NLS (NLSV40) at the N-terminus of spc105Δ313–638 did not rescue its ability to support the cell viability in the absence of wild-type Spc105 (figure 2b and bottom left). This dataset reveals that the proper localization of Spc105 at the KT is essential for its proper function. They also infer that the lack of nuclear localization cannot explain the lethality of the (spc105Δ313–638) mutant.

We confirmed these observations using the plasmid-shuffle assay (data not shown). Therefore, we hypothesized that the domain of Spc105 housed within amino acid 507–638 directly interacts with Kre28. Deletion of this domain abrogates the interaction resulting in delocalization of Spc105 from the KTs.

To biochemically confirm the results of the 2-hybrid and localization experiments, we immunoprecipitated GFP-labelled versions of Spc105 from strains expressing either Spc105Δ555::GFP (FL) or spc105Δ313–638::GFP (Δ313–638) and examined if both molecules interact with Kre28 (figure 2c). Immunoprecipitation followed by immunoblot analysis demonstrated that even though Spc105Δ555::GFP binds Kre28-5xFlag, the mutant of spc105Δ313–638::GFP is unable to do so, which indicates that a domain harboured within Spc105Δ507–638 is essential for its interaction with Kre28 and subsequently its recruitment at the KTs.

2.3. Localization of truncation mutants of Kre28 and their ability to support cell viability

Our yeast two-hybrid assay (figure 2a) indicated that the Spc105 interacting domain of Kre28 lies within amino acid 127–201 of Kre28. The predicted secondary structure of Kre28FL showed that the aforesaid region of Kre28 is helix rich and structured (figure 3a, http://bioinf.cs.ucl.ac.uk/psipred, also see electronic supplementary material, figure S3A). To check which domain of Kre28 is essential for its loading at theKT and interaction with Spc105, we created yeast strains that express GFP-fused Kre28 fragments from the ADH1 promoter. We examined their localization in a diploid yeast strain where one genomic copy of KRE28 is deleted, and the other allele is tagged with mCherry at its C-terminus (figure 3b). As expected, GFP-Kre28FL localizes at the KT. On the contrary, we could not detect the localization of GFP-kre28Δ313–182 or fragments with larger truncations at the KTs. When over-expressed from pRADH1, GFP-fused versions of Kre28FL and its truncations revealed a high cytoplasmic GFP signal (figure 3b). Therefore, we performed similar experiments with the SV40-NLS at the C-terminus of GFP-kre28Δ313–182. Even in this case, we did not see any KT localization (data not shown). It should be noted that the GFP tagged Kre28FL and its truncations were expressed at similar levels (figure 3c).

Does Kre28 delocalization affect the cell viability when yeast cells express the mutants in the absence of wild-type Kre28? To test this, we sporulated these diploid strains and isolated haploid spores. We observed that the segre- gants over-expressing truncated versions of kre28 rescued the deletion of endogenous KRE28 (kre28Δ). However, the segre- gants expressing truncated kre28 molecules from their native promoter (KRE28pr) could not complement genenic KRE28 deletion (figure 3d). These data indicate that full-length Kre28 is essential for binding with Spc105 and its interaction with the Mtw1 complex. However, truncated kre28 mutants with defective KT localization were able to sustain cell viability when over-expressed. We backcrossed viable spores with the parent strain (YEF473) to avoid background mutations. The segre- gants isolated from those crosses were subjected to further experiments.
2.4. Truncation of Kre28 significantly reduces the recruitment of Spc105 at the kinetochore

We observed slower colony growth among the segregants expressing only the kre28 truncation mutants (figures 3d and 5b). The slow growth suggested that these mutants have a high propensity of chromosome missegregation, and this will affect their viability. Among these mutants, we chose kre28Δ127–182 for further analysis because this is the smallest truncation that has a significant defect in KT localization. To check whether kre28Δ127–182 affects the biorientation of sister KTs, we tagged Ndc80 and Spc105 individually with mCherry and studied KT biorientation in kre28 truncation mutant (figure 4a,b). We did not find any noticeable defects in the bipolar attachment of metaphase KTs in this mutant (electronic supplementary material, figure S2F). However, we observed a significant decrease (approx. 61%) in Spc105 recruitment in the truncation mutants (figure 4b) with a much smaller change (approx. 24%) in Ndc80 localization (figure 4a).

To further characterize the correlation between Kre28 and Spc105 recruitment to the KT, we varied the amount of either Kre28 or Spc105 per the KTs by exploiting variable

