Mitotic spindle formation in the absence of Polo kinase

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Abstract
Mitosis is a fundamental process in every eukaryote, in which chromosomes are segregated into two daughter cells by the action of the microtubule (MT)-based spindle. Despite this common principle, genes essential for mitosis are variable among organisms. This indicates that the loss of essential genes or bypass-of-essentiality (BOE) occurred multiple times during evolution. While many BOE relationships have been recently revealed experimentally, the BOE of mitosis regulators (BOE-M) has been scarcely reported and how this occurs remains largely unknown. Here, by mutagenesis and subsequent evolutionary repair experiments, we isolated viable fission yeast strains that lacked the entire coding region of Polo-like kinase (Plk), a versatile essential mitotic kinase. The BOE of Plk was enabled by specific mutations in the downstream machinery, including the MT-nucleating γ-tubulin complex, and more surprisingly, through downregulation of glucose uptake, which is not readily connected to mitosis. The latter bypass was dependent on casein kinase I (CK1), which has not been considered as a major mitotic regulator. Our genetic and phenotypic data suggest that CK1 constitutes an alternative mechanism of MT nucleation, which is normally dominated by Plk. A similar relationship was observed in a human colon cancer cell line. Thus, our study shows that BOE-M can be achieved by simple genetic or environmental changes, consistent with the occurrence of BOE-M during evolution. Furthermore, the identification of BOE-M constitutes a powerful means to uncover a hitherto under-studied mechanism driving mitosis and also hints at the limitations and solutions for selecting chemotherapeutic compounds targeting mitosis.

Introduction
Different organisms have different sets of essential genes for their viability and propagation (1, 2). This indicates that most ‘essential’ genes are context-dependent and can become dispensable during evolution. Plausibly, a loss of essentiality is compensated for by manifestation of an alternative, currently ‘masked’, or far less active mechanism to ensure a similar cellular activity. Many experimental efforts have been made to recapitulate the molecular diversity found in nature (3). Large-scale systematic surveys have been recently conducted in budding and fission yeasts, in which a number of bypass-of-essentiality (BOE) events have been identified (4-7). In these studies, suppressors were screened for haploid strains in which an essential gene was experimentally disrupted. For 9—27% of the essential gene disruptants, a mutation or overexpression of other gene(s), or chromosomal gain makes the strain viable, indicating that essentiality depends on
genetic background and that BOE could indeed occur at a certain frequency. However, in most cases, the underlying mechanism remains unexplored. It is also unclear why BOE is rarely observed in certain processes such as mitotic cell division.

Mitotic cell division is controlled by many essential genes in a given cell type (8-10). Evolutionary evidence of BOE is clearly visible for this fundamentally critical biological process. One striking example is the centrosome, which is assembled by the action of many essential proteins in animal, fungal, and algal species, and plays a vital role in cell division and cellular motility (11). However, the centrosome and most of its components have been lost in land plants, and yet plant cells undergo spindle assembly and chromosome segregation at high fidelity (12). Kinetochore components, such as the CCAN complex, spindle microtubule (MT)-associated proteins, such as TPX2, augmin, and mitotic motors, and cell cycle regulators, such as anaphase-promoting complex/cyclosome are among other examples; they are not universally conserved or essential factors (12-15). Despite the evidence of BOE for almost all the genes involved in mitosis, only a limited number of cases can be found in experimental BOE screening. For example, two random BOE screenings in fission yeast encompassing 23 mitotic genes have identified only a single protein, Cnp20/CENP-T, despite the fact that >20% BOE has been observed for mitosis-unrelated genes (6, 7). The BOE of cnp20 is conceivable, as CENP-T functions in parallel with CENP-C for kinetochore assembly (16, 17). Another known BOE of mitosis regulators (BOE-M) in fission yeast is MT plus-end-directed kinesin-5/Cut7, which is required for bipolar spindle formation through force generation on spindle MTs. The viability of cut7Δ was restored when the opposing minus-end-directed kinesin-14/Pkl1 was simultaneously deleted. Thus, the balance of forces applied to spindle MTs is critical (18-20). However, many other essential mitotic genes have no apparent functionally redundant or counteracting factors, and whether these relationships are general mechanisms of BOE-M is unclear. Essential mitotic genes are potential targets of cancer chemotherapy (21); it is also important to understand the BOE that underlies drug resistance.

In this study, we found that the essentiality of the sole Polo-like kinase (Plk) in fission yeast (Plo1) can be bypassed. Plo1, similar to human Plk1, is assumed to be essential for spindle MT formation and spindle bipolarisation. However, these essential processes were restored in the absence of Plo1 by multiple independent mechanisms that increase MT nucleation and stabilisation, one of which involved a remarkably simple change in glucose concentration in the culture medium and depended on casein kinase I (CK1). Thus, our study uncovered an unexpected alternative mechanism of spindle MT formation and further implies that more BOE-M can be recapitulated in the experimental system.

Results

Viable yeast cells without Polo-like kinases in several genetic backgrounds

Our previous BOE screening randomly selected 93 genes on chromosome II, which encompassed 12 mitotic genes (7). BOE-M could not be detected in any of these genes. To further screen for BOE-M, we selected eight other mitotic genes (ark1, bir1, fta2, fta3, mis6, mis14, pic1, and plo1) and applied the same screening method. Seven days after plating and UV mutagenesis of spores of each disruptant, we found a growing haploid colony for plo1Δ, the sole Polo-like kinase (Plk) in Schizosaccharomyces pombe (Fig. 1A, first step). Plks play versatile roles in animal cell division, including centriole duplication.
(by Plk4), centrosome maturation, spindle assembly checkpoint satisfaction, and cytokinesis (by Plk1). It is also a possible target for cancer chemotherapy (21-23). The responsible suppressor mutation was identified through whole genome sequencing (WGS) followed by genetic crossing, which turned out to be ght5 (Fig. 1B, left, fourth line). In S. pombe, eight hexose transporters have been identified, which show different affinities to glucose; Ght5 is a major hexose transporter with the strongest affinity to glucose and plays a critical role in glucose uptake (24). This prompted us to test and found that the plo1Δ strain grows, albeit slower than the wild-type, when the glucose concentration of the medium was lower than 0.3% (Fig. 1B, left, third line, Fig. S1A).

Thus, plo1Δ became viable under low-glucose conditions. To obtain a full scope of suppressor mutations, we repeated the UV mutagenesis of plo1Δ spores on a larger scale, obtained multiple colonies, and determined the responsible mutations. Simultaneously, an ‘experimental evolution’ (EVO) (also called ‘evolutionary repair’ (3)) experiment was conducted for the plo1Δ strain in low-glucose medium (0.08%), in which serial dilution and saturation enriches the cells that have acquired beneficial mutations for proliferation (Fig. 1A, ‘1st EVO’). The faster growing strains obtained through this step were subjected to further evolutionary repair experiments in high-glucose medium (‘2nd EVO’) and at a different temperature (‘3rd EVO’).

