

# Probing Yeast Polarity with Acute, Reversible, Optogenetic Inhibition of Protein Function

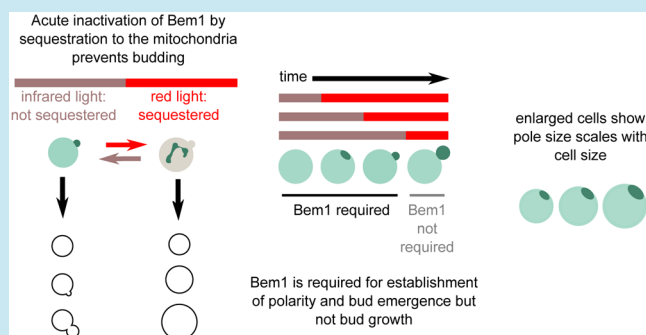
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## Supporting Information

**ABSTRACT:** We recently developed a technique for rapidly and reversibly inhibiting protein function through light-inducible sequestration of proteins away from their normal sites of action. Here, we adapt this method for inducible inactivation of Bem1, a scaffold protein involved in budding yeast polarity. We find that acute inhibition of Bem1 produces profound defects in cell polarization and cell viability that are not observed in *bem1Δ*. By disrupting Bem1 activity at specific points in the cell cycle, we demonstrate that Bem1 is essential for the establishment of polarity and bud emergence but is dispensable for the growth of an emerged bud. By taking advantage of the reversibility of Bem1 inactivation, we show that pole size scales with cell size, and that this scaling is dependent on the actin cytoskeleton. Our experiments reveal how rapid reversible inactivation of protein function complements traditional genetic approaches. This strategy should be widely applicable to other biological contexts.

**KEYWORDS:** cell polarity, budding yeast, Bem1, optogenetics, cell size



Polarization of cell signaling and growth is essential for a wide variety of cell types. Yeast cells must restrict their growth to a small region of the cell cortex in order to bud or mate. This process of polarization is fast, occurring over a period of about 30 min. In budding yeast, cell polarity is orchestrated by the small membrane-bound GTPase Cdc42. Active Cdc42 polarizes at the nascent bud site, where it directs assembly of actin cables that serve as tracks for polarized secretion of the machinery that builds the bud.<sup>1</sup> Cdc42 activation is both necessary<sup>2</sup> and sufficient<sup>2,3</sup> to specify polarity. Cdc42 activation is mediated by its guanine nucleotide exchange factor (GEF), Cdc24, and other regulatory proteins such as Bem1.<sup>4</sup>

Because polarization in yeast is fast, a full understanding of the process requires techniques for observation and perturbation that operate on a similarly rapid time scale. Traditionally, polarity signaling has been studied using loss-of-function experiments with slow, irreversible techniques like genetic knockouts or mutations. While these techniques are powerful, they have several limitations. First, constitutive genetic knockouts cannot be used to study essential genes. Second, it is impossible to use these constitutive perturbations to determine *when* a protein acts during a dynamic process. Third, cells are able to compensate for loss of a gene of interest over long time scales, either by upregulating related genes or paralogs<sup>5</sup> or through other processes like post-translational modifications or aneuploidy.<sup>6</sup> It takes many cell generation

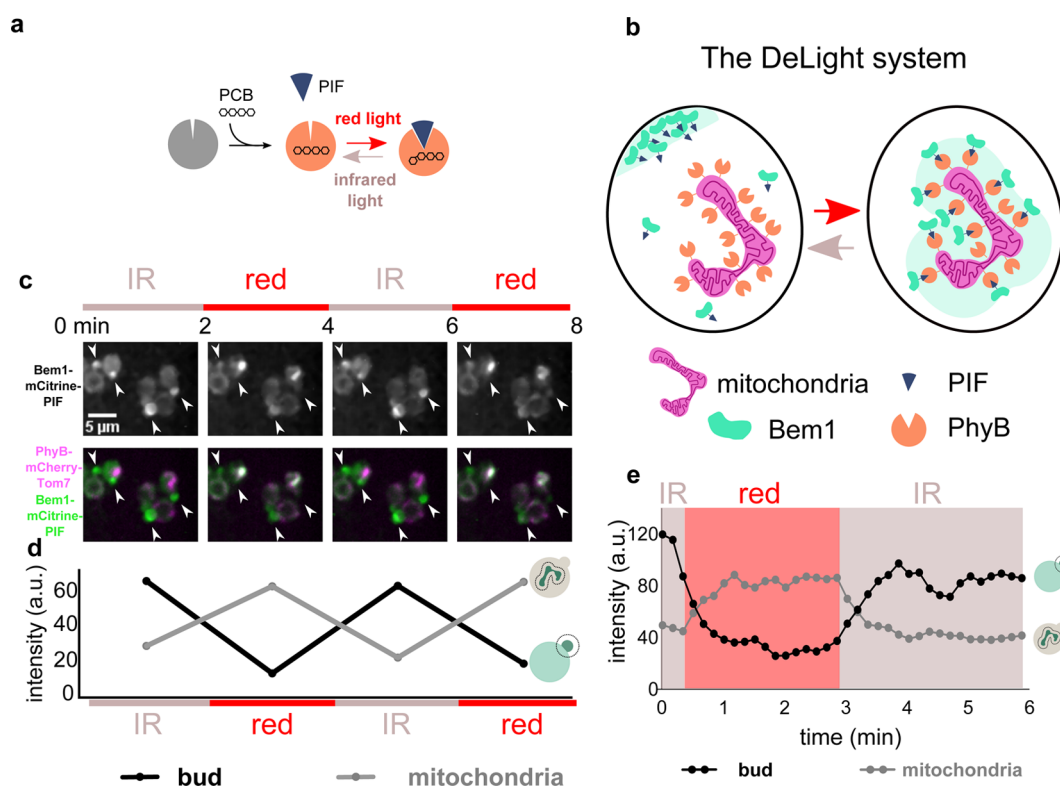
times to make a knockout strain, giving ample time for these adaptations to occur.