Figure 3. Truncation of Kre28 interferes with its localization to the kinetochore. (a) Secondary structure prediction for kre28127–201 (http://bioinf.cs.ucl.ac.uk/psipred). (b) Representative images of GFP fusions of Kre28FL and its truncated versions (kre28Δ127–181 or kre28Δ127–201) exogenously expressed by the ADH1 promoter. (c) Western blot assay with anti-GFP antibody on the lysates of the strains expressing Kre28FL or its truncated version from ADH1 promoter (ADH1pr, over-expression) or its native promoter (KRE28pr, expression from LEU2 locus). Image of Ponceau S stained blot is shown as loading control. Molecular weight of GFP-Kre28FL: 73.67 kDa, GFP-Kre28Δ127–182: 67.33 kDa, GFP-Kre28Δ1–201: 50.6 kDa, GFP-Kre28Δ1–127: 59.36 kDa. (d) Images of tetrad dissection plates for the heterozygous diploid strains. Genotypes are indicated above each photograph. kre28Δ spores expressing kre28 truncations are marked with red circles. The plate images on the left were taken after replica plating. Hence, the segregant colonies in those images look larger than the colonies on the right panel.
Figure 4. Kinetochores with truncated kre28 mutants form biorientation despite impaired Spc105 recruitment. (a) Left: representative micrographs of GFP-Kre28 (full length and truncation) and Ndc80-mCherry are shown, scale bar approximately 3.2 µm. Right: scatter plot of Ndc80-mCherry intensities (mean ± s.e.m) is shown for strains with Kre28FL and kre28Δ127–182. Unpaired t-test revealed p < 0.0001, indicated at the top. (b) Left: representative micrographs of GFP-Kre28 (full length and truncation) and Spc105-mCherry are shown, scale bar approximately 3.2 µm. Right: scatter plot of Spc105-mCherry intensities (mean ± s.e.m) is shown. According to unpaired t-test p < 0.0001, indicated at the top. (c) (i) Left: representative micrographs depict Spc105-GFP fluorescence from kinetochore cluster containing different amount of Kre28-mCherry, scale bar approximately 2.1 µm. Right: scatter plot where each grey circle represents the binned average number of Spc105-GFP molecules plotted against the average number of Kre28-mCherry molecules per bioriented kinetochore. Line in the plot indicates nonlinear regression, \( R^2 = 0.9774 \). (ii) Left: representative micrographs show Kre28-mCherry fluorescence from kinetochore cluster containing different amount of Spc105-GFP, scale bar approximately 2.1 µm. Right: scatter plot where each grey circle represents the binned average number of Kre28-mCherry molecules plotted against the average number of Spc105-GFP molecules per bioriented kinetochore. Line in the plot indicates nonlinear regression. \( R^2 = 0.9751 \). (iii) Left: representative micrographs depict Ndc80-GFP fluorescence from kinetochore cluster containing different amount of Kre28-mCherry, scale bar approximately 2.1 µm. Right: scatter plot where each grey circle represents the binned average number of Ndc80-GFP molecules plotted against the average number of Kre28-mCherry molecules per bioriented kinetochore. Line in the plot indicates nonlinear regression. \( R^2 = 0.9671 \). (d) Immunoblot assay with anti-GFP and anti-DsRed antibodies following RFP-trap assay on the cell lysates, flow through and elutes of indicated strains. Ponceau S staining of the membrane is shown as the loading control. Molecular weight of Spc105-mCherry, GFP-Kre28 and GFP-kre28Δ127–182 are approximately 132 kDa, approximately 74 kDa and 67 kDa, respectively. Normalized intensities of Spc105-mCherry and GFP-Kre28 for input and IP samples, which was calculated by ImageJ are depicted below (see Material and methods).
expression of the respective protein using the galactose-induced GALL promoter and quantified the amount of Spc105 and Kre28, respectively, per bioriented KT (figure 4c(i), (iii)). This quantification revealed that the amounts of KT localized Kre28 and Spc105 are mutually correlated. As the number of molecules of either protein per kinetochore increases, so does the number of molecules for the other. As expected, both numbers saturate at approximately 8 molecules per KT, which is close to the maximal number of Ndc80 complex molecules per yeast KT [27]. Given that Spc105 can localize to KTs even in strains over-expressing Kre28 fragments, these results strongly suggest that Kre28 positively contributes to Spc105 interactions with the Mtw1 complex. We also found that the number of Ndc80 molecules per KT was slightly lower in cells with KTs containing small numbers of Kre28 molecules (figure 4c(iii)). This result is consistent with our prior work, which showed that a reduction in Spc105 molecules per KT similarly lowers the number of Ndc80 molecules per KT [27].
Finally, we performed the immunoblot assay following RFP-trap experiments to assess the severity of impairment in the interaction between kre28127–182Δ and Spc105. We observed that the co-precipitation of kre28127–182Δ with Spc105-mCherry was significantly reduced (approx. 40%; figure 4d; see the figure legend and Material and methods section for details of band intensity calculation and normalization). Moreover, we detected the protein level of Spc105 was reduced in the presence of kre28127–182Δ (approx. 69%, figure 4d, panel of anti-DsRed blot). This set of observations imply that the coiled domain Kre28 containing amino acid residue 127–182 plays a significant role in the binding of Kre28 and Spc105. They also suggest that Kre28 plays a role in maintaining the stability of Spc105 protein.

2.5. Effect of Kre28 truncation on spindle assembly checkpoint and error correction pathway

Spc105 contains short linear interaction motifs known as MELT motifs that, when phosphorylated by the Mps1 kinase, serve as the scaffold for checkpoint components [17,18]. Moreover, the evolutionarily conserved RSSF motif present in the N-terminus of Spc105 acts as the primary binding motif of protein phosphatase I (PPI) that dephosphorylates the MELT repeats to silence the SAC [20]. Therefore, Spc105 delocalization may affect SAC activation and silencing.

