Supplementary Figure 1. She1 increases dwell time between individual steps of dynein. (a) Cartoon representation of the GFP-GST-dynein$_{1331}$-HALO-Qdot$^{525}$ used in the stepping assays. (b) Plots and fits of the cumulative survival probabilities of the N-terminal GFP and the C-terminal (motor) HALO tag-coupled Qdot$^{525}$. Note the large difference in photostability between the two fluorophores (see Methods; n ≥ 190 motors for each). (c) Histograms of dwell times (duration
between individual steps) for motor domain labeled (via Qdot\textsuperscript{525}) GST-dynein\textsubscript{331} in the absence or presence of She1, and with either 1 \( \mu \)M or 1 mM ATP, as indicated. The histograms were fit to a convolution of two exponential functions \([tk^2\exp(-kt)]\) with equal decay constants, \(k\), which reflects the number of steps taken per second\textsuperscript{1,2} (\(k \pm\) standard error of the fit is shown).
Supplementary Figure 2. *Cumulative probability functions used for determination of mean run length and dwell time values of dynein with and without She1 on control and subtilisin-treated microtubules.* Raw data (circles) and fits (dashed lines) are shown for run length (top) and dwell time (bottom) in the absence (green) and presence of She1 (red) on control (left) or subtilisin-treated microtubules (right; *n* ≥ 199 individual motors for each condition). Data were fit as previously described².
Supplementary Figure 3. Characterization of recombinant fragments used in the recruitment assays, and two-hybrid data. (a) Mean fluorescence intensity values (along with standard deviations) of microtubule-bound monomeric GFP-dynein<sub>331</sub> in the absence (magenta) or presence of ATP and vanadate (green). (b) Recombinant protein fragments used in the recruitment assays. With the exception of the dynein motor domain fragments, which were purified from yeast, all proteins were purified from E. coli (see Methods). (c) Cartoon representation
and reconstructed structural model of the $\text{SRS}_{\text{CC}}$-$\text{dynein}_{\text{MTBD}}$ fusion. Image was generated from a yeast model of the $\text{DYN1}$ MTBD (threaded into 3ERR$^3$) and 1SRY$^4$. (d) Mean fluorescence intensity values (along with standard deviations) of microtubule-bound GFP-$\text{SRS}_{\text{CC}}$-$\text{dynein}_{\text{MTBD}}$ (red) and GFP-$\text{dynein}_{\text{CC+MTBD}}$ (green; n ≥ 19 microtubules, and ≥ 151 µm of MT length for each condition) along with fits and resulting dissociation constants ($K_D$). (e) Two-hybrid assay demonstrating an interaction between $\text{dynein}_{\text{MTBD}}$ and $\text{She1}$ (see Methods). (f) Schematic representation of the experimental setup used for panel g. (g) Representative images depicting the inability of microtubule-bound $\text{She1}$ to recruit GFP-$\text{dynein}_{331^4\text{MTBD}}$ to microtubules. Images were acquired prior and subsequent to washing the chamber with motility buffer (see Methods; scale bar, 1 µm).
Supplementary Figure 4. **Comparison of various dynein MTBDs.** (a) Cartoon and homology models of the yeast dynein MTBD bound to α and β-tubulin in the high (grey) and low (tan) microtubule affinity states. The models were generated using one-to-one threading of the yeast DYN1 sequence into 3J1T (high affinity) and 3J1U (low affinity). CC1 and H1, which exhibit the largest differences between the two structures, are depicted as follows: CC1, red and pink, for high and low affinity states; H1, blue and cyan, for high and low affinity states, respectively. (b, left) Crystal structure of human dynein-2 (4RH7) docked onto microtubules (from 3J1T). (right) Homology model of the yeast MTBD (colored) – along with a short region of the CC (grey) – bound to α and β-tubulin in the high
microtubule affinity state. The residues are colored to reflect the degree of conservation between yeast and mouse primary sequence (see legend). (c) Mean fluorescence intensity values (along with standard deviations) of microtubule-bound GFP-dynein331 (green) and GFP-dynein331mMTBD (red; n ≥ 15 microtubules, and ≥ 68 µm of MT length for each condition) along with fits and resulting dissociation constants (K_D). Note the differences in apparent B_max values (4645 ± 763 A.U. for wild-type, and 2452 ± 517 A.U. for mMTBD; ± SE of fit) are likely a consequence of microtubule unbinding during the chamber washes (see Methods), and likely differences in microtubule dissociation rates between the two motor domains.