We have developed an acute, rapid, and reversible optogenetic technique<sup>7</sup> in order to overcome these limitations and study dynamic processes like yeast polarity. Our technique sequesters protein away from its normal site of action by trapping the soluble pool, building on previous work with light- or chemical-gated dimerizer techniques.<sup>8–12</sup> Tools like temperature-sensitive (*ts*) mutations<sup>13</sup> and degrons<sup>14–16</sup> can be used to study dynamic processes, but both have drawbacks. Generating *ts* alleles is time-consuming, and may not be possible for all genes. Degrons are significantly slower than our system, operating on a scale of minutes rather than seconds. Degron systems are reversible, but on a much slower time scale, limited by the kinetics of protein synthesis. Our system is beneficial in cases where a *ts* allele is difficult to generate or where extra speed is needed.

In this work, we investigate yeast polarity signaling using our previously developed optogenetic system.<sup>7</sup> In this case, we inactivate the protein by inducibly sequestering it away from its normal site of action, a technique we call depletion with light (DeLight). Using this tool, we find that inactivation of Bem1 by sequestration prevents polarization and budding, resulting in cells that grow isotropically until they burst. Next, we use the DeLight system to sequester Bem1 at different times during the

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**Figure 1.** The DeLight system can be used to inducibly sequester Bem1. (a) PhyB is a photoreceptor that interacts with a small molecule chromophore, PCB. PhyB interacts with phytochrome interacting factor (PIF) in the presence of red light and dissociates in the presence of infrared light. (b) By anchoring PhyB at the mitochondria of budding yeast, a PIF-tagged protein can be rapidly and reversibly sequestered away from its normal site of action using red and infrared light. We have named this approach depletion with light, or DeLight. (c) Images of PhyB-mCherry-Tom7 (anchor) and Bem1-mCititrine-PIF (polarity regulator) strains treated with PCB and exposed to red and infrared light, alternating on a 2 min interval; corresponds to Movie S1. White arrowheads indicate small buds or poles. Fluorescence images are a maximum intensity projection of z-stacks. Color bars indicate periods of exposure to red light (Bem1 sequestered at mitochondria) and infrared light (Bem1 released from mitochondria). (d) Intensity of Bem1-mCititrine-PIF measured at the bud *versus* mitochondria during red and infrared light exposure. (e) Timecourse of Bem1 sequestration and release, measured every 10 s.

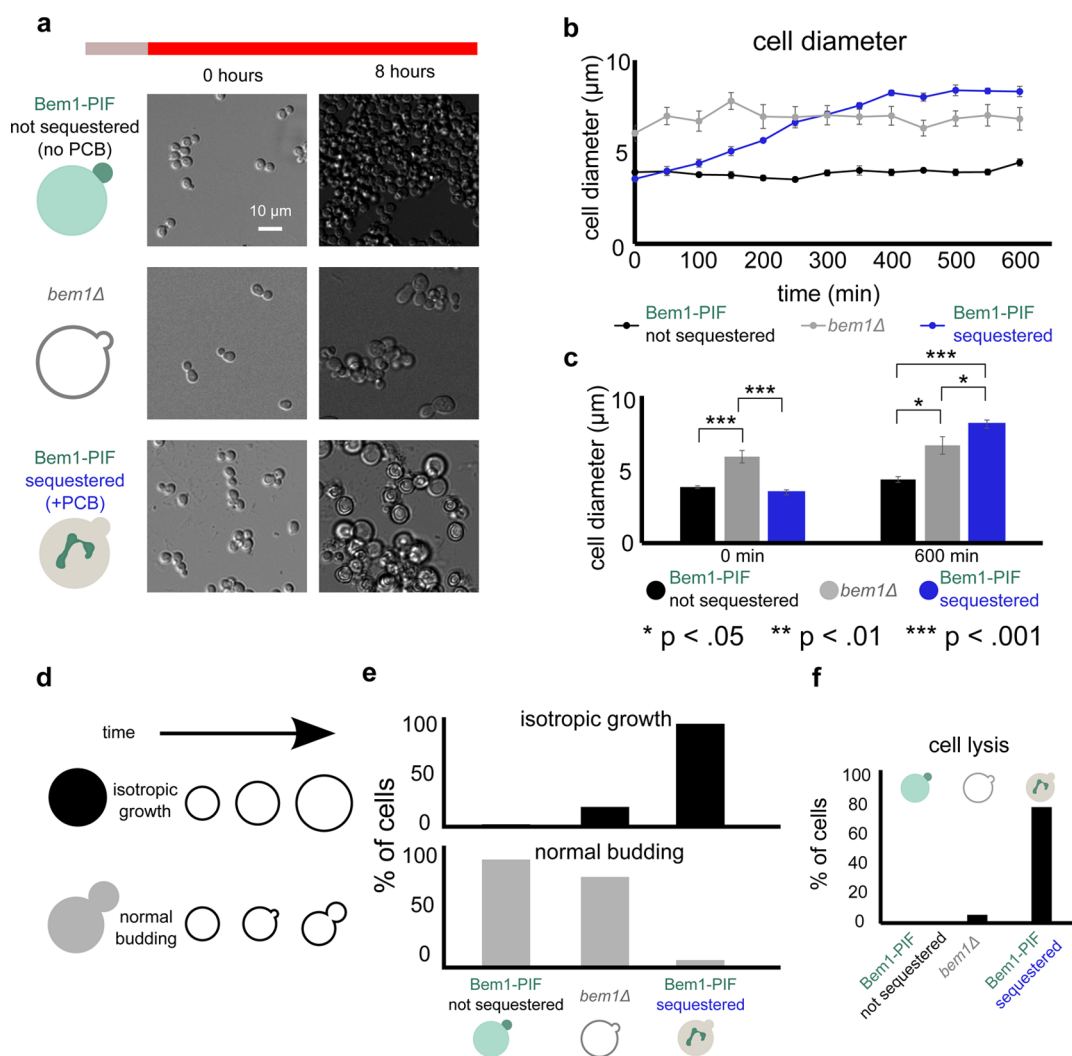
polarization process and find that Bem1 is required for polarization and bud emergence but not for bud growth. Finally, we use the enlarged cells generated by Bem1 sequestration to demonstrate that pole size scales with cell size.

