

Transcription Factor Condensates Mediate Clustering of *MET* Regulon and Enhancement in Gene Expression

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Abstract

Some transcription factors (TFs) can form liquid-liquid phase separated (LLPS) condensates. However, the functions of these TF condensates in 3D genome organization and gene regulation remain elusive. In response to methionine (met) starvation, budding yeast TF Met4 and a few co-activators, including Met32, induce a set of genes involved in met biosynthesis. Here, we show that the endogenous Met4 and Met32 form co-localized puncta-like structures in yeast nuclei upon met depletion. Recombinant Met4 and Met32 form mixed droplets with LLPS properties *in vitro*. In relation to chromatin, Met4 puncta co-localize with target genes, and at least a subset of these target genes are clustered in 3D in a Met4-dependent manner. A *MET3pr*-GFP reporter inserted near several native Met4 binding sites becomes co-localized with Met4 puncta and displays enhanced transcriptional activity. A Met4 variant with a partial truncation of an intrinsically disordered region (IDR) shows less puncta formation, and this mutant selectively reduces the reporter activity near Met4 binding sites to the basal level. Overall, these results support a model where Met4 and co-activators form condensates to bring multiple target genes into a vicinity with higher local TF concentrations, which facilitates a strong response to methionine depletion.

eLife assessment

This **important** study investigates the relationship between transcription factor condensate formation, transcription, and 3D gene clustering of the MET regulon in the model organism *S. cerevisiae*. The authors provide **solid** experimental evidence that transcription factor condensates enhance transcription of MET-regulated genes, but the evidence that nuclear condensates per se drive MET gene clustering is **incomplete** and would benefit from further experimental analyses. This paper will be of interest to molecular biologists working on chromatin and transcription, although its impact would be strengthened by revising the literature citations and including additional experimental work.

Introduction

Chromosomes form extensive 3D contacts that can occur over long linear genomic distances (Marenduzzo et al. 2007 [↗](#); Fazary et al. 2017 [↗](#); Misteli 2020 [↗](#)). A subset of these interactions, such as promoter-promoter or promoter-enhancer looping, are thought to play a central role in gene regulation (Mifsud et al. 2015 [↗](#); Schoenfelder and Fraser 2019 [↗](#)). In some cases, multiple genes come together in 3D space to form “multi-gene clusters” (Mir et al. 2018 [↗](#); Lim and Levine 2021 [↗](#); Zhu et al. 2021 [↗](#); Uyehara and Apostolou 2023 [↗](#)). These clusters often involve co-regulated genes responding to stress or other types of environmental cues, such as heat shock, starvation, virus infection, or developmental signals (Ferrier and Minguillón 2003 [↗](#); Sawado et al. 2003 [↗](#); McCord et al. 2011 [↗](#); Du and Bai 2017 [↗](#); Zhao and Faryabi 2023 [↗](#)). Evidence from imaging and genomic studies suggests that these gene clusters may be related to a phenomenon called “transcription factories”, where RNA polymerase II (Pol II), mediator, and nascent transcripts coalesce into distinct foci (Bartlett et al. 2006 [↗](#); Faro-Trindade and Cook 2006 [↗](#); Carter et al. 2008 [↗](#); Xu and Cook 2008 [↗](#); Sutherland and Bickmore 2009 [↗](#); Papantonis and Cook 2011 [↗](#); Rieder et al. 2012 [↗](#)). Proteomic analysis of transcription factories showed that they are also enriched with other components in the transcription pathway, including transcription factors (TFs), histone modification enzymes, and chromatin remodelers (Lyons et al. 2023 [↗](#)). Overall, these observations suggest that some co-regulated genes can physically cluster into sub-nuclear compartments with elevated local concentrations of transcription-related factors.

How multi-gene clusters are formed is not well understood. Since co-regulated genes are often activated by a common set of TFs (Kiran and Nagarajaram 2016 [↗](#); Mir et al. 2018 [↗](#); Tsai et al. 2019 [↗](#); Uyehara and Apostolou 2023 [↗](#)), such clustering may be mediated by the DNA-protein and protein-protein interactions of the TFs. Recent studies showed that TFs like Gcn4, OCT4, and TAZ, as well as some co-factors such as the mediator and histone-acetylation reader Brd4, tend to form biomolecular condensates through liquid-liquid phase separation (LLPS) (Whyte et al. 2013 [↗](#); Boija et al. 2018 [↗](#); Sabari et al. 2018 [↗](#); Gibson et al. 2019 [↗](#); Lu et al. 2020 [↗](#); Lee et al. 2021 [↗](#)). In LLPS, multivalent interactions among proteins, often through their intrinsically disordered regions (IDRs), allow them to condensate into liquid-like droplets that are separated from the aqueous phase (Banani et al. 2017 [↗](#); Dignon et al. 2020 [↗](#); Peng et al. 2020 [↗](#); Mitrea et al. 2022 [↗](#)). The existence of TF / co-factor condensates is supportive of the “transcription factory” model mentioned above where transcription-related factors are locally concentrated through LLPS. So far, LLPS of TFs and co-factors are often studied *in vitro* using purified systems or *in vivo* with artificial induction. How these proteins behave in a chromatin context at the endogenous concentration, and more specifically, if they contribute to multi-gene cluster formation, is not very clear.

Another unresolved issue related to multi-gene clusters is their functional role in gene regulation. Earlier studies of gene clusters found a positive correlation between clustering and gene expression level (Li et al. 2012 [↗](#); Whyte et al. 2013 [↗](#); Zhang and Bai 2016 [↗](#); Mir et al. 2018 [↗](#); Zhu et al. 2021 [↗](#); Zhao and Faryabi 2023 [↗](#)), but the mechanism of the phenomenon was not clear. This again may be related to TF / co-factor condensates: if a gene cluster is situated within a condensate, higher local concentrations of these factors can potentially enhance expression. This is proposed to be the case at super-enhancers, where the condensates of co-factors like Med1 and Brd4 are thought to drive the high transcriptional activities of the target genes. However, some reports studying TF condensates reveal a complex, sometimes contradictory, relation to gene expression, where the condensates are observed to have positive, negative, or neutral effects on transcription (Lu et al. 2020 [↗](#); Lee et al. 2021 [↗](#)). In a recent study of the *GAL* genes, for example, it is proposed that Gal4 condensation facilitates their recruitment to target genes but does not contribute to gene activation (Meeussen et al. 2023 [↗](#)). In general, the relation between TF condensates, multi-gene cluster, and gene regulation needs to be further studied.