Supplementary Figure 5. **Cumulative probability functions and resulting mean values for run length and dwell time values of GST-dynein^{mMTBD} with and without She1.** (a and b) Raw data (circles) and fits (dashed lines) are shown for run length (a) and dwell time (b) in the absence (green) and presence of indicated concentrations of She1 (yellow and red; n ≥ 147 individual motors for each condition). Data were fit as previously described\(^2\). (c and d) Mean run length (c) and dwell time (d) for GST-dynein\(_{331}^{mMTBD}\) in the presence of the indicated concentration of She1 (error bars, standard error). Dashed line indicates the mean run length or dwell time for wild-type (WT) GST-dynein\(_{331}\) in the absence of She1.
Supplementary Figure 6. Dynein relocalization to astral microtubules upon She1 overexpression requires the dynein MTBD, but not Pac1. (a) Cartoon representation of the two possible models to account for dynein relocalization upon She1 overexpression. The model on the left depicts a mechanism whereby the entire plus end targeting complex (composed of Dyn1, Pac1, Bik1 and Bim1; note that dynactin is not an obligate component of this complex) is required for the relocalization. Given the dispensable nature for the MTBD in plus end targeting, this would indicate an MTBD-independent mechanism. The model on the right depicts a mechanism whereby dynein microtubule binding activity via...
the MTBD is required. (b and c) Representative images of \textit{GAL1p:SHE1} cells expressing mTurquoise2-Tub1 (b), or mRuby2-Tub1 (c), and either Dyn1\textsubscript{\textsc{AMtbd}}-3YFP (b) or Dyn1-3GFP (c), the latter of which is deleted for \textit{PAC1}. Cells were grown to mid-log phase in SD media supplemented with raffinose (uninduced; “- galactose”) or galactose plus raffinose (induced for 3.5 hours; “+ galactose”) and then mounted on agarose pads for confocal fluorescence microscopy. Foci were identified in two-color movies and scored accordingly (see Methods; blue arrows, plus end foci; blue arrowheads, cortical foci; red arrowhead, dynamic cytoplasmic foci not associated with microtubules or spindle poles). Note the accumulation of Dyn1 near the spindle poles in \textit{pac1\textDelta \textit{GAL1p:SHE1}} cells grown in galactose-containing media (the same was observed in \textit{GAL1p:SHE1 PAC1} cells; not shown). Movies reveal these spots exhibit dynamic movements in a manner that is consistent with them localizing to short astral microtubules, and not the spindle poles themselves. Note that our data support the MTBD-dependent model, depicted in panel a, right.
Supplementary Figure 7. **In vivo assessment of dynein\textsuperscript{mM\textsubscript{TBD}} mutant function.** (a and b) The percentage of cells with the indicated spindle orientation phenotype (green, normal; blue, aligned along mother-bud axis, but not through the neck; red, improperly aligned) is plotted for the indicated yeast strains (WT, wild-type). Anaphase spindles were visualized using mRuby2-Tub1 (\(\alpha\)-tubulin). Strains were imaged after growth to mid-log phase in SD media supplemented with either (a) 2% galactose, or (b) 2% glucose, the former of which induces overexpression of She1 in \textit{GAL1p:SHE1} cells (scale bars, 2 \(\mu\)m; error bars,
standard error of proportion; n ≥ 17 and n ≥ 21 anaphase spindles for each strain in panels a and b, respectively). Note the higher prevalence of misoriented spindles in She1-overexpressing cells (GAL1p:SHE1) than in cells lacking dynein (dyn1Δ; *, p ≤ 0.015). This suggests that She1-overexpression disrupts other non-dynein-mediated spindle orientation processes (e.g., Kar9 pathway). (c) Representative fluorescence images of kar9Δ, hydroxyurea (HU)-arrested GFP-Tub1 (α-tubulin) expressing cells with the indicated SHE1 and DYN1 alleles (scale bars, 1 µm), along with kymographs depicting spindle movements over time (horizontal scale bars, 1 µm; vertical scale bars, 1 min). Dashed lines indicate the position of the bud neck in each example. Note the frequency with which the spindle traverses the bud neck in wild-type, but not mutant cells (green arrows; see Fig. 7g for quantitation). P-values were calculated using a two-tailed unpaired t test.
**Supplementary Table 1: Strains used in this study.**

