

iPAR: a new reporter for eukaryotic cytoplasmic protein aggregation

Sarah Lecinski¹, Chris MacDonald^{2,3}, Mark C. Leake^{1,2,3,4}

¹ School of Physics, Engineering and Technology, University of York, York, YO10 5DD, UK.

² Department of Biology, University of York, York, YO10 5DD, UK.

³ York Biomedical Research Institute, University of York, York, YO10 5DD, UK.

⁴ Correspondence to mark.leake@york.ac.uk

1. Abstract

Cells employ myriad regulatory mechanisms to maintain protein homeostasis, termed proteostasis, to ensure correct cellular function. Dysregulation of proteostasis, which is often induced by physiological stress and ageing, often results in protein aggregation in cells. These aggregated structures can perturb normal physiological function, compromising cell integrity and viability, a prime example being early onset of several neurodegenerative diseases. Understanding aggregate dynamics *in vivo* is therefore of strong interest for biomedicine and pharmacology. However, factors involved in formation, distribution and clearance of intracellular aggregates are poorly understood. Here, we report an improved methodology for production of fluorescent aggregates in model budding yeast which can be detected, tracked and quantified using fluorescence microscopy in live cells. This new openly-available technology, iPAR (inducible Protein Aggregation Reporter), involves monomeric fluorescent protein reporters fused to a Δ ssCPY* aggregation biomarker, with expression controlled under the copper-regulated *CUP1* promoter. Monomeric tags overcome challenges associated with non-physiological aggregation, whilst *CUP1* provides more precise control of protein production. We show that iPAR enables quantitative study of cytoplasmic aggregate kinetics and inheritance features *in vivo*. If suitably adapted, iPAR offers new potential for studying diseases in other model cellular systems.

Keywords: *Saccharomyces cerevisiae*, cytoplasmic aggregates, confocal microscopy, inheritance.

2. Introduction

Accumulation of misfolded protein aggregates is triggered by environmental stress conditions, which in turn compromise cell function. However, cells have evolved to respond to these changes to maintain metabolic function and ensure their survival. In eukaryotic cells, systems such as the temporal protein quality control (PQC) sustain the proteome and actively contribute to the detection of misfolded proteins (1, 2), their refolding mediated by chaperone proteins (2, 3). The degradation of damaged proteins is actively mediated by the ubiquitin-proteasome system (UPS) (4, 5) but not all proteins are recognised this way, and

other selective processes exist to degrade proteins, such as the autophagy pathway (6). Generally, these systems require acute control of the temporal and spatial dynamics of subcellular components for quality control *in vivo* to prevent or clear aggregates and maintain proteomic homeostasis (2, 3, 5).

When quality control responses and processes fail, misfolded proteins accumulate in the intracellular environment with a heterogeneous size distribution of aggregates (7, 8), consistent with diffusion-nucleation mechanisms of formation (9). This distribution of protein aggregates is harmful to the cell (10, 11), with endogenous protein aggregation effectively depleted from the cellular environment. Further toxicity is mediated by aggregation through perturbation of other functional proteins present in the crowded intracellular environment (12, 13). Ultimately, this can lead to pathogenic phenotypes (14, 15). Many neurodegenerative diseases (e.g. Parkinson's, Alzheimer's) are associated with a process which involves aggregation of amyloid result in packed beta-sheet structures and fibres (16-18), due in part to amyloid- β oligomerization (19). Other diseases such as cataracts (20) or Huntington's disease (21) result from the formation of amorphous aggregates (22, 23). Understanding the formation of such proteotoxic factors is crucial to elucidating underlying mechanisms associated with cellular malfunction and toxicity. Insight into the associated *in vivo* dynamics of these factors can also contribute to the development of new therapeutic methods.

The budding yeast, *Saccharomyces cerevisiae*, has been used to investigate several important processes affecting intracellular organisation which are highly conserved across all eukaryotes, including key survival mechanisms (24, 25), essential metabolic pathways such as DNA replication (26, 27), transcription (28, 29), membrane trafficking pathways (30-33), and PQC machinery for aggregate detection and clearance (3, 34, 35). In light of its excellent genetic tractability, and ease of cell culturing and optical imaging, we used budding yeast as a eukaryotic cellular model to investigate intracellular dynamics of aggregation. Various markers for aggregation use key conserved proteins present in yeast. Chaperone proteins are a good example of this; considered a first response against misfolded proteins, they are recruited at the site of misfolded proteins or aggregates to promote re-folding or initiate degradation pathways if necessary (36, 37). Current approaches to analysing and quantifying protein aggregates include optical microscopy with use of fluorescent biomarkers of aggregation, typically using chaperone proteins as reporters (e.g. HSP70, HSP40, HSP140) (20, 38-40). Additionally, fluorescent variants prone to form aggregates were developed, such as the thermosensitive mutant of Ubc9 (41) derived from a SUMO-conjugating enzyme and unable to properly fold in yeast cells (42).

Another common marker for aggregation used in *S. cerevisiae* is the engineered reporter Δ ssCPY*, a misfolded version of the vacuolar enzyme carboxypeptidase Y (CPY) which is prone to form aggregates and to be aberrantly localized to the cytoplasm (43, 44). This variant, derived from the native CPY (45, 46) carries a single amino acid mutation with a glycine substituted by an arginine at position 255 (G255R) (44, 47) (Figure 1). This mutation (labelled CPY*) is responsible for its misfolding, and when combined with an N-terminal truncated

signal peptide (Δ ss) results in aberrant localization of this misfolded protein to the cytoplasm. Tagging of Δ ssCPY* with enhanced GFP (EGFP) has been used as a model to uncover PQC (48-50) and sorting dynamics (51, 52), cellular perturbations and protein aggregation dynamics in stressed cells (53, 54). Studies have revealed that protein aggregate interactions and localization *in vivo* have a crucial role in established toxicity (54).

The Δ ssCPY* aggregation reporter is typically expressed from the endogenous *PRC1* promoter, which is problematic as this gene is metabolically regulated, for example being upregulated under certain stress conditions, like nutrient starvation (55, 56). As protein aggregation correlates with cellular abundance of proteins and local protein concentrations, and is often assessed under stress conditions, there are challenges in disentangling phenotypes associated with expression and aggregation in such experiments. Furthermore, EGFP, and indeed several other fluorescent protein tags, has the capacity to dimerize (57-59), which can also potentially introduce challenging artifacts when assessing the aggregation of tagged molecules.

To address the limitations of existing aggregation biomarkers, we present newly developed versions of Δ ssCPY* as reporters for cytoplasmic protein aggregates that are tagged with monomeric fluorescent proteins and are expressed under the control of an inducible promoter which minimises interference with other metabolic processes. These novel reagents, which we denote as an inducible Protein Aggregation Reporter (iPAR), are useful tools to study many aspects of stress induced aggregation.

For iPAR, we replaced the metabolically regulated endogenous promoter (*PRC1*) used to express Δ ssCPY* as reporter with the copper inducible promoter (*CUP1*). The fluorescent fusion tag EGFP was additionally mutated to its definitively monomeric version (mEGFP) to eradicate tag-induced oligomerization effects, improving the model compatibility for intracellular aggregation dynamics studies using optical tools. To enhance the utility of this new aggregation reporter reagent, we have used iPAR with a brighter green monomeric fluorescent protein mNeonGreen as well as the red fluorescent protein mScarlet-I. Using these tools with newly designed aggregate analysis techniques, we are able to quantify the induced aggregation of cytoplasmic factors following cell stress, and also to assess the capacity for mother cells to retain aggregates during the process of asymmetric cell division, during which other cellular organelles such as the nucleus and lytic vacuole are inherited in budding daughter cells. In this study, we comprehensively characterise iPAR using biochemical, cell biology and advanced optical methods, and make it openly accessible as a resource to the research community.

3. Material and Methods

3.1 Strains and plasmids used in the study

The yeast cell strains and plasmids used in this study are listed in Table 1 and 2, respectively and the oligonucleotides used in Table 3.

Parental strain	Genotype	Reference	Figure used
BY4742	MAT α , <i>his3Δ leu2Δ lys2Δ ura3Δ</i>	Brachmann, et al. 1998 (60)	Figs. 2-5
BY4741	MAT α , <i>his3Δ leu2Δ met15Δ ura3Δ</i>	Brachmann, et al. 1998 (60)	Fig. 5
BY4741	MAT α , <i>his3Δ leu2Δ met15Δ ura3Δ</i> <i>NRD1-mCherry::hgrB</i>	Shashkova, et al. 2021 (61)	Fig. 5

Table 1: List of background yeast strains used in this study.

Plasmid	Genotype	Reference	Figure Used
pLS190	pRS316 expressing Δ ssCPY*-EGFP from the <i>PRC1</i> promoter	Stolz and Wolf, 2012 (44)	Fig. 2
pCM695	pRS316 expressing -GFP from the <i>CUP1</i> promoter	Laidlaw et al., 2021 (62)	Fig. 2
pLS191	pRS316 expressing Δ ssCPY*-mEGFP from the <i>CUP1</i> promoter	This study	Figs. 2-5
pLS195	pRS316 expressing Δ ssCPY*-mScarlet-I from the <i>CUP1</i> promoter	This study	Fig. 4
pLS196	pRS316 expressing Δ ssCPY*-mNeonGreen from the <i>CUP1</i> promoter	This study	Fig. 4
pCM264	Mup1-EGFP from the <i>MUP1</i> promoter	MacDonald <i>et al.</i> , 2015 (63)	Supplementary Fig. 1

Table 2: List of plasmids used in this study.