We determined the whole genome sequences of several viable plo1Δ strains and confirmed the suppressor mutations by independently generating a double mutant with plo1Δ. In total, 16 genes were found to assist in the growth of the otherwise inviable plo1Δ strain (Fig. 1B; D; S1B, C). An example of the evolutionary repair process is shown in Fig. S1D. This strain acquired mutations in alp6 and aps1 during the 1st EVO but still possessed the benefit of plo1+ gene for strain fitness (Fig. S1D, left). However, additional mutations in mip1 and ahk1 during the 2nd EVO bypassed the requirement of Plo1, since adding back the plo1+ gene to the original locus did not further promote colony growth (Fig. S1D, right). Most of the responsible genes were categorised into three classes: Spt–Ada–Gcn5 acetyltransferase (SAGA) complex, glucose/PKA pathway, and MT regulators (Fig. 1D). The SAGA complex is a general regulator of transcription, possessing histone acetyltransferase activity, and affects the expression of many genes (25). We did not analyse this in the present study. The cAMP-PKA pathway is linked to glucose homeostasis in fission yeast. Glucose is detected by a receptor (Git3), and the G-protein complex (Gpa2, Git5, and Git11) is activated, which then activates adenylate cyclase (Cyr1) to produce cAMP (26). Eventually, cAMP releases the inhibitor Cgs1 from Pka1, converting Pka1 to its active form (27). Furthermore, yeast cells regulate glucose uptake by changing the localisation and transcriptional level of hexose transporters, including Ght5, depending on environmental conditions (24, 28, 29). There is a link between glucose/PKA and MT stabiliser CLASP (cytoplasmic linker-associated protein) during interphase (30). In our case, mutations in MT regulators (alp4, alp6, and asp1) and glucose/PKA-pathway genes additively supported the growth of plo1Δ (Fig. 1B).

**Monopolar spindles predominate during mitosis in the absence of Polo**

The major MT nucleator at the centrosome is the γ-tubulin ring complex (γ-TuRC), which consists of γ-tubulin and GCP subunits, including GCP2 (Alp4/Spc97) and GCP3 (Alp6/Spc98) (31). In animal cells, Plk1 is a critical regulator of mitosis, which is, in the early stage, required for γ-TuRC recruitment to the centrosome and thus centrosome
maturation; inhibition of Plk1 leads to monopolar spindle formation (22, 32). In fission yeast, cytokinesis/septation defects are most profoundly observed in \textit{plo1} mutants, whereas monopolar spindle formation has also been described (33-37). However, actual spindle dynamics have not been analysed for \textit{plo1}Δ in live imaging. To analyse spindle dynamics in the absence of Plk1, live imaging of mCherry-tubulin and a spindle pole body (SPB) marker, either Sad1\textsuperscript{SUN}-GFP or Alp\textsubscript{6}GCP3-GFP, was performed after \textit{plo1}Δ spor germation with spinning-disc confocal microscopy (Fig. 2A–E). The control cell assembled bipolar spindles immediately after the disappearance of interphase MT networks (Fig. 2A and D: time 0 corresponds to the onset of mitosis), and cell division was completed in \textasciitilde30 min. In contrast, 53 out of 56 \textit{plo1}Δ cells after spor germation were arrested with a monopolar spindle for >1 h (Fig. 2B, C, E, wherein stronger laser exposure was applied in Fig. 2C to visualise the faint MT signals).

We compared the phenotype with other known mutants that show monopolar spindle formation, including Cut12, which drives SPB insertion into the nuclear envelope (NE) (38, 39), Cut\textsuperscript{7kinesin-5}, which is required for anti-parallel MT crosslinking and sliding (40), and Cdc31\textsuperscript{cintrin}, which is required for SPB duplication (41). In the \textit{cut12-1} temperature-sensitive (ts) mutant, the lack of insertion of one of the duplicated SPBs causes partial breakage of the NE and detachment of an SPB from the NE (38, 42). In another study, \textit{Plo1} was shown to regulate the formation of the Sad1\textsuperscript{SUN} ring structure, which might be required for SPB insertion (43). We assessed the integrity of the NE in \textit{plo1}Δ. First, we tagged GFP to Pcp\textsubscript{1}\textsuperscript{PCNT}, a core SPB component (35, 44), in the \textit{plo1}Δ background. In contrast to the \textit{cut12-1} mutant, we always detected punctate Pcp\textsubscript{1}\textsuperscript{PCNT}-GFP signals (n = 20) at the pole of the monopolar spindle, suggesting that SPBs in the NE generate spindle MTs in the absence of Plo1 (Fig. S2A, B). Next, we conducted an NLS-GFP efflux assay, in which partial NE breakage because of SPB insertion error leads to nuclear GFP signal efflux into the cytoplasm (38). We first confirmed the efflux in the \textit{cut12-1} ts mutant; GFP started to leak out from the nucleus \textasciitilde10 min after interphase spindle disassembly at non-permissive temperatures in 20 out of 20 cells that assembled monopolar spindles (Fig. S2D, 20, 30, and 70 min; (38)). In contrast, \textit{plo1}Δ cells maintained GFP signals inside the nucleus during the early stage of mitosis, similar to the control strains (Fig. S2C). These data suggest that SPBs are properly inserted into the NE at the onset of spindle formation in the absence of Plo1. Notably, GFP efflux during mitotic arrest occurred in 16 out of 20 \textit{plo1}Δ cells (69 \textpm 18 min after spindle assembly), suggesting that NE was partially dissolved during prolonged arrest (Fig. S2C, 100 and 200 min).

Next, we isolated cut7\textsuperscript{Δ} and cdc31\textsuperscript{Δ} spores with Alp\textsubscript{6}GCP3-GFP (SBP) and mCherry-tubulin markers and germinated them in normal culture medium. As expected, monopolar spindles were prevalent in each sample, with only a single dot of Alp\textsubscript{6}GCP3-GFP detectable at the end of spindle MTs (Fig. 2F, G). However, Alp\textsubscript{6}GCP3-GFP signal intensity was significantly lower in the \textit{plo1}Δ spindles than in cut7\textsuperscript{Δ} or cdc31\textsuperscript{Δ} (Fig. 2H). Consistent with this phenotype, spindle MTs were dimmer in \textit{plo1}Δ (compare Fig. 2E and 2F/G) and \textit{plo1}Δ was sensitive to thiaobendazole (TBZ), an MT-destructing drug (Fig. S3A). Finally, we checked whether Cut\textsuperscript{7kinesin-5} localisation was defective in \textit{plo1}Δ, which would cause spindle monopolarisation. Cut\textsuperscript{7kinesin-5}-GFP accumulation at the SPB and spindle was delayed in the absence of \textit{plo1} (Fig. S3B, 0 min). However, the signals gradually recovered and reached a level comparable to the early prometaphase of control cells, at which spindle bipolarity was not recovered (Fig. S3C). Thus, it is unlikely that failure in
Cut7 kinesin-5 recruitment is the major cause of spindle monopolisation in plo1Δ. Rather, our data favour the model whereby decreased MT nucleation at SPB leads to spindle monopolisation in plo1Δ. Consistent with this notion, monopolar spindles have been observed in mutants of the γ-TuRC component (Alp4GCP3) (45).