We first studied SAC signalling in cells over-expressing kre284127–182Δ by treating cell cultures with the MT poison nocodazole and performing flow cytometry to quantify cellular DNA content. The flow cytometry revealed that SAC strength was not discernably different in strains expressing Kre28FL and kre28127–182Δ, as indicated by the arrest of the cell population with 2 N DNA content (figure 5a). However, we have previously shown that this assay cannot detect smaller defects in SAC and that of the error correction [6]. To examine the efficacy to detect smaller defects in SAC and that of the error correction process in kre28 mutants, we subjected them to a low dose of another MT poison, benomyl. At its dosage used in this assay, benomyl destabilizes MTs and forces yeast cells to rely on a combination of effective error correction and SAC signalling to ensure chromosome biorientation and accurate chromosome segregation [6]. The strains that express kre28 with larger truncations (Δ127–201, Δ1–201, and Δ1–126) demonstrated growth defects even in non-selective growth media (YPD). We also performed the benomyl-sensitivity assay using a strain where SV40-NLS is fused to Kre28127–182Δ and Spc105-mCherry which results in synthetic lethality [14]. Most strikingly, we observed that the stoichiometry of Kre28 to Spc105 is 1:1, whereas it was previously thought to be 2:1 [12].

Results of the yeast two-hybrid assays involving Kre28Δ1–201 and Spc105455–709Δ were consistent with previously published data from Yanagida lab [15,25]. However, it was unclear why the yeast two-hybrid assay did not work between Kre28Δ1–201 and spc105455–917Δ (spc105ΔC-RWD). It may be the case that CC + RWD (Spc105455–917) fusion with GAL4-binding domain (GBD_C1) did not fold in a way that they can interact with Kre28. On the other hand, it is also possible that RWD interfering with the binding of CC and Kre28 has an unknown physiological significance. Although we obtained yeast two-hybrid interaction between kre28Δ1–201 and spc105455–709Δ, we did not detect any KT localization of kre28Δ1–201. It was also not sufficient to support cell viability in the absence of wild-type Kre28. Much to our surprise, even after taking the data of Herzog lab into consideration, when we maintained the two binding sites of Spc105 (kre28Δ126–169Δ and kre28Δ229–259Δ) in our kre28 cassette (kre28Δ112Δ) [14], we did not see any KT localization. After considering these data, we can conclude that full-length Kre28 is essential for its localization and for full KT recruitment of Spc105.

Despite being an essential and conserved component of the KT, the structure of Kre28 remains unknown, and the absence of a structure prevents a clear understanding of the implications of our data. Therefore, to provide a structural context to our interaction-mapping and KT localization experiments, we used an implementation of Alphafold2 to predict structures of protein complexes using Google
### Table 1. Summary of observations of the experiments involving kre28 truncations.

| kre28 truncations | sufficiency for cell viability when expressed from \( KRE28 \) promoter | localization at the kinetochore when expressed from \( KRE28 \) promoter | interaction with Spc105 \( ^{455-917} \) | structured region within Spc105 |
|-------------------|-------------------------------------------------|-------------------------------------------------|---------------------------------|--------------------------------|
| FL                | yes                                             | yes                                             | yes                             | yes                            |
| \( \Delta 1-201 \) | no                                              | not done                                        | not done                        | not done                       |
| \( \Delta 127-182 \) (with and without NLS) | no                                              | not done                                        | not done                        | not done                       |
| \( \Delta 1-917 \) | no                                              | not done                                        | not done                        | not done                       |
| \( \Delta 1-320 \) (with and without NLS) | no                                              | not done                                        | not done                        | not done                       |

Note: SV, sensitivity to sensitivity to benomyl; V, viability when expressed from \( KRE28 \) promoter; MT, mating type; NLS, nuclear localization signal; FL, full length; C, truncation construct.

ColabFold (https://colab.research.google.com/github/sokrypton/ColabFold/blob/main/beta/AlphaFold2_advanced.ipynb [31]). Using ColabFold, we predicted the structure of a heterodimer of Kre28\(^{917} \) and the C-terminal region of Spc105\(^{455-917} \) (electronic supplementary material, figure S3B). The N-terminal boundary of Spc105 used in the structure prediction was chosen based on the start of the predicted model is also in agreement regarding the residues in Kre28 identified by a previous study to be critical in mediating its interaction with Spc105 [14]. In conclusion, the extensive interaction interface between the two proteins with involvement from residues over nearly the entire length of Kre28 predicted by the model can explain why we could not eliminate the interaction between Spc105 and Kre28 using extensive truncations in either protein.

Our data of Spc105 protein becoming destabilized in the \( knr28^{\Delta 127-182} \) mutant (figure 4l, panel of anti-DsRed blot) imply that impairment of Kre28-Spc105 interaction significantly affects Spc105 integrity of expression. This implication is in accordance with a previously reported study by Zhang et al. [32] where they depleted Zwint1 by RNAi and demonstrated that the cellular protein level of hSpc105 is depleted. A study in the fission yeast \( S. pombe \) revealed that KT proteins like Spc105 are surveilled by Bag102, the 26S proteasome, Ubc4, and the ubiquitin-ligases Ubx1 and San1 [33]. In that study, the authors suggested that cells employ this mechanism to maintain the homeostasis of nuclear components and genomic integrity. We came across another indirect evidence of protein quality regulation of Spc105 and Kre28 from Yong-Gonzalez and colleagues [34]. They showed that both of these proteins become sumoylated by Smc3-Smc6 complex and deleterious mutations in Smc5-Smc6 complex leads to chromosome loss [34]. Consistent with the above-mentioned studies, the Biggins lab also...
discovered Mub1/Ubr2 ubiquitin ligase complex to be a part of a quality control mechanism that monitors KT protein Dsn1 [35]. A similar mechanism probably controls Spc105 and/or Kre28 levels in S. cerevisiae.