3.2 Plasmid construction

The initial iPAR fusion construct *CUP1*- Δ ssCPY*-mEGFP was generated using several cloning steps. Initially, the parental plasmid encoding *PRC1*- Δ ssCPY*-EGFP (pLS190) was modified by site-directed mutagenesis using the S1 and S2 primers to incorporate the monomeric A206K mutation in EGFP (65). This template was then used to amplify Δ ssCPY* (oligos cm193 and cm194) and mEGFP (oligos S8 and S9) with compatible regions for Gibson Assembly (64). Δ ssCPY*-mEGFP was cloned between the *CUP1*-promoter and the *CYC1* terminator of pCM695 linearized with *EcoRI* and *HindIII* to generate pLS191 (*CUP1*- Δ ssCPY*-mEGFP). This assembly strategy using two PCRs introduced 5' *XhoI* and 3' *HindIII* restriction sites flanking mEGFP.

pLS191 was linearized with *XhoI* and *HindIII* and Gibson assembly used to exchange mEGFP with mScarlet-I (66) (using oligos S14 and S15, conserving *XhoI* and *HindIII* sites) and mNeonGreen (67) (using oligos S16 and S17) variants of iPAR (respectively denoted as plasmids pLS195 and pLS196).

Oligo Name	Sequence (5'-3')	Description
cm193	GATATTAAGAAAAACAACTGTAACG AATTCATGATCTCATTGCAAAGACCG	<i>CUP1</i> - ΔssCPY* - Forward primer - used to synthesise ΔssCPY* sequence for Gibson Assembly in pCM695
cm194	AGAATCGAGTTAAAAGGTATTGATTTT AAAGAAGATGGAAACGTTCTTGACAC	ΔssCPY*-EGFP - reverse primer - used to synthesise ΔssCPY* sequence for Gibson Assembly in pCM695
S1	CACACAATCTAAACTTTGAAAGATCC	EGFP - Forward primer - used to induce site directed mutagenesis (EGFP to mEGFP)
S2	CAGACAACCATACCTGTC	EGFP – Reverse primer – used to induce single nucleotide mutagenesis (EGFP to mEGFP)
S8	CCACGGTGGTTTCTCTTACTCGAGAG TAAAGGAGAAGAACTTTTCACTGG	Forward primer - <i>Xho</i> I site Gibson assembly for mEGFP
S9	CCAGATATTCTATGGCAAAGCTTTTATT TGTATAGTTCATCCATGCC	Reverse primer- <i>Hind</i> III site Gibson assembly for mEGFP
S5	GGTGTTCCTCAACTGTCGCCGCTGGT AAGG	ORF ΔssCPY* - Forward sequencing primer
S25	AACTAATTACATGATATCGACAAAGGA AAA	Reverse sequencing primer in the CPY terminator - for verification of the EGFP sequence
S3	GGCAGACAAACAAAAGAATGG	Forward sequencing primer in mEGFP sequence - used to verify mEGFP site-directed mutagenesis.
cm3	TGTATCAATTGCATTATAATATCTTCTT GT	Forward sequencing primer in the <i>CUP1</i> promoter - used to verify the ΔssCPY* sequence
S14	GGTGGTTTCTCTTACTCGAGATGGTG AGCAAGGG	Forward primer - <i>Xho</i> I site Gibson assembly for mScarlet-I
S15	CCAGATATTCTATGGCAAAGCTTCTAC TTGTACAGCTCGTCC	Reverse primer - <i>Hind</i> III site Gibson assembly for mScarlet-I
S16	CCACGGTGGTTTCTCTTACTCGAGAT GGTCTCAAAGGAGAGGCC	Forward primer - <i>Xho</i> I site Gibson assembly for mNeonGreen
S17	CCAGATATTCTATGGCAAAGCTTTTATT TATACAGCTCATCC	Reverse primer - <i>Hind</i> III site Gibson assembly for mNeonGreen

Table 3: Primers used for the construction of the initial iPAR fusion construct *CUP1*-ΔssCPY*-mEGFP and subsequent variants using mScarlet-I and mNeonGreen fluorescent proteins.

3.3 Site-directed mutagenesis

The NEB Q5® Site-Directed Mutagenesis Kit (part number: E0554S, New England Biolabs Inc.) was used to perform the mutation responsible for mEGFP following the manufacturer's protocol, with designed primers (S1 and S2, see Table 3) used at a concentration of 10 μM and the template DNA at a concentration between 1 to 25 ng/μl. The reaction mix was incubated for 5 min at room temperature before transformation.

3.4 Gel DNA extraction

To extract linearized plasmid backbones, gel DNA extraction was performed using the “QIAquick Gel extraction kit” (part number: 28706X4, QIAGEN, Ltd.), following the supplier’s instructions. In short, the DNA band of interest (cut from agarose gel following electrophoresis) was transferred into a sterile 1.5 ml Eppendorf tube. QG buffer was added to the tube to dissolve the gel (at a 3:1 volume proportion) and incubated for 10 min at 50°C. The sample was loaded onto a silica-membrane-based spin column (1.5 ml volume) and centrifuged at 13,000 rpm. After discarding the supernatant, the column was rinsed once with 100% isopropanol followed by a wash with PB buffer. A final elution was performed by loading 50 µl of either EB buffer (10 mM Tris.Cl, pH 8.5) centrifuged at 13,000 rpm into a clean, sterile 1.5 ml Eppendorf tube.

3.5 Cell culture

Single colony isolates from frozen stock following 24-48 h growth at 30°C were used to inoculate 5 ml liquid culture of either Yeast Extract-Peptone-Dextrose media (YPD: 2% glucose, 1% yeast extract, 2% bacto-peptone) or synthetic drop-out media lacking uracil (2% glucose, 1x yeast nitrogen base; 1x amino acid and base drop-out compositions (SD -URA, Formedium Ltd, UK), according to cell strains and selection requirements. Yeast cells were grown in the prepared liquid culture to mid-log phase ($OD_{600} = 0.4-0.6$) at 30°C before harvesting for imaging. A 100 mM copper sulphate stock solution was prepared, filter-sterilised with 0.22 µm diameter cut-off filters, and stored at room temperature. For the induction experiments, cells were first grown for 1-4 h in media containing 5 µM copper chelator bathocuproine sulfonate (BCS) before washing and incubation in media containing 100 µM copper sulphate to induce expression via the *CUP1* promotor (68). To promote the formation of aggregates, cells at the log phase were harvested, diluted to approximately $OD_{600} = 0.2$ and heat shocked for 2 h at either 37°C, 42°C or 30°C (the latter temperature being the control condition). The cells were then harvested and prepared for imaging with confocal microscopy.

3.6 Vacuole labelling

To label vacuoles, 0.8 µM FM4-64 was added to 1 ml of cell culture in YPD-rich media and incubated with shaking for 1 h. Cells were then washed two times with SC media then grown for a further 1 h chase period in SC media lacking dye. After incubation, samples were prepared for imaging.

3.7 Sample preparation for imaging

Imaging was performed in “tunnel” slides (69) using 22x22 mm glass coverslips (No. 1.5 BK7 Menzel-Glazer glass coverslips, Germany). To immobilize cells to the surface, 20 µl of 1 mg/ml Concanavalin A (ConA) was added to the tunnel slide. Excess ConA was rinsed with 200 µl of imaging media before 20 µl of cells were added, incubated for 5 min upside down in a humidified chamber to promote cell adhesion. Finally, any unbound cells were removed by washing with 200 µl of imaging media and sealed with fast-drying nail varnish before loading

on the microscope for imaging (70). Time-lapse experiments were performed in 35 mm glass-bottom dishes (Ibidi GmbH, Germany) with similar ConA coating methods adapted to the dishes support (71). 300 µl of 1 mg/ml of ConA were added to the dishes and incubated for 5 min then washed three times with sterile water. The dishes were then dried under a laminar flow hood ready for imaging. Typically, mid-log phase cells were diluted to OD₆₀₀ <0.1 before addition to the ConA coated dish and incubated for 5 min at room temperature. The dish was washed two times with imaging media to remove any unbound cells and finally topped with fresh media for imaging.

3.8 Confocal microscopy imaging

Cell strains were excited using 488 nm and 561 nm wavelength lasers on the LSM 880 Zeiss microscopes with a 1.4 NA (Nikon) objective. Intensity and gain were optimised and then maintained for each experiment. Green fluorescence (from mEGFP and mNeonGreen fluorophores) was imaged using 2% laser excitation power and red fluorescence (from the mScarlet-I fluorophore) with 1% power to minimise photobleaching. Detector digital gain was set to 1 with a scanning time of 1.23 seconds per frame. Z stack images to generate 3D movies of cells expressing aggregates were acquired with 0.33 µm thick sections across the sample covering 5-6 µm thickness. FM4-64 vacuolar staining was imaged with the 561 nm wavelength laser at 5% laser power using a bandpass emission filter range set to 578-731 nm. Timelapse imaging was performed by acquiring 10 min intervals of 3 µm thick section slices images over 90 min for optimal cytoplasmic volume visualisation during cell division (As describe in previous work (72)).