The DeLight system consists of two protein components, Phytochrome B (PhyB) and phytochrome interacting factor (PIF).<sup>17,18</sup> The light-sensitive protein PhyB is localized to a subcellular compartment where it can act as a light-regulated “anchor” for proteins fused to its binding partner, PIF. PhyB is only light responsive when bound to the small molecule chromophore phycocyanobilin (PCB), which can be delivered exogenously to yeast cells<sup>19</sup> (Figure 1a). When cells containing both protein components and PCB are exposed to red light, PhyB changes conformation, and the PIF-tagged protein is recruited to the anchor and depleted from the cytosol and plasma membrane. This interaction can be reversed with infrared light (Figure 1b).<sup>7</sup>

In budding yeast, several putative feedback loops modulate Cdc42 activation at the nascent bud site, including a positive feedback loop centering on the scaffold protein Bem1.<sup>20–22</sup> Bem1 binds to both Cdc24 (the GEF for Cdc42) as well as Cla4 (a p21 activated kinase), forming a trimeric complex. Cla4 and Bem1 can both bind to active Cdc42,<sup>23</sup> enriching the Bem1-GEF complex at sites of Cdc42 activity. This increased local GEF recruitment activates more Cdc42, which in turn recruits more Bem1 complex, forming a positive feedback loop

that amplifies Cdc42 signal. While this feedback loop appears to be important in activating Cdc42, the *bem1Δ* strain is viable, with relatively mild polarity and growth defects, suggesting that other parallel pathways may be able to compensate for loss of Bem1. Much of the argument for the importance of Bem1 stems from the observation that *bem1Δ* is synthetic lethal with *rsr1Δ*, a landmark protein involved in directing the location of the bud site, indicating that the Bem1-dependent positive feedback loop may be more important when cells must break symmetry.<sup>23</sup> However, recent evidence indicates that the *bem1Δ rsr1Δ* double mutant is viable in some genetic backgrounds.<sup>24</sup> Here we sought to revisit the role of Bem1 in yeast polarity using our DeLight system. We hoped to clarify if and when Bem1 function is required for polarity in a manner that would not be confounded by the compensation that may accompany long-term inactivation of Bem1.

We set out to establish that the DeLight system can be used to inducibly inactivate Bem1. For complete depletion of Bem1, every Bem1 protein in the cell must have a PIF tag. Therefore, we tagged Bem1 with mCititrine and PIF at the endogenous locus and verified that localization was unaffected by the PIF tag. As expected, Bem1-PIF polarizes to the nascent bud site and the cortex of growing buds (Figure S1). Next, we sought a localized PhyB that could efficiently sequester Bem1-PIF from the cytoplasm and the plasma membrane. We initially used a PhyB-mCherry-Htb2 construct from our previous work,<sup>7</sup>



**Figure 2.** Bem1 sequestration with DeLight prevents cells from budding. (a) *bem1Δ* and Bem1 DeLight strains (+PCB, Bem1-sequestered or –PCB, not sequestered) were pretreated with infrared light and then switched to red light to activate sequestration at  $t = 0$  and imaged in DIC for a period of 10 h; see Movie S2. (b) Cell diameter for each condition, measured every 5 min (error bars = standard error of mean, SEM; 10 cells measured for each condition at each time point). (c) Average cell diameters at the beginning and end of the experiment (error bars = SEM, 10 cells measured for each condition at each time point). (d) Schematic of phenotypes for cell scoring. (e) Quantification of movies from (b). Cells were scored for phenotypes based on the schematic in (d). (f) Quantification of cell lysis during the 10-h red light exposure. For panels (e) and (f), >100 cells were scored for each condition. Acute inhibition of Bem1 with DeLight produces more profound defects in budding and cell viability than genetic nulls of Bem1.

reasoning that sequestering Bem1 inside the nucleus would be most likely to remove it from its relevant effectors. However, persistent recruitment of mCitrine-PIF alone to the PhyB-mCherry-Htb2 construct slowed cell growth and altered cell size and morphology, possibly due to misregulation of histone function (Figure S2).

We sought a new PhyB-anchor that would provide efficient sequestration of PIF-tagged proteins without off-target effects. Sequestration at the mitochondria has previously been used to inactivate proteins.<sup>12,25</sup> The mitochondrial outer membrane protein Tom7, a component of the translocase of outer membrane (TOM) complex, has its N terminus facing the cytosol, making it suitable for fusion with PhyB, since PhyB only tolerates C-terminal fusions.<sup>26</sup> Our pADH-PhyB-mCherry-Tom7 construct expressed well, localized to mitochondria, and was well-tolerated by yeast cells (Figure 1c). Unlike the PhyB-Htb2 anchor strains, the PhyB-Tom7 anchor strains were unaffected by constitutive recruitment of mCitrine-PIF (Figure

S2). This result demonstrates that mitochondrial sequestration with PhyB-Tom7 is a good anchor for use with the DeLight system.