Modulation of MT nucleation and stability bypassed Polo essentiality

Two BOE strains had point mutations in alp4GCP2 and alp6GCP3. Double alp4-D440E plo1Δ and alp6-V664F plo1Δ strains recovered colony-formation ability in normal (high-glucose) medium (Fig. 1B). Thus, the essentiality of Plo1 was bypassed by a single specific mutation in the MT nucleating machinery. We also performed a spot test for single alp6-V664F and alp4-D440E mutants (Fig. 1C). alp6-V664F grew more slowly than the wild-type, whereas no difference in colony growth was observed for alp4-D440E.

We investigated whether the mutation in alp4 could restore γ-TuRC recruitment to the SPB in the absence of plo1. To address this, we isolated a double alp4-D440E plo1Δ mutant with Alp6GCP3-GFP and mCherry-tubulin markers (Fig. 3A). Quantification indicated that both GFP and mCherry signals were partially but significantly restored by the alp4-D440E mutation (Fig. 3B, C). In 50% of the cells (n = 46), spindle bipolarity was recovered after a delay and cytokinesis was completed (Fig. 3A, left), whereas monopolar states were persistent for >60 min in 30% of the cells (Fig. 3B, right), explaining the partial rescue of the viability by this specific mutant of alp4. Consistent with frequent spindle bipolarisation, GFP efflux in the viable alp4-D440E plo1Δ and ght5Δ plo1Δ strains was less frequently observed than in single plo1Δ (19% and 21%, n = 26 and 48) (Fig. S2E, F).

Asp1PPP5K/Vip1 is another MT-related factor and its mutation assisted in the growth of plo1Δ (Fig. 1B). Asp1PPP5K/Vip1 is known to have a kinase domain at the N-terminus and a phosphatase domain at the C-terminus, and the latter is required for MT destabilisation (46). Interestingly, two mutations acquired during experimental evolution were located at the C-terminus (Fig. 1D). The mutation did not affect colony growth in the presence of Plo1 (Fig. 1C). However, time-lapse imaging showed that the asp1-D507G plo1Δ strain exhibited more spindle MT signals than plo1Δ, indicating that mutations in the C-terminal domain of Asp1 cause spindle MT stabilisation (Fig. 3D–G). These data suggested that bypass of Plo1 essentiality is achieved by increasing MT stability and/or generation.

Glucose limitation bypasses Plo1 essentiality

The mechanism by which glucose limitation recovers the viability of plo1Δ is not readily explainable. Glucose reduction did not appear to change Plo1-GFP localisation. Both in high (3%) and low (0.08%) glucose media, Plo1-GFP was localised to SPBs from prophase to metaphase and delocalised at anaphase (Fig. S4A, B). To observe the process of mitosis, we followed Alp6GCP3-GFP and spindle MTs in double ght5Δ plo1Δ (Ght5 is a glucose transporter). Interestingly, Alp6GCP3-GFP accumulation at the SPB and spindle MT abundance were restored in the double mutant (Fig. 4A–D). MTs appeared to be more stable in ght5Δ, as incomplete disassembly of interphase MTs was often observed at the onset of mitosis, which reflected more total mCherry signals in the mutant than in the wild-type (arrows in Fig. 4B). Consistent with this observation, ght5Δ conferred resistance to TBZ (Fig. S3D).
Next, we tested the localisation of Mid1\textsuperscript{anillin}, which is recruited to the equatorial region during mitosis and defines the division site, depending on phosphorylation by Plo1 (47). We observed that Mid1 was not properly localised to the cortex in the viable ght5Δ plo1Δ strain, whereas the cortical localisation was normal in single ght5Δ (Fig. 4E–G). Consistent with this observation, the septum was mislocalised in ght5Δ plo1Δ, similar to mid1Δ (Fig. S4C–E). The results revealed that the septation error was not directly linked to the lethality of plo1Δ. Cdc7\textsuperscript{Hippo} is another downstream factor of Plo1; the SPB localisation of Cdc7\textsuperscript{Hippo} during metaphase, but not telophase, is impaired in the plo1 mutant (48). In the viable ght5Δ plo1Δ strain, Cdc7\textsuperscript{Hippo}-GFP localisation at metaphase SPB was not detectable (Fig. 4H, I). These results indicated that not all Plo1 downstream events, including the phosphorylation of the direct substrate, are restored by ght5 mutations.

**Casein kinase 1 (CK1) constitutes a masked mechanism for spindle bipolarisation**

Since proteins in the glucose-PKA pathway are not SPB- or spindle-associated, we hypothesised that other pathways are enhanced when glucose is limited, which promotes γ-TuRC localisation. To identify the effector proteins in such pathways, we performed a genetic screening, with the aim to acquire mutants that were synthetic lethal with double ght5Δ plo1Δ or pka1Δ plo1Δ. For this, we first transformed a plasmid containing the plo1\textsuperscript{+} gene in the double mutants, conducted mutagenesis, and selected the strains that could not lose the plasmid (Fig. 5A). A total of 13 mutants were identified that were synthetic lethal with either ght5Δ plo1Δ (seven strains) or pka1Δ plo1Δ (six strains). Possibly responsible genes were selected based on sequencing (e.g. dramatic amino acid changes, nonsense mutations, or identified in multiple strains). Synthetic lethality was confirmed for five genes (bub1, hhp1, iml1, mak1, and wis1) and one gene (sin1) by gene disruption and crossing with ght5Δ plo1Δ and pka1Δ plo1Δ, respectively. However, two mutants (iml1 and wis1) and one mutant (sin1) resulted in poor growth when singly combined with ght5Δ and pka1Δ, respectively. These were excluded from further analysis because the major basis of synthetic lethality may not involve the lack of Plo1 kinase. mak1 showed complex genetic interaction; while mak1Δ ght5Δ grew normally, synthetic lethality was revealed when mCherry-tubulin was introduced. In addition, the double mak1Δ pka1Δ grew poorly in the absence of mCherry-tubulin expression. Therefore, this gene was also excluded from further analyses. In contrast, triple disruption was not selected for two other genes, bub1 and hhp1 (Fig. S5A, B), whereas the double mutants with ght5Δ grew in a manner indistinguishable from the single ght5Δ even in the presence of mCherry-tubulin. We further confirmed the synthetic lethality of hhp1Δ with other PKA-pathway genes ght1Δ plo1Δ and pka1Δ plo1Δ (Fig. S5C, D). Thus, bub1 and hhp1 were essential for plo1Δ viability.