3.9 ImageJ image analysis

Confocal microscopy data were analysed using ImageJ/Fiji software (ImageJ 2.14.0/1.54f/Java 1.8.0_322) to extract fluorescence intensities from pre-defined segmentation outlines. Cell outlines were generated either manually using the ImageJ selection tool or in a semi-automated process using the Cell Magic Wand plugin (73). Fluorescent foci within each cell were detected using our bespoke ImageJ macro SegSpot allowing for the selection of a threshold method (within the range of inbuilt thresholding functions available in ImageJ) and object detection function within pre-defined cells outlines or regions of interest stored in ImageJ ROI Manager. Finally, pixel intensities and area parameters of the identified foci are extracted and displayed in an output table (See Supplementary Figure 4). Z stack images were visualized with the 3D project inbuilt ImageJ plugin.

4. Results

4.1 Construction of an inducible monomeric marker for cytoplasmic aggregation in budding yeast cells

The vacuolar hydrolase CPY trafficks through the biosynthetic pathway as an inactive precursor before activation in the yeast vacuole (74). A mutant version of CPY prone to aggregation, denoted CPY* (44), has been used in several previous studies as a model to

assess protein folding and regulatory control of misfolded proteins (74-77). The CPY* variant carries a single amino acid substitution of glycine for arginine at position 255 (G255R) near the enzymatic active site (Figure 1.A – 1.C). Deletion of the N-terminal signal peptide (Δ ss) of CPY inhibits entry to the secretory pathway and consequently the hydrolase mislocalizes to the cytoplasm (78). The Δ ssCPY* mutant, which aggregates in the cytoplasm, serves as a useful marker for protein aggregation (50, 52, 79, 80). However, the endogenous *PRC1* promoter (81) typically used to induce expression of this aggregate marker is metabolically regulated (55, 56); therefore, expression, and aggregation, often vary depending on the specific growth and stress conditions resulting in potential difficulties of interpretation.

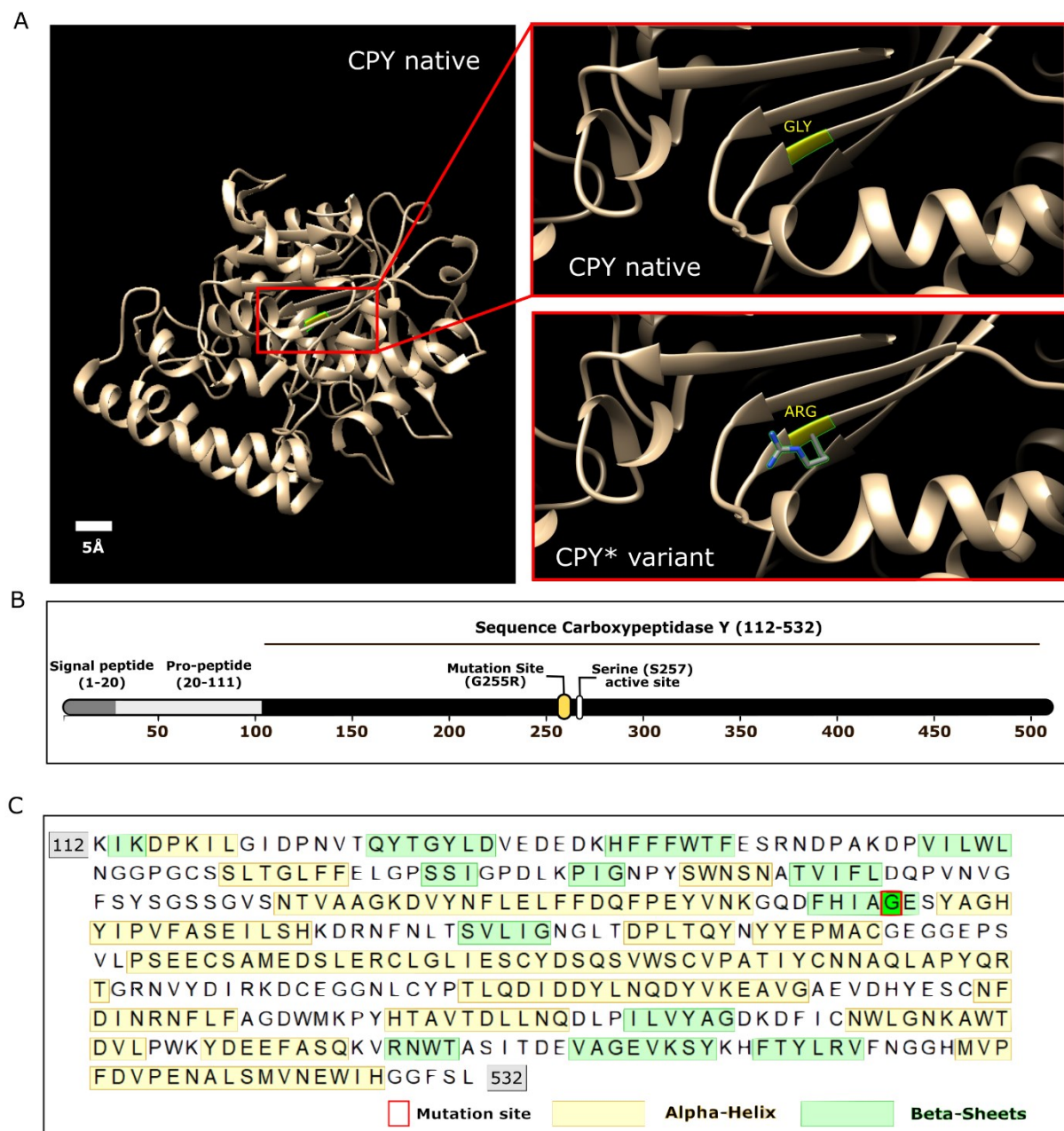


Figure 1: Modifications of CPY to enable its use as a reporter of cytoplasmic protein aggregation. A) Left; a 3D model of the native CPY structure. Right; zoom-in of the mutated region, showing Glycine

residue 255 in the native protein and the arginine substitution in the misfolded CPY* variant. Both amino acids are indicated in yellow. The 3D crystal structure of CPY (PDB ID: 1WPX) was visualised using Chimera software. B) CPY sequence showing G255R mutation site near the S257 active site, responsible for the protein unfolding and aggregative behaviour. C) Sequence for CPY, mutation site and native secondary structures. A red rectangle indicates the position of the mutation site (G255), alpha-helix regions in the native protein are shown in yellow and beta-sheets regions are displayed in green.

To overcome this limitation, we generated a fusion construct which expressed Δ ssCPY* from the copper inducible *CUP1* promoter (82) in the presence of 100 μ M copper sulphate (see Methods and schematic Figure 2.A), using definitive monomeric fluorescent protein tags (monomeric GFP in the first instance) to mitigate against issues associated with fluorescent protein oligomerization. To ensure GFP fluorescence was not affected by levels of copper used in this study, we confirmed that a titration from 0-200 μ M copper sulphate had no effect on a GFP-tagged methionine permease we previously used for membrane trafficking studies (63), assessed by flow cytometry compared to wild-type cells lacking a GFP-tagged protein used as background control (Supplementary Figures 1 and 2).

Copper-dependent expression levels of *CUP1*- Δ ssCPY*-mEGFP in budding yeast cells were characterized using confocal microscopy. Induction times from 1-5 h were imaged followed by image segmentation to extract the fluorescence intensity and the integrated pixel volume information of cells and aggregates. We found that expression of Δ ssCPY* could be rapidly induced in the presence of 100 μ M copper sulphate (Figure 2.B), with a strong increase observed after 1 h copper exposure, with slow rates of expression after 2 h and 3 h exposure and ultimate saturation of expression after approximately 4 h (Figure 2.B and C). At 5 h induction we noticed a small decrease in fluorescence intensities which we hypothesize was the result of active aggregates clearance pathways.

A 2 h copper incubation time was selected as a standard induction condition to express the Δ ssCPY*-mEGFP marker to generate a sufficient pool of protein aggregates for subsequent analysis. We noticed that after 2 h expression there was a reasonable level of expression and a number of aggregates forming in the cytoplasm (Figure 2.C).

