Core Fermentation (CoFe) granules focus coordinated glycolytic mRNA localization and translation to fuel glucose fermentation.
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SUMMARY
Glycolysis is a fundamental metabolic pathway for glucose catabolism across biology, and glycolytic enzymes are among the most abundant proteins in cells. Their expression at such levels provides a particular challenge. Here we demonstrate that the glycolytic mRNAs are localized to granules in yeast and human cells. Detailed live cell and smFISH studies in yeast show that the mRNAs are actively translated in granules, and this translation appears critical for the localization. Furthermore, this arrangement is likely to facilitate the higher level organization and control of the glycolytic pathway. Indeed, the degree of fermentation required by cells is intrinsically connected to the extent of mRNA localization to granules. On this basis, we term these granules, core fermentation (CoFe) granules; they appear to represent translation factories, allowing high-level coordinated enzyme synthesis for a critical metabolic pathway.

INTRODUCTION
The glycolytic pathway lies at the core of metabolic activity as a virtually ubiquitous biochemical pathway across living cells. The pathway serves both to supply energy and maintain levels of biochemical intermediates (Bar-Even et al., 2012). Multiple genes express a variety of isoforms for many glycolytic enzymes providing abundant scope for adaptable regulation (Masters et al., 1987; Oparina et al., 2013; Postmus et al., 2012; Warmoes and Locasale, 2014). The pathway was gradually pieced together by a succession of influential biochemists including Meyerhof, Embden, and Parnas (Bar-Even et al., 2012; Barnett, 2005; Schurr and Gozal, 2015). After these major biochemical breakthroughs, interest in central metabolism waned over a period where it was often perceived to perform mundane “housekeeping” functions (Bar-Even et al., 2012; Ray, 2010). More recently, the pathway and its regulation have received renewed interest for various reasons, including connections to cancer and cellular proliferation (Diaz-Ruiz et al., 2011; Gill et al., 2016), moonlighting activities of the glycolytic enzymes (Castello et al., 2015; Kim and Dang, 2005), and increased interest in central metabolism as a focus for metabolic engineering in a synthetic biology era (Lim and Jung, 2017).

In many aerobic cells, the pyruvate produced by glycolysis is transported to and oxidized in the mitochondria via respiration (Gray et al., 2014). However, under anaerobic conditions and in various aerobic cells, such as yeast, lymphocytes, and cancer cells, glucose is fermented through to ethanol or lactic acid. Hence, in these cells glycolysis serves as the major source of ATP and intermediates (Lunt and Vander Heiden, 2011). Indeed, the reduction of pyruvate to ethanol or lactic acid can be viewed as an extension of glycolysis.

Given the critical nature of the glycolytic pathway in energy production and cellular metabolism, it is unsurprising to find that the pathway is regulated by a myriad of different mechanisms. These include direct regulation of the enzymes via substrate and product concentration (Wegner et al., 2015), allosteric enzyme regulation by small molecules (Shen et al., 2016), and post-translational covalent modifications (Shenton and Grant, 2003; Tripodi et al., 2015). Aside from controls of enzymatic activity, other regulatory mechanisms act at the level of gene transcription (Chambers et al., 1995; Yeung et al., 2008), mRNA
processing/stability (Krieger and Ernst, 1994; Lunghi et al., 2015), protein stability (Benanti et al., 2007; Lu et al., 2014; Riera et al., 2003), translation (Daran-Lapujade et al., 2007; Man and Pilpel, 2007), and protein localization (Jin et al., 2017). Although clearly the glycolytic enzymes and the mRNAs that encode them can be regulated, they are often viewed as providing “housekeeping” functions. Indeed, in yeast, many of the glycolytic mRNAs are among the most abundant, heavily translated mRNAs in the cell. This raises obvious questions, such as how is this level of gene expression attained both at the transcriptional and post-transcriptional levels? Furthermore, how is this scale of gene expression coordinated across the pathway such that appropriate levels of enzyme are produced to generate a metabolic flux that is pertinent to the cellular conditions?

A number of recent observations have supplemented the understanding of glycolysis and the role of glycolytic enzymes in cells. For instance, it has become evident that a number of glycolytic enzymes “moonlight” as RNA binding proteins (Castello et al., 2015). Indeed, it has been suggested that many of the glycolytic proteins bind to glycolytic mRNAs to orchestrate control of the pathway (Matia-Gonzalez et al., 2015). In addition, the localization of two glycolytic mRNAs in yeast, PDC1 and ENO2, has been identified as important in their translation control and in the formation of mRNA processing bodies or P-bodies (PBs) after glucose starvation (Lui et al., 2014).

mRNA localization has been commonly considered as a means to generate localized sources of protein, with specific examples involved in cellular polarization identified across many biological systems—ASH1 mRNA in yeast (Long et al., 1997), Bicoid in Drosophila oocytes (Berleth et al., 1988), and Vg1 in Xenopus oocytes (Melton, 1987). In these cases, translationally repressed mRNAs are localized in a transit process involving motor proteins and cytoskeletal elements (Besse and Ephrussi, 2008). Another situation where translationally repressed mRNAs become localized is under stress conditions, where non-translated mRNAs can enter either PBs or stress granules (SGs) to play roles in mRNA degradation and/or storage (Hoyle and Ashe, 2008; Hubstenberger et al., 2017; Jain and Parker, 2013). More global assessments of mRNA localization suggest that the phenomena is widespread: large numbers of mRNA species are localized in Drosophila, neuronal cells, and yeast (Gadir et al., 2011; Lecuyer et al., 2007; Miyashiro et al., 1994; Pizzinga and Ashe, 2014; Zipor et al., 2009; Zivraj et al., 2010). Even so, mRNA localization is rarely thought to play a role in core housekeeping functions such as central metabolism.

In this study, we show that glycolytic mRNAs in yeast and human cells are specifically localized to granules. In yeast, we define the core fermentation (CoFe) granule, a core glycolytic mRNA granule where glycolytic mRNAs are colocalized and translated. Translation is a prerequisite for CoFe granules, and individual mRNA translation is required for localization. Finally, we show that the presence of mRNA granules correlates with the degree of glycolytic function required by the cell. We suggest that the localization of these mRNAs provides a means to generate the scale of protein expression required for such a critical pathway and permits rapid coordinated regulation or complex formation.

RESULTS
Glycolytic mRNAs localize to granules under active growth conditions

Previous work from our laboratory has highlighted that the glycolytic mRNAs, PDC1 and ENO2, encoding pyruvate decarboxylase and enolase, respectively, are translated in cytoplasmic granules (Lui et al., 2014) (Figure 1A). To evaluate whether glycolytic mRNAs in general are localized to these sites, we have utilized the m-TAG system, where elements of the MS2 bacteriophage are used to tether GFP to an mRNA to study its localization in live cells (Haim-Vilmovsky et al., 2011). Accordingly, MS2 stem loops were directly inserted into the 3'UTR sequences of the glycolytic genes at their endogenous genomic loci. The localization of the resulting mRNAs was then followed via coexpression of the MS2 coat protein-GFP fusion (MS2-CP-GFP). It should be noted that MS2-CP-GFP expression alone generates diffuse fluorescence throughout the cell (Figure 1A). In contrast, when MS2 stem loops are integrated into glycolytic mRNA 3'UTRs, the vast majority of the resulting mRNAs are observed in granules (Figure 1B). This includes mRNAs that encode enzymes at every step of the glycolytic pathway (Figure 1C). Notably, not all MS2-tagged mRNAs localize to granules of this kind (Lui et al., 2014; Simpson et al., 2014); for instance, two non-glycolytic mRNAs, GLO1 and PFK26, where the gene products are involved in the control of glycolysis, and the glycolytic mRNA, PYK2, are not observed in granules (Figure 1A). Previously PDC1 and ENO2 mRNAs were shown to localize to ~20 granules per cell (Lui et al., 2014), and here we show that 14 of the 15 tested glycolytic mRNAs localize to a similar number of granules (Figure 1D).
Figure 1. MS2-tagged glycolytic mRNAs are localized to granules in S. cerevisiae

(A) and (B) z-stacked images of strains expressing MS2-tagged mRNAs as labeled and the MS2 coat protein GFP fusion. Scale bar: 2 μm.

