- 1 AMP-activated protein kinase and adenylate kinase prevent the ATP catastrophe and
- 2 cytotoxic protein aggregation
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ABSTRACT

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ATP is the main source of chemical energy in all life and is maintained at several millimolar in eukaryotic cells. However, the mechanisms responsible for and physiological relevance of this high and stable concentration of ATP remain unclear. We herein demonstrate that AMP-activated protein kinase (AMPK) and adenylate kinase (ADK) cooperate to maintain cellular ATP levels regardless of glucose concentrations. Single cell imaging of ATP-reduced yeast mutants revealed that ATP concentrations in these mutants underwent stochastic and transient depletion of ATP repeatedly, which induced the cytotoxic aggregation of endogenous proteins and pathogenic proteins, such as huntingtin and α-synuclein. Moreover, pharmacological elevations in ATP levels in an ATP-reduced mutant prevented the accumulation of α-synuclein aggregates and its cytotoxicity. The removal of cytotoxic aggregates depended on proteasomes, and proteasomal activity cooperated with AMPK or ADK to resist proteotoxic stresses. The present results provide the first evidence to show that cellular ATP homeostasis ensures proteostasis and revealed that stochastic fluctuations in cellular ATP concentrations contribute to cytotoxic protein aggregation, implying that AMPK and ADK are important factors that prevent proteinopathies, such as neurodegenerative diseases.

INTRODUCTION

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Adenosine triphosphate (ATP) is the main energy currency used by all living organisms. Due to high demand, the turnover rate of ATP is estimated to be less than one minute in yeast and metazoans (Mortensen, Thaning et al., 2011, Takaine, Ueno et al., 2019). In addition to its role as energy currency, recent studies reported that ATP may influence the balance between the soluble and aggregated states of proteins, suggesting that proteostasis is maintained by energy-dependent chaperones and also by the property of ATP as a hydrotrope to solubilize proteins (Hayes, Peuchen et al., 2018, Patel, Malinovska et al., 2017). Recent proteomic analyses demonstrated that many cellular proteins are insoluble at submillimolar ATP concentrations and become soluble at ATP concentrations > 2 mM (Sridharan, Kurzawa et al., 2019); however, it currently remains unclear whether ATP-dependent protein solubilization/desolubilization play any significant roles in cell physiology. We recently established a reliable imaging technique to quantify intracellular ATP concentrations in single living yeast cells using the genetically encoded fluorescent ATP biosensor QUEEN (Yaginuma, Kawai et al., 2014), which enables long-term evaluations of ATP homeostasis in individual cells (Takaine et al., 2019). Our findings demonstrated that intracellular ATP concentrations did not vary within a yeast population grown in the same culture (Takaine et al., 2019), which was in contrast to the large variations observed in intracellular ATP concentrations within a bacterial cell population (Yaginuma et al., 2014). Moreover, intracellular ATP concentrations in individual living yeast cells were stably and robustly maintained at approximately 4 mM, irrespective of carbon sources and cell cycle stages, and temporal fluctuations in intracellular ATP concentrations were small (Takaine et al., 2019). Based on these findings, we hypothesized that an exceptionally robust mechanism may exist to precisely regulate ATP concentrations in eukaryotes. It also remains unclear why ATP is stably maintained at a markedly higher concentration than the K_m (Michaelis constant) required for the enzymatic activity of almost all ATPases (Edelman, Blumenthal et al., 1987), and the consequences associated with failed ATP homeostasis in living organisms have not yet been elucidated. The most promising candidate regulator of ATP homeostasis is AMP-activated protein kinase (AMPK). AMPK, which is activated by AMP and inhibited by ATP (Xiao, Heath et al., 2007), has long been regarded as an important regulator of the whole-body and cellular energy status in eukaryotes (Hardie, Schaffer et al., 2016). AMPK is activated by increases in the

AMP:ATP or ADP:ATP ratio (i.e., low-energy state), and regulates its downstream effectors

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by phosphorylation to redirect cell metabolism from an anabolic (ATP-consuming) state to catabolic (ATP-producing) state (Herzig & Shaw, 2017). In the budding yeast Saccharomyces cerevisiae, the sucrose non-fermenting 1 (Snf1) protein kinase complex is the sole AMPK. Similar to other AMPKs, the yeast Snf1 complex comprises three subunits: the catalytic α subunit (SNF1), scaffolding β subunit (SIP1, SIP2 or GAL83), and regulatory γ subunit (SNF4) (Ghillebert, Swinnen et al., 2011). The role of the Snf1 complex in adaptation to glucose limitations has been characterized in detail (Hedbacker & Carlson, 2008). In the presence of sufficient glucose in the medium, the Snf1 complex is inactive (Wilson, Hawley et al., 1996). When glucose concentrations decrease, the Snf1 complex is activated and phosphorylates the transcriptional repressor Mig1, which then triggers the transcription of numerous glucose-repressed genes (Carlson, 1999). However, the contribution of AMPK or the Snf1 complex to cellular ATP levels remains unknown. Other possible candidate regulators of ATP homeostasis include genes whose mutation leads to decreases in the cellular content of ATP. However, based on biochemical analyses of cell populations, few yeast mutants reduced ATP levels (Gauthier, Coulpier et al., 2008, Ljungdahl & Daignan-Fornier, 2012). Adenylate kinase (ADK) is a key enzyme that synthesizes ATP and AMP using two ADP molecules as substrates, and the null-mutant of ADK (adk1 Δ) was shown to have a reduced cellular ATP concentration (~70% of the wild type) (Gauthier et al., 2008). Bas1 is a transcription factor that is required for de novo purine synthesis and $bas 1\Delta$ has also a reduced ATP concentration (~50% of the wild type) (Gauthier et al., 2008). However, the regulation of ATP concentrations and the physiological consequences of reduced ATP levels in these mutants remain unclear, particularly at the single cell level. In the present study, we investigated the roles of AMPK, ADK, and Bas1 in ATP homeostasis using the QUEEN-based single cell ATP imaging technique. We demonstrated for the first time that AMPK is involved in the regulation of intracellular ATP concentrations, even under glucose-rich conditions. Furthermore, time-lapse ATP imaging revealed that cells lacking both AMPK and ADK frequently underwent transient ATP depletion, while ATP concentrations oscillated in those lacking Bas1. These ATP dynamics in the mutants were overlooked in previous biochemical studies. We found that some intrinsic proteins and aggregation-prone model proteins, including α -synuclein, which is responsible for Parkinson's disease, aggregated and were cytotoxic in all of the ATP-reduced mutants tested. The present results suggest that the stable maintenance of ATP is essential for proteostasis and imply that

an ATP crisis promotes proteinopathies, such as neurodegenerative diseases.