Does Kre28 act as a chaperone to stabilize the recruitment of Spc105 at the KT? Some of the previous studies argue against this hypothesis. In human KT, Zwint1 is dispensable for the interaction between hSpc105 and Mis12 complex [15,16]. While performing ex-vivo KT assembly experiments, Biggins lab showed that Ipl1 phosphorylation of Dsn1, which triggers outer KT assembly, also recruits Kre28, which should be specific to mitotic cells [36]. Contrarily, we observed a similar level of Kre28 at the KTs at every stage in the cell cycle, including the G1-S phase (unbudded and small budded cells, data not shown), which implies that Kre28 loading at the KT takes place at the same time point as loading of Spc105. Combining our observations with those from Herzog lab, we can conclude that kre28Δ−182 contributes to the main interaction between Kre28 and Spc105 and kre28Δ−259 contributes to interaction with the Mtw1 complex. However, full-length Kre28 is essential for proper binding with Spc105, their mutual recruitment and their activity at the KTs. Kre28 may also become phosphorylated by Ipl1, which can trigger its association with Spc105 and subsequently their loading at the KTs.

Does Kre28 have a specific function in SAC and error correction or during meiosis? The results of the functional assays (figure 5) clearly show that the delocalization of Kre28 from the KT impairs the processes. However, all these phenotypes may be linked with the delocalization of Spc105. A similar conclusion has been noted for Zwint1 [22,32,37]. Our experiments, Biggins lab showed that Ipl1 phosphorylation of Dsn1, which triggers outer KT assembly, also recruits Kre28, which should be specific to mitotic cells [36].

Contrarily, we observed a similar level of Kre28 at the KTs at every stage in the cell cycle, including the G1-S phase (unbudded and small budded cells, data not shown), which implies that Kre28 loading at the KT takes place at the same time point as loading of Spc105. Combining our observations with those from Herzog lab, we can conclude that kre28Δ−182 contributes to the main interaction between Kre28 and Spc105 and kre28Δ−259 contributes to interaction with the Mtw1 complex. However, full-length Kre28 is essential for proper binding with Spc105, their mutual recruitment and their activity at the KTs. Kre28 may also become phosphorylated by Ipl1, which can trigger its association with Spc105 and subsequently their loading at the KTs.

4. Material and methods

4.1. Plasmid and strain construction

The strains and plasmids used in this study are documented in tables 2 and 3, respectively. Yeast strains containing multiple genetic modifications were constructed by standard yeast genetic techniques. GFP (S65T) and mCherry fusion of proteins were used to localize KTs by fluorescence microscopy. The C-terminal tags like GFP, mCherry, 5xFlag and gene deletion cassettes like spc105Δ::NAT and kre28Δ::NAT were introduced at the endogenous locus through homologous recombination of PCR amplicons [38]. A 7-amino acid linker (sequence: ‘RFPGLIN’) bridges the tags (GFP, mCherry, or 5xFlag) from the C-termini of the tagged proteins. Earlier, we observed that the intensity of mCherry-tagged KT proteins varies significantly from one transformant to another for the same strain, due to inherent variability of the mCherry brightness. Therefore, to construct all the FRET strains with Ndc80, Stu2, Nsl1, Kre28 and Ask1-mCherry, we crossed a specific mCherry strain with haploid strains of all GFP-fused Spc105 alleles and sporulated the heterozygous diploids to obtain the desired segregants.

To construct the yeast strains with internally tagged Spc105 mentioned in electronic supplementary material, figure S1, and the strains with truncated Spc105 mentioned in figure 3, first we used BstElI digest of pRS305 chimera or Stul digest of pRS306 chimera of Spc105-GFP fusion alleles to transform AJY3278 (SPC105Δ::NAT) that was later sporulated to obtain the haploid segregants expressing only GFP fusion copy of Spc105. To construct some of these strains, first we deleted the genomic copy of SPC105 in a strain already supplemented with the wild-type SPC105 gene expressed from centromeric plasmid yCP50 (URA3). Then, the pRS315 chimera containing Spc105-GFP alleles were transformed in that strain. After that, the strains expressing only Spc105-GFP alleles were generated by negative selection for yCP50 on 5-FOA plates.

The construction of chimeras for over-expression of Kre28F1 or truncations was achieved by cloning N-terminal GFP tagged fusions of Kre28 in a centromeric plasmid pRS414, where KRE28 ORF is flanked by ADH1 (alcohol dehydrogenase 1) promoter and GFP ORF at the upstream and CYC (Cytochrome C) terminator at the downstream (obtained from Dr. Maria Duncan’s lab, department of Cell and Developmental Biology, University of Michigan). Kre28 fragments were cloned in BamHI- PstI sites. SV40- NLS was cloned within PstI- Sall. For wild-type expression, Kre28F1 and its truncations were cloned in BamHI- PstI of a pRS305 plasmid. They were expressed as N-terminal GFP tagged fusions by its own promoter and terminator. BstElI digests of these chimeras are transformed in AJY3298 (kre28Δ::NAT/KRE28-mCherry-Hyg) to check for their localization. To create diploid zygotes, two strains of a and α mating types are mixed with each other and spotted on YPD plate which was incubated for 3–4 h at 30°C. To induce sporulation, diploid yeast cells were grown in YPD overnight to stationary phase. Next day cells were pelleted down and resuspended with starvation media (0.1% yeast extract, 1% potassium acetate, 0.025% dextrose) and incubated 4–5 days at RT.