(C) Diagram of glucose fermentation to ethanol depicting the glycolytic mRNAs investigated in this study.

(D) A dotplot showing the variation in the number of granules per cell for each of the MS2-tagged strains above. n = 50. The mean ± SD are indicated for each strain.

(E) z-stacked images of smFISH performed on strains expressing MS2-tagged mRNAs and the MS2 coat protein GFP fusion. smFISH was performed for the canonical GPM1 gene (smGENE) or the MS2 stem loop sequence (smMS2). Scale bar: 3 μm.

(F) Beeswarm plot showing the proportion of smMS2 foci that colocalize with smGENE foci for a subset of strains expressing MS2-tagged glycolytic mRNAs. Each dot represents a single cell. n > 300.
Recent commentaries have highlighted the potential for an accumulation of fragments of mRNA carrying MS2 stem loops that can impact upon the interpretation of experiments using MS2 tethering systems (Garcia and Parker, 2015, 2016; Haimovich et al., 2016; Heinrich et al., 2017). It is possible that such fragments would accumulate at sites of mRNA degradation. To assess whether this is the case for the granules observed here, a range of approaches were taken. Firstly, it should be noted that all of the experiments presented are conducted on cells actively growing in nutrient replete media. Under these conditions in our experiments, PBs are largely absent (Lui et al., 2014), so the high-level accumulation of mRNA fragments at sites of mRNA decay seems unlikely. Secondly, although most of the glycolytic mRNAs tested are present in granules, the MS2 stem loops have a highly variable impact on the steady state level of the mRNAs: some mRNAs are stabilized, others destabilized, and some remain unchanged (Figure S1A). This profile is not consistent with MS2 fragments explaining the observed localization. Thirdly, the major mRNA species observed on Northern blots under active growth conditions for either the non-tagged or MS2-tagged ENO2 and PDC1 mRNAs were full-length mRNAs (Figure S1B). In contrast, in stressed cells where PBs are present, such as shortly after glucose depletion, degradation fragments for the MS2-tagged mRNAs comprise a high proportion of the total mRNA (Figure S1B). Fourthly, a subset of granule localizing glycolytic mRNAs has been tagged with a version of the MS2 system (termed the “version 6” system) where MS2 fragments do not accumulate (Tutucci et al., 2018). This new MS2 system reveals an identical granular pattern of glycolytic mRNA localization to the original MS2 system (Figure S1C). Finally, a single-molecule fluorescent in situ hybridization (smFISH) strategy where probes were targeted to either the MS2 stem loops or the body of the mRNA revealed greater than 75% signal overlap between these two probes (Figures 1E and 1F). This result suggests that the MS2 region of the mRNA reports the localization of full-length mRNAs. In addition, significant overlap is seen between the MS2-CP-GFP protein signal and either the MS2 RNA probe signal or mRNA body probe signal suggesting that the GFP signal also reports full-length mRNAs (Figure 1F). This overlap is observed despite the fact that the MS2-CP-GFP signal is only seen for those granules that exceed a specific intensity threshold, because, as reported previously, (Pizzinga et al., 2019) the MS2 live cell system predominantly detects multi-mRNA granules. Overall, the combination of different validatory analyses used show that glycolytic mRNAs evaluated using the m-TAG system localize to multi-mRNA granules.

However, insertion of MS2 stem loops could still alter some aspect of an MS2-tagged mRNAs fate. Therefore, in order to provide an independent assessment of the glycolytic mRNA localization, endogenous unmodified mRNAs were evaluated using smFISH. smFISH strategies commonly use ~30–50 fluorescently labeled oligonucleotides that are hybridized to mRNAs in fixed cells (Pizzinga et al., 2019; Tsanov et al., 2016). Because many glycolytic genes are present in yeast as multiple paralogues with very high levels of sequence identity, the use of smFISH to unambiguously study the localization of individual mRNA species is problematic. Therefore, sets of smFISH probes were designed to study the localization of mRNAs encoded by glycolytic genes that either lack paralogues or harbor substantial sequence differences to their paralogues. As a result, four different glycolytic mRNAs, GPM1, FBA1, TPI1, and PGK1, were analyzed and shown to localize specifically to granules (Figure 2A). In terms of the number of granules per cell, the smFISH data for endogenous mRNAs are entirely complementary to the live cell MS2-tagged mRNAs (cf. Figures 2B and 1D). One of the key advantages of the smFISH versus the m-TAG experiments is that for the smFISH data, single mRNA foci are visible. This allows an estimate of mRNA copy number per cell and the proportion of single mRNAs present in multi-mRNA granules (Pizzinga et al., 2019). From the data, it is clear that ~70% of the glycolytic mRNA molecules are present in large granules (Figure 2C). The results correlate well with live cell m-TAG data, where a similar fraction of the PDC1 and ENO2 mRNAs were previously found in multi-mRNA granules (Lui et al., 2014). For the non-glycolytic NPC2 mRNA, which is not localized to large multi-mRNA granules (Pizzinga et al., 2019), a speckled pattern is observed with a reduced, homogeneous fluorescent intensity for each speckle relative to the glycolytic mRNAs. The quantitation of fluorescent intensity profiles shows that the NPC2 mRNA is rarely localized to multi-mRNA granules, and even where NPC2 multi-mRNA granules can be identified, the proportion of mRNA present is very low (Figures 2A–2C). Overall, these data confirm that glycolytic mRNAs are housed in large cytoplasmic bodies or granules and, combined with our concurrent studies on translation factor mRNAs (Pizzinga et al., 2019), indicate that the localization observed with the MS2 system in actively growing cells can be representative of and meaningful to the localization observed for an untagged mRNA.

**Glycolytic mRNAs colocalize to the same RNA granules**

Our previous assessment of the ENO2 and PDC1 mRNAs suggested that these two mRNAs colocalize to the same set of granules (Lui et al., 2014) (Figure 3A) but are distinct from the translation factor mRNA...
granules we have recently described as factories for the production and inheritance of the translation machinery (Pizzinga et al., 2019). On this basis, we speculated that the colocalization of glycolytic mRNAs might generally allow a concerted production and/or regulation of the pathway of the glycolytic enzymes. More recent work highlights the potential for cotranslational assembly of components of the glycolytic pathway (Shiber et al., 2018), which again hints that actively translating glycolytic mRNAs might colocalize. In both the live cell and fixed cell mRNA localization experiments presented here, both the pattern and number of mRNA granules in a cell is remarkably similar across the different glycolytic mRNAs (cf. Figures 1Ca and 2A; cf. Figures 1Da and 2B). This similarity is consistent with a model where many of the glycolytic mRNAs colocalize to the same site.

In order to directly assess glycolytic mRNA colocalization, we made use of a PP7/MS2 system, which allows the simultaneous visualization of two mRNAs in the same live cell (Hocine et al., 2013; Lui et al., 2014; Pizzinga et al., 2019). A series of yeast strains were generated carrying PP7-tagged ENO2 mRNA and another MS2-tagged glycolytic mRNA. By coexpressing the MS2 and PP7 coat proteins fused to mCherry and GFP respectively, the localization of each MS2-tagged mRNA was compared directly with that of ENO2 mRNA in the same living cell. As previously shown (Lui et al., 2014), we observed a strong colocalization of PDC1 with ENO2 using this system (Figure 3A). Equally, for many of the glycolytic enzymes tested, a high degree of colocalization with the ENO2 mRNA pattern was observed (Figure 3A). Interestingly, however, despite the high sequence homology between ENO2 and ENO1 mRNAs, these mRNAs localized to discrete foci (Figure 3A).