112 **RESULTS** 113 AMPK is involved in the maintenance of cellular ATP levels 114 We recently developed a reliable monitoring system for cytoplasmic ATP concentrations in 115 living yeast cells using the ATP biosensor QUEEN (Takaine et al., 2019). We conducted a 116 more detailed examination of ATP dynamics in wild-type and mutant yeast cells using this 117 system in the present study. We investigated whether the deletion of SNF1, which encodes a catalytic subunit of AMPK, affects cellular ATP concentrations. In SNF1-null mutant $(snf1\Delta)$ 118 119 cells, cellular ATP levels were significantly lower than those in wild-type cells at all glucose concentrations tested (2, 0.2, and 0.05%) (Fig. 1A, B). It is important to note that even under 120 121 high glucose conditions (2% glucose), in which the Snf1 complex is considered to be inactive, 122 $snfl\Delta$ cells showed significantly reduced ATP levels. Moreover, the deletion of each gene 123 encoding the β subunit of AMPK (SIP1, SIP2, or GAL83) reduced cellular ATP levels to a 124 similar extent as the deletion of SNF1, suggesting that three subtypes of the Snf1 complex are 125 involved in maintaining ATP concentrations (Fig. 1C). On the other hand, the deletion of 126 MIG1, which encodes a major downstream target of the Snf1 complex in glucose repression, 127 had a negligible effect on ATP levels (Fig. 1D), suggesting that unknown factors other than 128 Mig1 primarily regulate ATP levels under the control of the Snf1 complex. Collectively, these 129 results demonstrated for the first time that AMPK/SNF1 affect cellular ATP concentrations 130 even under glucose-rich conditions. 131 132 Adenylate kinase ADK1 cooperates with AMPK to regulate ATP homeostasis 133 Adenylate kinase catalyzes the interconversion of adenine nucleotides (ATP+AMP \longleftrightarrow 134 2ADP), which is important for *de novo* adenine nucleotide synthesis and the balance of ATP, 135 ADP, and AMP. Previous biochemical studies reported that the deletion of the adenylate 136 kinase gene reduced ATP concentrations in mouse skeletal muscle cells and budding yeasts 137 (Gauthier et al., 2008, Janssen, Dzeja et al., 2000). We confirmed these findings using an 138 ATP imaging method: $adkl\Delta$ cells showed significantly lower QUEEN ratios than wild-type 139 cells on average in the presence of sufficient carbon sources (Fig. S1). 140 141 In addition to being a key enzyme in purine metabolism, adenylate kinase has also been 142 suggested to cooperate with AMPK in order to monitor the cellular energy state (Hardie, 143 Carling et al., 1998). Therefore, we compared ATP levels in $snfl\Delta \ adkl\Delta$ double mutant cells 144 with those in $snfl\Delta$, $adkl\Delta$, and wild-type cells (Fig. 2A, B). The data obtained were 145 represented by a dot plot that shows distribution characteristics in extensive detail. $snfl\Delta$ 146 $adkl\Delta$ cells had significantly lower average ATP levels than single mutant cells. We also

- found not only a general reduction, but also a huge variation in ATP levels in the $snfl\Delta$,
- 148 $adkl\Delta$, $snfl\Delta$ $adkl\Delta$ cell population, as indicated by the large coefficient of variance (CV)
- 149 (Fig. 2B). Furthermore, some cells had extremely low ATP levels in $snfl \Delta adkl \Delta$ (Fig. 2A,
- B). These results suggest that Adk1 and the Snf1 complex both synergistically contribute to
- 151 ATP homeostasis.

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- 153 A large pool of adenine nucleotide is important for maintaining cellular ATP
- 154 concentrations
- We also examined a $bas 1\Delta$ mutant, which is defective in the expression of genes responsible
- 156 for adenine biogenesis (Daignan-Fornier & Fink, 1992, Denis, Boucherie et al., 1998).
- 157 Consistent with previous biochemical findings (Gauthier et al., 2008), ATP levels quantified
- by QUEEN were reduced by $\sim 50\%$ in $bas 1\Delta$ cells (Fig. 2C, D). We found not only a general
- reduction, but also a large variation in ATP levels in the $bas1\Delta$ cell population, as indicated
- by the large CV (Fig. 2D). The decrease observed in ATP levels was due to reduced adenine
- biosynthesis because the addition of extra adenine to media partially restored ATP levels (Fig.
- 162 2D). These results suggest that the sufficient production of adenine nucleotides is essential for
- the stable maintenance of ATP levels. Moreover, the role of Bas1 in maintaining ATP levels
- appeared to be epistatic to that of Snf1 because $bas1\Delta snf1\Delta$ double mutant cells showed a
- similar distribution of ATP levels to $bas1\Delta$ cells (Fig. S2).
 - ATP levels temporally fluctuate in ATP mutant cells
- To investigate the mechanisms contributing to the large variations in ATP concentrations in
- $snfl\Delta adkl\Delta$ cells in more detail, we employed time-lapse ATP imaging (Fig. 3). We found
- that the QUEEN ratio often underwent a rapid decline followed by recovery in $snfl \Delta adkl \Delta$
- 171 cells (see 116 and 132 min in Fig. 3A, C, and Movie S1, and 180 and 356 min in Fig. 3B, D,
- and Movie S2). The sudden decrease in ATP concentrations (hereafter called "the ATP
- catastrophe") occurred within a few minutes without any sign and was rarely observed in
- wild-type cells (Takaine et al., 2019). The ATP catastrophe appeared to be a stochastic event
- and cell intrinsic: these events occurred independent of cell cycle stages or cell sizes (compare
- 176 Fig. 3C with D). Under some conditions, the QUEEN ratio did not recover after the ATP
- catastrophe and the cell died, as judged by the loss of QUEEN signals in the cell (Fig. S3).
- These results suggest that the large variations in ATP concentrations observed in $snfl \Delta adkl \Delta$
- cells were not simply due to a mixed population with different basal ATP levels, but were
- rather due to the stochastic ATP catastrophe in individual cells.

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Time-lapse imaging of $bas 1\Delta$ revealed oscillatory cycles in ATP concentrations (Figs. 3E and S4A, and Movies S3 and S4): ATP cycling in $bas1\Delta$ cells was slow (~35 min on average, Fig. S4B) and distinguishable from that in $snf1\Delta \ adk1\Delta$ cells; however, the common characteristics of these mutants were that the concentration of ATP repeatedly reached close to 0 mM. The ATP oscillation cycle was unsynchronized in the population and independent of cell cycle progression, suggesting a unique metabolic rhythm intrinsic to each cell. The oscillatory nature of ATP cycling in the $bas 1\Delta$ mutant may involve a transcription/translation cycle and will be described elsewhere. ATP homeostasis is required for preventing protein aggregation. We recently reported that cellular ATP concentrations are stably maintained at ~4 mM in budding yeast (Takaine et al., 2019) and herein demonstrated that in the Snf1/AMPK complex, Adk1 and Bas1 are required for the regulation of ATP homeostasis. However, the physiological importance of ATP homeostasis remains unknown. To clarify the significance of high ATP concentrations, we examined the global genetic interactions of $snfl\Delta$, $adkl\Delta$, and bas1∆ using CellMap ((Usaj, Tan et al., 2017), thecellmap.org). An in silico analysis identified genes involved in "protein folding/glycosylation" as common negative genetic interactors with $adk1\Delta$ and $bas1\Delta$ (Fig. 4A). Negative genetic interactors of ura6, a gene encoding uridylate kinase that also exhibits adenylate kinase activity, were enriched in the "protein folding/glycosylation" category (Fig. 4A). We also found that interactors of snfl were implicated in "protein folding/glycosylation. None of these mutants exhibited apparent genetic interactions with genes in the "metabolism" category (Fig. 4A). The same analysis using genetic and physical interactors provided similar results and showed that many interactors were enriched in the "protein turnover" category (Fig. S5). These results imply that although these three mutants regulate ATP with distinct mechanisms, all three have a common cellular function. To examine possible defects in protein folding and turnover (i.e., proteostasis), we challenged these mutants with various proteotoxic stresses. We found negligible growth defects in ATP mutants under normal growth conditions with 2% glucose at 30°C (control in Fig. 4B). suggesting that a high concentration of ATP is not necessary for cellular growth. However, the adk1 and bas1 mutants both exhibited severe growth defects with a high temperature of 40°C, 1 hour of heat shock at 55°C, or in the presence of 0.5 μg/ml of the glycosylation inhibitor tunicamycin or 2 mM H₂O₂. The SNF1 deletion increased the stress sensitivity of $adkl\Delta$ (Fig. 4B). This sensitivity to proteotoxic stress suggests that ATP homeostasis mutants