To determine whether our mitochondrial anchor was functional in sequestering Bem1-PIF, we added the PhyB-Tom7 construct to Bem1-mCitrine-PIF cells. After incubating cells with PCB to render the PhyB light responsive, we exposed the cells to successive rounds of red and infrared light, acquiring a 3D stack of images of Bem1-PIF between each light exposure. We observed reversible depletion of Bem1 from poles and reversible enrichment of Bem1-PIF at the mitochondria (Figure 1c,d, Movie S1). Both enrichment at the mitochondria and sequestration from the bud are rapid, with a  $t_{1/2}$  of roughly 30 s, and the interaction can be reversed on a similar time scale (Figure 1e).

Deletion of BEM1 generates enlarged cells displaying morphological abnormalities (Figure 2a) that grow more slowly than a wild-type strain.<sup>27</sup> Acute sequestration of Bem1

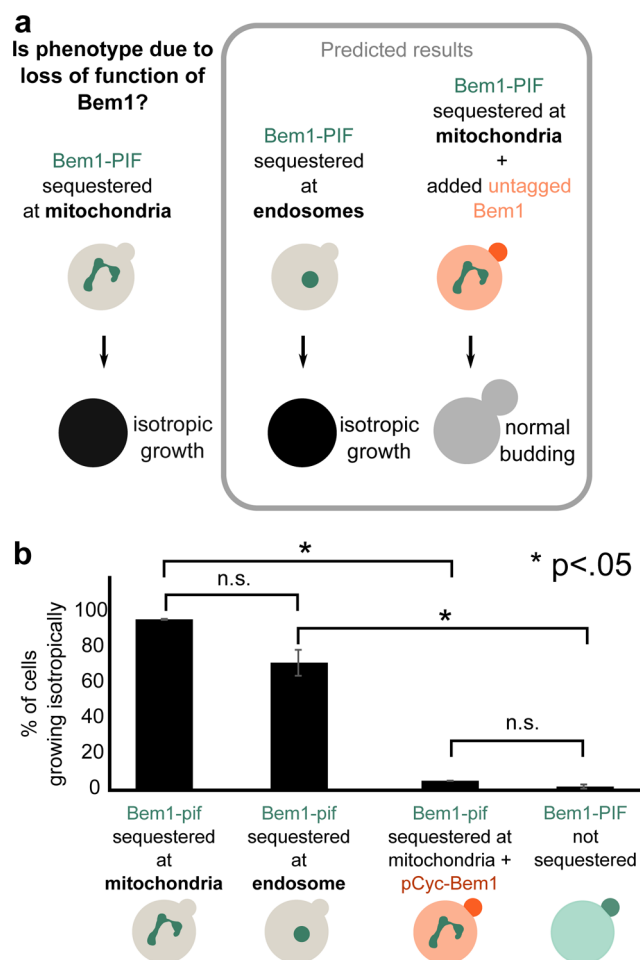
at the mitochondria generates a morphological phenotype that is significantly more profound than *bem1Δ*. Bem1-PIF sequestered cells fail to bud and instead grow isotropically until they are about twice as wide as control cells that have not been incubated with PCB and therefore do not sequester Bem1. These nonsequestered cells are indistinguishable from wild type cells (Figure S3). The Bem1 sequestered cells grow even larger than *bem1Δ* cells (Figure 2b,c, Movie S2). After 10 h of red light exposure, controls without PCB have an average diameter of  $4.4 \pm 0.2 \mu\text{m}$  (mean  $\pm$  SEM), whereas Bem1-PIF depleted cells have an average diameter of  $8.3 \pm 0.3 \mu\text{m}$ . In contrast, *bem1Δ* cells have an average diameter of  $6.8 \pm 0.6 \mu\text{m}$  (Figure 2c). If we assume the cell is a sphere, this corresponds to a 6.6 fold increase in volume for Bem1-PIF sequestered cells relative to control cells, compared to a 3.6 fold increase in volume for *bem1Δ* relative to the control.

95% of Bem1-PIF sequestered cells grow isotropically and do not form new buds after exposure to red light. In contrast, only 17% of *bem1Δ* cells grow isotropically (Figure 2e). In the acutely sequestered Bem1-PIF cells, this isotropic growth phenotype often causes the cells to lyse; 79% of cells lyse by 10 h of sequestration, as opposed to 6% of *bem1Δ* cells and 0% of unperturbed cells (Figure 2f).

In the above experiments, it is possible that the phenotypes observed upon Bem1-PIF recruitment result from a gain-of-function phenotype at the mitochondria rather than loss of Bem1 activity at polarity sites. We performed two sets of experiments to discriminate between these possibilities (Figure 3a). First, we investigated whether the phenotype depended on the intracellular site of Bem1 sequestration. If sequestration induced loss of function, recruitment to other organelles should generate a similar phenotype. As an additional test, we added an additional untagged (nonsequesterable) copy of Bem1. This should rescue the budding phenotype if it is due to loss of function. In contrast, if the budding phenotype reflects gain of function at the mitochondria, it would not be replicated by sequestering Bem1 to other anchors and would not be rescued by adding an additional wildtype copy of Bem1.