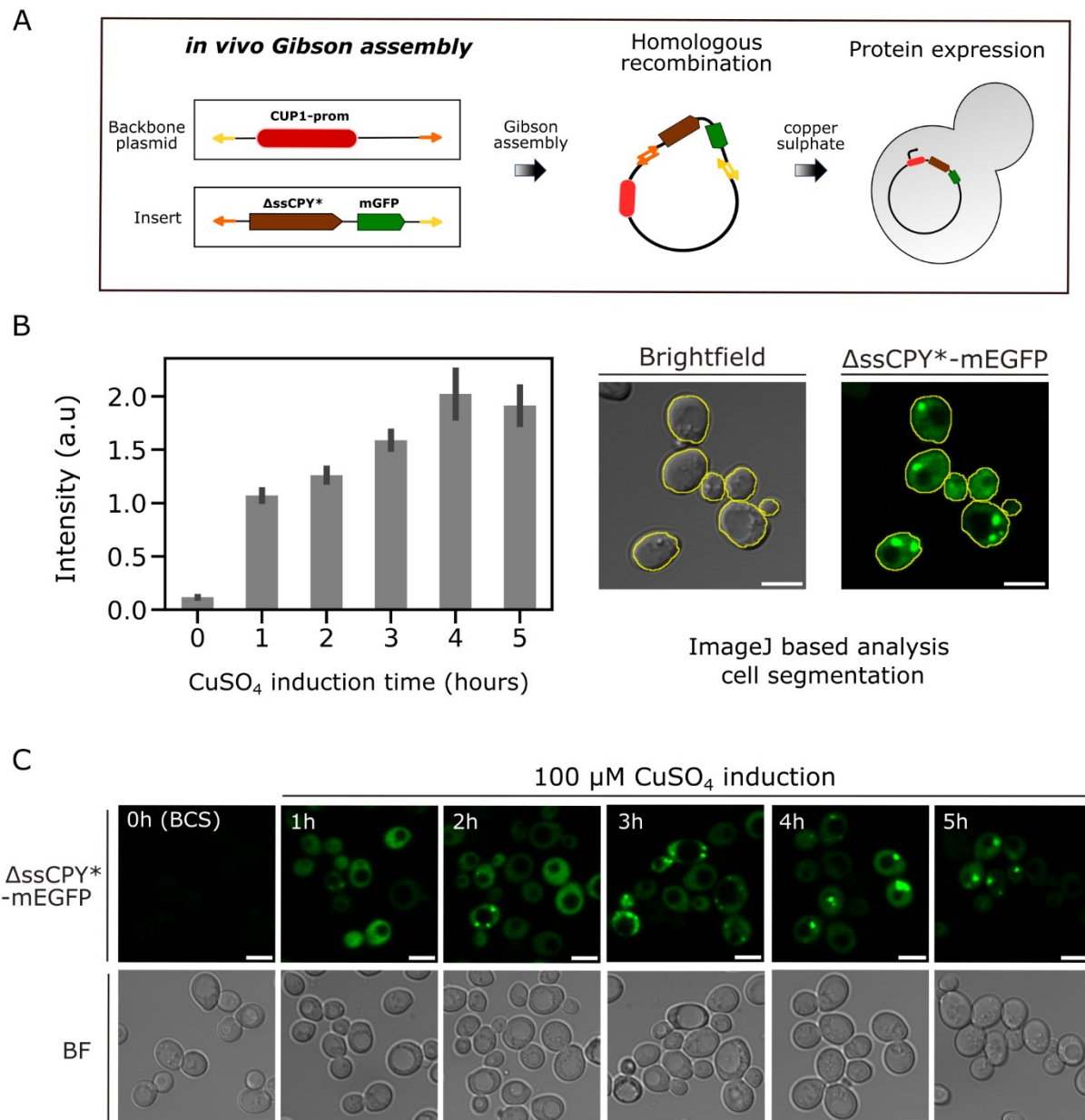


Figure 2: Induction of *CUP1* promoter by copper sulphate results in expression of protein aggregates, visible in confocal microscopy. A) Schematic representation of cloning strategy to produce copper-inducible cytoplasmic Δ ssCPY*-mEGFP aggregates. B) Bar plot for the fluorescence intensity of *CUP1*- Δ ssCPY*-mEGFP incubated in the copper chelator BCS (0 h) or following induction by 100 μ M copper sulphate, at 1 h, 2 h, 4 h and 5 h, n = 100 cells for each condition. SEM error bars represented. The micrographs on the right show cell segmentation using Cell Magic Wand ImageJ tool from brightfield image than was used to measure fluorescence intensity from the GFP channel in each cell. C) Fluorescence micrographs representing the Δ ssCPY*-mEGFP aggregation at different induction time points.

We then characterized the effect of temperature of cells expressing Δ ssCPY*-mEGFP following heat shock. As expected, cells grown for 1 h at 30°C exhibited very few aggregates,

however, shifts to heat stress conditions using temperatures of 37°C or 42°C resulted in visible reporter aggregation (Figure 3.A). There was a significant increase in cells following heat shock at both 37°C or 42°C in comparison to any cells at 30°C that had detectable aggregates of $\Delta ssCPY^*$ -mEGFP (Figure 3.B). A significant increase in number of aggregates was observed, in addition to the number of cells in which aggregates were detected, following heat stress (Figure 3.C).

Between 30°C and 37°C, we observed an increased number of aggregate-positive cells (defined as a cell which contains at least one detected fluorescent focus) by a factor of approximately 2.5, from an average of 19% (± 5.8 , s.d.) to 47% (± 9.4), corresponding to a Student's *t*-test *p* value of 7.59×10^{-5} (i.e., highly significant). Similarly, between 30°C and 42°C, the pool of aggregate-positive cells increased by a factor of approximately 3 from 19% (± 5.8) at 30°C to 59% (± 14.3) at 42°C with a significant *p* value of 5.00×10^{-3} . Although 42°C induced a greater number of aggregate foci across the population, we also detected elevated levels of cell death (Figure 3.A; arrows). Additionally, there was no significant increase in aggregate-positive cells by heat shocking at 42°C compared with 37°C (*p* = 0.261) (see Figure 3.B and Supplementary Table 1).

The total number of detected aggregates increased by a factor of 2.4 from 30°C to 37°C, and by a factor of 4.9 between 30°C and 42°C; and, although the number of aggregate-positive cells was similar between 37°C and 42°C, we still observed a significant increase in the number of aggregates detected (Figures 3.B,C and Supplementary Table 2). We subsequently used 2 h copper induction followed by 1 h heat shock at 37°C as our standard protocol, which we found to be sufficient to induce trackable $\Delta ssCPY^*$ -mEGFP aggregates without compromising the phenotype or viability of the cells.

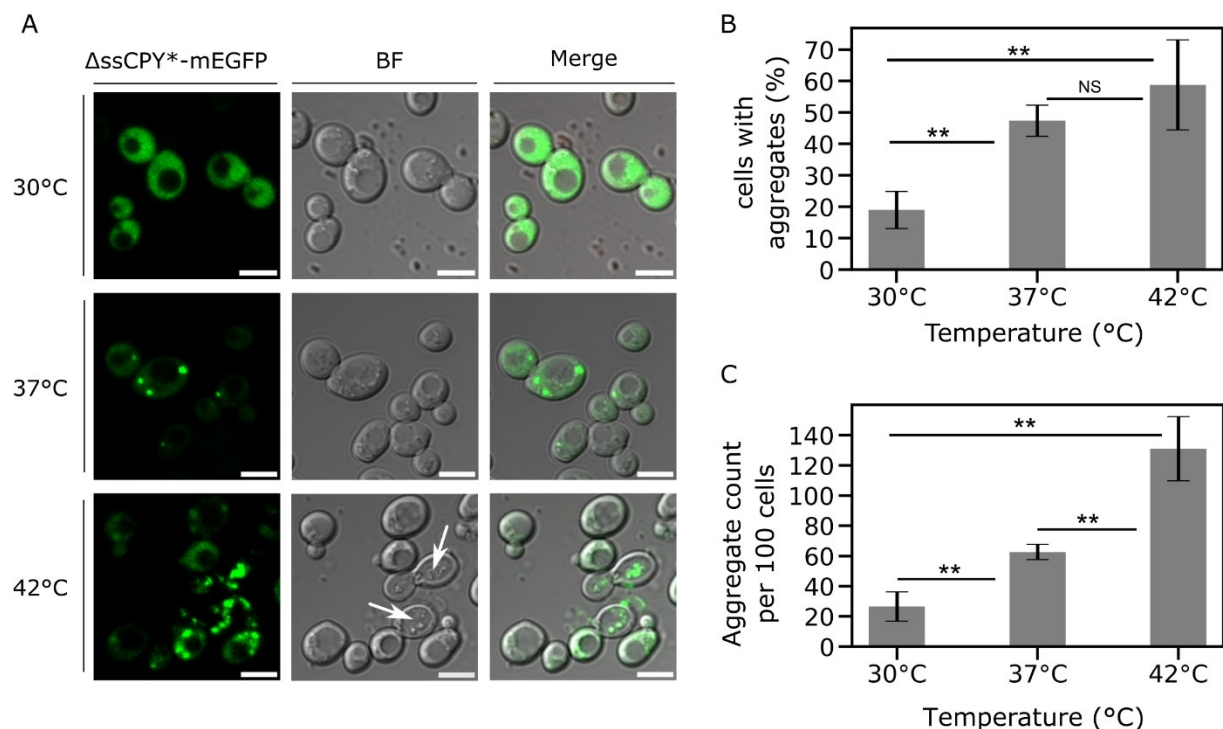


Figure 3: Short-term heat shock induces the formation of aggregates. A) Confocal micrographs from a representative cell population of yeast cells expressing the *CUP1*- Δ ssCPY*-mEGFP protein after induction with copper sulphate for 2 h followed by 1 h at either the initial growth temperature 30°C or the heat shock temperatures of 37°C and 42°C. White arrows indicate dead cells in the brightfield channel, which were not used in subsequent analysis. Scale bar: 5 μ m. B) Bar plot representing the percentage of cells which were positive for aggregates for cells exposed to the control 30°C, or the 37°C and 42°C heat shock. Non-significance is indicated by a Student's *t*-test *p* value ≥ 0.05 , the double asterisk indicates a *p* value < 0.05 . C) Bar plot showing the number of aggregates detected and counted in the cell population, bringing it to *n* = 100 cells in total. See also Supplementary Table 2.