217 are defective in some aspects of proteostasis. We found that all four mutants tested contained 218 significantly increased numbers of Hsp104-GFP foci, a marker of protein aggregation 219 (Josefson, Andersson et al., 2017) (Fig. 4C, D). In contrast to Hsp104-GFP foci, Pab1-GFP, a 220 marker of stress granule (SG) assembly (Hoyle, Castelli et al., 2007), did not form foci in 221 ATP mutants, suggesting that protein aggregation and SG assembly are regulated in a distinct 222 manner (Fig. 4D). These analyses identified abnormal protein aggregation as a common 223 defect associated with ATP homeostasis mutants for the first time. 224 225 The transient depletion of ATP leads to the formation of protein aggregates 226 To examine whether ATP depletion triggers protein aggregation in living yeast, we artificially 227 depleted cellular ATP levels by replacing glucose with 2-deoxyglucose (2DG), a strong 228 inhibitor of glycolysis, in the media and monitored protein aggregation using Hsp104-GFP as 229 a marker of protein aggregation (Josefson et al., 2017) in wild-type cells. We previously 230 showed that ATP levels were almost completely depleted 2 minutes after the 2DG treatment 231 (Takaine et al., 2019). Within 15 min of the 2DG treatment, more than 20% cells contained 232 Hsp104-GFP foci indicative of protein aggregation (Fig. 5A, B). In contrast to ATP, the concentration of which recovers to normal levels within 1 min of glucose refeeding (Takaine 233 234 et al., 2019), the dissolution kinetics of Hsp104-GFP were significantly slower (up to hours, 235 Fig. 5A, B). 236 To further confirm whether a high concentration of ATP is required for protein solubility, 237 we also tested the Ubc9-ts protein, a model protein that is prone to aggregate (Kaganovich, 238 Kopito et al., 2008), and found that ATP depletion by the 2DG treatment triggered Ubc9-ts 239 protein aggregation (Fig. 5C). Thus, not only Hsp104-GFP-positive intrinsic proteins, but also 240 extrinsic model proteins aggregate after ATP depletion. 241 242 SG are assembled in budding yeast cells after stressful conditions, such as glucose depletion 243 (Hoyle et al., 2007). In contrast to the formation of Hsp104-GFP foci, ATP depletion after the 244 2DG treatment did not instantly trigger the formation of SG (Fig. S6). Consistent with recent 245 findings (Jain, Wheeler et al., 2016), the present results suggest that SG formation requires 246 ATP. We also noted that Hsp104-GFP foci and SG did not co-localize, suggesting that these 247 structures are derived from distinct mechanisms (Fig. S6). Thus, the artificial depletion of 248 ATP may trigger abnormal protein aggregation in living yeast cells.

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ATP homeostasis is required for the protection of cells from cytotoxicity caused by protein aggregation Protein aggregation is often associated with neurodegenerative diseases, such as Alzheimer's, Huntington's, and Parkinson's diseases (Eftekharzadeh, Hyman et al., 2016). Mitochondrial failure has also been associated with many neurodegenerative diseases; however, it currently remains unclear whether energy failure causes protein aggregation because mitochondria also produce cytotoxic reactive oxygen species (ROS). (Bhat, Dar et al., 2015, Pathak, Berthet et al., 2013). The abnormal aggregation of α -synuclein has been implicated in Parkinson's disease (Lashuel, Overk et al., 2013). To clarify whether ATP is preventing the formation of cytotoxic protein aggregation, we examined the toxicity of α-synuclein-GFP (Syn-GFP) in budding yeast. As reported previously, the expression of Syn-GFP exhibited negligible toxicity against wild-type yeast when expressed under the inducible *GAL1* promotor (Fig. 6A) (Outeiro & Lindquist, 2003, Sharma, Brandis et al., 2006, Wijayanti, Watanabe et al., 2015). However, $snfl\Delta$, $adkl\Delta$, and $basl\Delta$ are hypersensitive to the expression of Syn-GFP (Fig. 6A). We also found that $rpn4\Delta$, which encodes a key transcription factor for proteasomal subunits (Xie & Varshavsky, 2001), was very sensitive to Syn-GFP (Fig. 6A), which is consistent with the concept that Syn-GFP is degraded in the ubiquitin-proteasomal pathway in yeast (Tofaris, Kim et al., 2011, Wijayanti et al., 2015). We then visualized the cellular localization of Syn-GFP. Consistent with previous findings (Willingham, Outeiro et al., 2003), Syn-GFP expressed in yeast mainly localized to the plasma membrane (Fig. 6B, C). In addition to the plasma membrane, we found that Syn-GFP localized to punctated structures, most likely corresponding to protein aggregation (Fig. 6B, C). Punctated structures were not as obvious in $rpn4\Delta$ cells defective in proteasomes, suggesting that the accumulation of Syn-GFP puncta was not simply due to defective degradation. To investigate whether a high concentration of ATP protects cells from toxic protein aggregation, we added extra adenine to the medium (Fig. 6D). A previous study demonstrated that the addition of 300 µM adenine to the medium increased ATP concentrations from 4 to 5.5 mM in wild-type cells and from 1 to 4 mM in bas 1 \Delta cells (Gauthier et al., 2008) (similar results are shown in Fig. 2D), but induced little or no changes in $adk1\Delta$ cells (from 2 to 2 mM) (Gauthier et al., 2008). Consistent with our hypothesis, the addition of adenine reduced Syn-GFP toxicity and aggregation in $bas 1\Delta$, but not $adk 1\Delta$ cells (Fig. 6D). Thus, a high concentration of ATP prevented Syn-GFP aggregation and toxicity.

We examined another model protein involved in neurodegenerative diseases. PolyQ

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containing the huntingtin protein tends to aggregate and has been implicated in Huntington's disease (Jiang, Poirier et al., 2005). We investigated the toxicity of Htt103Q, a mutant form of the huntingtin protein, which tends to aggregate and cause cytotoxicity in yeast (Meriin, Zhang et al., 2002). Consistent with the concept that a high concentration of ATP prevents protein aggregation, the ATP homeostasis mutants $snfl\Delta$, $adkl\Delta$, $snfl\Delta$ $adkl\Delta$, and $basl\Delta$ were very sensitive to Htt103Q expression (Fig. S7). Proteasomes are essential for the removal of protein aggregates caused by ATP depletion Protein aggregation caused by ATP depletion was cytotoxic (Figs. 4B and 6A) and was not easily dissolved after ATP repletion (Fig. 5). To identify a pathway that is essential for the removal of aggregates, we examined the involvement of proteasomes and autophagy. The deletion of RPN4, which encodes a transcription factor of proteasomal genes (Xie & Varshavsky, 2001), down-regulated proteasomal activity (Kruegel, Robison et al., 2011) and resulted in synthetic growth defects with $adkl\Delta$, $snfl\Delta$ bas $l\Delta$ at a high temperature of 38°C and in the presence of H₂O₂ (Fig. 6E). In contrast to proteasomes, autophagy did not appear to have genetic interactions with the above mutants (Fig. 6E). The deletion of an essential component of the autophagic pathway, ATGI did not affect the sensitivity of $adk1\Delta$, $snf1\Delta$, $bas 1\Delta$ to a high temperature of 38°C or to H_2O_2 (Fig. 6E). We also did not observe the accumulation of Hsp104-GFP foci in the autophagy mutants atg1, atg8, and atg13 (not shown). To investigate the involvement of proteasomes in the removal of protein aggregates after the transient depletion of ATP, we pretreated cells with the proteasomal inhibitor MG132 or DMSO and examined the kinetics for the formation of Hsp104-GFP foci after the 2DG treatment (Fig. 6F) using the drug-sensitive yeast strain Y13206 (Piotrowski, Li et al., 2017). Under both conditions, more than 90% of cells exhibited Hsp104-GFP foci within 30 min of the 2DG treatment. More than two-thirds of Hsp104-GFP foci dissolved in the DMSO control. while less than one-third dissolved in MG132-treated samples, indicating that proteasomes are required for the dissolution process (Fig. 6F).