4.2. Yeast two-hybrid assay

We performed yeast two-hybrid experiments by co-transforming both of prey (pGAD_C1) and bait (pGBD_C1) chimera in strain AJY3802 (PFP69A) [39]. Then, we streaked two of the transformants for each prey-bait pair on synthetic dextrose plates of histidine dropout (-HIS) and dropout of histidine and adenine (-HIS-ADE). Plates were incubated in 32°C for at least 3 days.

4.3. Benomyl-sensitivity assay

This experiment was performed as described previously [6,26]. Starting from 0.1 OD600 of log-phase cultures, we prepared 10-fold serial dilutions and frogged or spotted them on YPD and YPD containing 20 µg ml−1 and 30 µg ml−1 benomyl. At least two biological replicates were used, and spotting was repeated twice for each set of experiments. The plates were incubated at 32°C and pictures were taken
Table 2. Strains used in this study.

| strain (AJY#) | genotype | background |
|---------------|----------|------------|
| 2987          | SPC105::GFP-, KANMX6, NDC80::mCherry- KANMX6 | YEF473     |
| 3711          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), NDC80::mCherry- KANMX6 | YEF473     |
| 3712          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), NDC80::mCherry- KANMX6 | YEF473     |
| 3713          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (LEU2), NDC80::mCherry- KANMX6 | YEF473     |
| 3714          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (LEU2), NDC80::mCherry- KANMX6 | YEF473     |
| 3795          | spc105::GFP, SPC105 (HIS3), NDC80::mCherry-KANMX6 | YEF473     |
| 3796          | spc105::GFP, SPC105 (HIS3), NDC80::mCherry- KANMX6 | YEF473     |
| 3435          | spc105::GFP::NAT, Spc105<sup>222</sup>::GFP (LEU2), NDC80::mCherry- KANMX6 | YEF473     |
| 3513          | spc105::GFP::NAT, Spc105<sup>222</sup>::GFP (LEU2), STU2-mCherry-NAT | YEF473     |
| 3639          | spc105::GFP::NAT, SPC105 (HIS3), STU2-mCherry-NAT | BIY4743    |
| 3641          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (LEU2), STU2-mCherry-NAT | YEF473     |
| 3709          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), STU2-mCherry-NAT | YEF473     |
| 3710          | Spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), STU2-mCherry-NAT | YEF473     |
| 3735          | SPC105::GFP- KANMX6, STU2-mCherry-NAT | YEF473     |
| 3736          | SPC105::GFP-KANMX6, STU2-mCherry-NAT | YEF473     |
| 3212          | spc105::GFP::NAT, SPC105 (HIS3), Spc105<sup>222</sup>::mCherry (CEN, LEU2) | BYI4743    |
| 3215          | spc105::GFP::NAT, SPC105 (HIS3), Spc105<sup>222</sup>::mCherry (CEN, LEU2) | BYI4743    |
| 3217          | spc105::GFP::NAT, Spc105<sup>222</sup>::GFP (URA3), Spc105<sup>222</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3218          | spc105::GFP::NAT, Spc105<sup>222</sup>::GFP (URA3), Spc105<sup>222</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3219          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (URA3), Spc105<sup>222</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3220          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (URA3), Spc105<sup>222</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3801          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (URA3), Spc105<sup>455</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3799          | SPC105::GFP-KANMX6, Spc105<sup>709</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3800          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (URA3), Spc105<sup>709</sup>::mCherry (CEN, LEU2) | YEF473     |
| 3658          | spc105::GFP::NAT, SPC105 (HIS3), GAL1::pr-mCherry-NUF2 (KANMX6) | BYI4743    |
| 3659          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (LEU2), GAL1::pr-mCherry-NUF2 (KANMX6) | YEF473     |
| 3660          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), GAL1::pr-mCherry-NUF2 (KANMX6) | YEF473     |
| 4171          | SPC105::GFP-KANMX6, NSL1::mCherry-TRP1 | YEF473     |
| 4172          | SPC105::GFP-KANMX6, NSL1::mCherry-TRP1 | YEF473     |
| 4175          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), NSL1::mCherry-TRP1 | YEF473     |
| 4176          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (LEU2), NSL1::mCherry-TRP1 | YEF473     |
| 3760          | spc105::GFP::NAT, Spc105<sup>222</sup>::GFP (LEU2), ASK1::mCherry-NAT | YEF473     |
| 3794          | spc105::GFP::NAT, SPC105 (HIS3), ASK1::mCherry-NAT | YEF473     |
| 3107          | GFP(S65T)-NDC80, KRE28::mCherry-HYG | YEF473     |
| 2991          | NDC80::GFP-KANMX6, KRE28::mCherry-HYG | YEF473     |
| 2993          | SPC24::GFP-KANMX6, KRE28::mCherry-HYG | YEF473     |
| 3160          | KRE28::GFP-KANMX6/KRE28::mCherry-HYG | YEF473     |
| 3206          | KRE28::GFP-KANMX6/KRE28::mCherry-HYG | YEF473     |
| 2986          | SPC105::GFP-KANMX6, KRE28::mCherry-HYG | YEF473     |
| 3221          | spc105::GFP::NAT, Spc105<sup>222</sup>::GFP (CEN, LEU2), KRE28::mCherry-HYG | YEF473     |
| 2977          | spc105::GFP::NAT, Spc105<sup>709</sup>::GFP (CEN, LEU2), KRE28::mCherry-Hyg | YEF473     |
| 2982          | spc105::GFP::NAT, Spc105<sup>455</sup>::GFP (CEN, LEU2), KRE28::mCherry-HYG | YEF473     |
| 3802          | trp1-901, leu2-3,112 ura3-52, his3-200 gal4-4a, gal80-4A, GAL2-ADE2 LYS2::GAL1-HIS3, met2::GAL7-lacZ | —          |
| 3278          | spc105::GFP::NAT/SPC105 | YEF473     |
| 5022          | spc105::GFP::NAT/SPC105, Spc105<sup>455</sup>::GFP (CEN, LEU2) | YEF473     |
| 5023          | spc105::GFP::NAT/SPC105, spc105<sup>455</sup>::GFP (CEN, LEU2) | YEF473     |