In order to corroborate the colocalization observed using the PP7/ MS2 systems, we assessed pairwise colocalization of endogenous unmodified glycolytic mRNAs using smFISH (Figure 4A). Because, in contrast to the MS2 system where only multi-mRNA granules can be followed, the smFISH technique detects all of the probed granules we have recently described as factories for the production and inheritance of the translation machinery (Pizzinga et al., 2019). On this basis, we speculated that the colocalization of glycolytic mRNAs might generally allow a concerted production and/or regulation of the pathway of the glycolytic enzymes. More recent work highlights the potential for cotranslational assembly of components of the glycolytic pathway (Shiber et al., 2018), which again hints that actively translating glycolytic mRNAs might colocalize. In both the live cell and fixed cell mRNA localization experiments presented here, both the pattern and number of mRNA granules in a cell is remarkably similar across the different glycolytic mRNAs (cf. Figures 1C and 2A; cf. Figures 1D and 2B). This similarity is consistent with a model where many of the glycolytic mRNAs colocalize to the same site.
mRNA present in a cell, the signal is very congested. This congestion is particularly apparent for glycolytic mRNAs, which are typically estimated to be present at >100 copies per cell (Lahtvee et al., 2017). So, in order to objectively measure pairwise colocalization, a computational strategy was developed. In short, the distance between the centroid of a granule for one mRNA species was measured relative to the centroid of the nearest neighboring granule for the other mRNA species. A spot was deemed to colocalize if its centroid was within the sphere of a spot in the opposite channel (Figure 4B). Using this method, a significant proportion (~50%–60%) of the glycolytic mRNA granules were deemed to overlap with one another (Figure 4C). Because each mRNA is present in ~20 multi-mRNA foci per cell (Figure 2B) with a similar number of single mRNA foci, we were concerned that high levels of colocalization could simply stem from the proportion of cytosolic space occupied by the mRNA foci. Therefore, to control for this, we established a Monte Carlo simulation model (Fletcher et al., 2010), where the position of real foci were randomized within the cell accounting for vacuole space and cross-compared with randomized simulated foci for a second mRNA. From this analysis it is clear that relative to the simulated model, the various tested glycolytic mRNAs display significant colocalization (Figure 4C). Although these smFISH results mirror the colocalization observed using the MS2 and PP7 stem loop systems (Figure 3), the scale of colocalization appears lower. One possible reason for the lower colocalization reflects the extra sensitivity of the smFISH technique in detecting single mRNA foci, which may not colocalize to the same extent as multi-mRNA granules. To explore this, we considered only the most intense foci in the smFISH data, which likely represent multi-mRNA containing foci. This analysis revealed a positive shift in the level of colocalization by ~10%–20% (Figure 4C). Importantly, simulated controls using these brighter, larger spots did not display this shift in colocalization (Figure 4C).

Overall, the smFISH results combined with the live cell studies reveal that in optimally growing yeast there is a high-degree of colocalization to multi-mRNA granules for the majority of glycolytic mRNAs tested. Not

Figure 3. Glycolytic mRNAs colocalize to granules in actively growing cells
(A) and (B) z-stacked images show the localization of various MS2-tagged mRNAs (via coexpression of the MS2-CP-mCh fusion) relative to the ENO2-PP7 mRNA (visualized using coexpression of the PP7-CP-GFP fusion). The percentage of observable tagged mRNA colocalizing with the PP7-tagged mRNA is indicated ± SD. Scale bars: 2 μm.
every localized glycolytic mRNA colocalizes, for instance the ENO1 mRNA does not, but the vast majority do. These colocalized mRNAs encode enzymes that catalyze most of the reactions that are required for glucose fermentation to ethanol; therefore, we have termed the granules "core fermentation" mRNA granules or CoFe granules.

Glycolytic mRNA granules do not reside on the ER or mitochondria

A number of studies have described the role of organelles in the localization of specific mRNAs in yeast (Fundakowski et al., 2012; Gadir et al., 2011). Given data showing that mRNAs encoding non-organellar proteins can also be enriched with organelles (Jan et al., 2014), it is plausible that the localization of glycolytic mRNAs to multi-mRNA granules could be occurring on specific organelles such as the ER or mitochondria. Although the profile of the glycolytic mRNAs does not necessarily match the known localization pattern for these organelles, we could not rule out that the CoFe granules are somehow anchored or stabilized by interactions with these organelles. In order to test this idea, the localization of the glycolytic mRNAs was assessed in strains carrying the fluorescent tagged organelle markers Sec63p (ER) and Cox4p (mitochondria) (Figure S2). For both markers little, if any, evidence of colocalization was observed. Instead, the glycolytic mRNA granules appear largely separated from either the ER or the mitochondria, suggesting that their structure and localization are not reliant upon either of these organelles.

mRNA translation both occurs in and is required for localization to CoFe granules

Previously, we have shown that in contrast to most mRNA containing granules, which carry translationally repressed mRNAs, the granules housing the glycolytic mRNAs PDC1 and ENO2 are sites where these mRNAs are translated (Lui et al., 2014). A variety of experiments supported this hypothesis, including data from FRAP assays where newly synthesized unbleached protein accumulated in the granules (Lui et al., 2014). In addition, treatment of cells with cycloheximide, which freezes ribosomes on mRNA, caused an increase in the number of mRNA granules. Furthermore, the quantification of ribosome-associated mRNA relative to granule-associated mRNA showed that even though 95% of these mRNAs are translated, at least 70% are present in granules. Finally, under polysome run-off conditions a rapid coalescence of the
Figure 5. mRNAs translation in CoFe granules is required for localization

(A) Schematic of TRICK reporter system. Ribosomes on translated RNAs “knock off” the PP7-CP-GFP fusion, whereas on untranslated RNAs the coat protein remains bound.

(B) z-stacked images of TRICK-tagged mRNAs coexpressing the MS2-CP-mCh fusion and the PP7-CP-GFP fusion, in + and – glucose. Scale bars: 3 μm.

(C) % of mCherry only granules

(D) * STOP codon (sc)

(E) PDC1

(F) pPDC1

(G) mRNA granules per cell

(H) Relative fold change
mRNA granules to form PBs was observed, suggesting that prior to the stress the mRNAs were in a distinct granule, being actively translated.

To extend this analysis further, a technique called TRICK (translating RNA imaging by coat protein knockoff) was used, which allows visualization of translation in live cells (Halstead et al., 2015; Pizzinga et al., 2019). This technique relies upon observations that a PP7 coat protein fusion bound to PP7 stem loops upstream of the STOP codon is displaced under active translation conditions, whereas the MS2 coat protein fusion tethered downstream of the STOP codon remains associated (Halstead et al., 2015) (Figure 5A). For PDC1 mRNA under active growth conditions, most granules observed only carry the MS2-CP-mCherry (MS2-CP-mCh) fusion protein (Figures 5B and 5C). In contrast, after a 10-min glucose depletion to elicit a robust and global inhibition of protein synthesis (Ashe et al., 2000), both MS2-CP-mCh and PP7-CP-GFP colocalize to mRNA granules to form PBs was observed, suggesting that prior to the stress the mRNAs were in a distinct granule, being actively translated.