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DISCUSSION In the present study, we demonstrated for the first time that the Snf1 complex, budding yeast AMPK, is required for the stable maintenance of cellular ATP concentrations (ATP homeostasis) in collaboration with the adenylate kinase Adk1 (Fig. 7A). This function of the Snf1 complex in ATP homeostasis is independent of glucose concentrations in the medium and Mig1, the major transcriptional repressor involved in glucose repression (Fig. 1); therefore, this is distinct from its well-characterized role in adaptation to glucose limitations. The activity of the Snf1 kinase complex may be sharply tuned depending on the intracellular concentrations of adenine nucleotides or other metabolites indicative of cellular energy to prevent a rapid ATP catastrophe (Fig. 3), even in the presence of sufficient amounts of glucose. It is important to note that the reductions observed in intracellular ATP concentrations in $snfl \Delta$ cells in the presence of glucose were overlooked in previous biochemical analyses, again demonstrating the usefulness of QUEEN-based ATP imaging. Since the deletion of BASI caused the greatest reduction in ATP levels and is epistatic to $snfl\Delta$, a large pool size of adenine nucleotide is a prerequisite for ATP homeostasis. This assumption is reasonable because the pool size of recyclable ATP restricts ATP concentrations based on the rapid turnover rate of ATP. Bas1 maintains the pool size of ATP by balancing ATP synthesis and irreversible decreases, such as incorporation into RNA and DNA (following conversion to deoxy-ATP), degradation, and excretion in rapidly proliferating yeasts. We also showed that key regulators of ATP homeostasis play roles in preventing cytotoxic protein aggregation in budding yeast (Fig. 7B). The common feature associated with these mutants is reduced ATP levels, suggesting that high ATP levels are essential for protein solubilization. A recent study suggested that the requirement for ATP changes depending on its concentration. At concentrations lower than 0.5 mM, ATP mainly serves as a substrate for enzymes, such as protein kinases and heat shock protein chaperones, whereas at concentrations higher than 2 mM, ATP may exert solubilizing effects on disordered proteins (Sridharan et al., 2019). ATP homeostasis may also be required to constantly drive proteasomal protein degradation, which requires large amounts of ATP (Benaroudi, Zwickl et al., 2003, Tanaka, Waxman et al., 1983).

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Previous biochemical measurements indicated that although ATP levels were lower in adk1 and bas I mutants than in the wild type, these mutants still had ATP concentrations that were higher than 2 mM (Gauthier et al., 2008), which does not directly explain the accumulation of protein aggregates in these mutants (Fig. 4) because most proteins are expected to be soluble at > 2 mM ATP (Sridharan et al., 2019). The visualization of ATP dynamics in living cells at the single cell level revealed that ATP undergoes transient depletion repeatedly (the ATP catastrophe) in AMPK and ADK mutants, which explains how aggregation-prone proteins aggregate and cause cytotoxicity. A similar, but distinct instability of ATP was observed in bas 1 \Delta, further confirming that the ATP catastrophe causes protein aggregation. Although severe ATP depletion in these mutants was gradually recovered by as yet uncharacterized negative feedback regulation, the duration period of ~15 min with reduced ATP levels may induce some proteins to form aggregates that last for generations. Many neurodegenerative diseases, such as Alzheimer's, Huntington's, and Parkinson's diseases, are associated with protein aggregation (Eisele, Monteiro et al., 2015, Josefson et al., 2017). On the other hand, mitochondrial dysfunction and accompanying energy failure in nerve cells may result in many types of neurodegenerative diseases based on a large body of evidence (Haelterman, Yoon et al., 2014, Pathak et al., 2013). For example, previous studies demonstrated that ATP levels in the brain were decreased in patients with early Parkinson's disease (Mochel, N'Guyen et al., 2012) and also that ATP synthesis in the brain was not properly regulated in patients with early Huntington's disease (Hattingen, Magerkurth et al., 2009) and in the corresponding mouse model (Mochel, Durant et al., 2012). Therefore, protein aggregation induced by the ATP catastrophe, as revealed in the present study, may link energy failure and protein aggregation, providing a comprehensive insight into the onset of neurodegenerative diseases. Further studies are warranted to clarify whether the ATP catastrophe also occurs in the neurons of patients at risk of neurodegenerative diseases or in the elderly. However, neither biochemical measurements nor mass spectrometry is capable of detecting the ATP catastrophe because of their insufficient time and space resolution. Thus, an ATP imaging approach using the yeast model system will be the first choice for elucidating the molecular mechanisms underlying ATP homeostasis and ATP catastrophe-induced protein aggregation. A recent study reported that the activation of AMPK by metformin ameliorated the progression of Huntington's disease in a mouse model (Arnoux, Willam et al., 2018), and the potential therapeutic use of metformin for neurodegenerative diseases is being discussed

(Rotermund, Machetanz et al., 2018). Furthermore, the involvement of ATP and ADKs in preventing the manifestation of Parkinson's disease in mouse models and patients has been proposed (Garcia-Esparcia, Hernandez-Ortega et al., 2015, Nakano, Imamura et al., 2017). Protein aggregation induced by the ATP catastrophe may be a general mechanism for the development of proteinopathies. The present study study using ATP imaging revealed a physiological consequence of a failure in ATP homeostasis in living cells for the first time and suggests that ATP homeostasis has potential as a target for preventing/treating neurodegenerative diseases.

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MATERIALS AND METHODS Yeast strains and plasmids The budding yeast strains and plasmids used in the present study are listed in Supplementary Tables S1 and S2, respectively. These strains were constructed by a PCR-based method (Janke, Magiera et al., 2004) and genetic crosses. The yeast knockout strain collection was originally purchased from GE Healthcare (cat. #YSC1053). Media and cell culture The standard technique for the yeast culture and manipulation was used (Guthrie & Fink, 2002). Synthetic medium (SC) was prepared according to the recipe of Hanscho et al. (Hanscho, Ruckerbauer et al., 2012). 2-Deoxy-D-glucose (2DG), tunicamycin, and MG132 were purchased from FUJIFILM Wako (cat. # 046-06483, 202-08241, and 139-18451, respectively). Tunicamycin and MG132 were dissolved in dimethylsulfoxide (DMSO) to make stock solutions (5 mg/ml and 42 mM, respectively). Cells were grown to the mid-log phase at 30°C in SC before imaging unless otherwise noted. Microscopy Cells expressing Hsp104-GFP or GFP-Ubc9ts were concentrated by centrifugation and sandwiched between a slide and coverslip (No. 1.5 thickness, Matsunami, Osaka, Japan). Immobilized cells were imaged using an inverted fluorescent microscope (Eclipse Ti-E, Nikon) equipped with an Apo TIRF 100× Oil DIC N2/NA 1.49 objective lens and electron-multiplying charge-coupled device camera (iXon3 DU897E-CS0-#BV80, Andor) at approximately 25°C. The Hsp104-GFP and GFP-Ubc9ts fluorescent signal was collected from stacks of 11 z-sections spaced by 0.5 µm, and the maximum projections of the optical sections were shown in Figs. 4, 5, and S6. Cells expressing QUEEN were immobilized on a concanavalin A-coated 35-mm glass-bottomed dish (#3971-035, No. 1.5 thickness, IWAKI). The dish was filled with an excess amount of medium (4.5–5 ml) against the cell volume to minimize changes in the chemical compositions of the medium during observations. QUEEN fluorescence was acquired as previously described (Takaine et al., 2019). Cells expressing α-synuclein-GFP were immobilized on a slide glass as described above, and the fluorescence signal was collected from a single z-plane using an inverted fluorescent microscope (Eclipse Ti2-E, Nikon, Tokyo, Japan) equipped with a CFI Plan Apoλ100× Oil DIC/NA1.45 objective

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lens and CMOS image sensor (DS-Qi2, Nikon). Images of cells were acquired from several fields of view for each experimental condition. Data analysis Numerical data were plotted using KaleidaGraph software ver. 4.5.1 (Synergy Software). Means, SDs, and p values were calculated using Excel software (Microsoft, WA, US). Significance between two sets of data was tested using the unpaired one-tailed Welch's t-test unless otherwise noted, and was indicated by an asterisk or p value. The horizontal bar in the dot plot indicates the average of each population. All measurements were repeated at least twice to confirm reproducibility. ATP concentrations in yeast cells were estimated using QUEEN-based ratiometric imaging, as previously described (Takaine, 2019), (Takaine et al., 2019). The QUEEN ratio is proportional to ATP concentrations and pseudo-colored to reflect its value throughout the present study. The mean QUEEN ratio inside of a cell represents the intracellular ATP concentration of the cell. The autocorrelation functions (ACF) of oscillations in the QUEEN ratio were calculated using R studio software ver. 3.4.1 (R Core Team, 2017). The apparent period of oscillation was estimated from the positive second peak of the correlation coefficient, which was outside the 95% confidence interval and significant (p < 0.05), rejecting the assumption that there is no correlation.