Upon met depletion (-met) in budding yeast, Met4 is recruited by several sequence-specific DNA-binding co-activators, including Cbf1, Met31, and Met32, to bind and rapidly activate a subset of genes (Thomas et al. 1992 [\[1\]](#); Lee et al. 2010 [\[2\]](#); Ouni et al. 2010 [\[3\]](#); Carrillo et al. 2012 [\[4\]](#)). This activator complex is further stabilized with Met28 (Kuras et al. 1997 [\[5\]](#)). In a previous study, we carried out a screen to probe long-distance gene regulation that affected *MET3* promoter (*MET3pr*) activity by inserting an insulated *MET3pr*-GFP reporter into many genomic loci and measuring GFP intensity in -met (Du et al. 2017 [\[6\]](#)). At most locations, the reporter has comparable induction level; however, it shows enhanced activity at a small subset of loci, which were defined as “transcriptional hotspots” (Du et al. 2017 [\[6\]](#)). These hotspots tend to be located near endogenous Met4 target genes, and 3C assay detects physical proximity among the hotspots. These results lead to a hypothesis that hotspots occur at loci where *MET* genes cluster, and such clustering and transcriptional elevation may be both mediated by the condensates of related TFs.

In this study, we test the hypothesis above using a combination of biochemistry, genetics, genomics, and imaging approaches. We demonstrate that Met4 and Met32 form co-localized puncta in the yeast nuclei upon met depletion. Purified Met32 forms condensate with LLPS properties *in vitro*, with which Met4 can merge. In relation to chromatin, we show that the loci that are associated with the Met TFs co-localize with the Met4 puncta, and at least a subset of Met4 target genes cluster in 3D nuclear space in a Met4-dependent manner. Functionally, the *MET3pr*-GFP reporter inserted near the endogenous Met4 target sites shows higher enrichment of Met TFs / Pol II and enhanced transcriptional activity (hotspots). A Met4 IDR truncation that reduces Met4 puncta formation selectively affects the reporter activity at hotspots. Collectively, our results support a model where TFs in the met response pathway form LLPS condensates, organize multiple target genes in their vicinity, and enhance gene expression by creating a local environment with elevated concentrations of transcription-related factors. This may represent a common strategy for cells to concentrate limited resources to mount a strong response to stress.

Results

The trans-activator Met4 and its co-factor Met32 form puncta in the nucleus

Our previous study presented evidence that the “hotspots” for *MET3pr* transcription may form 3D clusters, but the mechanisms of clustering and transcriptional elevation are unknown. Given the recent reports on TF condensation (Sabari et al. 2018 [\[7\]](#); Lu et al. 2020 [\[8\]](#); Zhang et al. 2022 [\[9\]](#); Okada et al. 2023 [\[10\]](#)), we hypothesize that these phenomena may be related to condensates formed by TFs in the *MET* pathway. To test this possibility, we labeled the endogenous Met4 protein with GFP and visualized Met4 in live cells under Airyscan confocal microscopy (**Materials and Methods**; see **Table S1** for strain list). We observed nuclear-localized Met4-GFP under both the repressed (+met) and activated (-met) conditions, with the latter showing higher intensity. Importantly, the Met4-GFP displays puncta-like structures, especially under -met (**Fig. 1A** [\[11\]](#) & **S1A**). As controls, we constructed strains expressing a nuclear-localized free GFP or GFP fused to various chromatin-associated factors, including TFs Cbf1 and Reb1, and Sth1, a subunit of RSC nucleosome remodeler, which also bind chromatin with sequence specificity (Cai and Davis 1990 [\[12\]](#); Ju et al. 1990 [\[13\]](#); Donovan et al. 2023 [\[14\]](#)). The GFP signals appear more evenly distributed in these control strains (**Fig. 1A** [\[11\]](#)). To evaluate the distribution more quantitatively, we calculated the coefficient of variance (CV) of the fluorescent intensities among the pixels inside each nuclei (**Materials and Methods**, **Table S2**). Uneven distribution of the GFP molecules would result in large variations in the pixel intensities and higher CVs. The CVs of Met4-GFP in both ±met conditions are significantly higher than free GFP and other nuclear proteins (**Fig. 1B** [\[11\]](#)). Since the strains in **Fig. 1A** [\[11\]](#) express GFP at different levels, which directly affect CV, we also calculated the Fano number (variance divided by the mean) to control for this variable. Met4-GFP in the -met

condition has the largest Fano number (**Fig. 1C** & **S1B**), which indicates that in the activated state it forms the most punctate structures. These data support the notion that Met4 forms condensates in the nuclei upon activation.

We next carried out the same experiment with Met32, the DNA binding co-factor of Met4. We imaged cells containing endogenous Met32 labeled with mCherry and compared its distribution to free mCherry. Unlike Met4, Met32-mCherry is barely visible in +met, and it is significantly upregulated in -met (**Fig. 1D** & **S1C**). In the latter condition, Met32-mCherry also shows uneven distribution with high CV and Fano number (**Fig. 1E, F** & **S1D**). We also imaged Met31-mCherry, but its concentration is too low to be detected. When Met4-GFP and Met32-mCherry are tagged in the same strain, their fluorescent signals largely overlap (**Fig. 1G** & **S1E**). To quantify the overlap, we extracted the GFP and mCherry fluorescence intensities for all the pixels inside each nucleus and calculated their correlation coefficient. As a control, we carried out the same analysis with co-expressed Sth1-GFP and Met32-mCherry, which appear to be less co-localized (**Fig. 1G**). Indeed, the Pearson correlation between Met4 and Met32 fluorescence on average was 0.75, much higher than that between Sth1 and Met32 ($R = 0.19$) (**Fig. 1H**). Overall, the puncta-like structures of Met4 and Met32 and their co-localization support the possibility that these two factors co-condensate under the -met condition.