To expand the utility of the iPAR reagent, the mEGFP fluorescent tag was flanked with unique cutting sites (5' *Hind*III and 3' *Xho*I sites) to enable interchangeability and future extension of the construct library (Figure 4.A). We used this strategy to create iPAR variant *CUP1*- Δ ssCPY*-mNeonGreen and *CUP1*- Δ ssCPY*-mScarlet-I, which we found also formed inducible aggregates following the optimised protocol described above in a similar manner (Figure 4.B).

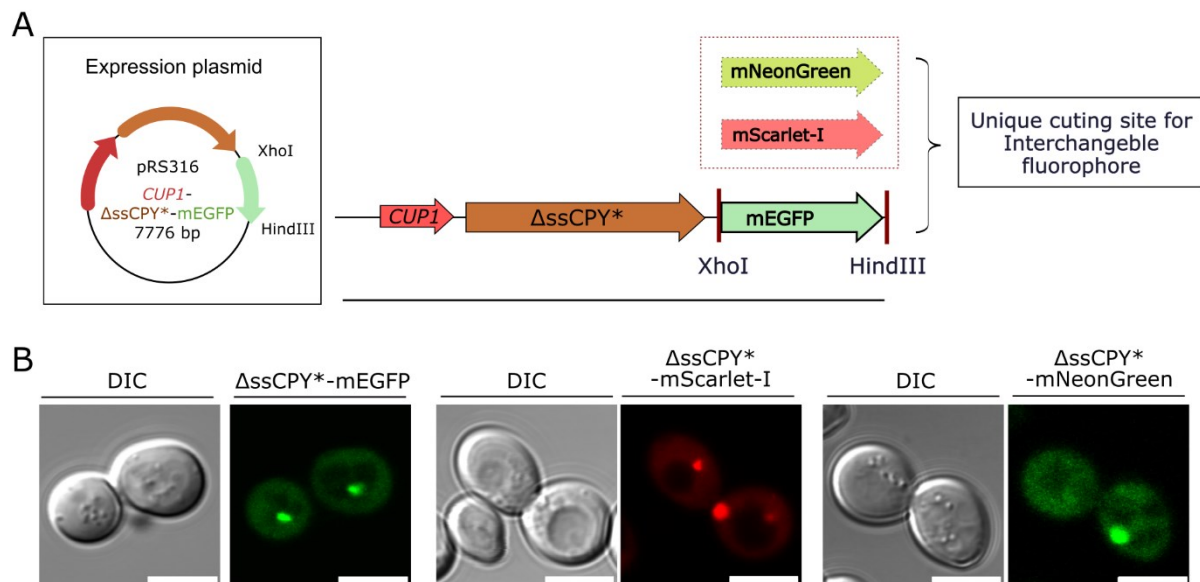


Figure 4: iPAR enables interchangeable monomeric fluorescent proteins to be used for reporting on protein aggregation inside the cytoplasm of living yeast cells. A) Schematic of the expression plasmid constructed for *CUP1*- Δ ssCPY*-mEGFP, the fluorophore with *Hind*III and *Xho*I cutting sites used to facilitate the exchange of fluorescent markers. B) From left to right, micrographs with differential interference contrast (DIC) and fluorescence channel for *CUP1*- Δ ssCPY* in pRS316 with the mEGFP, mScarlet-I and mNeonGreen fluorescent proteins shown respectively.

4.2 Cytoplasmic aggregates and localization in time and space in budding yeast

We performed further characterization of iPAR as a reporter for cytoplasmic aggregation, focussing on spatiotemporal dynamics of newly formed aggregates. We first investigated the number of aggregates and their spatial distributions between mother and daughter cells. Figure 5.A shows the analysis focused on budding cells, where mother and daughter cell images were independently segmented using our bespoke *SegSpot* macro coded for ImageJ

which enabled thresholding and object detection of fluorescent foci (see Methods and Supplementary Figure 4). The area and intensity of fluorescent foci were automatically extracted by this macro, and their values plotted (Figure 5.B). Jitter plots revealed that the mean foci areas measured in mother cells were approximately twice as large as those measured in daughter cells, with a mean focus area of $0.987 (\pm 0.744) \mu\text{m}^2$ measured in mother cells vs $0.393 (\pm 0.291) \mu\text{m}^2$ for daughters (Figure 5.B: left plot and Supplementary Table 3). Conversely, mother cells contained aggregates of higher measured volume with a measured mean fluorescence intensity significantly higher than daughter cells, corresponding to a measured integrated intensity (measured in arbitrary units A.U.) of 52,375 A.U. (± 8512) vs 37,138 A.U. (± 9785) respectively (right plot of Figure 5.B, see also Supplementary Table 3). We note that the distribution of numbers of aggregate foci in both cell types is heterogeneous yet more pronounced in mother cells (Figure 5.B), which was also reflected by high standard deviation values. These results suggest a polarity behaviour of formation/clearance of $\Delta\text{ssCPY}^*\text{-mEGFP}$ during cellular growth resulting in statistically different sizes of aggregates between two cells which are dividing (the older cells displaying larger aggregates with higher intensities than those of the emerging daughter buds).