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Molecular and Cellular Regulation, Gunma University, Japan.

AUTHOR CONTRIBUTIONS

- 456 M.T. and S.Y. conceived and designed the project. M.T. conducted experiments and the data
- analysis. H.I. provided a key reagent and expertise. M.T. and S.Y wrote the manuscript with
- 458 input from H.I.

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CONFLICT OF INTEREST

The authors declare no competing interests.

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FIGURE LEGENDS

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- Fig. 1 AMPK is involved in the maintenance of cellular ATP levels in non-starving cells
- (A) Visualization of intracellular ATP levels in wild-type (WT) and $snfl\Delta$ cells using the
- ATP sensor QUEEN. Cells were grown to the mid-log phase in SC-H medium containing the
- indicated concentrations of glucose and then imaged. White scale bar = 5 μ m. (B) $snfl\Delta$ cells
- had lower ATP levels than wild-type cells. The mean QUEEN ratio inside a single cell (mean
- 605 QUEEN ratio/cell), which represents the intracellular ATP level of the cell, was quantified for
- each cell from the ratio image. Data are the mean of the cell population \pm 1SD (error bar)
- normalized to wild-type cells. N = 105–193 cells were scored. Asterisks indicate a significant
- difference from WT (p < 0.05). (C) ATP levels in single-deletion mutants of AMPK β
- subunits. Data are the mean of the cell population \pm 1SD (error bar) normalized to wild-type
- 610 cells. N = 177–223 cells were scored. Asterisks indicate a significant difference between the
- two strains (*, p < 0.05; **, p < 0.01). N.S., no significance. (D) $mig 1\Delta$ cells had slightly
- lower ATP levels than wild-type cells. Data are the mean of the cell population \pm 1SD (error
- bar) normalized to wild-type cells. N = 190–231 cells were scored. Asterisks indicate a
- significant difference between the two strains.
- Fig. 2 Interconversion and active synthesis of adenine nucleotides are important for
- 617 ATP homeostasis

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- 618 (A) Adk1 and Snf1 synergistically control cellular ATP levels. QUEEN ratio images of ATP
- 619 homeostasis mutant cells grown in the medium containing 2% glucose. The asterisk indicates
- an example of cells with significantly reduced ATP levels. (B) Mean QUEEN ratios of cells
- were translated to ATP concentrations and shown in a dot plot. The horizontal bar indicates
- the mean of each population. Asterisks indicate p values versus WT: $*=1.5\times10^{-59}$,
- **= 2.3×10^{-34} , ***= 2.9×10^{-117} . CV: coefficient of variance. N = 134–276 cells were scored.
- 624 (C) QUEEN ratio images of $bas 1\Delta$ cells grown in the 2% glucose medium. Growth in media
- supplemented with 0.11 mg/ml adenine partially restored the low ATP phenotype of $bas 1\Delta$.
- 626 (D) The ATP concentrations of cells shown in (C) were plotted. Asterisks indicate p values:
- *=2.3×10⁻¹⁶⁰, **=8.8×10⁻¹² (versus WT in glucose), ***=3.6×10⁻²⁰ (versus *bas1*Δ in glucose).
- 628 CV: coefficient of variance. N = 186–296 cells were scored.
- 630 Fig. 3 Temporal fluctuations in ATP levels in $snf1\Delta adk1\Delta$ and $bas1\Delta$ cells
- 631 (A, C) Time-lapse imaging of QUEEN in $snfl\Delta \ adkl\Delta$ cells in the 2% glucose medium.
- Images at the representative time points were shown. The QUEEN ratio decreased twice
- 633 (indicated by arrowheads) within a short interval. See also Movie S1. Data were converted

634 into ATP concentrations and plotted in (C). (B, D) Another example of time-lapse imaging of QUEEN in $snfl\Delta adkl\Delta$ cells in the 2% glucose medium. The QUEEN ratio decreased twice 635 636 (indicated by arrowheads) with a long interval. See also Movie S2. Data were converted into 637 ATP concentrations and plotted in (D). (E) Time-lapse imaging of QUEEN in $bas 1\Delta$ in the 638 2% glucose medium. The ATP concentrations of the mother (cell-1) and daughter (cell-2) 639 were plotted at the bottom. Images at the representative time points were shown on the top. 640 Note that the QUEEN ratio is synchronized until cells undergo separation at the time point of 641 76 min indicated by an arrow. After separation, each cell has a unique periodic cycle of ATP. 642 The movie is available in Movie S3. White scale bar = $5 \mu m$. 643 644 Fig. 4 ATP homeostasis is required to prevent protein aggregation 645 (A) Functional landscape of known interactors of ATP mutants. Negative genetic interactors 646 of the indicated gene were derived from the SGD database (https://www.yeastgenome.org/) 647 (Cherry, Hong et al., 2012) and overlaid on a functional map based on the global genetic 648 interaction network of the yeast genome (Baryshnikova, 2016, Usaj et al., 2017). URA6 649 encodes an uridylate kinase that is essential for viability, which also exhibits adenylate kinase 650 activity. (B) Each strain of the indicated genotype was serially diluted (five-fold), spotted on 651 the SC + 2% glucose medium, and grown under the indicated stress conditions. Photos were 652 taken after 2-3 days. (C) Formation of Hsp104-GFP foci in ATP homeostasis mutants. The 653 GFP signal (inverted grayscale) was imaged in the log phase culture of the indicated mutant 654 cells expressing Hsp104-GFP at 35°C. (D) Quantification of data shown in (C). Data from 655 similar experiments using strains expressing Pab1-GFP, instead of Hsp104-GFP, were also 656 plotted. Values are the mean \pm 1SD (error bars). Asterisks indicate a significant difference from WT (p < 0.05) (N=3-4). White scale bar = 5 μ m. 657 658 659 Fig. 5 ATP depletion triggers protein aggregation in living yeast cells 660 (A) The formation of Hsp104-GFP foci after ATP depletion. Wild-type cells expressing 661 Hsp104-GFP were grown to the log phase at 35°C in the medium containing 2% glucose. At 662 the time point of 0 min, the medium was replaced with 20 mM 2DG (red) or 2% glucose (as a 663 control; blue). Cells were released back to media containing 2% glucose at the time point of 664 50 min. Cells were imaged at the indicated time points, classified, and scored according to the 665 number of Hsp104-GFP foci. Values are the mean (N = 3). Asterisks indicate a significant

difference from the 2% glucose treatment (p < 0.05). (B) Representative images of cells

GFP-Ubc9-ts under an inducible *GAL* promoter were grown in the medium containing 2%

analyzed in (A). (C) Formation of Ubc9-ts foci after ATP depletion. Cells expressing

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- galactose (SC-gal) at 33°C, and the medium was then exchanged with 2DG or SC-gal. After
- 30 min, cells were imaged and scored for the number of GFP-Ubc9-ts foci. Representative
- images (inverted grayscale) are shown on the left and summarized on the right. Values are the
- mean \pm 1SD (error bars) (N = 4). White scale bar = 5 μ m.