Met32 forms condensates with LLPS properties *in vitro*

LLPS is proposed to be a mechanism that leads to TF condensation (Banani et al. 2017; Boeynaems et al. 2018). To test if this mechanism applies to Met4 and Met32, we first evaluated their propensity to phase separate using the FuzDrop algorithm (Hatos et al. 2022). All the Met activators, including Met4, Met31, Met32, and Met28, are predicted to have higher LLPS probabilities than GFP and mCherry (**Fig. 2A**). Among these TFs, Met4 and Met32 have particularly high scores, which suggests they function as droplet drivers that can spontaneously phase separate (Vendruscolo and Fuxreiter 2022). In addition, both factors are predicted to have large unstructured domains, a feature known to promote LLPS (**Fig. S2A & B**).

We next purified Met4 and Met32 to study their condensation properties *in vitro*. Like many other recombinant TFs, Met4 is prone to degradation. We managed to extract full length Met4 tagged with 6xHis and MBP on the N- and C-terminal, respectively, although some degradation products could not be avoided (**Fig. 2B**) (**Materials and Methods**). We imaged the purified Met4 labeled with Alexa488 in various buffers. Met4 remains diffusive up to 5 μ M at physiological salt concentrations and forms aggregates at higher density or lower salt concentrations (**Fig. 2C**). Although these results suggest that Met4 may not inherently undergo LLPS, we cannot rule out the potential effects from impurities (degradation products) and tags on our observations. In contrast, we managed to purify high-quality Met32 fused with mCherry (**Fig. 2B**). At the physiological salt concentration with no crowding agents, Met32-mCherry forms condensates at various concentrations from 1 to 20 μ M (**Fig. 2C** & **S2C**). These condensates show properties consistent with LLPS such as droplet fusion (**Fig. 2D**) and fluorescent recovery after photobleaching (FRAP) (**Fig. 2E**). The Met32-mCherry condensates have a 64 ± 9.7 s half recovery time, comparable to well-established LLPS proteins such as FUS (5-120 seconds), DDX4 (3-200 seconds), and hnRNP A1 (5-120 seconds) (Nott et al. 2015; Shin et al. 2017; McSwiggen et al. 2019).

To further test potential Met4 / Met32 co-condensation, we added purified Met4 and Met32 together in 150 mM NaCl buffer and imaged in both Alexa488 and mCherry channels. The two proteins are mixed into droplets in solution (**Fig. 2F**). In these co-localized droplets, Met32-mCherry still undergoes FRAP recovery with a slightly slower rate (83 ± 11 s half recovery time) (**Fig. S2D**). These results are consistent with the idea that Met4 and Met32 can co-condense through a LLPS mechanism *in vitro*, which may contribute to the Met4 / Met32 puncta formation *in vivo*.