To investigate the consequences of Bem1 sequestration to a different organelle, we recruited Bem1 to the endosome using PhyB-mCherry-Snf7. This anchor, developed in our previous work, efficiently sequesters PIF-tagged protein but is not fully reversible, making it less suitable for more complex time-varying experiments, but appropriate for comparison with PhyB-mCherry-Tom7 in this case.<sup>7</sup> Sequestration to the endosome via PhyB-Snf7 produced a budding phenotype similar to the one generated by sequestration of Bem1 to the mitochondria via PhyB-Tom7. As with mitochondrial sequestration of Bem1, the majority of cells (70%) did not form new buds after sequestering Bem1 at the endosome and instead grew isotropically until they burst (Figure 3b, Movie S3). The phenotypes are qualitatively similar, but the magnitude of the defect is somewhat smaller for Snf7 sequestration of Bem1-PIF, possibly due to differing expression levels or accessibility to PIF between Snf7 and Tom7.

To further validate that Bem1 sequestration to the mitochondria causes a loss of protein function, we tested the ability of an additional untagged copy of Bem1 to rescue the phenotype. We performed Bem1-PIF sequestration to the mitochondria in the presence of an additional untagged copy of Bem1 driven by the pCyc promoter. The nonsequesterable copy of Bem1 rescued the phenotype, with only 4% of cells growing isotropically, compared to 95% for cells in which



**Figure 3.** The Bem1 sequestration phenotype is due to loss of function of Bem1. (a) The phenotype generated by sequestration of Bem1-PIF to the mitochondria could arise in two ways: gain of function of Bem1 at the mitochondria or loss of function due to sequestration away from its normal site of action. These two models can be distinguished by recruiting Bem1 to another organelle (endosome) and by adding a nonsequesterable copy of Bem1. Predicted results are shown for Bem1 loss of function (gain of function predicts opposite phenotypes). (b) Quantification of phenotypes from movies (Movie S3). >100 cells were scored for each condition. The phenotypes are consistent with mitochondrial sequestration inducing acute loss of Bem1 function.

sequestration was active and in which all copies of Bem1 had the PIF tag (Figure 3b, Movie S3). The rescued cells grew at a similar rate to unperturbed cells, with an average time between buds of  $112 \pm 6$  min for the rescued cells and  $108 \pm 2$  min for the unperturbed cells. In contrast, the *bem1Δ* cells had a time between buds of  $133 \pm 18$  min.

While these results argue that the Bem1 sequestration phenotype is likely to be due to loss of function of Bem1, it could also be due to sequestration of other Bem1 binding partners. For example, Cdc24, the Cdc42 GEF, binds Bem1 and could be sequestered along with Bem1 at the mitochondria. In this case, we would expect an additional copy of Cdc24 to rescue the phenotype, similar to the rescue we observe for adding additional wildtype Bem1 (Figure 3b). However, adding an additional copy of Cdc24 fails to rescue the growth and polarity phenotypes; these cells behave similarly to the Bem1 sequestered cells without added Cdc24 (Figure S4).



Taken together, our observations that (1) the Bem1 sequestration phenotype does not depend on the site of sequestration and (2) the Bem1 sequestration phenotype can be rescued by nonsequesterable Bem1 argue that the observed phenotype is due to acute loss of function of Bem1 rather than gain of function at the anchor location. Because this acute phenotype is stronger than the *bem1Δ* phenotype, our data suggest that the *bem1Δ* strain may have undergone compensation, either by genetic changes or post-transcriptional regulation. Similar isotropic growth phenotypes have been observed in cases where cells cannot polarize secretion, specifically with temperature-sensitive alleles of myosin, tropomyosin, and formins.<sup>28,29</sup>

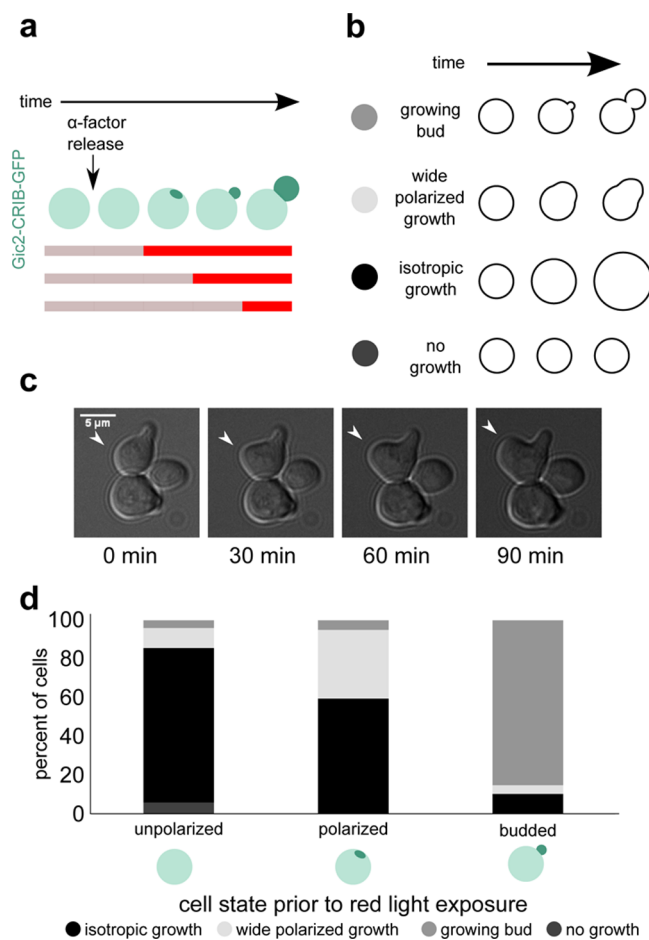
Acute sequestration provides a platform to examine the effects of inactivating Bem1 before compensation can occur and may help to clarify conflicting data regarding the role of Bem1. We find that Bem1 is essential for polarity, not just redundant with other components. Previous work indicated that Bem1 was only essential in a symmetry-breaking background, where the landmark protein Rsr1 was deleted.<sup>23</sup> Our results indicate that Bem1 is crucial even when cells still contain landmark proteins.