| Strain     | Genotype                                                                 | Yeast background | Source     |
|------------|---------------------------------------------------------------------------|------------------|------------|
| SMY193     | Mata kar9Δ::KANr GFP-TUB1::LEU2 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY263     | Mata dyn1ΔMTBD::3YFP::TRP1 TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY272     | Mata dyn1ΔMTBD::3YFP::TRP1 HIS3::GAL1p::SHE1 TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY293     | Mata TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY295     | Mata dyn1Δ::TRP1 TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY628     | Mata dyn1ΔMTBD::3YFP::TRP1 she1Δ::HIS3 kar9Δ::KANr GFP-TUB1::LEU2 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY714     | Mata dyn1ΔMTBD::3YFP::TRP1 kar9Δ::KANr GFP-TUB1::LEU2 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY950     | Mata DY1N1-3YFP::TRP1 JNM1-3mCherry::HIS3 she1Δ::URA3 TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY1006    | Mata GAL1p:ZZ-TEV-6xHis-GFP-3XHA-fSNAPgs-dyn1Δ31::gsDHA::KANr prb1Δ pep4Δ3::His5 his3-11,15 ura3-52 leu2-3,112 ade2-1 trp1 | W303             | This study |
| SMY1008    | Mata GAL1p:ZZ-TEV-6xHis-GFP-3XHA-GST-dyn1Δ31::gsDHA::KANr prb1Δ pep4Δ3::His5 his3-11,15 ura3-52 leu2-3,112 ade2-1 trp1 | W303             | This study |
| SMY1049    | Mata GAL1p:ZZ-TEV-6xHis-GFP-3XHA-fSNAPgs-dyn1Δ31::gsDHA::KANr prb1Δ pep4Δ3::His5 his3-11,15 ura3-52 leu2-3,112 ade2-1 trp1 | W303             | This study |
| SMY1051    | Mata GAL1p:ZZ-TEV-6xHis-GFP-3XHA-GST-dyn1Δ31::gsDHA::KANr prb1Δ pep4Δ3::His5 his3-11,15 ura3-52 leu2-3,112 ade2-1 trp1 | W303             | This study |
| SMY1377    | Mata GAL1p:ZZ-TEV-6xHis-GFP-3XHA-fSNAPgs-dyn1Δ31::gsDHA::KANr prb1Δ pep4Δ3::His5 his3-11,15 ura3-52 leu2-3,112 ade2-1 trp1 | W303             | This study |
| SMY1427    | Mata dyn1ΔMTBD::3YFP::TRP1 KANr::GAL1p::SHE1 TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY1475    | Mata dyn1ΔMTBD::3YFP::TRP1 KANr::GAL1p::SHE1 TUB1+3'UTR::HPH::His3p:mTurquoise2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY1522    | Mata DY1N1-3GFPP::TRP1 KANr::GAL1p::SHE1 pac1Δ::HIS3 TUB1+3'UTR::HPH::His3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
| SMY1554    | Mata kar9Δ::KANr she1Δ::HIS3 GFP-TUB1::LEU2 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63 | YEF473           | This study |
This study