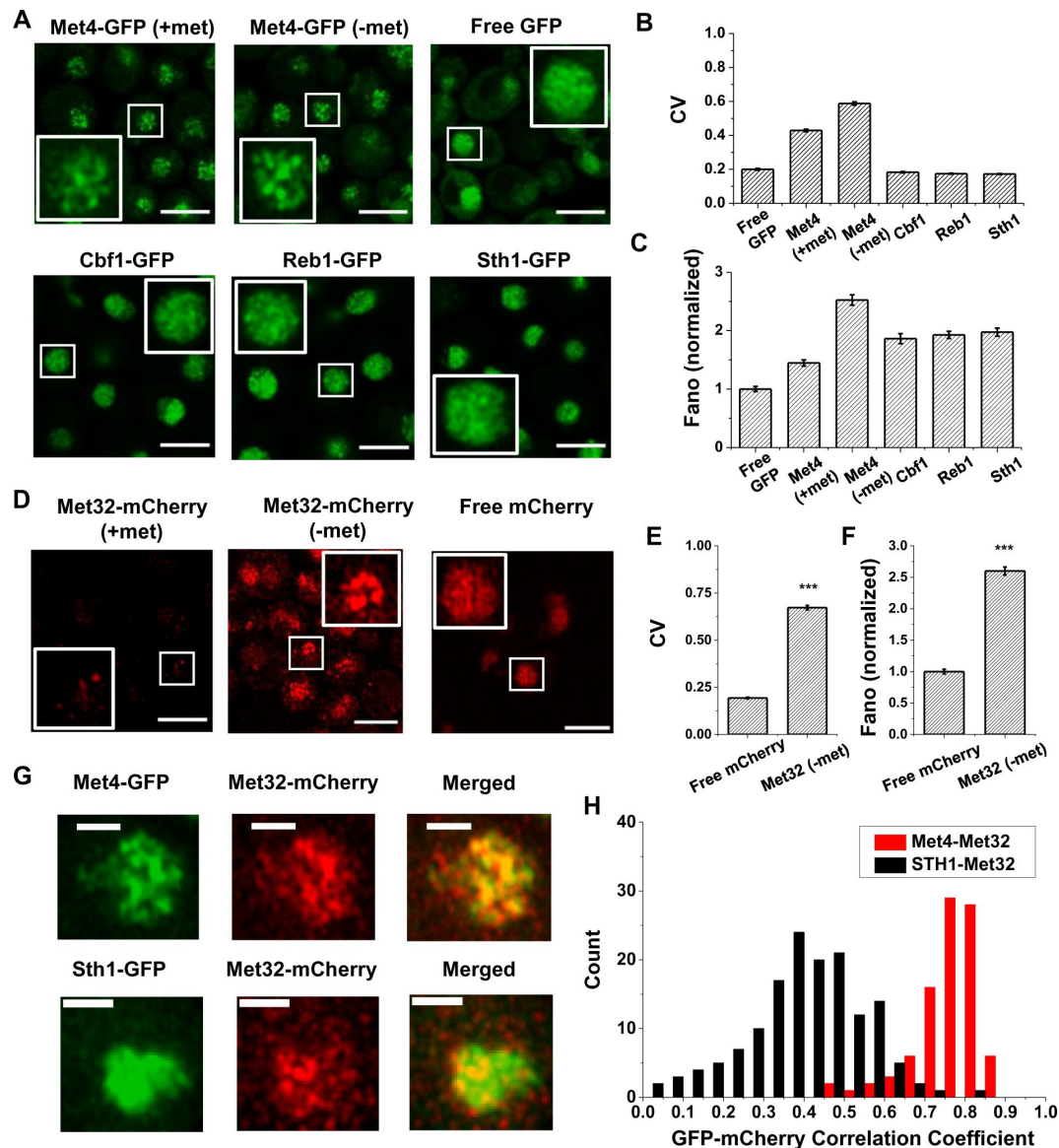


Figure 1.

The trans-activator Met4 and its co-factor Met32 form puncta in the nucleus.

A) Representative images of Met4-GFP in $\pm$ methionine (met) conditions. Control cells expressing free GFP (driven by the *HOpr*), Cbf1-GFP, Reb1-GFP, and Sth1-GFP cells are shown with similar contrasts. Scale bars represent 4 μ m (same as in D & G). **B & C**) The mean coefficient of variance (CV) and Fano numbers of nuclear pixel intensities for different versions of GFPs in panel A. The Fano numbers are normalized so that the free GFP has a Fano number 1. **D**) Representative images of strains expressing Met32-mCherry in $\pm$ met conditions. Control strain expressing free mCherry (driven by the *HOpr*) is shown with similar contrasts. **E & F**) The mean CV and normalized Fano number of pixel intensities of Met32-mCherry vs free mCherry. **G**) Representative images of strains co-expressing Met4-GFP and Met32-mCherry (top row), or Sth1-GFP and Met32-mCherry (bottom row). **H**) A histogram of Pearson correlation coefficient between co-expressed Met4-GFP and Met32-mCherry signals (red bars $N=93$) or Sth1-GFP and Met32-mCherry signals (black bars $N=149$) in each cell.

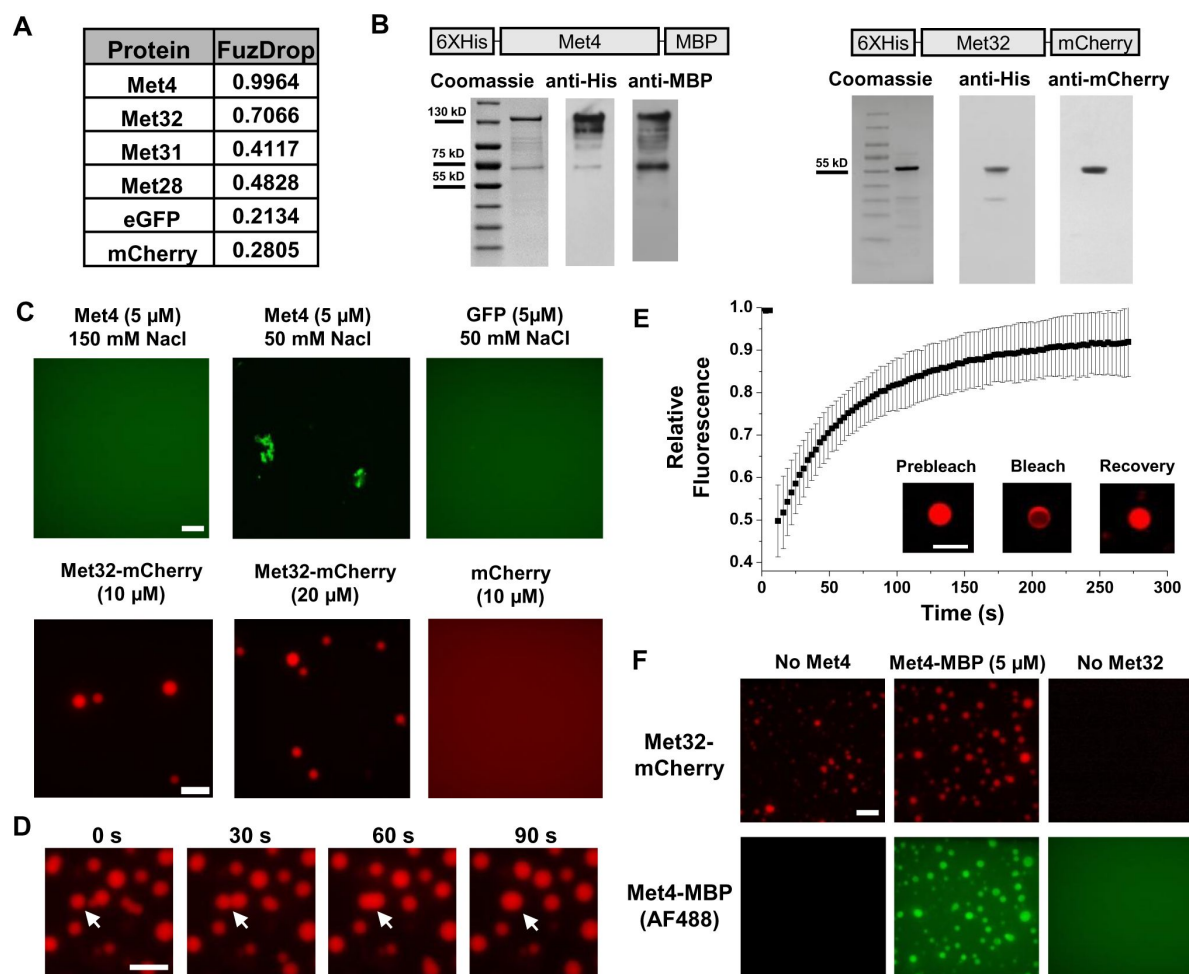


Figure 2.