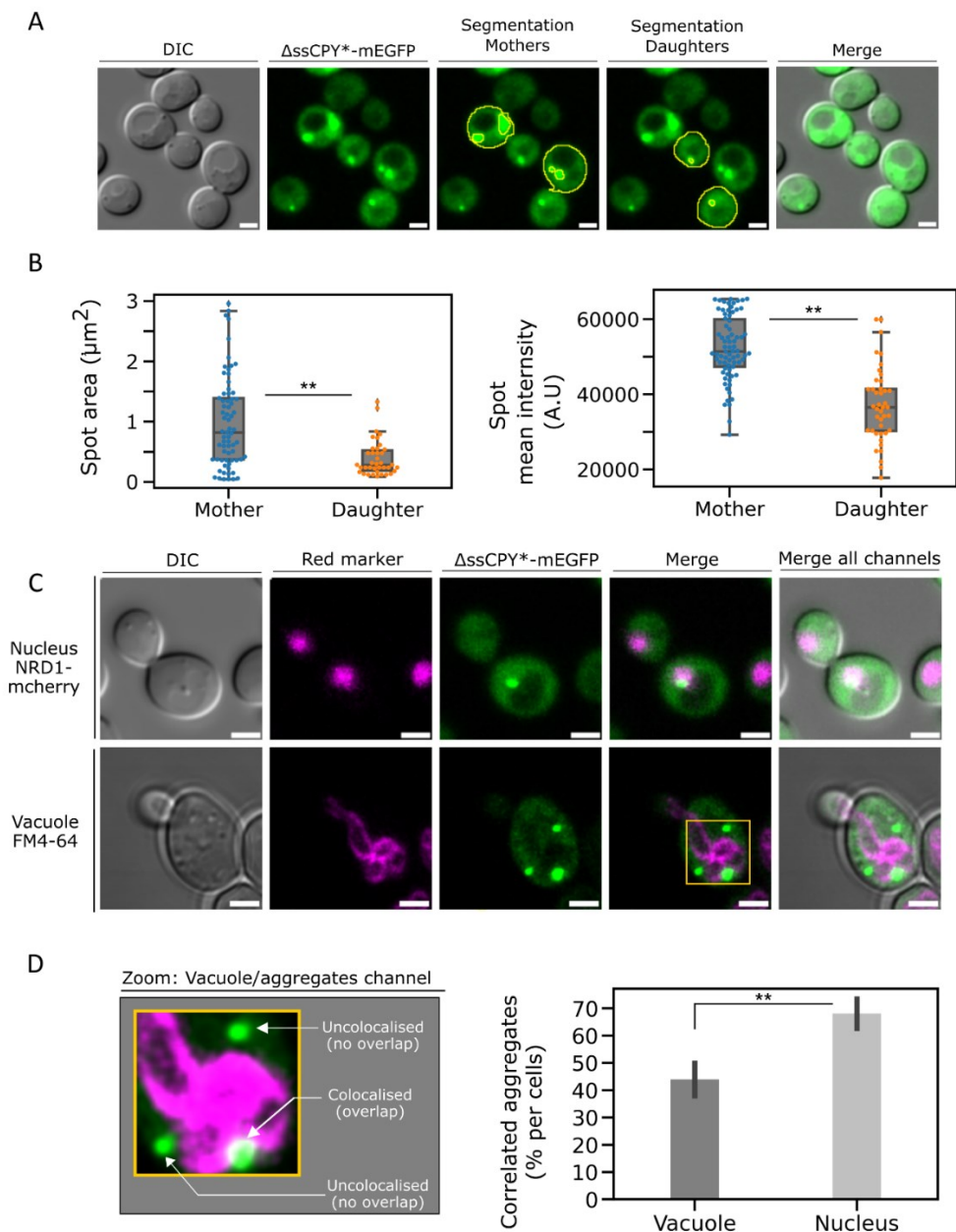


Figure 5: Protein aggregates localize specifically to vacuolar and nuclear compartments. A) Semi-automated segmentation (a combination of the ImageJ selection tool and our bespoke automated macro processing) of mother cells and daughter cells to characterize fluorescent foci. From left to right: DIC image of the cell, fluorescence channel, segmentation of the mother cells, of the daughter cells and merge of the fluorescence channel with the DIC. Scale bar: 2 μm . B) Characterization of aggregate foci, jitter plot of the detected foci area between mother cell and daughter cells. On the right, jitter plot of the intensity measured in each fluorescent focus identified. Outlier detection and removal was performed using standard interquartile methods (83, 84). C) Fluorescence micrographs of dual label strain for simultaneous observation of aggregates and key cellular compartments. Top row shows the nucleus labelled by nuclear reporter Nrd1-mCherry background strain, bottom row shows the vacuole labelled with FM4-64, which mark the vacuole location. Micrographs showing the brightfield, the red channel with the marked compartment of interest, the green channel with the iPAR aggregate reporter and the merge of both fluorescence channels along the brightfield. Scale bar: 5 μm .

We then sought to verify whether iPAR indicated any qualitatively similar spatiotemporal behaviour as reported previously for other cytoplasmic aggregation reporters (54, 85, 86). For example, Δ ssCPY* aggregates were previously shown to be localized in JUNQ and IPOD (87), while inclusion bodies have been observed near the nucleus (88) and the vacuole (89). To elucidate whether our copper sulphate induced Δ ssCPY*-mEGFP colocalized near the membrane of either of the nucleus or the vacuole, we constructed dual colour cell strains including a fluorescent red tag as a reporter for the location of the nucleus or the vacuole. Figure 5.C shows the resulting dual colour images of representative live cells, the top row showing Nrd1-mCherry (90, 91) marking the nucleus, the bottom row showing using FM4-64 pulse-chased labelling to mark the vacuole (see Methods), both simultaneously expressed with Δ ssCPY*-mEGFP.

We quantified the proportion of aggregates present in each cellular compartment, by assessing the proximity/colocalization of both colours (micrographs in Figure 6.D) and found that a mean of approximately 44% of aggregates colocalized with the vacuole compartment and 68% with the nucleus (Figure 5.D). This result is broadly consistent with earlier observations that a significant number of aggregates appear to localize both near the nucleus or vacuole (73). The higher percentage of aggregates identified as being associated with the nucleus may indicate that aggregates preferentially sequester into JUNQ inclusion bodies.

We also acquired 3D data to visualize the patterns of aggregate spatial expressions inside the entire volume of the cell (Supplementary Figure 3 and Supplementary Videos 3-6). 3D projections of cells expressing iPAR, including labels of either the vacuole or nucleus, further confirmed the presence of cytoplasmic aggregates, appearing preferentially in the mother cells and confirming localization in regions that are in the close vicinity (i.e. in putative contact) to the nucleus and vacuole to within our optical resolution limit of approximately 250 nm.

Finally, we performed time-course experiments during cell division with the dual label strain detailed above. In both cases, as a cell divides, we observed protein aggregates sequestered in the mother cell (Figure 6 and Supplementary Videos 1 and 2). We observed that both vacuoles and nuclei were inherited into budding daughter cells whilst aggregates were retained in the mother cells. This observation reinforces the hypothesis that there is a diffusion barrier between mother and daughter cells during cell division (92-94). The sequestration of misfolded cytoplasmic proteins has been reported previously as being a highly conserved quality control process which is crucial to cellular rejuvenation (95-99); the presence of Δ ssCPY* associated with both JUNQ and IPOD inclusion bodies suggests a potential cellular recognition and cellular response for clearance and degradation.

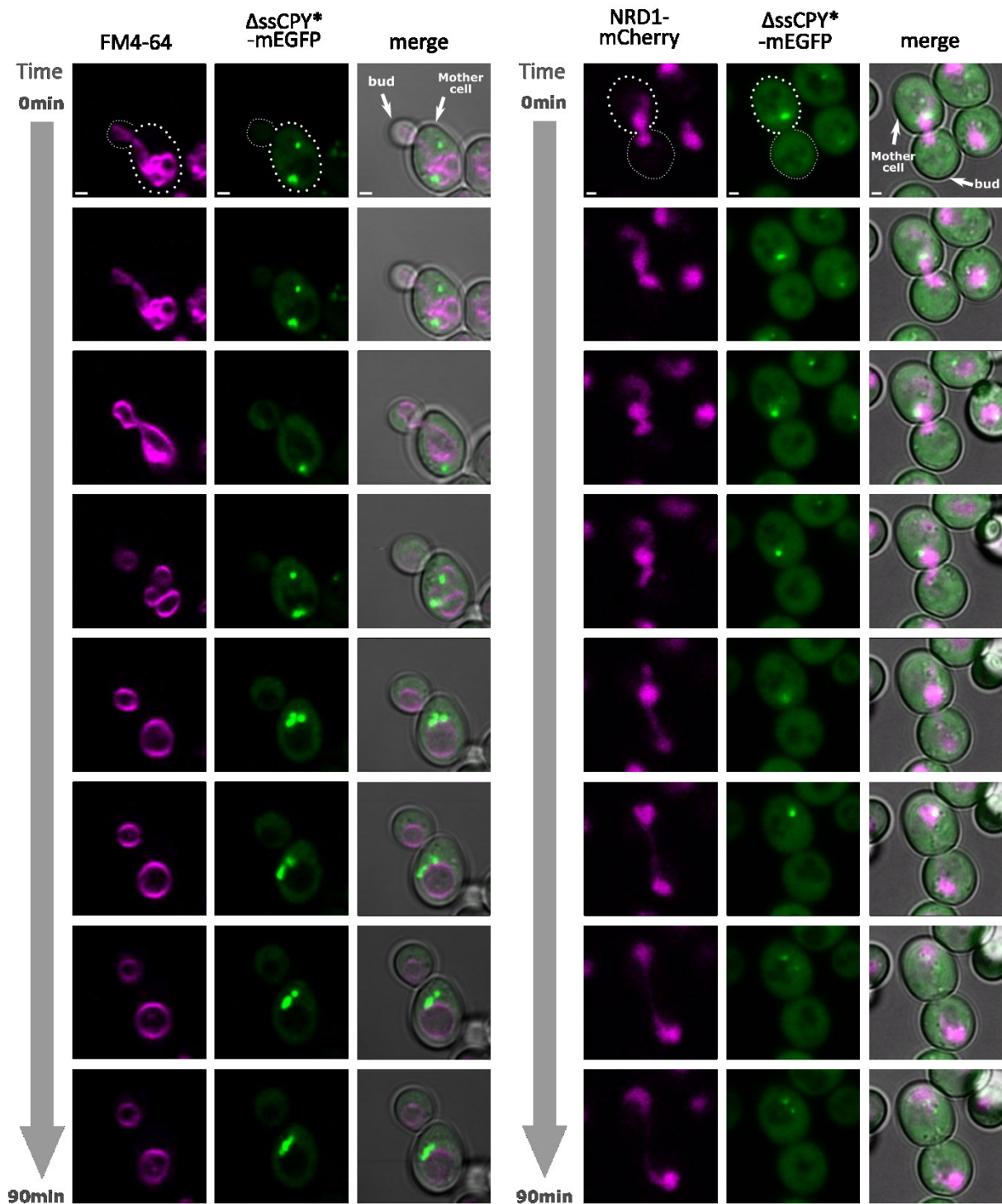


Figure 6: Protein aggregates are localized near to the vacuole and nucleus during cell division. Cells expressing the Δ ssCPY*-mEGFP trackable aggregates (generated after 2 h copper sulphate induction including 1 h heat shock at 37°C) in combination with either Nrd1-mCherry expressed in the nucleus or a WT background strain labelled with FM4-64 at the vacuole, imaged over 90 min during cell division. Micrographs show the red channel for those two markers of interest, the green channel of the imaged aggregate marker and the merge of both fluorescence channels along the brightfield. White arrows indicate the position of the tracked aggregates. Scale bar: 1 μ m.