Another key question is whether translation of a molecule of mRNA is a requirement for entry into the granule. In order to address this question, we selected the PDC1 mRNA and sought to limit its translation, then assess the impact on localization. More specifically, we adopted two different strategies toward reducing PDC1 mRNA translation. In the first approach a STOP codon was inserted immediately downstream of the translation START codon (PDC1-sc) (Figure 5D). We reasoned that this would severely reduce the number of ribosomes associated with this mRNA and significantly increase the pool of non-translated PDC1 mRNA. As a second strategy, a stem loop was inserted into the mRNA downstream of the translation START codon (PDC1-sl) (Figure 5G). Introduction of this well-characterized stem loop (ΔG value of ~41 kcal/mol) has previously been shown to reduce translation of specific mRNAs by limiting scanning of the 43S preinitiation complex through to the AUG Start codon (Palam et al., 2011; Pizzinga et al., 2019; Vattem and Wek, 2004). In strains carrying these altered PDC1 mRNAs, mRNA localization was followed relative to the non-modified mRNA using the MS2 system.

Introduction of either the STOP codon or the stem loop structure into the PDC1 mRNA dramatically reduced the number of PDC1-MS2 mRNA granules: decreasing from ~20 granules per cell to less than 5 (Figures 5E–5G). Coincident with this effect on the number of mRNA granules in the cell, both strategies used to limit PDC1 mRNA translation also resulted in reduced mRNA levels (Figure 5H). Insertion of the STOP codon caused an ~8-fold reduction in PDC1-MS2 mRNA, whereas stem loop insertion reduced mRNA levels ~2-fold. The reduction of mRNA caused by the introduction of the STOP codon is consistent with premature STOP codons leading to nonsense mediated mRNA decay (Hagan et al., 1995). The impact of the stem loop on PDC1 mRNA levels is not as pronounced as the STOP codon insertion, and it is a little surprising that this insertion leads to mRNA destabilization, as this same stem loop has been inserted into a number of mRNAs without impacting upon overall mRNA levels (Palam et al., 2011; Pizzinga et al., 2019; Vattem and Wek, 2004). This suggests that the context of a stem loop in the 5’UTR of an mRNA is important in determining to what extent the insertion impacts upon the fate of the mRNA. These results highlight the intimate connection between the translation of an mRNA and its stability and add to many observations showing that a reduction in translation can lead to mRNA destabilization (Roy and Jacobson, 2013).
Figure 6. CoFe granule number varies with quality of carbon source

(A) z-stacked images of PDC1-MS2 or ENO1-MS2 mRNA in strains coexpressing the MS2-CP-GFP fusion, grown in SC media with either 2% glucose, 2% raffinose, 2% galactose, or 3% ethanol. Scale bar: 2 μm.

(B) Quantification of mRNA granules per cell for PDC1 and ENO1 mRNA in strains grown in the different carbon sources (Gluc = Glucose, Raff = Raffinose, Gal = Galactose, EtOH = Ethanol). n = 50. Error bars are ±SD. *p < 0.005 relative to other columns from a 1-way ANOVA and the Tukey’s HSD test.

(C) Graph representing the gene expression ratio of PDC1 or ENO1 in strains (yMK1586 and yMK2468) grown in the different carbon sources. The gene expression ratio was calculated according to the Pfaffl method. This approach...
Figure 6. Continued

considers the PCR efficiency for the different genes and is expressed as the change in target gene levels (PDC1 and ENO1) between glucose conditions and the different carbon sources over the change in reference gene levels (ACT1) between glucose conditions and the different carbon sources (Pfaffl, 2001). (Gluc = Glucose, Raff = Raffinose, Gal = Galactose, EtOH = Ethanol). Error bars are ± SD.

(D) z-stacked images of FBA1, GPM1, and TPI1 mRNAs probed using smFISH in DAPI-stained wild-type strains. Strains were grown in SC media with either 2% glucose or 3% ethanol. Scale bar: 1 μm.

(E) Graph representing the number of mRNA molecules within each smFISH foci in cells grown in either glucose (black) or ethanol (EtOH; magenta). Foci were separated into either all foci or multi-mRNA foci dependent upon the number of mRNAs predicted to reside within that foci (see Methods). Box and line represent the interquartile range and the median, respectively.

(F) Quantification of total mRNA molecules per cell in cells grown in either glucose (black) or ethanol (EtOH; magenta), as measured by smFISH. Box and line represent the interquartile range and the median, respectively.

(G) Quantification of the number of multi-mRNA containing foci per cell in cells grown in either glucose (black) or ethanol (EtOH; magenta). Box and line represent the interquartile range and the median, respectively.

A surprising observation was made when the localization of the PB marker Dcp2p was assessed in cells bearing either the PDC1-sc or PDC1-sl mRNAs. In both strains, Dcp2p was constitutively present in PBs even in unstressed cells, and the PDC1-sc and PDC1-sl mRNAs colocalized with these bodies (Figures 5E and 5F). This result is especially intriguing as generally PBs are barely visible unless cells are stressed in some way (Lui et al., 2014). Yet in these unstressed cells, just a single point mutation to introduce a STOP codon into one mRNA species is sufficient to induce PB formation. This result is also interesting with regard to the controversy surrounding MS2 tagging. The specific introduction of a mutation that inhibits translation changes the mRNA localization pattern dramatically and causes PB formation. Therefore, if the RNA granules observed for non-mutated glycolytic mRNAs during exponential growth (Figure 1C) were due to the accumulation of RNA fragments carrying the MS2 stem loops, we would expect a similar colocalization with PB markers. However, we do not observe such colocalization with PB markers in unstressed cells (Lui et al., 2014; Pizzinga et al., 2019).

Overall, these results highlight that in keeping with many observations over the years it is difficult to alter the translation of an mRNA without affecting its stability (Mugridge et al., 2018; Roy and Jacobson, 2013). However, the results do suggest that as well as translation occurring in granules, inhibiting translation of individual glycolytic mRNAs changes the fate of those mRNAs so that instead of entering the granules for translation, alternative mRNA fates become apparent, such as relocation to PBs.

Glycolytic mRNA localization varies according to the level of fermentation

In order to understand the potential physiological role the CoFe granules play, experiments were undertaken where yeast were grown on a range of carbon sources selected based upon the pathways required for carbon source metabolism. For example, although yeast cells ferment glucose to ethanol even under aerobic conditions, for other carbon sources the degree of fermentation varies. Yeast cells grown on ethanol as the sole carbon source derive their energy from respiration and only require the glycolytic enzymes for gluconeogenesis. Raffinose is catabolized initially via the action of the secreted enzymes invertase (Suc2p) and α-galactosidase (Mel1p). These enzymes yield the monosaccharides glucose, fructose and galactose, which are readily available for fermentation via the glycolytic enzymes (Barnett, 1976). Equally, yeast that are pre-adapted to galactose express enzymes of the Leloir pathway, allowing fermentation of this sugar via entry into the glycolytic pathway (Timson, 2007).

Microscopic analysis revealed that yeast cells grown on glucose, raffinose, or galactose harbored approximately 10–20 granules of either PDC1 mRNA or ENO1 mRNA per cell, whereas cells grown on ethanol harbored significantly fewer granules (Figures 6A and 6B). In terms of the levels of the PDC1 and ENO1 mRNAs, these also vary with carbon source, consistent with glucose representing the preferred yeast carbon source. For both mRNAs, glucose grown cells harbor significantly more glycolytic mRNA than cells grown on most other carbon sources (Figure 6C).