Met32 forms condensates with LLPS properties *in vitro*.

A) Probabilities of individual Met TFs (Met4, Met32, Met31, Met28), GFP, and mCherry to undergo liquid-liquid phase separation from a published prediction program (Fuzdrop). **B)** Coomassie staining and western blots of purified Met4 and Met32 proteins. **C)** Protein aggregate and droplet formation observed for Met4-MBP and Met32-mCherry fusion proteins. Met4 and GFP are 5 μ M, Met32-mCherry and mCherry are 10 μ M in 20 mM HEPES pH 7.5, 150 mM NaCl buffer. Met4 and GFP are in 20 mM HEPES pH 7.5, 150 mM NaCl or 50 mM NaCl. Met4 is labeled with Alexa 488 (AF488). Scale bar represent 10 μ m (same as in D-F). **D)** Merging of Met32-mCherry droplets. Met32-mCherry is at 30 μ M in 20 mM HEPES pH 7.5, 150 mM NaCl. **E)** Fluorescence recovery after photobleaching (FRAP) data for 20 μ M Met32 droplets. The intensity data was collected every 3 seconds for 270 seconds and normalized to percent bleaching. Inset: Representative images of Met32 FRAP. **F)** Colocalization of Met32-mCherry droplets (10 μ M) with Met4-MBP (5 μ M) in 150 mM NaCl.

Met4-activated genes co-localize with Met4 puncta

Since Met32 is a sequence-specific DNA binder, the Met4 / Met32 condensates can potentially contact multiple target genes. This leads to our model in **Fig. 3A**, which predicts that 1) at least some Met4 activated genes should be co-localized with Met4 condensates, and 2) multiple target genes should cluster in 3D. To test these predictions, we first performed chromatin immunoprecipitation with sequencing (ChIP-seq) of related TFs (Met4, Met32, Met28, Cbf1), as well as RNA-seq, to identify genes directly targeted by Met4. As expected, the bindings of Met4, Met32, and Met28 are strongly induced by met depletion, while Cbf1, an abundant pioneer factor (Kuras et al. 1996; Blaiseau and Thomas 1998; Ouni et al. 2010; Carrillo et al. 2012; Yan et al. 2018; Donovan et al. 2019), shows constitutive binding (**Fig. 3B & C**, **S3A & B**). Under the induced condition, we identified 34 regions co-bound by all four TFs (**Fig. 3C**, **Table S3**), and these binding sites potentially regulate 46 genes (some sites are located in divergent promoters). Met depletion for 2 hrs causes wide-spread changes in the mRNA level with both up- and down-regulation (**Fig. 3D**, **S3C**), reflecting the fundamental role of methionine in metabolism (Thomas and Surdin-Kerjan 1997). Importantly, 40 out of the 46 genes co-bound by all four TFs show significant increased expression under the -met condition (**Fig. 3D**, **Table S4**), indicating that they are direct targets of Met4 and its co-activators.

To determine if the Met4 target genes are co-localized with the Met4 puncta, we inserted 196X tetO repeats near two of these genes, *MET13* and *MET6*, which allow us to visualize these loci through their association with tetR-mCherry. We also created control strains with the same array inserted near *PUT1* and the mating locus *MAT*, two loci with no Met4 association. These chromatin dots were imaged together with Met4-GFP under $\pm$ met conditions. Visually, *MET13* and *MET6* loci have higher probabilities to be co-localized with Met4 puncta in -met compared to *MAT* and *PUT1* (**Fig. 3E**). To test this more quantitatively, we aligned these images with the chromatin dots at the center and averaged the surrounding GFP signals (**Materials and Methods**). For all four loci, the averaged GFP intensities are higher near the center because the distant pixels are more likely to be outside nuclei boundaries, where Met4 signals are absent (**Fig. 3F & G**). In +met condition, Met4-GFP has low intensities. Upon activation, we can see localized GFP dots with heightened peak intensities at the *MET13* and *MET6* loci in comparison to controls (**Fig. 3F & G**). These results indicate that at least some Met4 target genes are associated with Met4 condensates upon activation, presumably through sequence-specific co-binders like Met32.

Met4-dependent clustering of *MET* genes upon induction

We next tested the second prediction of the model in **Fig. 3A**, i.e., Met4 condensates mediate the 3D clustering of target genes. We first carried out a Hi-C measurement under -met condition, which did not show any significant interactions among Met4 targets (Li et al. 2023). Given the limitations of Hi-C in detecting fine-scale interactions (Oluwadare et al. 2019), we used an assay recently developed in our lab, Methyltransferase Targeting-based chromosome Architecture Capture (MTAC) (Li et al. 2023), to identify potential chromosomal interactions with selected Met4 targets (**Materials and Methods**). In this method, a 256X LacO array is inserted near the gene of interest (viewpoint, or VP) to recruit LacI fused with DNA methyltransferase M.CviPI, which methylates proximal cytosines in the “GC” context in *cis* and in *trans* (**Fig. 4A**). Physical proximity to the VP can thus be evaluated based on the enrichment of the methylation signal in comparison to a control strain containing the same LacI-M.CviPI but no LacO array.

In a previous study (Li et al. 2023), we performed MTAC assays with a VP near the *MET6* gene under $\pm$ met condition. This VP 3kb away from *MET6* captures highly-specific inter-chromosomal interactions with four other Met4-targeted genes under -met, but not +met, condition. To further illustrate this point, we carried out MTAC with *MET13* as the VP (**Fig. 4B**). Under +met condition, MTAC only detects interactions from the adjacent local chromatin (**Fig. 4C**, **S4A**). When the same strain was switched to the -met condition, MTAC captures two far-*cis* interactions (intra-