5. Discussion

We have developed iPAR, an improved reporter for highly precise quantification of cytoplasmic protein aggregation in model budding yeast *Saccharomyces cerevisiae*. By replacing the metabolically regulated *PRC1* promoter with the copper sulphate inducible *CUP1* promoter and introducing definitively monomeric fluorescent tags, iPAR enables precise control of protein expression in growing cells with minimized interference from the fluorescent tag in the aggregation process. These modifications make iPAR a better choice of reporter for stress-related studies and for investigating the dynamics of aggregation, compared to heat shock protein biomarkers of aggregation which use non-monomeric GFP (83). We first characterized iPAR by measuring the expression response of Δ ssCPY*-mEGFP to 100 μ M copper sulphate, indicating that a 2 h standard induction time was optimal to produce a strong fluorescence signal of protein aggregates. We then tested the effects of heat shock on aggregation following inducible expression. At 37°C, we measured a strong increase in aggregate-positive cells (greater than twice as many cells that contain protein aggregates compared to cells incubated at the 30°C no-stress control condition). At 42°C, we observed a similar number of aggregate-positive cells, but we detected a higher total number of aggregates across a population of cells as well as a higher number of aggregates per cell. However, the physiological cell phenotype of 42°C was visibly impaired in several instances, including abnormal morphology and dead cells, consistent with cell metabolic malfunction resulting in an increase in cytoplasmic aggregation. Therefore, we did not select this temperature in subsequent investigations using iPAR.

We verified that induced aggregates localize to the nucleus and vacuole JUNQ and IPOD compartments respectively, as reported from previous studies using existing aggregation reporters. We performed time lapse imaging to quantify the extent of inheritance of the vacuoles and nuclei during asymmetric cell division of iPAR yeast cells in real time, showing that these intracellular organelles are inherited to daughter cells whilst proteotoxic aggregates are retained in the mother cell (see Figure 6 and Supplementary Videos 1 and 2). These real time, simultaneous, same-cell observations are consistent with earlier reports using separate imaging of organelles and aggregates across different cells (86, 95). In budding yeast cells, the presence of multiple inclusion bodies typically observed during osmotic stress were shown previously to be further sequestered in targeted cellular locations (86, 100). Aggregates may be actively recognized by cells and sequestered in the mother cell volume, additionally, physicochemical properties such as local viscosity (101) and the molecular crowding at the junction between the two cells can potentially influence aggregate localization, as suggested by the results of our previous study (72) on the investigation sub-cellular crowding dynamics.

In summary, iPAR offers a robust, improved capability to report on cytoplasmic protein aggregation. We have made the plasmids that encode three fluorescently-tagged variants openly available as a research resource to the scientific community to, we hope, contribute to a wide range of future scientific studies. More generally, our new iPAR technology, if

suitably adapted to different eukaryotic model systems, may help to address several relevant ageing studies and diseases in which protein aggregation is a known or hypothesized factor.

6. Ethics

The presented work did not require the use human subject or animals and therefore was not subjected to a welfare committee.

7. Data accessibility

Raw data can be openly accessible from Zenodo, DOI: 10.5281/zenodo.10468170 (<https://zenodo.org/records/10468171>).

Segmentation analysis code can be openly accessible from <https://github.com/york-biophysics/ImageJ-Macros> (file name: SegSpot.ijm)

The Plasmid construct and cloning maps presented in this article were submitted to Addgene genomic bank under the following ID: 83606 (catalogue number: #212197) and accessible online from <https://www.addgene.org/212197/>.

8. Use of Artificial Intelligence (AI) and AI-assisted technologies

The work presented was not generated using Artificial intelligence tools.

9. Authors' contributions

S.L.: data curation, formal analysis, investigation, methodology, visualization, writing original draft and writing; C.M.: funding acquisition, conceptualization, methodology, resources, supervision, visualization and writing; M.C.L.: funding acquisition, conceptualization, methodology, project administration, resources, supervision and writing.

10. Conflict of interest declaration

The authors declare no competing interests.

11. Acknowledgements

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Supplementary Information

iPAR: a new reporter for eukaryotic cytoplasmic protein aggregation

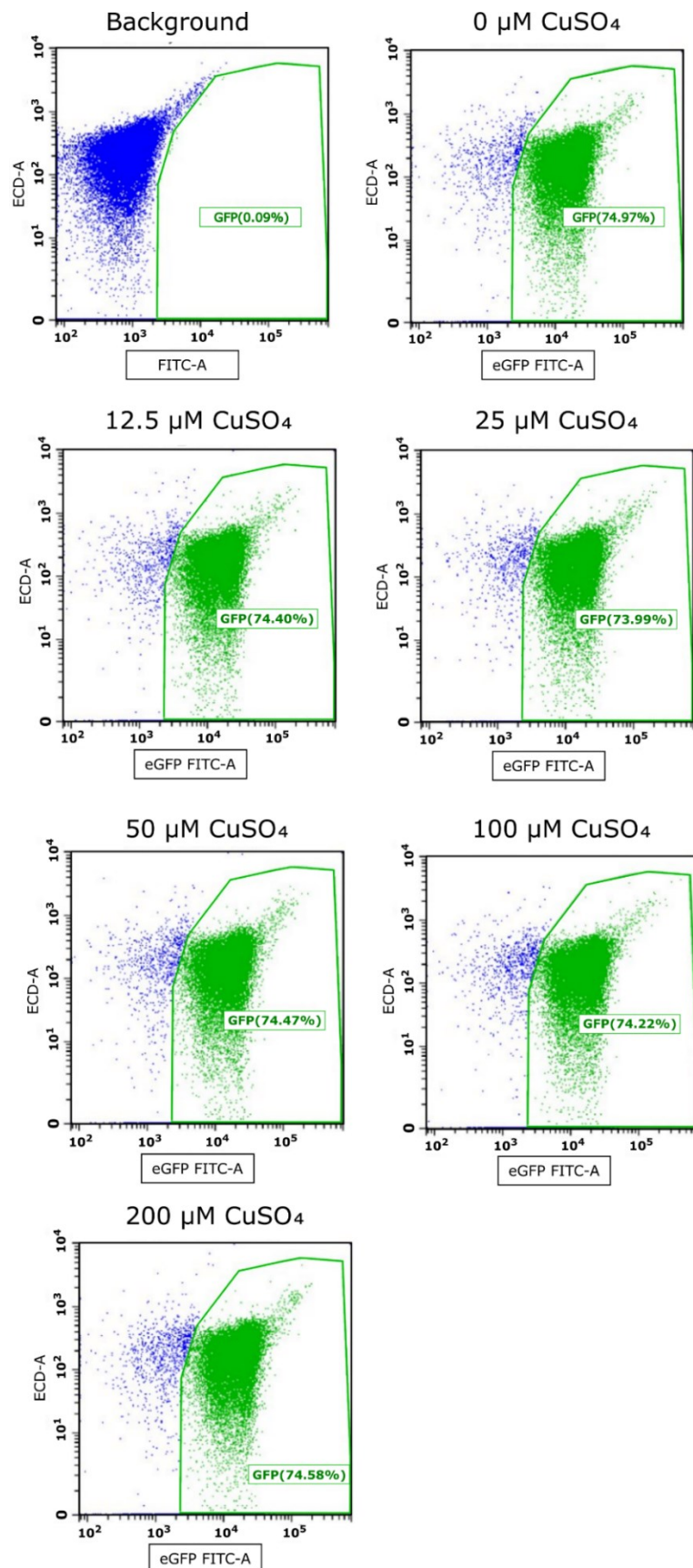
Sarah Lecinski¹, Chris MacDonald^{2,3}, Mark C. Leake^{1,2,3,4}

¹ School of Physics, Engineering and Technology, University of York, York, YO10 5DD, UK.

² Department of Biology, University of York, York, YO10 5DD, UK.

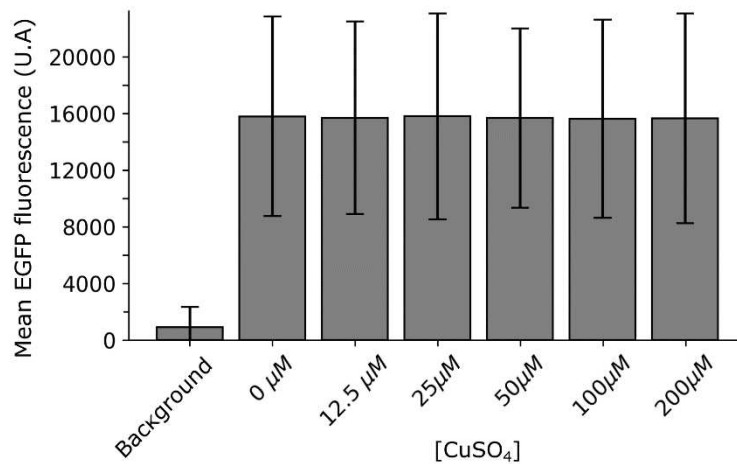
³ York Biomedical Research Institute, University of York, York, YO10 5DD, UK.