Similar observations were made when the localization of the FBA1, GPM1, and TPI1 mRNAs was investigated for cells grown on either glucose or ethanol using smFISH (Figure 6D). For yeast grown on either carbon source multi-mRNA granules were observed (Figures 6D and 6E), and, consistent with qRT-PCR results above, the number of mRNA molecules per cell was reduced for the ethanol grown cells (Figure 6F). However, despite this, the number of mRNA molecules per multi-RNA granule was similar for yeast grown on...
either carbon source (Figure 6E), suggesting some form of regulated recruitment to the RNA granule. Perhaps the most striking difference in the localization of the mRNAs was the number of multi-mRNA granules, with ethanol leading to dramatically reduced numbers of RNA granules (Figure 6G), consistent with MS2 experiments performed previously (Figure 6A).

Therefore, these results show that both the level of glycolytic mRNAs and prevalence of CoFe granules vary depending on the carbon sources. In particular, the presence of CoFe granules appears to correlate with a requirement for glycolytic flux to utilize the provided carbon source, with growth on a respiratory carbon source causing a dramatic reduction in the number of granules. Overall, the data are consistent with a view that the localization of glycolytic mRNAs to CoFe granules represents a strategy allowing high-level coordinated production of glycolytic enzymes in translation factories.

Glycolytic mRNA granules are also evident in human cells

In order to assess whether a similar organization of glycolytic mRNAs might exist in higher eukaryotic cells, smFISH analysis was conducted for four different glycolytic mRNAs in HeLa cells: two enolase mRNAs (ENO1 and ENO2), a lactate dehydrogenase mRNA (LDHA), and a phosphofructokinase mRNA (PFKM). For all four of the selected mRNAs, variation in mRNA signal was observed both in terms of particle size and intensity (Figure 7A, data not shown). In particular, large intense mRNA foci were observed for the glycolytic mRNAs (Figure 7) that were not present for other highly expressed mRNAs such as ACTB mRNA (Figure S3). This suggests that granules harboring multiple mRNAs can also be a feature for glycolytic mRNAs in higher eukaryotic cells. Similar observations were made in other cell lines such as HFF-1 cells and SH-SY5Y cells (data not shown). This opens up the possibility that glycolytic mRNAs might be coordinately localized in higher cells. Therefore, a multichannel smFISH approach was taken. Here, evidence for a specific colocalization of the ENO2 and PFKM mRNAs was obtained (Figures 7B and 7C). Although the degree of colocalization is not as comprehensive as observed in yeast, these data do show that in actively growing human tissue culture cells, glycolytic mRNAs can be localized to granules and that these granules can contain more than one type of glycolytic mRNA.

DISCUSSION

mRNA localization serves critical functions in the expression of proteins at specific loci within cells and in the response to stress in terms of PB and SG formation (Pizzinga and Ashe, 2014). In this study, we suggest that mRNA localization to granules can co-ordinate whole pathways of metabolism. We use a combination of live cell experiments and smFISH to show that glycolytic mRNAs localize to granules in yeast and human cells. In stark contrast to mRNAs localizing to PBs, SGs, or transport granules, in yeast the glycolytic mRNAs are translated in CoFe granules, and their translation is a requirement for localization.

Recent evidence suggests that liquid-liquid phase separation (LLPS) within cells produces membraneless compartments or biological condensates where enzymatic reactions and processes can occur. For instance, in the nucleolus, rRNA is produced via numerous highly complex reactions (Brangwynne et al., 2011), whereas in the centrosome microtubule nucleation occurs (Zwicker et al., 2014). The CoFe granules described here conform to many of the properties of phase-separated condensates: they are dynamic, can be observed to fuse, and are disrupted by low concentrations of 1,6-hexanediol (Lui et al., 2014). Therefore, our data suggest that translated glycolytic mRNAs are present in such biological condensates where molecular processes are not only maintained but might actually be enhanced (Kojima and Takayama, 2018).

Enhanced translation of mRNA is therefore one possible explanation as to why the glycolytic mRNAs would be localized within granules. Previous observations from our lab have shown that up to 95% of the glycolytic mRNAs are translated (Lui et al., 2014). In addition, the glycolytic mRNAs are among the most abundant in the cell and so may require rather specific mechanisms to maintain their high rates of translation. Equally LLPS has previously been associated with altered efficiency of a host of biological processes and enzymes (Zhou et al., 2008), so translation may prove to represent another example of such a process, especially where the coordinated generation of high volumes of glycolytic enzyme may be important.

Another possible rationale for localized mRNA translation is to aid the formation of multi-protein complexes. Many of the glycolytic enzymes are present in multimeric complexes. For example, almost all of the glycolytic enzymes function as multimers: in yeast the phosphofructokinase enzyme is present as an octamer (Schwock et al., 2004), phosphoglycerate mutase and pyruvate kinase are tetramers (Jurica et al.,
1998; Rigden et al., 1998), and enolase is dimeric (Sims et al., 2006). Co-translation of individual mRNAs at the same site within cells could therefore aid the formation and productive folding pathways for these complexes. We have not formally shown that different mRNA species are cotranslated at these sites. However, in previous work we have shown that for two different glycolytic mRNA species over 90% of these mRNAs are associated with heavy polysomes, whereas 60%–70% of these mRNAs are associated with the mRNA granules (Lui et al., 2014). The fact that most glycolytic mRNAs, including those that are coassociated in granules, are being actively translated, is suggestive that cotranslation at these granules is also occurring.

Figure 7. Human glycolytic mRNAs are present in granules and can colocalize

(A) Scatterplot of particles detected in z-stacked smFISH images of HeLa cells using probes to the mRNAs indicated. Images from three biological replicates were analyzed using the ImageJ ComDet plugin, which generates readouts of particle size (measured in pixels where each pixel = 45 × 45 nm) and the fluorescent intensity of particles.

(B) Single z-slice smFISH images of HeLa cells using probes to the mRNAs indicated. Scale bar: 10 μm. Insets magnified x2.

(C) Histogram showing the percentage of colocalized mRNA particles calculated using ComDet analysis of z-slices from three biological replicates. The significance across the various combinations was calculated using one-way ANOVA and the Tukey’s HSD test. ENO2 versus PFKM (p value <0.05 shown by asterisk) is significantly different to ENO2 versus ENO1, ENO1 versus PFKM, or ENO2 versus LDHA. Error bars are ±SD.
A range of precedents exist for the cotranslational production of complexes across various biological systems (Halbach et al., 2009; Kamenova et al., 2019; Shiber et al., 2018; Wells et al., 2015), whereas a systematic analysis in *Schizosaccharomyces pombe* suggests that cotranslational production of protein complexes is widespread, with a substantial fraction of proteins copurifying with mRNAs that encode interacting proteins (Duncan and Mata, 2011). Notably, in recent work characterizing the propensity for cotranslational folding in several different protein complexes in *Saccharomyces cerevisiae*, the PFK1 and PFK2 phosphofructokinase mRNAs were identified as key examples where the translated products are cotranslationally assembled or folded (Shiber et al., 2018).

It has also been shown that, as well as forming multimeric single enzyme complexes, various different glycolytic enzymes can be compartmentalized into much larger complexes (Masters, 1991). A variety of observations suggest that the physical compartmentalization of glycolysis is advantageous. For instance, in protozoan organisms such as *Trypanosoma* and *Leishmania*, a specific membrane-bound organelle called the glycosome has evolved, which is thought to provide these pathogens a scope for regulating metabolic activity (Haanstra et al., 2016). Furthermore, in human cells, such as skeletal muscle cells, neurons, and erythrocytes, glycolytic enzymes can be organized as complexes coordinated either on membranes or the cytoskeleton (Knoll and Walsh, 1992; Puchulu-Campanella et al., 2013). Moreover, a glycolytic metabolon has also been described in yeast (Masters, 1991), and it is thought to be stabilized by various weak interactions with actin (Araiza-Olivera et al., 2013). These multi-enzyme complexes are likely to promote both the channelling of metabolites from one enzyme to the next, as well as the reduction of potentially toxic intermediates. More recent work in yeast has shown that although glycolytic enzymes are broadly cytosolic under non-stress conditions, they can coalesce into “G-bodies” in response to hypoxic stress (Jin et al., 2017). Overall, therefore, the coproduction of the glycolytic enzymes at the same site by virtue of coordinated mRNA localization could promote the cotranslational formation of some of these higher order complexes of enzymes. Although, it should be noted that our own data suggest that under active growth conditions fluorescent-protein tagged forms of the glycolytic enzymes are generally found throughout the cytosol (Lui et al., 2014).