To identify the lethal event caused by these mutations, we observed live cells of the triple disruptants, bub1Δ ght5Δ plo1Δ and hhp1Δ ght5Δ plo1Δ. For this, we selected each triple disruptant that possessed the Plo1-GFP multicopy plasmid. Viable cells were cultured in non-selective medium, by which cells naturally lose the plasmid at a certain probability. Time-lapse images were then acquired. We analysed the cells that no longer had Plo1-GFP signals, as these cells represent triple gene disruptants. As a control, we prepared double ght5Δ plo1Δ possessing the Plo1-GFP plasmid and performed the identical ‘plasmid loss’ culture. In the control strain that had no GFP signals, monopolar
spindles were converted into bipolar spindles within 30 min, followed by entry into anaphase, in >60% cells, as expected (Fig. 5B and E). In contrast, in triple bub1Δ ght5Δ plo1Δ, anaphase began even when spindles were still monopolar in 13 out of 43 cells (Fig. 5C). This phenotype explains the lethality of the strain and is consistent with the fact that Bub1 is an integral component of the spindle assembly checkpoint, which prevents premature anaphase entry (49). In contrast, in hhp1Δ ght5Δ plo1Δ, >80% of cells were arrested in monopolar states for >30 min, and spindle bipolarisation and anaphase entry were scarcely observed, similar to the plo1Δ strain in the normal medium (Fig. 5D and E). We concluded that the lethality of hhp1Δ ght5Δ plo1Δ comes from a defect in spindle bipolarisation, similar to plo1Δ in the normal medium.

hhp1 encodes casein kinase I (CK1), which is distributed throughout the cell and is enriched at the SPB (50, 51). Hhp1^{CK1} is involved in a variety of cellular processes, such as DNA repair, ubiquitination-dependent regulation of septation initiation, DNA recombination and cohesin removal during meiosis (50, 52-54). However, to the best of our knowledge, Hhp1^{CK1} has not been directly linked to spindle function in fission yeast.

To investigate the basis of the unexpected genetic interaction, we first tested whether Hhp1^{CK1} expression/localisation was altered by ght5 disruption. To this end, we tagged GFP to Hhp1^{CK1} in the wild-type and ght5Δ backgrounds. Time-lapse mitosis imaging and GFP intensity quantification indicated that Hhp1^{CK1} localisation was unchanged, but the overall abundance became more variable and on average slightly increased in the absence of ght5 (96 ± 23, 120 ± 63 [AU, ± SD], n = 30 each). However, the level of hhp1 mRNA was not elevated by a glucose reduction, suggesting that post-transcriptional regulation underlies the increased Hhp1 in the cell (Fig. S5E). Next, we tested whether the upregulation of Hhp1^{CK1} is necessary for the bypass of Plo1. We selected the hhp1Δ ght5Δ plo1Δ triple disruptant that possesses the Hhp1^{CK1}-GFP multicopy plasmid, and conducted a plasmid-loss experiment. In this experiment, GFP signal intensity served as an indicator of intracellular levels of the Hhp1^{CK1} protein. Time-lapse imaging and subsequent image analysis showed that the level of the Hhp1^{CK1} protein was overall correlated with the efficiency of spindle bipolarisation (Fig. 5G, 5H [grey vs. green]). However, the impact of the slight increase in Hhp1^{CK1} observed in ght5Δ was marginal; when we compared the time required for monopolar-to-bipolar conversion, we observed a slight and statistically nonsignificant decrease (Fig. 5H [grey vs. orange]). Thus, a moderate increase in Hhp1^{CK1} facilitates bipolar spindle formation in the absence of Plo1 and Ght5, although it may not be a prerequisite for BOE.

Next, we investigated whether ectopic expression of Hhp1^{CK1} was sufficient for the recovery of plo1Δ viability. We tested the expression of Hhp1^{CK1} by two different promoters on the multicopy plasmid, but we could not obtain data that reproducibly showed that Hhp1^{CK1} expression restored plo1Δ colonies (Fig. S5F). In addition, Hhp1^{CK1} expression from the plasmid did not enhance the growth of alp6-V664F plo1Δ or alp4-D440E plo1Δ, which was viable on its own but had slower growth than the wild-type (Fig. S5F). Thus, an increase in Hhp1^{CK1} levels alone did not increase the fitness of plo1Δ and was insufficient for the bypass of Plo1 essentiality.

Finally, we observed spindle dynamics in the hhp1 single disruptant. Most of the cells (99%) assembled bipolar spindles, and mitosis proceeded comparably to the wild-type. However, among the 272 cells monitored, we found that 3 cells (1%) formed monopolar spindles; this was not observed in our imaging of control Hhp1^{CK1}+ cells (N
>336) (Fig. 5I). Furthermore, hhp1Δ ght5Δ was more sensitive to TBZ than ght5Δ (Fig. 3SD). Thus, Hhp1CK1 has a very mild, almost negligible level of contribution to MT stability and bipolar spindle assembly in the presence of Plo1, but becomes essential in the absence of Plo1. In S. pombe, hhp2Δ also encodes CK1 (52). Therefore, we selected the hhp1Δ hhp2Δ double disruptant expressing mCherry-tubulin and Alp6GFP, and performed time-lapse microscopy. Interestingly, monopolar spindles appeared at a much higher frequency than single hhp1Δ (29%, n = 79) (Fig. 5I). Other phenotypes, such as undeveloped spindle MTs, were also observed in the double disruptant (Fig. 5I, right). We further determined if hhp2Δ would be synthetically lethal with three viable plo1Δ strains (ght5Δ plo1Δ, git1Δ plo1Δ, and pka1Δ plo1Δ). Unlike hhp1Δ, no strains showed synthetic lethality with hhp2Δ. Thus, Hhp1CK1 and Hhp2CK1 were not completely redundant for bypass-related functions, which corroborates the previous report that they are different in subcellular localisation and abundance (51).

**Masked contribution of CK1 to spindle formation in a human colon cancer cell line**

Among the four Plks in mammals, Plk1 is required for centrosome maturation and bipolar spindle formation in many cell types and is thus most analogous to S. pombe Plo1. There are also several CK1 family members in mammals. As CK1δ is localised at the centrosome (55, 56), we tested whether CK1δ constitutes the masked mechanism behind Plk1 in human cells (Fig. 6). The treatment of a human colon cancer line (HCT116) with a low concentration (3 nM) of Plk1 inhibitor BI2536 resulted in a slightly higher frequency of monopolar spindle appearance in early prometaphase (Fig. 6A and B). PF670462, an inhibitor of CK1δ/ε (57, 58), did not increase the number of monopolar spindles. However, when both inhibitors were simultaneously treated, 36% of the cells first assembled monopolar spindles (Fig. 6A and B). The monopolar spindles were eventually converted to bipolar spindles; however, this process required >30 min in ~20% of the cells when two compounds were simultaneously added (Fig. 6C). These results highlight the importance of CK1, perhaps CK1δ, in spindle bipolarisation in human colon cancer cells, when Plk1 function is partially impaired.