⁴ Correspondence to mark.leake@york.ac.uk

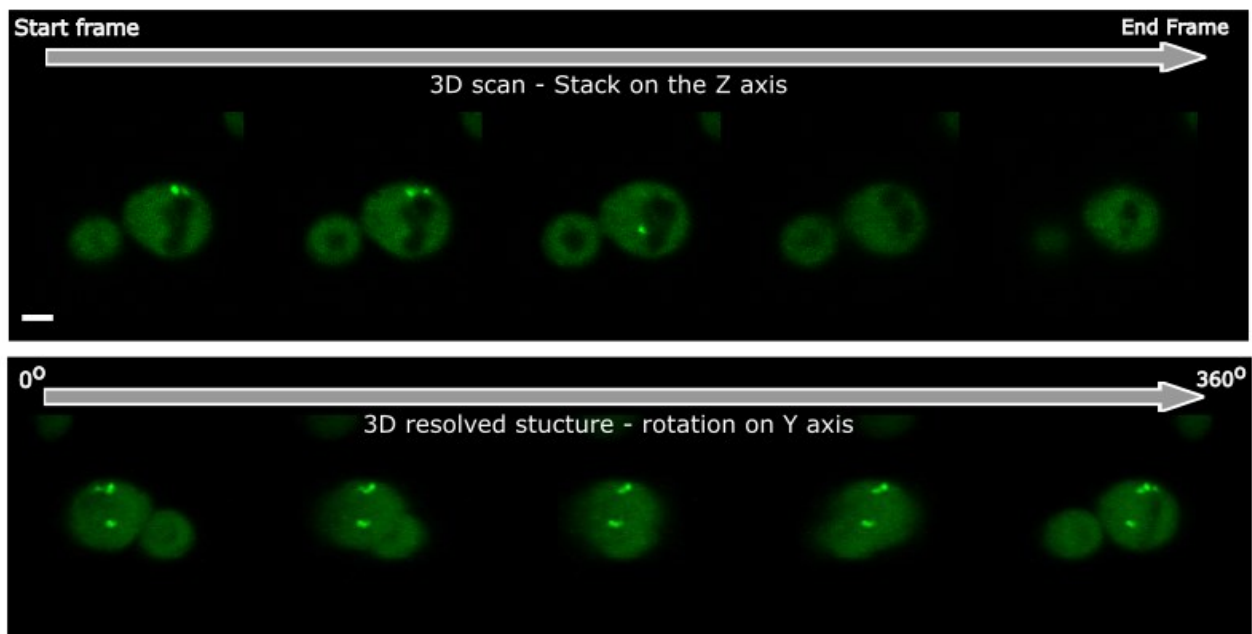


Supplementary Figure 1: Flow cytometry can quantify fluorescence of cells with high throughput, following induction by copper sulphate. Scatter plot representing the presence of fluorescent positive

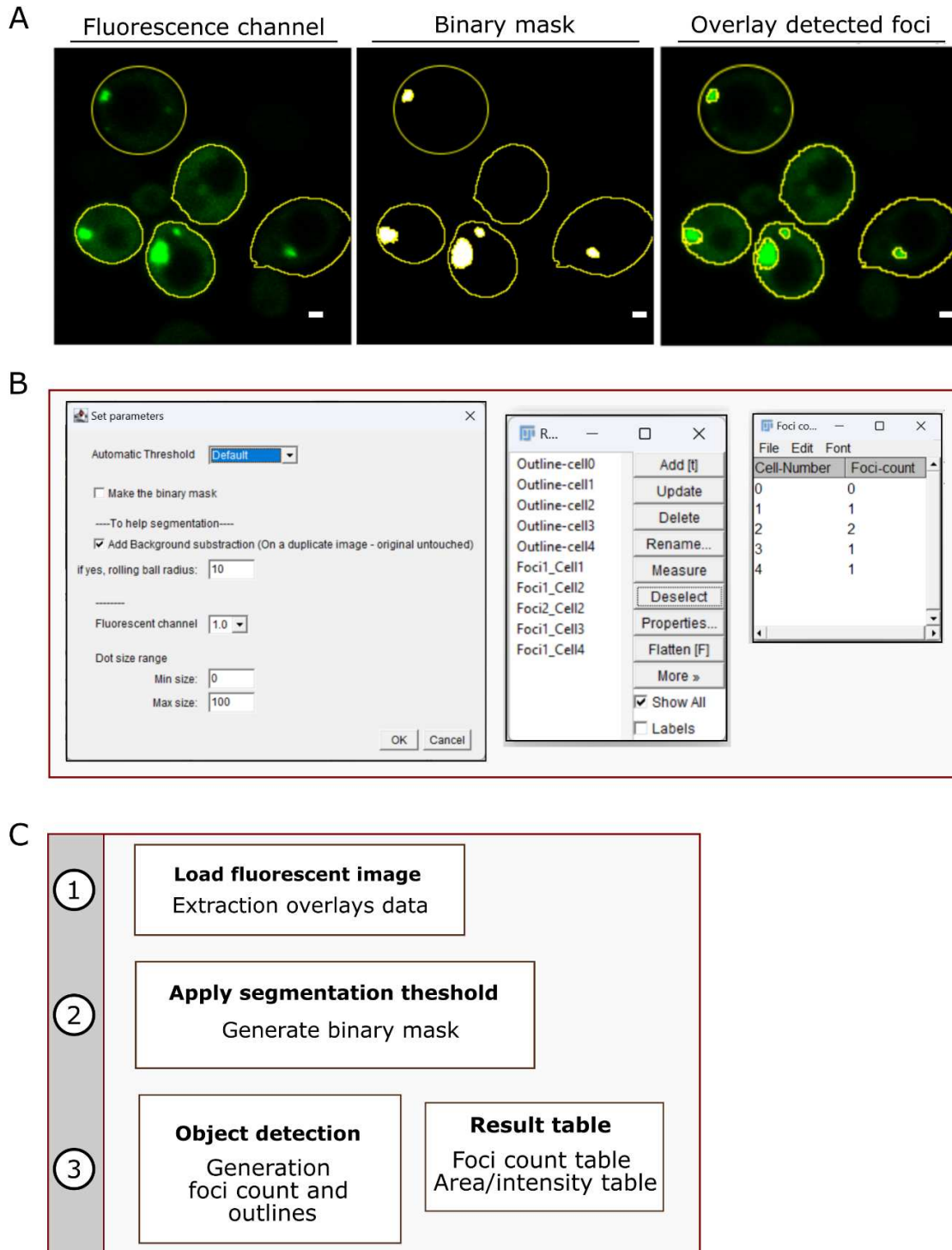
cells in the cell population analysed by flow cytometry. The background non-fluorescent strain was used to calibrate the presence of non-fluorescent cells, in blue colour. Positive cells expressing Mup1-EGFP are visualised in green and the calculated percentage of EGFP positive is indicated in green.



Supplementary Figure 2: Flow cytometry indicates relative insensitivity to fluorescence brightness for different concentrations of copper sulphate. Box plot representing the mean fluorescence measured in cell population expressing Mup1-EGFP in the presence of different copper sulphate concentrations. Error bar SEM. Number of cells $n \geq 10^4$.



Supplementary Figure 3: 3D visualization can be used to determine the localization of protein aggregates throughout the full cell volume. Top micrograph: Z stack of protein aggregates for iPAR reporter using mEGFP, 0.33 μm thickness between frames, scales bar: 2 μm. Bottom micrographs: reconstituted 3D volume of the strain, from the Z stack displayed above and using the ImageJ inbuilt 3D project function. See also Supplementary Video 3.



Supplementary Figure 4: ImageJ macro enables automated, objectivity foci quantification for live cell image data. A) Visual generated segmentation via ImageJ object detection function on the generated binary mask. B) Macro user interface and output sport count table. C) Schematic for the foci detection macro workflow.

	30°C	37°C	42°C
% aggregates positive cells	18.98173	47.39776	58.7738
Standard deviation of %	5.871798	4.943365	16.57905
T-test p value (30 vs 37°C)	7.59 x 10 ⁻⁵		
T-test p value (37 vs 42°C)		0.261	
T-test p value (30 vs 42°C)	5.0 x 10 ⁻³		

Supplementary Table 1: Heat shock and aggregates positive cells

Statistical properties including the p-value taken from Student's *t*-test corresponding to bar plot in Figure3.B

	30°C	37°C	42°C
mean foci count/100cells	26.502	62.644	131.025
Standard deviation of %	9.705	5.035	18.457
T-test p value (30 vs 3°C)	0.000167		
T-test p value (37 vs 42°C)		0.00021	
T-test p value (30 vs 42°C)	2.74 x 10 ⁻⁵		

Supplementary Table 2: Heat shock and aggregates counts

Statistical properties including the *t*-test p value corresponding to bar plot in Figure 3.C

	Mother cells	Daughter cells
Foci mean area	0.987	0.393
Foci median area	0.822	0.284
Standard deviation foci area	0.744	0.291
Foci mean intensity	52375.69	37138.98
Foci median intensity	51452.68	36550.59
Standard deviation foci intensity	8512.51	9785.60
T-test p value (foci area)	8.60 x 10 ⁻⁶	
T-test p value (Foci intensity)	3 x 10 ⁻¹⁴	

Supplementary Table 3: Comparison aggregates area and intensity properties between mother cells and daughter cells.

Statistical properties including the *t*-test p value corresponding to Jitter plots in Figure 5.B

	vacuole	Nucleus
average correlation percentage (%)	43.94737	68.11404
Standard deviation	36.34713	32.67815
T-test p value	0.0057	

Supplementary Table 4: Correlation aggregates and key subcellular compartments

Statistical properties including the *t* test p-value corresponding to Jitter plots in Figure 5.D