Next, we used the temporal control of DeLight to investigate when Bem1 is required for polarization of cell growth. Cells were arrested with  $\alpha$ -factor to generate a synchronized population of unpolarized cells. Consistent with previous reports,<sup>30</sup> most cells polarize the Cdc42 activity reporter Gic2-CRIB within an hour of release by  $\alpha$ -factor washout. (Figure S5). Cells were exposed to infrared light (no sequestration) until 45, 60, or 75 min after release, then switched to red light to sequester Bem1 to the mitochondria (Figure 4a). Cells were scored for polarity by imaging the Gic2-CRIB activity reporter prior to switching to red light. Unpolarized cells were strongly affected by Bem1-PIF sequestration; 80% of cells that were unpolarized when switched to red light did not subsequently develop polarity and instead grew isotropically (Figure 4d). Cells that were polarized but had not yet formed a bud were also affected, with 60% of cells growing isotropically following Bem1 sequestration. However, cells that had undergone bud emergence before Bem1-PIF sequestration were largely unaffected, with only 11% of cells growing isotropically.

In addition to the isotropic growth phenotype, we also observed cells that showed polarized growth but with abnormal morphology; the region of growth was much wider than a normal bud, and lacked the constriction seen in a bud neck (Figure 4c). 11% of cells that were unpolarized at the time of Bem1-PIF sequestration had the wide polarized growth phenotype. In cells that were already polarized but had not yet budded, 36% percent of cells showed wide polarized growth. In cells that had already formed a bud, the occurrence of wide polarized growth was much lower, with only 4.5% of cells showing this phenotype (Figure 4d).

From these data, we conclude that Bem1 is required for the establishment of polarity as well as initiation of bud growth. In contrast, Bem1 is largely dispensable for bud growth once the bud has emerged.

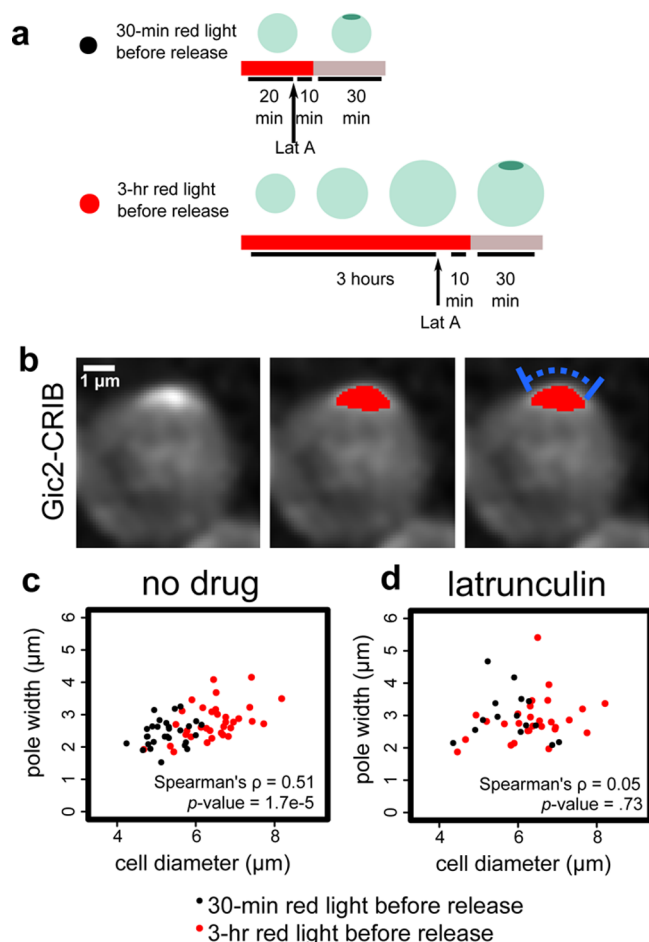
Because acute loss of Bem1 yields cells that grow isotropically and attain much larger size than control cells, we reasoned that we could use these large cells to investigate whether the size of the pole scales with the size of the cell. The pole is shaped by diffusion of activated Cdc42, active transport to the pole, and endocytosis.<sup>31</sup> Many mutations generate



**Figure 4.** Bem1 localization is required for polarization and bud emergence but not bud growth. Here we use temporal control of DeLight to determine when Bem1 function is required for budding. (a) Schematic of experimental setup. The colored bar indicates periods of red and infrared light exposure. (b) Phenotypes for cell scoring. (c) Example of wide polarized growth phenotype. (d) Percent of cells that exhibit each phenotype, binned by polarization state at the time of red light exposure. Polarization state is scored using the Gic2-CRIB reporter. Bem1 is required in the early phase of the polarization process (pole formation and bud emergence) but is dispensable for bud growth following emergence.

enlarged yeast cells, including Bem1; *bem1Δ* cells are in the largest 5% of mutants in the yeast whole-genome deletion collection,<sup>32</sup> and pole size could be measured in these backgrounds with persistently altered cell size. However, the reversibility of our system provides a major advantage: we use depletion of Bem1-PIF to generate large cells, but once they are moved to infrared light, they recover wild-type Bem1 activity and can polarize (Figure S6). This allows us to measure pole size in isogenic cells that are large or small. In addition, cells incubated in red light are large on average, but significant variation exists in the population, allowing comparison between large and small cells that have been treated identically.