- 674 Fig. 6 Aggregation and cytotoxicity of α-synuclein depends on ATP homeostasis
- 675 (A) Each strain of the indicated genotype was transformed with an expression vector carrying
- 676 α-synuclein-GFP or GFP, serially diluted (five-fold), spotted on SC + 2% galactose plates,
- and then grown at 30°C for 3 days. (B) The localization of α-synuclein-GFP in ATP mutants.
- 678 Cells were grown on galactose plates at 30°C for more than 42 h and then imaged.
- Representative images of α-synuclein-GFP (inverted grayscale) are shown. (C) Quantification
- of the data shown in (B). Cells were classified and scored for the localization pattern of
- α -synuclein-GFP. The percentage of cells showing α -synuclein-GFP foci are plotted. Data are
- the mean \pm 1SD (error bars) from 3–6 independent observations. N = 33–380 cells were
- scored in each measurement. P values versus WT are shown. N.S., no significance (p value >
- 684 0.05). (D) (top) Each strain of the indicated genotype was transformed with an expression
- vector carrying α-synuclein-GFP and grown on galactose plates containing 0 mM (–
- Adenine) or 0.3 mM (+ Adenine) adenine at 30°C for 3 days. (bottom) Cells were grown on
- galactose plates in the absence or presence of adenine at 30°C for 41-45 h and then imaged.
- The percentage of cells showing α -synuclein-GFP foci were plotted. Data are the mean \pm 1SD
- (error bars) from 5 independent transformants. N = 53-258 cells were scored in each
- 690 measurement. P values versus "- Adenine" are shown. (E) Each strain of the indicated
- 691 genotype was serially diluted (five-fold), spotted on the SC + 2% glucose medium, and grown
- under the indicated stress conditions. (F) Cells of the drug-sensitive strain Y13206 were
- grown to the log phase at 37°C in the medium containing 2% glucose and supplemented with
- 694 0.1% DMSO or 0.1% DMSO plus 42 μ M MG132 at t = -30 min. At t = 0 min, these cells
- were washed and released in the medium containing 20 mM 2DG \pm MG132, and the cells
- were then washed and released again in the medium containing 2% glucose \pm MG132. Cells
- were imaged at the indicated time points and scored for the number of Hsp104-GFP foci.
- Data are the mean \pm 1SD (error bars). Asterisks indicate a significant difference from DMSO
- 699 (p < 0.02) (N=3).

- 701 Fig. 7 Models for ATP homeostasis and its role in proteostasis
- 702 (A) Schematic summary of the roles of Snf1, Adk1, and Bas1 in ATP homeostasis. (B) A
- schematic model for ATP homeostasis preventing cytotoxic protein aggregation.

SUPPLEMENTARY INFORMATION

- 705 Supplementary figure legends
- Fig. S1 Adenylate kinase Adk1 is involved in the maintenance of cellular ATP
- 707 concentrations

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- 708 (A) QUEEN ratio images of wild-type and $adk1\Delta$ cells grown in 2% glucose or 2% galactose.
- 709 White scale bar = $5 \mu m$. (B) Quantification of data shown in (A). The mean QUEEN ratio/cell
- 710 was quantified for each cell from ratio images. Values are the mean of the cell population \pm
- 711 1SD (error bar). N = 182–236 cells were scored. Asterisks indicate a significant difference
- 712 from WT (p < 0.05).
- 714 Fig. S2 ATP levels in $bas1\Delta snf1\Delta$ cells
- 715 (A) QUEEN ratio images of wild-type, $bas 1\Delta$, $snf 1\Delta$, and $bas 1\Delta snf 1\Delta$ cells. (B)
- Quantification of data shown in (A). The mean QUEEN ratio/cell was quantified for each cell
- from ratio images. Data are shown as a dot plot. The horizontal bar in the plot indicates the
- mean of each population. Significance between two sets of data was tested using the unpaired
- two-tailed Welch's *t*-test and indicated by *p* values. N.S., no significance (*p*-value > 0.05).
- 721 Fig. S3 Time-lapse imaging of QUEEN in $snf1\Delta$ adk 1Δ cells
- 722 Time-lapse imaging of QUEEN in $snfl\Delta adkl\Delta$ cells in the 2% glucose medium. An example
- of $snfl\Delta \ adkl\Delta$ cell showing an irreversible decrease in the QUEEN ratio (indicated by an
- arrowhead). In this case, the cell eventually died (indicated by an arrow). Images at the
- representative time points were shown. The ATP concentration was plotted at the bottom.
- 726 White scale bar = $5 \mu m$.
- 728 Fig. S4. Oscillatory behavior of the ATP concentration visualized in *bas1*Δ cells
- 729 (A) Another example of $bas 1\Delta$ cells showing an oscillating QUEEN ratio. In this case,
- 730 cytokinesis had just been completed at t = 0 min. ATP concentrations in cell-1 and cell-2 were
- 731 plotted at the bottom. See also Movie S4. White scale bar = $5 \mu m$. (B) Autocorrelation
- function of the OUEEN ratio calculated from the data on cell-1 in Fig. 3E. Blue dotted lines
- 733 indicate the 95% confidence interval. An arrow indicates the second peak of the correlation
- and corresponds to the apparent period.
- 736 Fig. S5 *In silico* analysis of interactors of ATP mutants
- Genetic and physical interactors of the indicated genes were derived from the SGD database
- 738 (https://www.yeastgenome.org/) (Cherry et al., 2012) and overlaid on a functional map based

on the global genetic interaction network of the yeast genome (Baryshnikova, 2016, Usaj et al., 2017). $\it{URA6}$ encodes an uridylate kinase essential for viability, which also exhibits adenylate kinase activity.
Fig. S6 Simultaneous observation of Hsp104 and Pab1 foci Wild-type cells expressing Hsp104-GFP and Pab1-RedStar2 were grown to the log phase at 37°C in the medium containing 2% glucose. Cells were washed and released either in the medium containing 20 mM 2DG (top) or in the medium lacking glucose (bottom), and then imaged after 30 and 60 min. White scale bar = 5 μ m.
Fig. S7 Cytotoxicity of polyQ containing the huntingtin protein in wild-type and ATP-mutant yeast cells
Each strain of the indicated genotype was transformed with an expression vector carrying
Htt103Q and grown on glucose ((-), no induction) or raffinose ((+), leaky expression) plates at 30°C for 3 days.