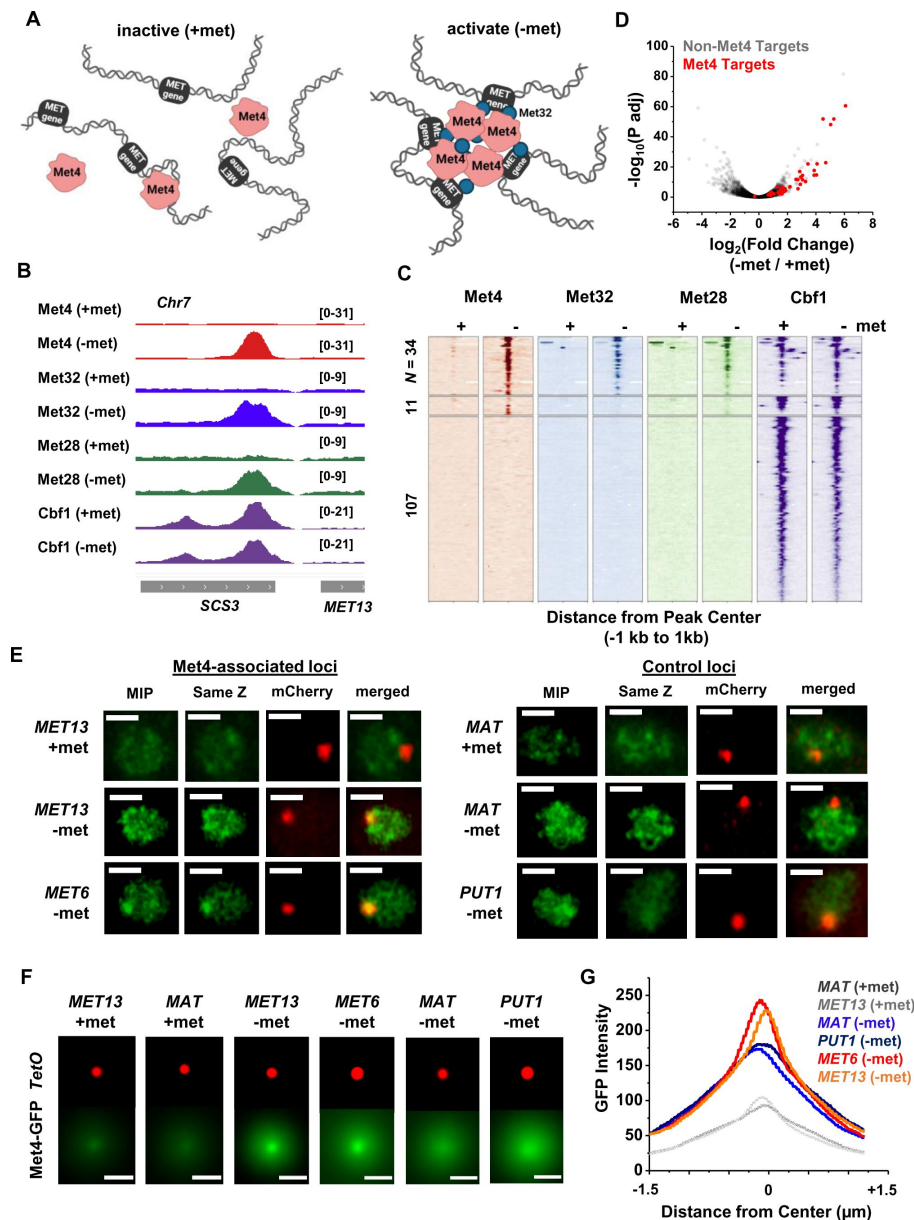


Figure 3.

Met4-activated genes co-localize with Met4 puncta.

A) Model of Met4 condensate in the chromatin context. Upon met depletion, Met TFs may form condensates that interact with multiple target genes, leading to the 3D clustering of these co-regulated genes. **B**) Examples of ChIP-seq signals of Met TFs (Met4, Met32, Met28 and Cbf1) in $\pm$ met conditions. The Met TF co-binding peak shown here is near the *MET13* gene. **C**) Heatmaps of Met4, Met32, Met28, and Cbf1 ChIP-seq peaks in $\pm$ met conditions. The peaks were clustered into ones enriched with all four Met TFs ($N = 34$), enriched with Met4 and Cbf1 ($N = 11$), and enriched with Cbf1 only ($N = 107$). **D**) Volcano plot of RNA-seq data comparing mRNA levels in $\pm$ met conditions. Most genes near Met TFs co-binding peaks are strongly induced by met depletion (red dots). **E**) Single nuclei images of yeast cells expressing Met4-GFP and TetR-mCherry with a TetO array integrated near *MET13* / *MET6* (Met4 targets), or *MAT* / *PUT1* (non-targeted control). Images were taken with 14 z-stacks with step size 0.4 μ m. "MIP" and "same Z" show Met4-GFP images with either maximum intensity projection among all z stacks, or with a single stack at the same z plane as "mCherry", where the mCherry labeled TetO array shows the highest intensity. "merged" is the merged image of the "same Z" and the "mCherry". Scale bars represent 1 μ m (same as in F). **F**) Average mCherry and GFP intensities centered at the mCherry labeled TetO arrays located at different genomic sites. **G**) The GFP intensity profile across the dot center in panel E. The GFP intensity near the dot center is significantly higher for *MET13* / *MET6* loci.