(Continued.)
Table 2. (Continued.)

| strain (AJY#) | genotype | background |
|---------------|----------|------------|
| 5024          | spc105Δ::NAT/SPC105, spc105Δ::GFP (CEN,LEU2) | YEF473 |
| 5025          | spc105Δ::NAT/SPC105, spc105Δ::GFP (CEN,LEU2) | YEF473 |
| 5026          | spc105Δ::NAT/SPC105, spc105Δ::GFP (CEN,LEU2) | YEF473 |
| 6275          | spc105Δ::NAT/SPC105, NLS+spc105Δ::GFP (CEN,LEU2) | YEF473 |
| 6273          | spc105Δ::NAT1, Spc105Δ::GFP (CEN,LEU2), KRE28-5xFlag-KANMX6 | YEF473 |
| 6274          | spc105Δ::GFP (CEN,LEU2), KRE28-5xFlag-KANMX6 | YEF473 |
| 3298          | kre28Δ::NAT/KRE28-mCherry-HYG | YEF473 |
| 3386          | KRE28-mCherry-HYG/kre28Δ::NAT, ADH1pr-GFP-kre28-Δ127-183 (CEN, TRP1) | YEF473 |
| 3387          | KRE28-mCherry-HYG/kre28Δ::NAT, ADH1pr-GFP-kre28-Δ127-201 (CEN, TRP1) | YEF473 |
| 3390          | KRE28-mCherry-HYG/kre28Δ::NAT, ADH1pr-GFP-KRE28Δ1 (CEN, TRP1) | YEF473 |
| 3391          | kre28Δ::NAT, ADH1pr-GFP-KRE28Δ1 (CEN, TRP1) | YEF473 |
| 3407          | kre28Δ::NAT, ADH1pr-GFP-KRE28Δ1 (CEN, TRP1) | YEF473 |
| 3408          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ127-183 (CEN, TRP1) | YEF473 |
| 3409          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ127-201 (CEN, TRP1) | YEF473 |
| 3410          | KRE28-mCh-Hyg, ADH1pr-GFP-kre28-Δ127-183 (CEN, TRP1) | YEF473 |
| 3411          | KRE28-mCh-Hyg, ADH1pr-GFP-kre28-Δ127-201 (CEN, TRP1) | YEF473 |
| 3471          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ11-201 (CEN, TRP1) | YEF473 |
| 3472          | KRE28-mCherry-HYG, ADH1pr-GFP-kre28-Δ11-201 (CEN, TRP1) | YEF473 |
| 3473          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ11-127 (CEN, TRP1) | YEF473 |
| 3474          | KRE28-mCherry-HYG, ADH1pr-GFP-kre28-Δ11-127 (CEN, TRP1) | YEF473 |
| 4951          | mad2Δ::TRP1 | YEF473 |
| 4786          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ128-183+Δ13 (CEN, TRP1) | YEF473 |
| 4787          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ128-183+Δ13 (CEN, TRP1) | YEF473 |
| 4788          | KRE28-mCherry-HYG, ADH1pr-GFP-kre28-Δ128-183+Δ13 (CEN, TRP1) | YEF473 |
| 4660          | sgo1Δ::Kan | YEF473 |
| 3477          | KRE28-mCherry-HYG/ kre28Δ::NAT, KRE28pr-GFP-Kre28Δ1-Tr(KRE28) (LEU2) | YEF473 |
| 3494          | KRE28-mCherry-HYG/kre28Δ::NAT, KRE28pr-GFP-kre28-Δ127-183-Tr(KRE28) (LEU2) | YEF473 |
| 3495          | KRE28-mCherry-HYG/kre28Δ::NAT, KRE28pr-GFP-kre28-Δ11-201-Tr(KRE28) (LEU2) | YEF473 |
| 3421          | kre28Δ::NAT, ADH1pr-GFP-KRE28Δ1, NDC80-mCherry-KANMX6 | YEF473 |
| 3423          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ127-183, NDC80-mCherry-KANMX6 | YEF473 |
| 3483          | kre28Δ::NAT, ADH1pr-GFP-KRE28Δ1 (CEN, TRP1), SPC105-mCherry-HIS3 | YEF473 |
| 3484          | kre28Δ::NAT, ADH1pr-GFP-kre28-Δ127-183 (CEN, TRP1), SPC105-mCherry-HIS3 | YEF473 |
| 3101          | NAT- GALLpr-SPC105-GFP-KAN, KRE28-mCherry-HYG | YEF473 |
| 3201          | NAT- GALLpr-KRE28-mCherry-HYG, SPC105-GFP-KAN | YEF473 |
| 3202          | NAT- GALLpr-KRE28-mCherry-HYG, NDC80-GFP-KAN | YEF473 |