Table S1. Strains used in the present study

Name	Genotype	Source	Figure
MTY3008	snf1∆::kanMX6 leu2∆0 lys2∆0 ura3∆0	Lab stock	4B, 6A-C,
			6E, S7
MTY3015	his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	Lab stock	4B, 6A-E,
			S7
MTY3049	adk1Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	Lab stock	4B, 6A-E,
			S7
MTY3118	bas1Δ::kanMX6 leu2Δ0 lys2Δ0 ura3Δ0	Lab stock	4B, 6A-E,
			S7
MTY3143	$his3\Delta 1:: 2 \times pRS303 - P_{TEF} - QUEEN - 2m - T_{CYCI} leu 2\Delta 0 lys 2\Delta 0 ura 3\Delta 0$	This study	S1
MTY3149	$adk1\Delta :: kanMX6\ his3\Delta 1 ::\ 2 \times pRS303 - P_{TEF} - QUEEN - 2m - T_{CYCI}\ leu \\ 2\Delta 0\ lys \\ 2\Delta 0$	This study	S1
	ura3Δ0		
MTY3264	$his3\Delta 1:: 3 \times pRS303 - P_{TEF} - QUEEN - 2m - T_{CYCI} leu 2\Delta 0 lys 2\Delta 0 ura 3\Delta 0$	Takaine et	2A, B, S2
	MYO1-3mCherry-hphMX6	al., 2019	
MTY3270	$bas1\Delta::kanMX6\;his3\Delta1::\;3\times pRS303-P_{TEF}-QUEEN-2m-T_{CYCI}\;leu2\Delta0\;lys2\Delta0$	This study	2C, D, 3E,
	ura3A0 MYO1-3mCherry-hphMX6		S2, S4
MTY3293	$adk1\Delta$:: $kanMX6\ his3\Delta1$:: $3 \times pRS303 - P_{TEF} - QUEEN - 2m - T_{CYCI}\ leu 2\Delta0\ lys 2\Delta0$	This study	2A, B
	ura3\Delta 0 MYO1-3mCherry-hphMX6		
MTY3355	$adk1\Delta$:: $kanMX6\ snf1\Delta$:: $natNT2\ his3\Delta1$:: $3 \times pRS303$ - P_{TEF} - $QUEEN$ - $2m$ - T_{CYC1}	This study	2A, B,
	leu2Δ0 lys2Δ0 ura3Δ0 MYO1-3mCherry-hphMX6		3A-D, S3
MTY3371	$snf1\Delta$:: $kanMX6\ his3\Delta1$:: $3 \times pRS303 - P_{TEF} - QUEEN - 2m - T_{CYCI}\ leu 2\Delta0\ lys 2\Delta0$	This study	2A, B, S2
	ura3A0 MYO1-3mCherry-hphMX6		
MTY3412	adk1Δ::kanMX6 snf1Δ::natNT2 leu2Δ0 lys2Δ0 ura3Δ0	This study	4B, 6A-E,
			S7
MTY3420	Hsp104-yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4C, D, 5A,
			В
MTY3421	snf1∆::kanMX6 Hsp104-yeGFP-hphNT1 his3∆1 leu2∆0 lys2∆0 ura3∆0	This study	4C, D
MTY3422	bas1Δ::kanMX6 Hsp104-yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4C, D
MTY3424	adk1Δ::kanMX6 Hsp104-yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4C-D
MTY3425	adk1Δ::kanMX6 snf1Δ::natNT2 Hsp104-yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0	This study	4C, D
	ura3Δ0		
MTY3489	atg1Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	Lab stock	6E
MTY3493	snq2Δ::KILeu2; pdr3Δ::KIura3; pdr1Δ::natMX4; can1Δ::STE2prSp_his5 lyp1Δ	From Y.	Parental

	$his 3\Delta 1\ leu 2\Delta 0\ ura 3\Delta 0\ met 15\Delta 0\ LYS 2+$	Ohya,	strain of
		(Piotrowski	MTY3501
		et al., 2017)	
MTY3501	$Hsp104$ -yeGFP-hphNT1 $snq2\Delta$::KILeu2; $pdr3\Delta$::KIura3; $pdr1\Delta$::natMX4;	This study	6F
	$can1\Delta$:: $STE2prSp_his5\ lyp1\Delta\ his3\Delta 1\ leu2\Delta 0\ ura3\Delta 0\ met15\Delta 0\ LYS2+$		
MTY3503	Pab1- yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4D
MTY3504	Hsp104-yeGFP-hphNT1 Pab1-RedStar2-natNT2 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	S6
MTY3505	snf1Δ::kanMX6 Pab1- yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4D
MTY3506	adk1Δ::kanMX6 Pab1- yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4D
MTY3507	adk1Δ::kanMX6 snf1Δ::natNT2 Pab1- yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0	This study	4D
	ura3Δ0		
MTY3508	bas1Δ::kanMX6 Pab1- yeGFP-hphNT1 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	4D
MTY3513	$atg1\Delta$:: $natNT2$ $adk1\Delta$:: $kanMX6$ $his3\Delta1$ $leu2\Delta0$ $lys2\Delta0$ $ura3\Delta0$	This study	6E
MTY3514	$atg1\Delta$:: $natNT2$ $snf1\Delta$:: $kanMX6$ $his3\Delta1$ $leu2\Delta0$ $lys2\Delta0$ $ura3\Delta0$	This study	6E
MTY3515	atg1Δ::natNT2 bas1Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	6E
MTY3516	rpn4Δ::natNT2 adk1Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	6E
MTY3517	rpn4Δ::natNT2 snf1Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	6E
MTY3518	rpn4Δ::natNT2 bas1Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	This study	6E
MTY3525	rpn4Δ::kanMX6 his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0	Lab stock	6A-C, 6E

Table S2. Plasmids used in the present study

Name	Structure	Source	Purpose
MTP3091	pESC-Leu-GFP-Ubc9-ts	Judith Frydman (pJF1089)	Fig. 5C
MTP3088	p426 103Q GAL	Addgene (cat.# 1188)	Fig. S7
MTP3108	pYES2-α-synuclein-GFP	H. Takagi (Wijayanti et al., 2015)	Fig. 6A-D
MTP3090	pYES2-GFP	K. Ohashi	Fig. 6A

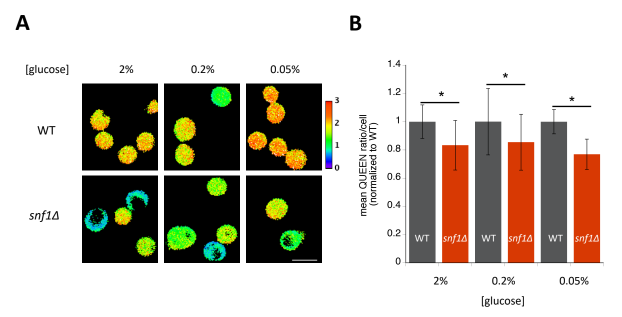
758 Captions for supplementary movies 759 Movie S1 760 Time-lapse imaging of QUEEN in $snfl\Delta adkl\Delta$ cells in the 2% glucose medium. 761 Corresponding to the data shown in Fig. 3A. The QUEEN ratio decreased twice (116 and 132) 762 min) within a short interval. White scale bar = $5 \mu m$. 763 764 **Movie S2** 765 Another example of $snfl\Delta adkl\Delta$ cells showing a sudden decrease in the QUEEN ratio. 766 Corresponding to the data shown in Fig. 3B. The QUEEN ratio decreased twice (180 and 356 767 min) with a long interval. White scale bar = $5 \mu m$. 768 769 Movie S3 770 Oscillatory behavior of the QUEEN ratio in $bas 1\Delta$ cells. Corresponding to the data shown in 771 Fig. 3E. Left, the QUEEN ratio image; middle, Myo1-mCherry (inverted grayscale image); right, bright field image. Images were taken every 4 min. White scale bar = $5 \mu m$. 772 773 774 **Movie S4** 775 Another example of $bas I\Delta$ cells showing an oscillating QUEEN ratio. Corresponding to the 776 data shown in Supplementary Fig. S4A. Left, the QUEEN ratio image; right, bright field 777 image. In this case, cytokinesis had already been completed at t = 0 min. Images were taken

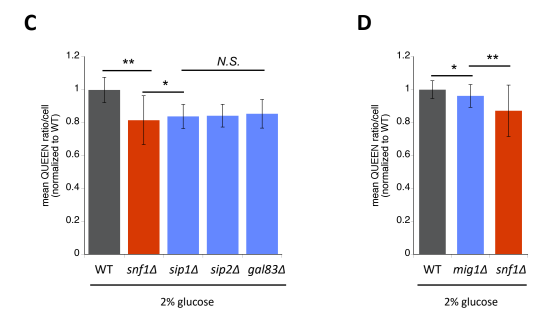
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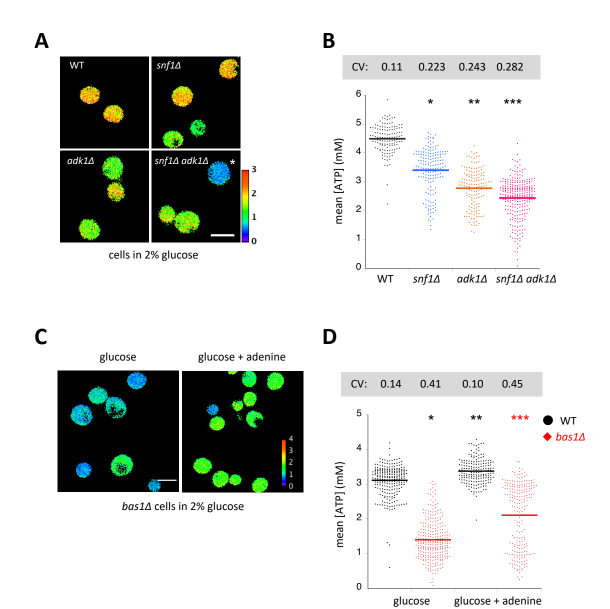
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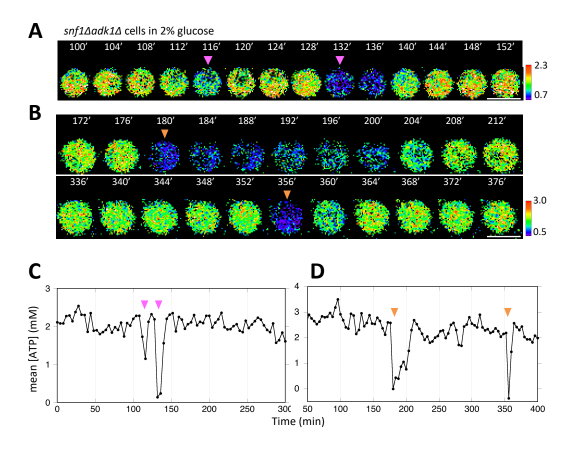
780