To generate enlarged cells, Bem1-PIF PhyB-Tom7 cells with the Gic2-CRIB reporter were exposed to red light for 3 h, which is sufficiently long to significantly enlarge the cells but not cause cell lysis (Figure 5a). After switching to infrared light, most cells polarized within 5 min, confirming reversibility (Figure S6). As a control for the Bem1 sequestration and release, Bem1-PIF PhyB-Tom7 cells were exposed to red light



**Figure 5.** Pole width scales with cell size in an actin-dependent manner. We use the reversibility of DeLight to investigate how pole size scales with cell size. (a) We treat Bem1 DeLight cells with red light for 3 h to generate enlarged cells, or 30 min to retain wild-type cell size, and then release Bem1 to allow cells to polarize. (b) Example of pole measurement. Image shown is a maximum intensity projection of a z-stack of CRIB-GFP, a Cdc42 activity reporter. Red overlay is the thresholded pole. Blue bar indicates pole width measurement. (c) and (d) Pole width *vs* cell diameter with no drug (c) *versus* latrunculin, an actin depolymerizing drug (d). Correlation is indicated by the Spearman's  $\rho$  value. Black points are measurements from 30 min cells and red points measurements from 3-h cells (enlarged).  $n = 63$  and  $n = 44$  for no drug and latrunculin, respectively.

for 30 min, not long enough to significantly alter cell size, and then switched to infrared light. In this context, cells were not enlarged, and most cells polarized within 5 min after switching to infrared light.

We measured pole width in each cell population by tracing the perimeter of the cell from end to end of the thresholded pole, based on the Gic2-CRIB reporter (Figure 5b). Particularly in Latrunculin A treated cells, and to a lesser degree in wildtype cells, the poles are quite dynamic, and it is not always clear which time point represents a stable pole to measure. Therefore, for each cell, we selected the time point with the brightest pole intensity for width measurement. We tested other time point selection criteria and found that each method generated similar results (Figure S7). In the pooled population of enlarged and normal-sized cells, the width of the pole is correlated with cell diameter (Spearman's  $\rho = 0.51$ ,  $p$ -value  $< 0.001$ ). Importantly, for cells with similar diameters, there is

no significant difference in pole size between the two red light durations ( $t$  test,  $p$ -value = 0.97, cells between 5.5 and 6  $\mu\text{m}$  diameter). Since pole width is thought to be governed by active transport and endocytosis at the pole,<sup>31</sup> we reasoned that pole width scaling with cell size would require actin polymer. Indeed, in cells treated with Latrunculin A to depolymerize actin, the correlation between cell diameter and pole size is lost (Spearman's  $\rho = 0.05$ ,  $p$ -value = 0.73) (Figure 5c). The enlarged cells generated by our light-gated perturbation were essential to observing pole width scaling; there is not enough endogenous variation in cell size to detect a correlation (Spearman's  $\rho = 0.31$ ,  $p$ -value = 0.10).

In summary, we have refined the DeLight system, generating a new anchor for protein sequestration, and applied this system to inactivate Bem1 rapidly and reversibly. Acute sequestration of Bem1 prevents budding, and generates a phenotype significantly stronger than genetic deletion of Bem1. We used the DeLight system's speed to probe the temporal regulation of polarity by Bem1 and found that Bem1 is necessary for polarization and bud emergence, but not for bud growth. Finally, we used the reversibility of the DeLight system to uncover pole size scaling with cell size.

While significant previous work has established the importance of the signaling surrounding Bem1 in yeast polarity, the viability of the *bem1* $\Delta$  strain indicated that Bem1 was not essential for polarization unless it was combined with mutations in other polarity components such as Rsr1. Here, we show that acute depletion of Bem1 prevents cells from polarizing, suggesting that Bem1 is essential for polarization under these conditions. Compensation over the relatively slow timescale of generating the *bem1* $\Delta$  mutant likely enables survival of the knockout strain. Our acute perturbation permits analysis of the cells before compensation can occur.

We show that the requirement for cytoplasmic Bem1 is restricted to the polarization and bud emergence phases and that sequestering Bem1 once the bud has emerged does not prevent further bud growth. This result could extend our current understanding of cell cycle regulation of bud growth. Cdc42 is activated in late G1 and must polarize in order for the cell to form a bud. It continues to polarize through bud emergence until G2, when growth in the bud switches from polarized to isotropic. In G2, Bem1 and Cdc24 lose their localization to the bud tip, and active Cdc42 disperses to the entire bud cortex, suggesting that polarization of these signaling components is important for polarized growth but not for isotropic growth of the bud.<sup>33</sup> This transition is known as the apical-isotropic switch. Importantly, previous experiments have primarily relied on observation of protein localization, rather than perturbation of protein function during specific phases of this multistage process. Techniques for assaying necessity of Bem1 and other polarity proteins before and after the apical-isotropic switch have been lacking. Because we do not use any cell cycle reporters, our results only establish that Bem1 localization is necessary before but not after bud emergence, but lack the time resolution to further clarify whether Bem1 specifically acts at the apical-isotropic switch. In future work, our system could be used in conjunction with cell cycle reporters or mutants to determine whether the change in Bem1 necessity occurs concurrently with the switch to isotropic growth.

Our finding that pole size scales with cell size raises several intriguing questions. What is the mechanism by which cells scale their pole size? Does increased pole size lead to increased

daughter cell size, and if so, for how many generations does this altered cell size persist? More generally, the inducibly isotropically growing cells generated here could be used in other analyses of scaling in budding yeast.

Broadly, we have demonstrated the application of the light-gated sequestration system to a dynamic and rapid cellular process, cell polarity. The DeLight system is generally applicable to a wide range of signaling cascades. Previously, we have applied the system to the mitotic cyclin Clb2<sup>7</sup>, and we envision that it can be used to probe the function of many other proteins in dynamic processes ranging from cell cycle regulation to yeast polarity and beyond.