SMY1664  Mata/α [pGBK7-53]::TRP1 [pRS315:SHE1p:GAL4-AD-SHE1]::LEU2  LYS2::GAL1uas–Gal1tata:HIS3 GAL2uas–Gal2tata:ADE2/ade2-101 MEL1/MEL1 trp1-901/trp1-901 leu2-3,112/leu2-3,112 ura3-52::URA3::Mel1tata:AUR1::MEL1 uas::URA3::GAL1uas–Gal1tata:LacZ his3-200/his3-200 gal4Δ/gal4Δ galo8Δ/gal80Δ met  Y2HGold/ Y187  This study

SMY1665  Mata/α [pGBK7:SRS-SRSCC-dyneinmTBD]::TRP1 [pGADT7-T]::LEU2 LYS2::GAL1uas–Gal1tata:HIS3 GAL2uas–Gal2tata:ADE2/ade2-101 MEL1/MEL1 trp1-901/trp1-901 leu2-3,112/leu2-3,112 ura3-52::URA3::Mel1uas–Mel1tata:AUR1::MEL1 uas::URA3::GAL1uas–Gal1tata:LacZ his3-200/his3-200 gal4Δ/gal4Δ galo8Δ/gal80Δ met  Y2HGold/ Y187  This study

SMY1666  Mata/α [pGBK7:SRS-SRSCC-dyneinmTBD]::TRP1 [pRS315:SHE1p:GAL4-AD-SHE1]::LEU2 LYS2::GAL1uas–Gal1tata:HIS3 GAL2uas–Gal2tata:ADE2/ade2-101 MEL1/MEL1 trp1-901/trp1-901 leu2-3,112/leu2-3,112 ura3-52::URA3::Mel1uas–Mel1tata:AUR1::MEL1 uas::URA3::GAL1uas–Gal1tata:LacZ his3-200/his3-200 gal4Δ/gal4Δ galo8Δ/gal80Δ met  Y2HGold/ Y187  This study

SMY1669  Mata/α [pGBK7:SRS-dyneinmTBD]::TRP1 [pGADT7-T]::LEU2 LYS2::GAL1uas–Gal1tata:HIS3 GAL2uas–Gal2tata:ADE2/ade2-101 MEL1/MEL1 trp1-901/trp1-901 leu2-3,112/leu2-3,112 ura3-52::URA3::Mel1uas–Mel1tata:AUR1::MEL1 uas::URA3::GAL1uas–Gal1tata:LacZ his3-200/his3-200 gal4Δ/gal4Δ galo8Δ/gal80Δ met  Y2HGold/ Y187  This study

SMY1670  Mata/α [pGBK7:SRS-dyneinmTBD]::TRP1 [pRS315:SHE1p:GAL4-AD-SHE1]::LEU2 LYS2::GAL1uas–Gal1tata:HIS3 GAL2uas–Gal2tata:ADE2/ade2-101 MEL1/MEL1 trp1-901/trp1-901 leu2-3,112/leu2-3,112 ura3-52::URA3::Mel1uas–Mel1tata:AUR1::MEL1 uas::URA3::GAL1uas–Gal1tata:LacZ his3-200/his3-200 gal4Δ/gal4Δ galo8Δ/gal80Δ met  Y2HGold/ Y187  This study

SMY1707  Mata/α dyn1Δ::HIS3 KAN::GAL1p:SHE1 TUB1+3'UTR::LEU2:HIS3p:mRuby2-TUB1 ura3-52 lys2-801 leu2-Δ1 his3-Δ200 trp1-Δ63  YEF473  This study

SMY1709  Mata/α [pGBK7-53]::TRP1 [pGADT7-T]::LEU2 LYS2::GAL1uas–Gal1tata:HIS3 GAL2uas–Gal2tata:ADE2/ade2-101 MEL1/MEL1 trp1-901/trp1-901 leu2-3,112/leu2-3,112 ura3-52::URA3::Mel1uas–Mel1tata:AUR1::MEL1 uas::URA3::GAL1uas–Gal1tata:LacZ his3-200/his3-200 gal4Δ/gal4Δ galo8Δ/gal80Δ met  Y2HGold/ Y187  This study
*rMTBD (MTBD obtained from rat DHC) is identical to mMTBD in amino acid sequence.*
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