Another point worth reflecting upon when considering the role of the CoFe granules is that several glycolytic enzymes have extra-glycolytic or “moonlighting” functions outside of their role in glycolysis. For example, many of the glycolytic enzymes have been identified as RNA-binding proteins that appear to interact with their own mRNA (Castello et al., 2015; Matia-Gonzalez et al., 2015). In addition, yeast enolase is important for both the mitochondrial import of tRNA\(^{12}_{\text{CUU}}\) (Entelis et al., 2006) and for vacuolar H\(^+\)-ATPase function (Lu et al., 2007). Many further moonlighting functions of glycolytic enzymes, including nuclear functions in transcription, DNA replication/repair, and histone modification have been described (Boukouris et al., 2016). One possible explanation for the presence of CoFe granules could be that they serve as a focus for the coordinated high-level production of the glycolytic machinery en masse, whereas individual translated glycolytic mRNAs outside of factories could provide the capacity for moonlighting protein activities.

One intriguing observation made during the course of our studies is that although the yeast ENO1 mRNA is observed to localize to granules, it does not appear to colocalize with the ENO2 mRNA in CoFe granules. Several possible non-mutually exclusive explanations could account for this. Firstly, the expression levels of the enolase isoforms vary greatly depending upon the fermentation/respiration status; Eno2p represents the predominant polypeptide in fermenting cells, whereas the distribution is much more even in stationary phase or respiring cells (Entian et al., 1987). So the discrete localization of the two mRNAs could contribute to these expression differences. Secondly, since the enolase enzyme is dimeric, and both Eno2p and Eno1p homodimers, as well as the heterodimer have been described (Holland et al., 1982), the discrete localization of the ENO1 and ENO2 mRNAs could serve to regulate the relative proportion of these different complexes via cotranslational homodimer formation. The rationale for requiring homodimers seems unlikely to reside in their enzymatic function, as the various forms display very similar enzyme kinetics (Holland et al., 1982). Therefore, an alternative possibility relates to the enolase moonlighting functions, with Eno2p playing a more active role than Eno1p in the targeting of the nuclear-encoded tRNA\(^{12}_{\text{CUU}}\) isoacceptor to mitochondria (Entelis et al., 2006).

Overall, glycolysis is perhaps the most fundamental of all biological pathways. The enzymes of the pathway are regulated at almost every level, and paralogues have evolved distinct functions. The pathology of many disease conditions is intimately connected to the glycolytic pathway. For instance, aerobic glycolysis serves...
as a hallmark of many malignant cancers and the surrounding stroma, which can serve as a negative prog-
nostic indicator due to increased resistance to therapy (Lee and Yoon, 2015; Ngo et al., 2015). The identi-
fication and characterization of factories for the production of glycolytic proteins can only serve to increase understanding of the functions, regulation, and possibility for genetic adjustment of this key metabolic pathway.

Limitations of the study
Although we have interpreted our data to mean that cotranslation of different colocalized mRNAs is occur-
bring in translation factories, due to technical limitations we have not directly shown the translation of two or more mRNAs at the same site. However, in this study and previous work, we have shown translation of single mRNA species occurs at these sites. We have shown that multiple mRNAs colocalize to the sites and that for any single glycolytic mRNA, over 90% of the mRNA is engaged with heavy polysomes. We have interpreted these results to mean that cotranslation of different mRNAs is indeed occurring at the same site.

Resource availability
Lead contact
Further information and requests for resources and reagents should be directed to the Lead Contact, Mark Ashe (mark.p.ashe@manchester.ac.uk).

Materials availability
Strains and plasmids generated as part of this study are available upon request.

Data and code availability
Code for bespoke yeast cell smFISH analysis is available at the GitHub repository github.com/CPBS/MoralesPolanco.

METHODS
All methods can be found in the accompanying Transparent methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2021.102069.

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AUTHOR CONTRIBUTIONS
All authors generated reagents, performed experiments, evaluated results, and generated figures. FMP, CB, JL, JC, and MPA conceived the study and designed experiments, whereas MPA wrote the manuscript. All authors contributed to the discussion and evaluation of the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Core Fermentation (CoFe) granules focus coordinated glycolytic mRNA localization and translation to fuel glucose fermentation

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Figure S1. mRNA levels and the use of lower affinity MS2 stem loops support the premise that the glycolytic mRNAs are localized to RNA granules. Related to Figure 1. (A) Graph representing the relative fold change of MS2-tagged mRNAs relative to untagged mRNA levels. Error bars represent ± SE. (B) Northern blots of ENO2 and PDC1 mRNA in glucose replete and starved conditions in untagged strains or strains bearing the MS2 tag. (C) z-stacked epifluorescent images of CDC19, ENO1 and ENO2 mRNAs tagged with both the MS2v6 and MS2v5 stem loop systems co-expressing the relevant MS2-CP-GFP fusion. MS2v5 images are the same as those shown in Figure 1. ENO1 and CDC19 MS2v6 constructs contain 24x stem loops, ENO2 contains 12x stem loops. Scale bar: 2 μm.
Figure S2. Glycolytic mRNA granules do not localize to ER or mitochondria. Related to Figures 3 and 4. z-stacked epifluorescent images of ENO1 and PDC1 mRNAs tagged with MS2 stem loops, visualized via co-expressed pMCP-GFP, with either an endoplasmic reticulum marker, Sec63p-CFP (A) or a mitochondrial marker, Cox4p-RFP, (B). Scale bars 2 µm.
Figure S3. ACTB mRNA localizes diffusely throughout the cytoplasm. Related to Figure 7. (A) z-stacked epifluorescent images of HeLa cells. Cells were stained with DAPI and hybridized with either ACTB specific smFISH probes (top) or no probe as a control (bottom). Scale bar 10 µm. (B) Scatter plot of ACTB smFISH foci size and intensity, as detected by ComDet reveals very little evidence for large mRNA granules.
TRANSPARENT METHODS

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Yeast growth conditions
Yeast experiments were performed in the *Saccharomyces cerevisiae* strain yMK467 - a derivative of W303-1A (SCR_003093). Unless stated otherwise, experiments were performed after strains were grown in synthetic complete (SC) media with 2% glucose at 30°C to exponential phase. Strains used in this study are listed in Table I. For live-cell microscopy, cells were pelleted at 500xg for 3 minutes at 30°C, resuspended in pre-warmed (30°C) media lacking methionine and incubated for 30 min to induce expression of the pCP-GFP/mCh fusions prior to imaging. For growth on alternative carbon sources SC media was supplemented with 2% raffinose, 2% galactose or 3% ethanol. For stress conditions, cells were incubated in media lacking glucose for 10 minutes.

Human cell-line growth conditions
Human cell-line experiments were performed in HeLa cells (CVCL_0030), grown in Dulbecco's modified Eagle's medium supplemented with 10% fetal bovine serum at 37°C.