**Discussion**

This study represents a rare example of the experimental BOE of genes required for mitosis. The BOE occurrence in Plo1 was unexpected, as it has been recognised as a versatile, essential kinase in mitosis not only in animal cells but also in fission yeast. However, there is evolutionary evidence supporting that this gene can be deleterious; for example, plants have lost Plks, whereas the ancestral algae possess Plks (59). In our initial BOE screening using plo1Δ spores, only one viable strain was recovered, in which the gene encoding the glucose transporter Ght5 was lost through a deletion event. Subsequent evolutionary repair (experimental evolution) experiments led to the identification of more mutations, many of which restored viability of plo1Δ without the ght5 mutation. Thus, the initial mutagenesis-based screen was not sensitive enough to capture all the possible BOE. More BOE may be uncovered in the yeast system, including BOE-M, by applying more sensitive methods, or simply by increasing the screen scale.

Plo1 loss can be rendered non-lethal by mutations in several genes, some of which were unrelated to each other and not associated with spindle functions at first glance.
However, this is in accordance with many previous examples of BOE or evolutionary repair in the laboratory, where compensatory mutations are often found in genes outside of the perturbed functional module (3). Subsequent analysis suggested that bypass mutations converge into a common outcome: the increase in spindle MTs. This was achieved by multiple direct and indirect mechanisms, such as mutations in an MT destabiliser and MT nucleator, or through glucose starvation. In contrast, the septum phenotype was not rescued in a viable strain. Thus, although multiple defects have been identified in the plo1 mutants, MT formation is directly linked to viability. In a broader sense, BOE analysis could be used to distinguish between essential and non-essential functions of an essential gene.

The bypass of Plo1 essentiality by glucose reduction in the medium is intriguing from multiple perspectives. First, it illustrates the non-automatic nature of gene essentiality (1). If the low-glucose medium was used as the standard yeast culture medium, then Plo1 would have been assigned as a non-essential gene in *S. pombe*. Second, a change in available nutrients occurs, perhaps frequently, in the natural yeast habitat. The decrease in available glucose allows the yeast to lose a critical mitotic kinase and develop an alternative mechanism. The data support the theory that environmental change combined with gene mutations drives molecular diversity, namely, variation in genes required for an essential process (3). Third, the change in fundamental metabolism alters the expression of many genes (60), offering a unique ‘genetic background’ that is not achieved by mutations in a few mitotic genes. In the case of plo1Δ, a critical factor for survival was Hhp1^{CK1}. Because Hhp1^{CK1} is SPB-associated, it is possible that critical Plo1 substrates (such as γ-TuRC or its associated factor) are phosphorylated by Hhp1^{CK1}. However, CK1 is unlikely the sole element of BOE based on glucose repression and other factors should be also involved, as Hhp1^{CK1} overexpression alone was not sufficient to restore the viability of plo1Δ (Fig. 7). Interestingly, *S. cerevisiae* Hrr25^{CK1} can phosphorylate and activate the γ-tubulin complex *in vitro* and this phosphorylation is required for *in vivo* γ-tubulin functions (61). Whether Hrr25^{CK1} constitutes a masked mechanism of Cdc5^{pkl} in *S. cerevisiae* is an intriguing question for future investigation.

BOE, or synthetic viability, is a critical challenge in cancer chemotherapy because of the emergence of resistance (62). Plk inhibitors have been recognised as promising antitumor drugs (21, 23). However, our study suggests that there may be resistant cells involving CK1 and that double inhibition of Plk1 and CK1Δ may be more suitable for mitotic cell perturbation.

### Materials and methods

**Yeast strains and media**

We followed Takeda et al. (2019) for the yeast culture and gene disruption. Complete medium YE5S contained 1% yeast extract and 3% (w/v) glucose, supplemented with adenine, leucine, histidine, uracil, and lysine, whereas glucose was reduced to 0.02% in the low-glucose medium. Synthetic PMG and EMM media were used when selection was based on adenine, leucine or uracil. Cells were cultured at 32°C (plate) or 30°C (liquid), unless other temperatures are indicated. Sporulation was induced in an SPA plate. Gene disruption, site-directed mutagenesis, and GFP/mCherry tagging were performed using the standard one-step replacement method (i.e. homologous recombination). Transformation was performed using the conventional LiAc/PEG method (63), and
targeted integration was confirmed by PCR. Selection of the strain after random spore or
transformation was based on leucine, uracil, G418 (100 µg/mL), hygromycin (50 µg/mL),
clonNAT (100 µg/mL), or blasticidin S deaminase (37.5 µg/mL). The strains, plasmids,
and primers used in this study are listed in Tables S1, S2, and S3, respectively.

**Human cell culture**
The human colon cancer-derived HCT116 line, in which the endogenous TubG1 gene (γ-
tubulin) was tagged with mClover, was cultured in McCoy’s 5A medium (Gibco)
supplemented with 10% serum and 1% antibiotics (64). Plk1 inhibitor BI2536 (3 nM) and
CK1 inhibitor PF670462 (2 µM) were treated for 24–30 h prior to imaging, whereas 30
nM BI2536 was added at the beginning of imaging. MTs were stained with 50 nM SiR-
tubulin for 2 h prior to imaging; this concentration of SiR-tubulin did not significantly
affect MT growth and nucleation in this cell line (64).

**BOE screening and confirmation**
We followed the method described by Takeda et al. (2019). The initial screening
encompassing eight mitotic gene disruptants was performed for 1 × 10^7 spores, and plo1
screening was repeated with a larger scale (8 × 10^7 spores). One or 11 viable colonies
were obtained for plo1 in the respective screening, for which PCR confirmed that the
strain was indeed deleted from the entire plo1 open reading frame (ORF). One of the
viable plo1Δ strains was crossed with the wild-type (975 or 972), followed by sporulation
on an SPA plate. The spores were incubated in YE5S with G418 (100 µg/mL) and
cycloheximide (100 µg/mL) plates. The viable colonies were subjected to WGS (i.e. bulk
segregant analysis). Analysis of the genome sequence of the strain identified a unique ~7
kb deletion on chromosome III, in which three genes, SPCC1235.17, SPCC1235.18, and
ght5, were included. By crossing other lines with the Ght5-GFP integrant strain (24), we
observed a strong genetic link for six of the strains (one strain was sterile and was not
analysed further). Sequencing of the three strains verified that all had mutations or
deletions in the ght5 locus; one showed a mutation from TGG to TAG that introduced a
premature stop codon in ght5 and another showed Gln153 to Pro substitution of ght5. To
confirm this suppression, the ght5 gene was independently deleted by homologous
recombination, and the growth of ght5Δ plo1Δ cells was investigated using a spot test.
WGS of the other three plo1Δ revertants identified other suppressor candidates (the other
line was sterile and was not further analysed). One strain had TT instead of AC in the
ngg1 coding region, which introduces a premature stop codon. Ngg1 (also called Ada3)
is a component of the SAGA complex (65). To investigate whether the mutations in ngg1
rescue plo1 lethal, a genetic linkage test was conducted with the rec6 gene, which is
located close to ngg1. The other two revertants had mutations in the cyr1 gene (encoding
adenylate cyclase, a component of the PKA pathway). Nucleotide insertion in cyr1 of one
strain caused a frameshift that induced a premature stop codon, whereas the other strain
had an amino acid change from Asn to Lys. Therefore, the plo1Δ rescue ability by pka1
or cyr1 deletion was tested. The strains lacking both pka1 and plo1 or cyr1 and plo1 grew
on 3% glucose medium, albeit slower than the wild-type.