after 2 (YPD) to 3 (YPD + Benomyl) days. For space limitations, we showed only YPD + Benomyl plates. For all spotting assays with benomyl, we used YEF473 or strain expressing Spc105Δ::GFP as wide-type (positive control) and mad2Δ or sgo1Δ as negative controls. Before spotting, we grew the strains expressing truncated kre28 or Kre28Δ1 control in synthetic dextrose media (Sd-Trp) without Tryptophan to culture only cells carrying the pRS414 chimera. As shown in the plate images, we also used strains expressing Kre28Δ1-mCherry along with truncated kre28 as controls which did not show any discernable difference in growth, compared to wild-type.

4.4. Microscopy and image acquisition and analyses

A Nikon Ti-E inverted microscope with a 1.4 NA, 100X, oil-immersion objective was used for experiments mentioned in the paper [40]. A ten-plane Z-stack was acquired (200 nm separation between adjacent planes). To measure Ndc80 and Spc105-mCherry, an extra 1.5x opto-var lens was used. We measured total fluorescence of KT clusters (16 KTs in metaphase) by integrating the intensities over a 6 × 6 region centred on the maximum intensity pixel. We used median intensity of pixels immediately surrounding the 6 × 6 area to correct for background.
fluorescence. The calculation was performed using semi-automated MATLAB programs as described earlier [41]. FRET, high-resolution colocalization, fluorescence distribution analyses and analyses of the images were performed as previously described [7,8,27,40,42]. While measuring proximity ratio, we considered any value below 0.10 as no FRET (mean of the data marked as black), range between 0.10 and 0.3 as medium to low FRET (average of the data marked as red) and any values above 0.3 as high FRET (mean of the data marked as red).

### 4.5. Titration of Kre28 and Spc105 proteins levels and quantification of Kre28, Spc105 and Ndc80 intensities

We grew the strains with prGALL-SPC105 or prGALL-KRE28 in the presence of raffinose (2%). On the day of the assay, we supplemented the media with variable amounts of galactose as discussed previously (2%, 1.5%, 0.5%, 0.2%, 0.1% and 0.05%) [27]. We determined the number of Kre28, Spc105 and Ndc80 from their intensities as reported previously.
[27]. We first deduced the fluorescence intensities of Kre28-mCherry, Ndc80-GFP and Spc105-GFP from bioriented KTs. We used AJY939 (Ndc80-GFP, Spc25-mCherry) as a reference to obtain the intensities for known number of Ndc80 and Spc25 molecules at the bioriented KTs. AJY939 was cultured under same imaging conditions as the experimental strains, and the calibration data were acquired throughout the duration of this study. This calibration accounted for alteration in the microscope and imaging technique set-up over time. We used the values of Ndc80-GFP and Spc25-mcherry to determine the number of molecules of Spc105-GFP and Kre28-mCherry that were loaded in the bioriented KTs.

4.6. Preparation of cell lysates and western blot assay

To prepare cell lysates, log-phase cells (OD₆₀₀ 2.0) were pelleted, resuspended in sample buffer (2% SDS, 1% 2-mercaptoethanol), boiled and lysed by glass-bead mechanical disruption [7]. The lysates were collected after centrifugation. After separating the proteins by 10% SDS-PAGE, samples were transferred to nitrocellulose or PVDF blocked with 5% skimmed milk in 1x phosphate-buffered saline-Tween (PBS-T, 137 mM NaCl, 10 mM phosphate, 2.7 mM KCl, 0.05% Tween 20). We probed the blot with primary and fluorescent secondary antibodies. Mouse α-GFP antibody was from Santa Cruz Biotechnology (1 : 2000, GFP(B-2):sc-9996). Peroxidase conjugated α-mouse IgG (1 : 5000; Sigma, A-4416) treated with ECL (Thermo Scientific) was used to develop the western blot.

4.7. GFP-trap and RFP-trap assay to pull down Spc105 and immunoblot assay

We used AJY6273 and AJY6274 for GFP-trap experiments. As mentioned in the strain list, AJY6273 expresses Spc105ΔKIRKL from a centromeric plasmid (pRS315) and genomic SPC105 allele is deleted. The truncation of 313–638 affects the cell viability; hence in AJY6274, the genomic SPC105 allele was intact and spc105ΔKIRKLΔα-Flag was expressed from a centromeric plasmid (pRS315). We grew both the strains in synthetic dextrose media devoid of leucine (SD-LEU) to maintain the centromeric plasmid before harvesting them for cell lysis. For RFP-trap assays, we used AJY3483 and AJY3484 (see strain list for detailed information on their genetics). We grew the strains in SD-TRP (synthetic dextrose devoid of tryptophan) media till late log-phase before harvesting the cells. We lysed the cells by glass-beads in the presence of buffer H 0.15 (25 mM HEPES of pH 8.0, 2.0 mM MgCl₂, 0.1 mM EDTA of pH 8.0, 0.5 mM EGTA-KOH of pH 8.0, 15% glycerol, 0.1% IGEPAL-CA-630 and 150 mM KCl), supplemented with 0.2 mM PMSE protease inhibitor cocktail and phosphatase inhibitor cocktails [43]. We isolated clear lysates of the strains and from there, we incubated equal amount of lysates with pre-equilibrated beads of GFP-trap (Chromotek, gta-20) or RFP-Trap (Chromotek, rta-20) overnight. Next day, we washed the beads with post IP wash buffer (composition as mentioned above) with and without 2 mM Dithiothreitol, before boiling them in the presence of 1xSDS loading buffer.