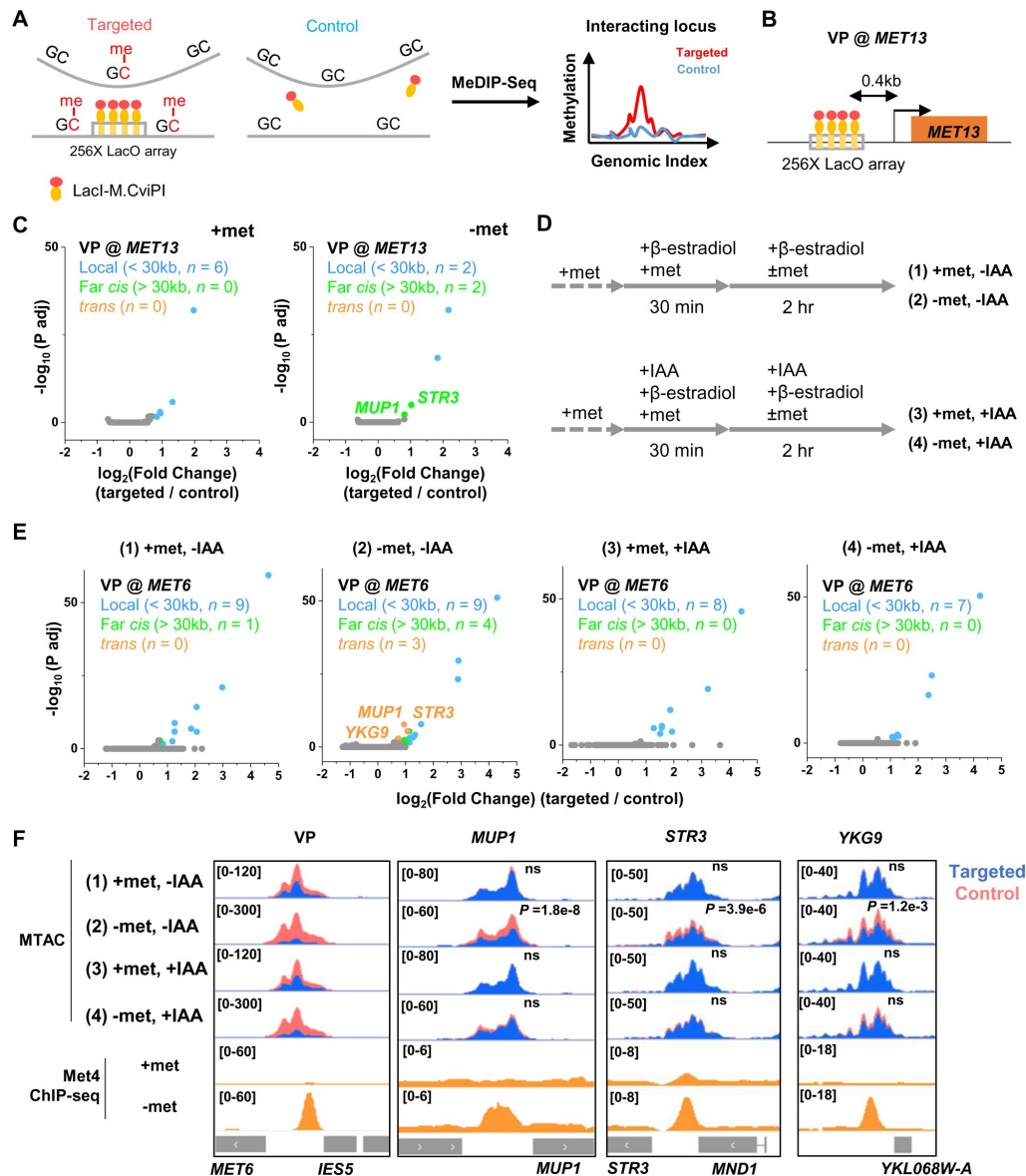


Figure 4.

Met4-dependent clustering of *MET* genes upon induction.

A MTAC workflow. In a "targeted" MTAC strain, a LacO array is integrated into a genomic locus (viewpoint, VP) and recruits LacI-M.CviPI, an ectopic DNA methyltransferase that methylates the cytosine in a "GC" dinucleotide in proximal DNA. LacI-M.CviPI is also expressed in a control strain with no LacO array insertion (background methylation). Methylation in these two strains is detected by ChIP, and methylation level in nucleosome-depleted regions (NDRs) are compared in targeted vs control strains. Significantly higher methylation in the targeted strain indicates proximity to the VP. **B** VP design of the *MET13* locus. **C** Volcano plot of MTAC signals with *MET13* as VP in $\pm$ met conditions. Each dot represents an individual NDR, and colored dots are the NDRs that show proximity to the VP (significantly higher methylation in the targeted vs control strains). Local (intra-chromosomal interactions within 30kb), far-cis (intra-chromosomal interactions over 30kb), and trans (inter-chromosomal interactions) are shown in blue, green, and orange. Same as below. **D** Design of Met4 depletion assay. Met4 is depleted by auxin-degron system in $\pm$ met conditions, resulting in four conditions: (1) +met, -IAA, (2) -met, -IAA, (3) +met, +IAA, (4) -met, +IAA. β -estradiol is added in all conditions to induce the expression of LacI-M.CviPI. **E** Volcano plot of MTAC signals with *MET6* as VP for the four conditions in panel D. Note that long-distance interactions are detected in condition (2) but are largely absent in other three conditions. **F** MTAC and Met4 ChIP-seq data at the *MET6* locus (VP) and *MUP1*, *STR3* and *YKG9* as interacting regions of the VP. MTAC signal is shown in the four conditions in panel D. The ChIP enrichment of Met4 is shown in $\pm$ met conditions. *P*, FDR-adjusted *P* value, Wald test by DESeq2. ns, non-significant.

chromosomal interactions that are over 30kb apart) with two promoter regions near Met4 targeted genes (*STR3*, *MUP1*) (Fig. 4C, S4A). Interestingly, these two regions were also found to interact with the *MET6* VP (Li et al. 2023). Taken together, these data support the clustering of a subset of Met4-targeted genes upon activation.

The model in Fig. 3A implies that the clustering of *MET* genes is mediated by Met4 / Met32. To test this idea, we carried out MTAC using the *MET6* VP ± acute auxin-induced degradation of Met4 ± met (Fig. S4B & C). For the -met experiment, auxin (IAA) was added 30 mins prior to the met depletion so that Met4 activation would be largely eliminated (Fig. 4D). This would also prevent the accumulation of Met32, as it relies on Met4 for induction (Menant et al. 2006; Lee et al. 2010; Carrillo et al. 2012). Consistent with our previous observations, very few long-distance interactions were found in the +met repressed condition, regardless of the presence or absence of Met4 (Fig. 4E, (1) and (3)). Upon met depletion without IAA addition, we observed significant *trans* (inter-chromosomal) interactions between the *MET6* locus with three other Met4-targeted genes (*MUP1*, *STR3*, *YKG9*) (Fig. 4E, (2)). Importantly, these interactions disappear when Met4 was degraded (Fig. 4E, (4)). These data show that the presence of Met4, and likely Met32, is required for *MET* gene clustering. This is consistent with the model in Fig. 3A that clustering is mediated by Met4 / Met32 condensates.