**Whole genome sequencing**
Approximately 2 × 10^8 cells were harvested, and their genomic DNA was extracted using
Dr. GenTLE (Takara). The genomic DNA was sequenced by BGI, Novogene, or the gene
sequencing facility at Nagoya University, where the read length of the sequence was 150 bp or 81 bp. The FASTQ file was modified and mapped using CLC genomic workbench software. The ends (5 bp from the 5′ end and 1 bp from the 3′ end) of each fragment were trimmed, and these fragments were then filtered for a minimum sequencing quality of 30 datasets to reduce the error ratio. The fragments were aligned to the *S. pombe* genome reference and formatted as BAM files. IGV was used to visualise the BAM file and check the mutation site. Insertion, deletion, single nucleotide variation (SNV), and large indels were investigated. All the WGS data are available at Sequence Read Archive (SRA) at NCBI (accession number: PRJNA768628).

**Experimental evolution**

Experimental evolution was conducted with serial dilution and saturation (66). Saturated culture after passage (10 mL) was 1,000-fold diluted for the next passage. After culturing for 150–200 generations, cell cultures were spread onto agar plates, and the fastest growing colonies were selected for each culture. In the first round, six *plo1Δ* colonies were inoculated into 0.08% glucose liquid medium with G418 (10 µg/mL; in order to avoid microbial contamination) and cultured at 30°C. Eventually, six independent colonies were picked up (hereafter called ‘1st EVO’ strains). They showed faster colony growth on 0.08% glucose agar plate than the original *plo1Δ* strain. Three of them also formed small colonies on the normal 3% glucose medium. These strains were further subjected to experimental evolution using 3% glucose medium with G418 (10 µg/mL) at 30°C (two or three independent cultures). In the end, eight ‘evolved’ strains (i.e. the fastest growing strains) were obtained as *plo1Δ* second-round evolution strains (hereafter called 2nd EVO). Another round of experimental evolution (3rd EVO) was carried out for *git1Δ plo1Δ* and *pkα1Δ plo1Δ* strains at 36°C (two and six independent cultures, respectively), as these formed tiny colonies at 36°C, whereas the *plo1Δ* strain did not grow even in the low-glucose medium at this high temperature. We determined the genome sequences and identified the specific mutations in five out of six strains from 1st EVO, including three strains for which 2nd EVO selection was carried out, and six out of eight strains from 2nd EVO. On average, we found 2.2 mutations introduced in the open reading frame. Three out of the five 1st EVO strains had a point mutation in SAGA complex genes (*spt20*, *sgf73*, and *spt7*). Mutations in glucose transporters or PKA-pathway genes (*ght5*, *git5*, *git1*, and *ght8*) were found in four of the six 2nd EVO strains. From the 3rd EVO, we sequenced the three fastest growing clones and found *asp1* mutations in two strains and *tral* (SAGA) in the third strain.

**Synthetic lethality screening**

Double disruptants *ght5Δ plo1Δ* and *pkα1Δ plo1Δ* were transformed with the Plo1 plasmid (*ura4*+). Cells (5 × 10⁴ – 1 × 10⁶ cells) were mutagenised with UV (150 or 200 × 100 µJ/CM²; UVP Crosslinker CL-1000) and plated onto a PMG (*ura4–*) plate. After 6 d, the colonies were replica plated onto YE5S plates. After 2 d, the colonies were replica plated onto 5-FOA and PMG (*ura4–*) plates. Cells that could not survive in the absence of the plasmid were isolated (inviable colonies after replica plating onto 2 mg/mL 5-FOA-containing plates). For one of the obtained strains, the genomic DNA library (pTN-L1, NBRP) was transformed into the strain, and the colonies found on 5-FOA plates were isolated. The plasmid extracted from the colonies was sequenced and found to encode *hpt1*. However, this method did not work well for other strains. Therefore, for the other
12 strains, whole genome sequences were determined and candidate mutations (nonsynonymous and frameshift) were inspected by independent deletion and crossing.

**Microscopy**

Exponentially growing yeast cells or spores were attached to a 1 μg/mL lectin-coated glass plate for >10 min at 32°C (67). Live imaging was performed with a spinning-disc confocal microscope (Nikon Ti; 100× 1.45 NA) with a perfect focus system. The cells were maintained at 32°C in a stage-top heater. Images were taken using an ImagEM CCD camera (Hamamatsu) every 1–3 min for exponentially growing cells or every 1 or 2 min for germinating spores (z-stacks: 1 μm × 5 sections). Spores were pre-incubated for 10–12 h in 3% YE5S medium before imaging. For imaging that involved plasmid loss, cells were pre-incubated overnight in PMG without uracil or leucine and exponentially growing cells were transferred to 3% YE5S medium for ~24 h prior to microscopy.

Human cell live imaging was performed with a spinning-disc confocal microscope (Nikon Ti; 40× 1.45 NA) with a perfect focus system. Images were captured using an ImagEM CCD camera (Hamamatsu). The cells were maintained at 37°C in a stage-top heater, where CO₂ was supplied. Z-stack images (3 μm × 5 sections) were taken every 3 min. Images were analysed using FIJI, and the data were plotted using GraphPad Prism software. The signal intensity of mCherry-tubulin on the spindle and Hhp1-GFP in late G2 was measured after maximum projection, whereas a single focal plane was selected for Alp6-GFP and Cut7-GFP measurements. Two Alp6-GFP signals on the bipolar spindles were summed to obtain the total intensity. The maximum projection images are presented in the figures.

**Real time PCR**

Exponentially growing yeast cells were lysed with zymolyase (50 U per 1 × 10⁶ cells), and total RNA samples were prepared using the RNeasy Plant Mini kit (Qiagen), using the yeast protocol described in the manufacturer’s instructions of RNeasy Mini Kit (Qiagen). To eliminate genomic DNA contamination, an additional DNase treatment was performed using an RNase-free DNase Set (Qiagen). PrimeScript II (Takara) was used for the reverse transcription. Real-time PCR was performed and analysed using Step One Plus (Applied Biosystems) with SYBR Green Master Mix (Applied Biosystems). The primer set designed at the C-terminus of act1 was used as the control (24). The copy number of hhp1 mRNA relative to act1 was calculated from a standard curve drawn using serial dilutions of cDNA as the templates.