METHOD DETAILS

Yeast strain and plasmid construction
MS2 and PP7 tagged strains were generated as previously described (Haim-Vilmovsky and Gerst, 2011; Hocine et al., 2013; Tutucci et al., 2018), using plasmid reagents generously donated by Jeff Gerst and Robert Singer. Dual MS2 and PP7 tagged strains were generated by crossing the single tagged haploid strains. Subsequent diploid strains were selected, sporulated and the appropriate dual tagged haploid strains were verified by PCR. Expression of PP7 and MS2 coat protein-fluorescent protein fusions was driven from the inducible MET25 promoter on plasmids that have been described previously (Haim-Vilmovsky and Gerst, 2011; Hocine et al., 2013; Tutucci et al., 2018; Lui et al., 2014). The strain harbouring a premature stop codon in the *PDC1* ORF was generated via recombination of a mutant PCR product generated from the *PDC1-MS2* tagged strain. More specifically, oligonucleotides with a specific mutation in the upstream primer were used to amplify a *PDC1::HIS5::MS2* cassette from genomic DNA prepared from an intermediate strain in the *PDC1-MS2* m-TAG procedure (Haim-Vilmovsky and Gerst, 2011). The mutation introduces a premature STOP codon in the *PDC1* ORF. The *PDC1::HIS5::MS2* cassette was then transformed and recombined into the *PDC1* locus of the yMK467 strain. Removal of the *HIS5* marker was carried out using a Cre recombinase strategy as previously described (Haim-Vilmovsky and Gerst, 2011). TRICK strains were generated using a similar approach to MS2 or
PP7 tagging, but using a DNA template developed for TRICK in yeast (Pizzinga et al., 2019). For generation of the yEPlac195-PDC1 (pPDC1-MS2) plasmid, PDC1-MS2 was amplified from genomic DNA of yMK1586 and cloned into the pGEM-T Easy vector and subsequently cloned into yEPlac195 using SphI and SacI restriction enzymes. A stem loop sequence (Vattem and Wek, 2004) was inserted upstream of the start codon in PDC1 using Gibson assembly (Gibson et al., 2009) to generate plasmid yEPlac195-PDC1-SL (pPDC1-MS2 (sl)).

**Single molecule fluorescent in situ hybridisation (smFISH)**

For yeast cultures, smFISH was performed as previously described (Pizzinga et al., 2019). In brief, exponential-phase yeast were fixed in 4% EM-grade formaldehyde (15714-S; Electron Microscopy Sciences) for 45 min, then spheroplasted and permeabilized with 70% ethanol. Gene-specific 20nt antisense oligonucleotides were designed with a 59nt Flap sequence, to which fluorescently labelled oligonucleotides were annealed (Pizzinga et al., 2019; Tsanov et al., 2016). The conjugated fluorophores included Alexa Fluor 488, Alexa Fluor 546, ATTO 590 and Alexa Fluor 648. After careful titration of the probe, 20 pmol fluorescently labelled smFISH probe was found to generate optimal signal relative to background when added to the cells in hybridization buffer (10 mg E. coli tRNA, 2 mM Ribonucleoside Vanadyl Complex, 200 μg/ml BSA, 10% dextran sulfate, 10% formamide, and 2× SSC in nuclease-free water). This hybridization buffer – probe mix was then added to cells and incubated overnight at 37°C, with gentle agitation. Cells were then washed in 10% formamide and 2× SSC and adhered to 0.01% poly-L-lysine–coated coverslips before mounting in ProLong diamond antifade mounting solution with DAPI (Life Technologies Cat# P36970).

For human cell experiments, HeLa cells were seeded in Dulbecco's modified Eagle's medium supplemented with 10% fetal bovine serum onto 13 mm laminin coated coverslips in sterile 24-well plates, then were fixed in methanol for 10 min at -20°C. Fixed cells were washed in 10% Formamide, 2x SSC buffer in nuclease-free water for 30 min at room temperature, then hybridized probes (as above) were added at a concentration of 25 nM for Cy7-conjugated probes and 75 nM for Cy5-conjugated probes at 37°C overnight. Cells were then washed and mounted (Tsanov et al., 2016). An extensive series of experiments was undertaken to optimise the probe concentration and fixation method such that the no signal was detected in the various channels when a particular probe was absent.

**Live-cell fluorescent microscopy**

All yeast live-cell epifluorescent microscopy was performed on a Nikon Eclipse E600 or a Delta Vision microscope (Applied Precision) equipped with a CoolSnap HQ camera (Photometrics),
using a 100x/1.40 NA oil plan Apo objective. Fluorescent parameters for each fluorophore are as follows; GFP (excitation-490/20 nm, emission-535/50 nm); mCherry (excitation-572/35 nm, emission-632/60 nm); and CFP (excitation-436/10 nm, emission-465/30). For routine live-cell imaging, exponentially growing cells were viewed on poly-L-lysine coated glass slides and images were taken with a z-spacing of 0.2μm. Images were acquired using Softworx 1.1 (Applied Precision), or Metamorph (Molecular Devices) software and processed using Image J software package (National Institute of Health, NIH).

**Fixed-cell fluorescent microscopy**

Images of human cells were acquired on an Olympus IX83 inverted microscope using Lumencor LED excitation, a 100x objective and the Penta filter set. The images were collected using a Retiga R6 (Qimaging) CCD camera with a z-optical spacing of 0.2 μm. Raw images were then deconvolved using the Huygens Professional software (Scientific Volume Imaging).

Yeast smFISH images were collected on a Leica TCS SP8 AOBS inverted gSTED microscope using a 100x/1.40 Plan APO objective and 1x confocal zoom, as described previously (Pizzi nga et al., 2019). DAPI staining was detected using a photon multiplying tube with a blue diode 405nm laser (5%). Confocal images of smFISH signals were collected using hybrid detectors with the following detection mirror settings; Alexa Fluor 488 498-536nm; Alexa Fluor 546 556-637nm (5 to 35μs gating); ATTO 590 603-637nm; Alexa Fluor 647 657-765nm using the 488nm (60%), 550nm (60%), 593nm (60%) and 646nm (60%) excitation laser lines, respectively. Images were collected sequentially in 200nm z-sections. Acquired images were subsequently deconvolved and background subtracted using Huygens Professional (Scientific Volume Imaging).

**Quantitative RT-PCR (qRT-PCR)**

RNA preparations were carried out using Trizol as described by the manufacturer (Thermofisher scientific, Cat#15596026), followed by isopropanol precipitation then treatment with Turbo DNase (Thermofisher scientific, Cat#AM2238). qRT-PCR was performed in a two-step manner using a ProtoScript First Strand cDNA synthesis kit (New England Biolabs, Cat#E6300S) and iQ SYBR Green Supermix (Bio-Rad, Cat#1708880) according to manufacturer’s instructions. Reactions were performed using 100ng of cDNA. iTaq Universal SYBR Green One Step Kit (Bio-Rad, Cat#1725150) was used to carry out one-step qRT-PCR and reactions were performed using 300ng of RNA. A CFx Connect Real-Time system was used to run reactions. Samples were run in triplicate and normalized to *ACT1* mRNA, and the fold change was calculated using either the 2^-ΔΔCq or the Pfaffl method (Livak and Schmittgen, 2001; Pfaffl, 2001).
QUANTIFICATION AND STATISTICAL ANALYSIS

Quantification of microscopy and statistics.

For quantification of granule numbers per cell from live cell experiments, 50 cells were counted for each strain. For quantification of overlapping MS2 and PP7 signal in double-tagged strains or TRICK strains, 100 granules were considered for each strain over three biological repeats. GraphPad Prism 6 (GraphPad Software, Inc.) was used to produce the graphs and to calculate the standard deviation, indicated by error bars. Two-way ANOVA was performed using GraphPad Prism 6. * denotes a P value < 0.0001.