## MATERIALS AND METHODS

### Yeast Strain Construction and Growth Conditions.

Standard yeast strain construction methods were used throughout. All strains were constructed in congenic w303 (MATa *his3-11,15 trp1-1 leu2-3 ura3-1 ade2-1*). PhyB-mCherry-Snf7 and PhyB-mCherry-Htb2 were constructed in our previous work.<sup>7</sup> To select a new anchor for light-induced protein sequestration, we used the yeast GFP localization collection (yeastgfp.yeastgenome.org)<sup>34</sup> to identify proteins that localized to the mitochondria and were expressed at high levels. This candidate list of proteins was narrowed by excluding those with overexpression phenotypes, since the PhyB-anchor constructs are expressed on top of the endogenous protein. Finally, we reviewed sequence and topology information for each protein to identify proteins for which the N-terminus was facing the cytosol, because PhyB only tolerates fusions on its C-terminus. From this list, Tom7 was the most favorable candidate. Tom7 was cloned from the genome and inserted in the PhyB-mCherry backbone using Gibson enzymatic assembly.<sup>35</sup> Endogenous Bem1 was tagged at the C-terminus with mCitrine and PIF using an optimized PCR-based tagging cassette<sup>36</sup> modified with a PIF addition. The pCyc-Gic2-CRIB-2xGFP (KAN) plasmid was a gift from Jessica Walter. Additional copies of pCyc-Bem1 and pCyc-Cdc24 were added using integrating plasmids at the TRP locus.

For all time-lapse microscopy experiments, cells growing exponentially in synthetic defined complete medium (Sunrise) were vortexed and plated on Concanavalin A coated coverslip-bottom 384-well plates (Brooks), which were then washed to remove floating cells. For optogenetic experiments, cells were incubated with a final concentration of 31.25  $\mu$ M (0.25  $\mu$ L of 12.5 mM stock in 100  $\mu$ L culture) PCB (purified ourselves<sup>37,38</sup> or purchased from Santa Cruz Biotechnology, Inc.) for 2–4 h, then pelleted and resuspended in fresh media without PCB before plating. For  $\alpha$ -factor synchronization and release experiments, cells were incubated with 100  $\mu$ M  $\alpha$ -factor (Zymo) for 90 min with or without PCB, washed, and then released into fresh synthetic medium.

**Microscopy and Optogenetic Control.** The experiments in Figures 1, 4, and 5 were performed on two spinning disk confocal microscopes, both based on a Nikon Eclipse Ti inverted microscope with a motorized stage (ASI). The first microscope used a Diskovery spinning disk confocal equipped with a 100  $\mu$ m pinhole disk (Spectral Applied Research), 405, 440, 488, 514, and 561 nm laser wavelengths (LMMS, Spectral Applied Research), a 60x 1.49NA TIRF Apo objective (Nikon), and a Zyla sCMOS camera (Andor). The second microscope used a Yokogawa CSU-X1 spinning disk confocal, an MLC400B monolithic laser combiner with laser lines at 405,

488, 561, and 640 nm (Agilent), a 60x 1.4NA Plan Apo VC objective (Nikon), and a Clara interline CCD (Andor).

The experiments in Figures 2 and 3 were performed on a Nikon Eclipse Ti inverted microscope equipped with a Lambda XL Broad Spectrum Light Source (Sutter), a 20x 0.75NA Plan Apo objective (Nikon) or a 60x 1.4NA Plan Apo VC objective (Nikon), FITC and TRITC filter cubes (Chroma), and a Clara interline CCD (Andor). A motorized stage (ASI) allowed automated sampling of multiple XY positions.

For all experiments, microscope settings, dichroic positions, filters, shutters, and cameras were controlled by NIS Elements (Nikon).

For optogenetic experiments, red and infrared light were provided by placing filters on top of the condenser in the brightfield lightpath<sup>18</sup> (650 nm 20 nm-bandpass filter, Edmund Optics, near-IR RG9 glass filter, Newport). To provide constant illumination of the sample, the transmitted light shutter was left continuously open during imaging. In order to illuminate multiple wells of a 384-well black glass-bottom imaging plate (Brooks), the condenser was lowered as close as possible to the plate, and the field and aperture diaphragms were opened fully. In this configuration, nine wells received uniform simultaneous illumination, allowing imaging of multiple strains and controls in the same experiment.

Fluorescence images were computationally denoised in collaboration with John Sedat (UCSF), using an algorithm built into the Priism image analysis package.<sup>39</sup>

All image analysis was performed in Fiji.<sup>40</sup> Intensity measurements in Figure 1 were performed on background-subtracted images using manually drawn ROIs. The Cell Counter (Kurt De Vos, University of Sheffield) plugin was used for manual scoring and counting of cell types. Pole widths were measured in Fiji. First, the image background was subtracted using Fiji's built-in rolling-ball background subtraction algorithm. Next, cytoplasmic background was measured in a hand-drawn ROI and then subtracted from the whole image. The maximum pixel intensity was measured and the image was thresholded using 25% of the maximum intensity value. Then, the arc around the cell periphery from end to end of the pole was measured using the curved line tool. The time point for with the highest maximum intensity value was used for each cell. This process was semiautomated with a custom-written macro.

Data analysis, statistical analysis, and graphing were performed in Microsoft Excel and R ([www.R-project.org/](http://www.R-project.org/)).<sup>41</sup>

## ASSOCIATED CONTENT

### Supporting Information

Figures S1–S7 and Movies S1–S3 are available as described in the text. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acssynbio.5b00053.

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### Notes

The authors declare no competing financial interest.

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