**Statics**

All statistical analyses were performed using the GraphPad Prism software. Two-way ANOVAs were applied between the two groups in Fig. 3B, 3C, 3G, 4C, 4D, and 5F, and a two-way ANOVA with multiple comparisons using Tukey’s test to compare the four groups in Fig. 2H and a one-way ANOVA with multiple comparisons using Tukey’s test to compare the four groups (Fig. 5H). The data distribution was assumed to be normal, but this was not formally tested. Established P values were denoted as follows: P > 0.05 (ns), P < 0.05 (*), P < 0.01 (**), P < 0.001 (***) and P < 0.0001 (****). Error bars in the graph represent the SEM or SD of each group. Experiments were performed twice or more, and the data from one experimental set were presented after quantitative analysis,
except for Fig. 4C, 4D, 5G, 5H, and 6C, where the data from multiple experiments were combined because of insufficient sample numbers in a single experiment.

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Figure 1. Isolation of viable plo1Δ strains

(A) Experimental procedure to isolate plo1Δ strains. The yeast spores in which plo1 gene was replaced by drug-resistant cassette were mutagenised by UV and plated onto the drug-containing medium. The haploid colonies that appeared after several days represent the plo1Δ strains. The whole genome sequence (WGS) was determined to map the responsible suppressor mutations, while a few strains were subjected to experimental evolution (EVO), where serial dilution and saturation accumulated fitness-increasing mutations. EVO was repeated three times in different conditions, and suppressor mutations were determined by WGS. (B) Viable plo1Δ strains obtained by indicated suppressor mutations. Cells (5,000, 1,000, and 200) were spotted onto normal YESS plates, except for the third row, where glucose (glc) concentration in the medium was reduced to 0.08% (YESS, 4 d, 32°C). (C) Single mutants of alp4-D440E, alp6-V664F, and asp1-D507G. Cells (5,000, 1,000, and 200) were spotted onto normal YESS plates and incubated for 3 d at 32°C. (D) List of suppressor mutations for plo1Δ. The ‘WGS’ column indicates at which step in (A) the mutation was identified. ‘Rescue’ column indicates whether the indicated mutation alone bypassed the essentiality of Plo1. The colony grew extremely poorly for the spt20Δ plo1Δ (marked with −).
Figure 2. Monopolar spindle formation with reduced microtubules (MTs) and γ-TuRC localisation in plo1Δ
(A, B) Live imaging of the control and plo1Δ strains expressing Sad1SUN-GFP and mCherry-tubulin. The first mitotic phase after spore germination was imaged. (C) Mitotic spindles of control and plo1Δ strains expressing Sad1SUN-GFP and mCherry-tubulin with longer exposure time. (D, E) Live imaging of the control and plo1Δ strains expressing Alp6(ΔC99)-GFP and mCherry-tubulin. (F, G) Monopolar spindles of the cut7Δ and cdc31Δ strains after germination. (H) Quantification of Alp6(ΔC99)-GFP intensity during mitosis. The signal intensities from 5 to 8 min were compared between the plo1Δ and wild-type (WT), cut7Δ, or cdc31Δ (****, p < 0.0001; *, p = 0.0226). Error bars indicate the standard error of the mean (SEM). Time 0 (min) was set at the onset of spindle formation.
Figure 3. Microtubule (MT) nucleation and spindle bipolarisation were rescued by a point mutation in a γ-TuRC subunit or MT destabiliser

(A) (Left) Spindle bipolarisation after a prolonged monopolar state by a specific mutation in the alp4-GCCP2 gene. Equatorial MTs during telophase were also recovered (arrow at 78 min). (Right) Failure in spindle bipolarisation. (B, C) Partial recovery of Alp6GCCP1-GFP and MT intensities by a specific mutation in the alp4GCCP2 gene. The signal intensities from 12 to 15 min were compared between plo1Δ and alp4-Δ440E plo1Δ; Alp6GCCP1-GFP intensity (**, *p* = 0.0049), and MT intensity (**, *p* = 0.0017). (D–G) Partial recovery of MT intensities by a mutation in the asp1PPPSK/Vip1 gene. MT intensity from 12 to 15 min was compared between plo1Δ with asp1-D507G plo1Δ (***, *p* < 0.0001). In all the graphs, error bars indicate the standard error of the mean (SEM). Time 0 is set at the onset of spindle formation. Time 0 (min) is set at the onset of spindle formation.
Figure 4. γ-TuRC localization was restored by mutations in a glucose transporter in the absence of Plo1
(A) Live imaging of the plo1Δ strain expressing Alp6\textsuperscript{(GFP)}-GFP and mCherry-tubulin. The first mitotic phase after spore germination was imaged. (B) Live imaging of the ght5Δ plo1Δ strain expressing Alp6\textsuperscript{(GFP)}-GFP and mCherry-tubulin. Mitosis in the exponentially growing phase was imaged. Arrows indicate interphase microtubules (MTs) that remain during spindle assembly. (C, D) Quantification of Alp6\textsuperscript{(GFP)}-GFP and MT intensities during mitosis. The signal intensities were compared between plo1Δ with ght5Δ plo1Δ: Alp6\textsuperscript{(GFP)}-GFP intensity from 7 to 10 min (****, p < 0.0001) and MT intensity from 12 to 15 min (****, p < 0.0001). Error bars indicate the standard error of the mean (SEM). Time 0 is set at the onset of spindle formation. Control data is identical to that in Fig. 2I and J. The increase in MT intensity during the early mitotic stage in ght5Δ plo1Δ is due to the incomplete disassembly of interphase MTs (see arrows in B). (E–G) Equatorial accumulation of Mid1\textsubscript{1min}–GFP is not restored in the viable ght5Δ plo1Δ strain. (H, I) Spindle pole body (SPB) localization of Cdc7\textsubscript{Bip}–GFP at metaphase is not restored in the viable ght5Δ plo1Δ strain. Arrows indicated GFP signals at SPBs. Time 0 (min) is set at the onset of spindle formation.
Figure 5. Hhp1<sup>KK1</sup> becomes essential for spindle formation in the absence of Plo1