every 3 min. White scale bar = $5 \mu m$.









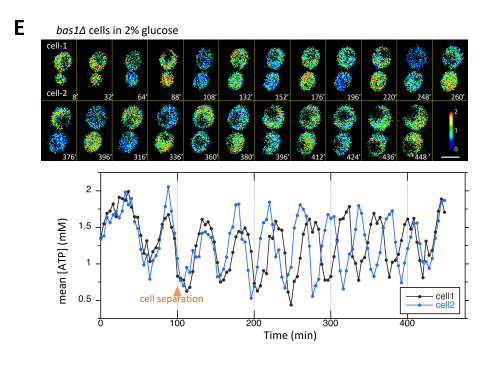
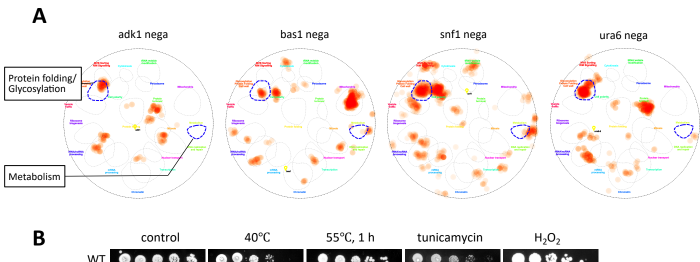
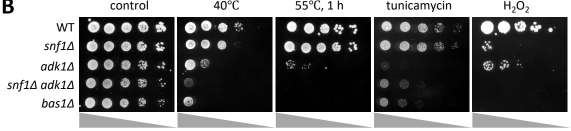
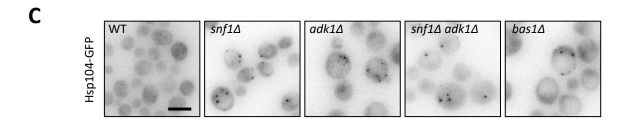


Fig. 4







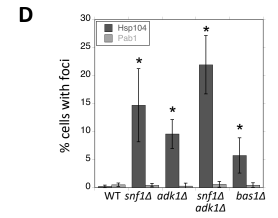
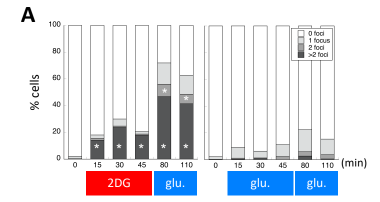
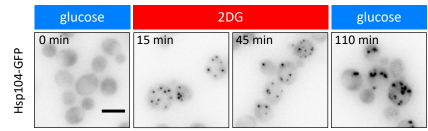
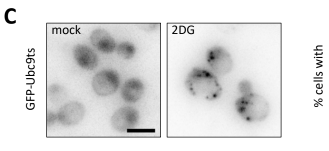


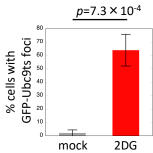
Fig. 5

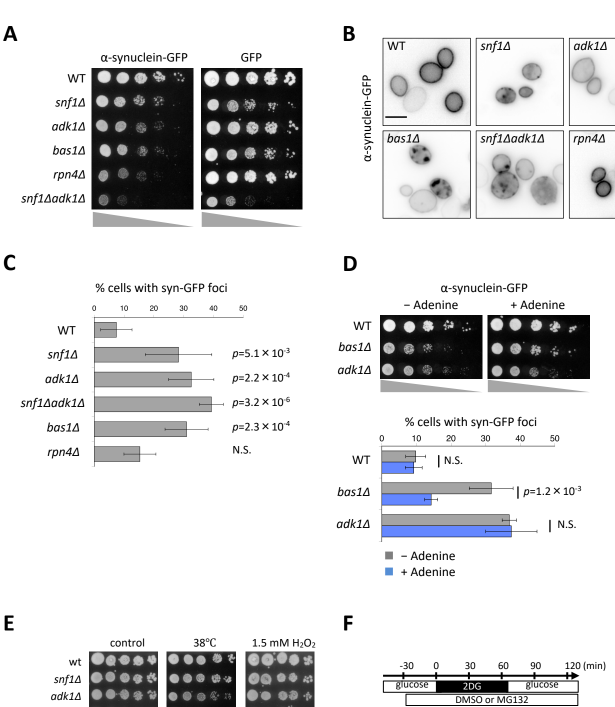


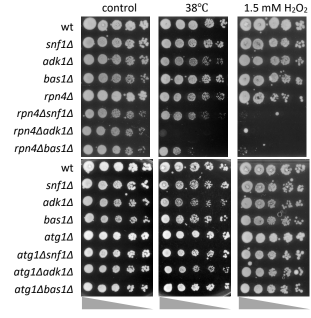
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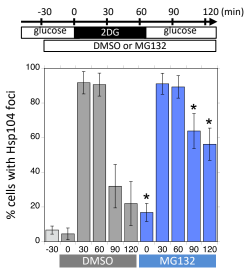


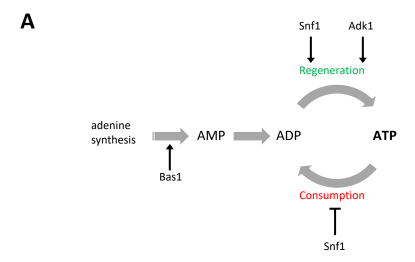


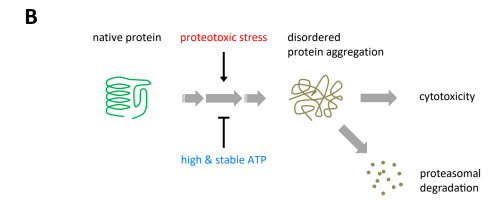


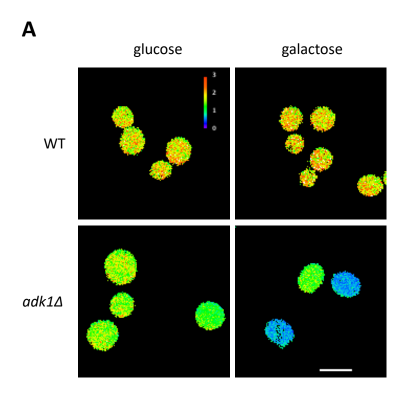


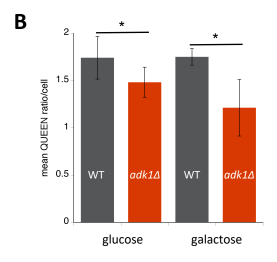


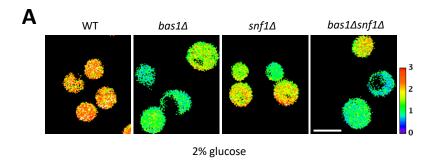












В