After subjecting the proteins through SDS-PAGE, we transferred them to nitrocellulose membrane which we blocked with 5% skimmed milk in 1x phosphate-buffered saline-Tween (PBS-T, 137 mM NaCl, 10 mM phosphate, 2.7 mM KCl, 0.05% Tween 20). We probed the blot with Mouse anti-GFP (JL8, Living Colors, Takara, 1 : 3000) or Mouse anti-Ds-Red (Santa Cruz Biotechnology, sc-390909, 1 : 2000), Mouse α-Flag M2 (Sigma-Aldrich, 1 : 5000) and HRP conjugated secondary anti-Mouse (Sigma-Aldrich, 1 : 10,000). ECL (Thermo Scientific) treatment was used to develop the western blot. We exposed the anti-Ds-Red blots to X-ray films. Anti-GFP and anti-Flag blots were imaged by C600 imager (Azure Biosystems). We calculated the band intensities by ImageJ in RFP-TRAP assay. Then, we normalized both Spc105 and Kre28 intensities of the truncation mutants with those of the full-length proteins. After that, we normalized Kre28 IP intensities with that of normalized Kre28 input values. Following that, we divided the Kre28 IP intensities with Spc105 IP intensities. We mentioned those values as fold difference at the bottom of the immunoblot panels (figure 4d).

4.8. Flow cytometry

We performed flow cytometry as described previously [6,26]. For strains expressing Kre28-FL or truncated kre28, we started with overnight inoculums grown in Sd-Trp and shifted to YPD to grow till early to mid-log phase before supplementing the media with Nocodazole (final concentration 15 µg ml⁻¹) or DMSO control. We collected cell samples at 0, 1, 2, 3 h post-drug treatment and fixed them with 70% ethanol before storing them at 4°C overnight. Next day, after removing the Ethanol, treated the samples with bovine pancreatic RNase (Millipore Sigma, final concentration 170 ng ml⁻¹) at 37°C for 6 h overnight in RNase buffer (10 mM Tris pH 8.0, 15 mM NaCl). After that, we removed the RNase and resuspended the cells in 1X PBS. We treated the samples with propidium iodide (Millipore Sigma, final concentration 5 mg ml⁻¹ in PBS) for at least 1 h at room temperature before analysing them using the LSR Fortessa machine (BD Biosciences) in Biomedical research core facility, University of Michigan medical school. We analysed and organized the data using FlowJo software (FlowJo_V10.7.1_CL).

4.9. Nuclear localization signal mapping

We performed NLS mapping by pasting the amino acid sequences of Spc105 and Kne28 in the website of http://nls-mapper.iab.keio.ac.jp/cgi-bin/NLS_Mapper_form.cgi and keeping the cut-off score to 5.0 [44,45]. It scored SV40-NLS (TPPKKKRKVA, monopartite) as 15.5. We observed score of 9.0 (monopartite) for Spc105Δ337–345 (SSNKRRKLD) and 6.9 (bipartite) for Spc105Δ917 (NTLKREYKLNEEVEKVN–IRGKIRKL). We did not find any NLS for Kne28 without setting the cut-off score to 4.0. Kne28Δ207–234 and Kne28Δ286–317 displayed NLS scores of 4.2 and 4.0 (bipartite), respectively.

4.10. Prediction of coiled-coil domains in Kre28 and spc105ΔKIRKL

We predicted the CC of full-length Kne28 and spc105ΔKIRKL by inserting their amino acid sequences on the website of https://embnet.vital-it.ch/software/COILS_form.html [46]. We used MTIDK matrix for the prediction. We downloaded the output in postscript format and further configured by adobe illustrator.
4.11. Prediction of spc105551–917 - Kre28 interaction interface by Colabfold

We used Colabfold to predict the complex formed by interaction of Kre28 and spc105551–917 (https://colab.research.google.com/github/sokrypton/ColabFold/blob/main/alpha/AlphaFold2_advanced.ipynb, [31]). The following parameters were used for the structure prediction: msa_method: mmseqs2, pair_mode: unpaired, pair_cov = 50, pair_qid = 20, rank_by = pTMscore, num_models = 5, use_ptm = True, max_recurrents = 3, tol = 0, num_samples = 1, subsample_msa = True, num_relax = None. We processed the figures of the predicted structure by PyMOL 2.5 (https://pymol.org/2/), Schrödinger, LLC. The pLDDT confidence matrix was depicted by spectrum b mapping in PyMOL. Pair Alignment Error matrix was prepared by Colabfold.

4.12. Statistical analysis

We analysed the data and assembled the graphs by GraphPad Prism 8 software. We performed unpaired t-test (Mann–Whitney test) and one-way ANOVA analyses to check the statistical significances of the data. The p-values are mentioned on the top of the graph.

Data accessibility. This article has no additional data.

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