Yeast smFISH images were processed and analysed using FISH-quant (Mueller et al., 2013) and FindFoci (Herbert et al., 2014) to identify spot position and size and provide spot enhancement via dual Gaussian filtering. To account for differences in smFISH signal intensity between fluorophores and experiments, different intensity thresholds for each channel/image were determined manually. However, the same thresholds were applied to all cells in that image. Cell outlines were automatically generated using a modified version of the CellProfiler (Carpenter et al., 2006) pipeline provided with FISH-quant that utilizes background cytoplasmic DAPI staining rather than brightfield images to determine cytoplasmic cell boundaries. Spot colocalization and other foci characteristics were assessed and quantified using custom scripts in python to scrape data from FISHquant into long format and R for more detailed analysis. Human cell line smFISH images were analyzed using the Image J ComDet plugin, which generates values for particle size (area) in pixels where each pixel = 45x45 nm and particle intensity in arbitrary units.

Colocalization analysis in yeast cells was performed by pairing spots between channels based on spot centroid distance in 3D space (Eliscovich et al., 2017). Spots were deemed to colocalize if the 3D distance between centroids was less than the radius of either of the two spots, i.e. if the centroid of one spot existed within the radius of another spot. mRNA quantitation was performed using Gaussian fitting, as described previously (Pizzinga et al., 2019). To account for stochasticity in initial fitting parameters, this fitting was repeated 1,000 times and averaged. Simulated controls were based on the Monte Carlo simulation method (Fletcher et al., 2010). Real spots identified using smFISH sampled using varying spot characteristics, such as size in x, y and z, and these were arbitrarily positioned within a simulated volume typical of a yeast cell as measured using the custom CellProfiler pipeline. This volume includes RNA depleted regions of the cell such as the vacuole. A variable number of spots were selected for the simulation of each mRNA dependent upon the average number of foci observed per cell for that particular mRNA. The colocalization of these randomly positioned foci was subsequently processed using the same script outlined above, and iterated 1,000 times per pairwise comparison. For the human cell line
colocalisation analysis, the ComDet plugin was used on ImageJ which takes a similar centroid-centroid overlap approach to that described above (Katrukha 2020).

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| EM-grade formaldehyde | Electron Microscopy Sciences | Cat#15714-S |
| ProLong Diamond Antifade Mountant | Life Technologies | Cat#P36970 |
| **Critical Commercial Assays** | | |
| Turbo DNase | Thermofisher Scientific | Cat#15596026 |
| iQ SYBR Green Supermix | Bio-Rad | Cat#1708880 |
| iTaq Universal SYBR Green One Step Kit | Bio-Rad | Cat#1725150 |
| Stellaris® FISH Probes, Human ACTB with Quasar® 570 Dye | Biosearch Technologies | Cat#VSMF-2002-5 |
| **Experimental Models: Organisms/Strains** | | |
| yMK467 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 | (Campbell et al., 2005) | |
| yMK807 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 | (Campbell et al., 2005) | |
| yMK1577 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 ENO2-MS2L p[MS2-GFPp3 HIs3] | (Lui et al., 2014) | |
| yMK1586 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PDC1-MS2L p[MS2-GFPp3 HIs3] | (Lui et al., 2014) | |
| yMK1651 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PDC1-MS2L DCP2-CFP p[MS2-GFPp3 HIs3] | (Lui et al., 2014) | |
| yMK2257 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 ENO2-PP7L PDC1-MS2L p[MS2-mCh3 HIs3] p[PP7-GFPp2 URA3] | (Lui et al., 2014) | |
| yMK2412 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PFK1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2413 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PYK2-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2415 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PFK2-MS2L (5 loops) p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2416 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 FBA1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2429 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 | This Study | NA |
| yMK2430 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 TPI1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2431 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 GLK1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2447 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 ADH1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2452 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PDC1-MS2L (sc) DCP2-CFP p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2480 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PFK2-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2535 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 PGK1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2468 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 ENO1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK580 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 HXK1-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
| yMK2582 MATa ADE2 his3-11,15 leu2-3,112 trp1-1 ura3-1 CDC19-MS2L p[MS2-GFPp3 HIs3] | This Study | NA |
This Study NA

yMK2699 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2494 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2588 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2600 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2602 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2603 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2604 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2697 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2700 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2705 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK2738 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3162 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3176 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3397 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK5000 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK5001 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK5002 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK5003MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3471 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3472 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3473 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

yMK3474 MATa ADE2 his3-1,15 leu2-3,112 trp1-1 ura3-1

This Study NA

Oligonucleotides

X-Flap Alexa Fluor 546: 5’Alex546N-CACGTAGTCGAGCTCGAAACCTAGGAGG

Integrated DNA Technologies NA

-3’AlexF546N

Y-Flap Alexa Fluor 488: 5’Alex488N-AATGCATGTCGAGCTCGAAACCTAGGAGG

Integrated DNA Technologies NA

-3’AlexF546N
| DNA Sequence                                                                 | Source                          | Notes     |
|------------------------------------------------------------------------------|---------------------------------|-----------|
| 3'AlexF488N                                                                  |                                 | NA        |
| Z-Flap Alexa Fluor 647: 5'Alex647N-CTTATAGGGCATGGATGCTAGAAGCTGG-3'AlexF647N  | Integrated DNA Technologies     | NA        |
| Y-Flap ATTO 590: 5'ATTO590N-AATGCATGTCGACGAGGTCCGAGTGTAA-3'ATTO590N           | Integrated DNA Technologies     | NA        |
| X-Flap ATTO 590: 5'ATTO590N-CACTGAGTCCAGCTCGAAACTTAGGAGG-3'ATTO590N           | Integrated DNA Technologies     | NA        |
| Z-Flap ATTO 590: 5'ATTO590N-CTTATAGGGCATGGATGCTAGAAGCTGG-3'ATTO590N           | Integrated DNA Technologies     | NA        |
| Recombinant DNA                                                              |                                 | NA        |
| yEPLac195-PDC1                                                               | This Study                      | NA        |
| yEPLac195-PDC1-SL                                                            | This Study                      | NA        |
| pMCP-GFP                                                                     | (Haim-Vilmovsky et al., 2011)   | pMS2-CP-GFP(x3) |
| 12xMS2v5                                                                     | (Haim-Vilmovsky et al., 2011)   | pLOXHi5MS2L |
| pMCP-mCh                                                                      | (Lui et al., 2014)              | NA        |
| pPP7CP-GFP                                                                   | (Hocine et al., 2013)           | Addgene Plasmid #45931 |
| 24xPP7SL                                                                     | (Hocine et al., 2013)           | Addgene Plasmid #45163 |
| 12xMS2v6-loxP                                                                | (Tutucci et al., 2018)          | Addgene Plasmid #104392 |
| 24xMS2v6-loxP                                                                | (Tutucci et al., 2018)          | Addgene Plasmid #104393 |
| MCPv6-NLS                                                                    | (Tutucci et al., 2018)          | Addgene Plasmid #104394 |
| Software and Algorithms                                                      |                                 |           |
| Softworx v1.1                                                                 | Applied Precision               |           |
| Metamorph                                                                    | Molecular Devices               |           |
| FISHQuant                                                                    | (Mueller et al., 2013)          |           |
| FindFoci                                                                     | (Herbert et al., 2014)          |           |
| CellProfiler 3.0                                                             | (Carpenter et al., 2006)        |           |
| Prism 6                                                                      | GraphPad                        |           |
| Huygens Professional | Scientific Volume Imaging | [https://svi.nl/HomePage](https://svi.nl/HomePage) |
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