(A) Schematic presentation of the synthetic lethal screening. The strain possessing Plo1 plasmid (ura4+) is sensitive to 5-FOA and therefore, does not grow. The strain that cannot grow specifically on the 5-FOA plate should have a mutation that is synthetical lethal with ght5A plo1A or plaA plo1A. The genome sequences of these strains were determined ("WGS"). (B–D) Plasmid-loss experiment. The indicated double or triple disruptants transformed with Plo1-GFP plasmid were grown. Time-lapse imaging was performed and mitotic cells with or without Plo1-GFP signals were analysed. (E) Frequency of mitotic phenotypes (in the absence of Plo1-GFP). (F) MT intensity decreased in the absence of Hhp1<sup>KK1</sup>(***, p <0.0001). (G) Plasmid-loss experiment using a plo1A ght5A hhp1A triple disruptant and multicopy Hhp1-GFP plasmid (leu<sup>+1</sup>). Time spent with monopolar spindles was plotted for each cell. GFP intensity (arbitrary unit) corresponds to the amount of Hhp1 in a cell. Mean intensities (± SD) of Hhp1-GFP signals in the wild-type background and ght5A background were indicated by grey and orange bars, respectively. In this analysis, the mean background intensity of the parental strain that had no GFP expression (80.3 AU, n = 31) was subtracted from the Hhp1-GFP intensity value. (H) Time required for monopolar-bipolar conversion. Three bars correspond to the samples with different GFP intensities described in (G). Error bars represent SD (*, p = 0.0458; ***, p =0.0001; ****, p <0.0001), and ns stands for “not significant.” (I) A total of 29% of the hhp1A hhp2A cells (n = 79) and 1% of the hhp1A cells (n = 272) assembled monopolar spindles, whereas this never occurred in control cells (n = 336). A lack of spindle MTs was also observed in the double disruptant (right). Time 0 (min) was set at the onset of spindle formation.
Figure 6. Synthetic monopolar spindle phenotype by partial inhibition of Polo-like kinase (Plk1) and casein kinase I (CK1)δ in human colon cancer cells

(A) Mitosis of the HCT116 cell line in the presence of Plk1 and/or CK1 inhibitors (BL2536 for Plk1, PF670462 for CK1). Green, β-tubulin-mClover (endogenously tagged); Red, Sir-tubulin. (B) Frequency of monopolar spindles (monopolar state for ≥10 min). (C) Duration of nuclear envelope breakdown (NEBD)-to-bipolar spindle formation. Time 0 (min) is set at the onset of spindle formation.
Figure 7: How Plo1 essentiality is bypassed
An increase in spindle microtubules (MTs) is the key to bypassing Plo1. This can be achieved by mutations in MT-associated proteins or nucleators (1) or global change in glucose metabolism (2), which involves casein kinase I and other unknown factors.
Figure S1. The bypass of essentiality (BOE) of Plo1

(A) Colony spotting on YES5 (+ G418, CHX) plates with various glucose concentrations. Spores (7,500, 1,500, 300, or 60) were spotted and incubated for 5 d at 32°C (expected drug-resistant cell numbers: 1,875, 375, 75, and 15, respectively). (B) Disruptant of PKA (pka1Δ) or adenylate cyclase (cyr1Δ) bypassed Plo1 essentiality. Colonies were grown on YES5 plates for 4 d at 32°C. (C) An example of suppressor mutation confirmation based on linkage test. nggl and rec6 genes are closely located on chromosome II. Double plo1Δ rec6Δ was never obtained after crossing rec6Δ (marked with hygromycin resistance) with a surviving plo1Δ strain (G418 resistant) in which a mutation was identified in nggl. (D) Experimental evolution (EVO) of plo1Δ strain was performed sequentially, as described in Fig. 1A. After each round, we isolated a few of the fastest growing colonies. We converted plo1Δ to plo1+ and tested if this enhanced the colony growth. Return of Plo1 helped colony growth of a strain obtained after the 1st round of evolution (left, strain #3), whereas a further evolved strain was insensitive to Plo1 (right, strain #3-3). Two glucose concentrations were tested (3% and 0.08%). Wild-type was used as the control.
Figure S2. Nuclear envelope appears to be intact in plo1Δ
(A, B) Live imaging of plo1Δ and cut12-1 strains expressing Pcp1<sup>pericentrin-GFP</sup> and mCherry-tubulin. cut12-1 cells were incubated at 36°C for 3 h before imaging. (C–F) Live imaging of plo1Δ, cut12-1, ght5Δ plo1Δ, and alp4-D440E plo1Δ strains expressing NLS-GFP-β-Gal and mCherry-tubulin. cut12-1 cells were incubated at 36°C for 3 h before imaging. Time 0 (min) was set at the onset of spindle formation.
Figure S3. plo1Δ deletion is sensitive to thiabendazole (TBZ) but does not critically affect Cut7kinetin-5 localisation

(A) In the presence of 10 μg/mL TBZ, plo1Δ failed to form colonies even in low glucose medium. Spores (10,000, 2,000, and 400) were spotted and colonies were formed on 0.08% glucose, G418-containing plates for 5 d at 32°C (expected G418-resistant cell numbers: 5,000, 1,000, and 200 cells, respectively). (B, C) Failure in spindle bipolarisation even when Cut7kinetin-5-GFP accumulated on the SPB at a normal level after a delay in plo1Δ. (D) ght5Δ confers resistance to TBZ, whereas hhp1Δ suppresses the effect. Cells (5,000, 1,000, and 200 cells) spotted and cultured in YE5S medium for 3 d (0 and 15 μg/mL TBZ plates) or 6 d (25 and 30 μg/mL TBZ plates) at 32°C. Time 0 (min) is set at the onset of spindle formation.
Figure S4. Septation defect of plo1Δ is not rescued by ght5 Δ deletion

(A, B) Live imaging of plo1Δ strains expressing Plo1-GFP and mCherry-tubulin. Cells were incubated in the YE5S medium containing 3% or 0.08% glucose. Arrows indicate Plo1-GFP signals at SPBs. Time 0 (min) was set at the onset of spindle formation. (C) Bright field images of wild-type (WT), mid1Δ, and ght5Δ plo1Δ. (D) Quantification of multi-septated cells. A total of 21% of mid1Δ cells and 26% of ght5Δ plo1Δ cells had multiple septations, whereas this was never observed in control cells. (E) Quantification of the septation site. Relative position of the septum along the long axis of the cell was determined.
Figure S5. ght5Δ plo1Δ is synthetic lethal with hhp1Δ, but Hhp1CK1 overexpression alone does not increase the colony growth of plo1Δ.

(A–D) Confirmation of synthetic lethality between the indicated strains by random spore spreading followed by replica-plating onto drug-containing plates. The bub1 and hhp1 ORFs were replaced with ClonNAT-resistant or BSD-resistant cassettes. The plo1 ORF was replaced with a G418-resistant cassette. ght5, git1, and pka1 were replaced with a hygromycin-resistant cassette. Colony formation of heterozygous diploids (red arrowheads) indicates the functionality of the plate. (E) Relative hhp1 mRNA levels in wild-type (WT), ght5Δ, and pka1Δ. Data from two independent experiments were plotted. (F) Investigation of the rescue ability by Hhp1 overexpression in the indicated strains. The calmodulin promoter (cam1-pr) (67) and the native promoter of hhp1+ gene were used. Hhp1 was expressed in the plo1Δ, alp4-D440E plo1Δ, and alp6-V664F plo1Δ strains. Cells (10,000, 5,000, and 1,000) were spotted and cultured in YES5 with G418 for 5 d at 32°C.