Regions near Met TF binding sites constitute *MET* “transcriptional hotspot” in haploid yeast

We next investigated the functional role of Met4 condensates in gene expression. In our previous screen, we identified the *MET13* locus on chr7 as a “transcriptional hotspot”: when the *S.kud MET3pr*-GFP reporter was inserted into *MET13* or neighboring genes, it showed higher GFP expression than most of the other insertion sites upon induction (Du et al. 2017). Combined with the observation that *MET13* locus co-localizes with the Met4 puncta (Fig. 3E-G), this suggests a potential link between the higher reporter activity and Met4 / Met32 condensates. However, the initial screen was performed in diploids, while all the experiments above were carried out in haploids. We therefore first tested if *MET13* also functions as a transcriptional hotspot in haploid yeast. We again used the *S.kud MET3pr* for this test, as the sequence of this promoter deviates significantly with the endogenous *MET3pr*, allowing them to be differentiated in PCR (Du et al. 2017).

We created two haploid strains with the *S.kud MET3pr*-GFP reporter inserted near the *MET13* gene (*CWC23* and *RSM23*) and measured its transcriptional profile (Fig. 5A) (Materials and Methods). For comparison, we also inserted the same reporter into two non-hotspot basal loci *ATG36* and *LDS2* (Fig. 5A). The GFP intensities from the reporter at *CWC23* and *RSM23* loci are ~50% higher (P -value < 1e-4) compared to the control under the -met condition (Fig. 5B). Reporter GFP mRNA levels, as well as Pol II associated with the GFP ORF, are also significantly higher in the former strains (Fig. 5C & D). These results are consistent with our previous observations in diploid cells that the *MET13* locus promotes the transcriptional activity of the *MET3pr*.

Given the model in Fig. 3A, we reasoned that the enhanced activities of the GFP reporter may be due to increased local concentrations of TFs, which likely lead to higher TF occupancies. To test this idea, we performed ChIP of Met4 and Met32 followed by qPCR on the *S.kud MET3pr* at the hotspot vs basal genomic locations because it contains a Met32 motif. Consistent with our expectation, we observed higher enrichment of Met4 and Met32 over the *S.kud MET3pr* at hotspot locations (Fig. 5E & F). Taken together, these findings confirm the existence of “hotspot” loci in haploid yeast and suggest that it may be caused by enhanced binding of Met32 and Met4.

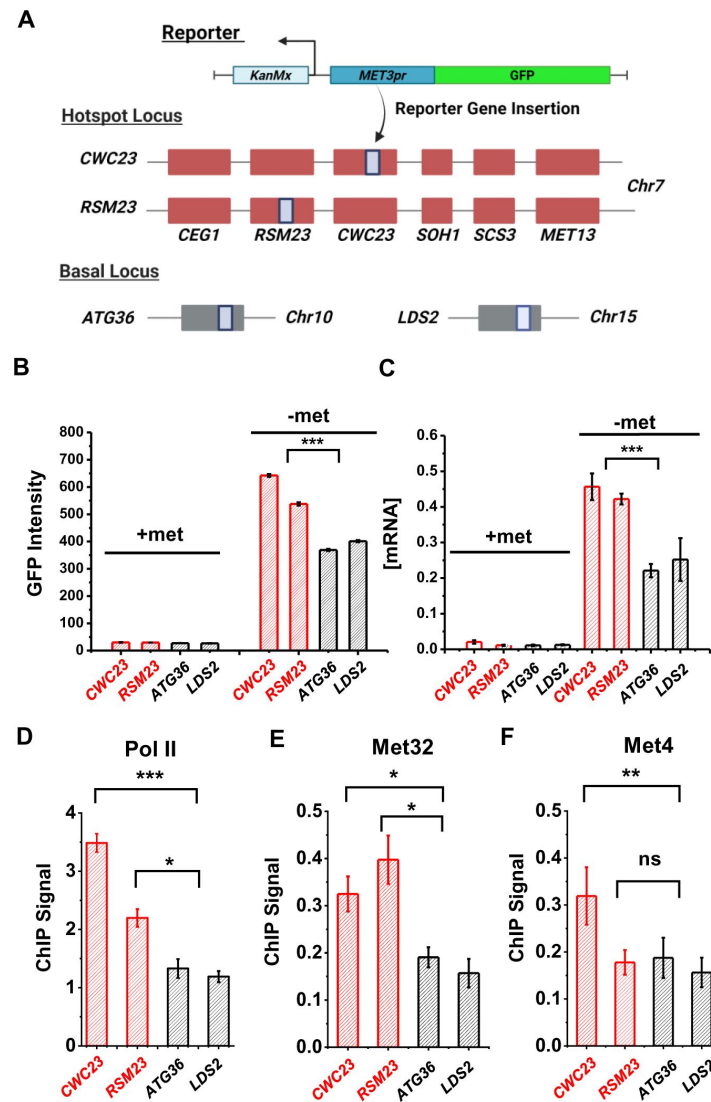


Figure 5.

Characterization of a *MET* “transcriptional hotspot” in haploid yeast.

A) Schematic of strains constructed with *S.kud* *MET3pr*-GFP reporter inserted near the *MET13* transcriptional hotspot (*CWC23*, *RSM23*) and “basal” loci (*ATG36*, *LDS2*). **B & C**) Mean cellular GFP fluorescent intensities and GFP mRNA measured by qRT-PCR in the four strains above in $\pm$ met conditions. **D**) ChIP-qPCR of Rpb1 over the GFP ORF in the four strains above in -met. **E & F**) ChIP-qPCR of Met32 and Met4 over the *S.kud* *MET3pr* in the four strains above in -met. The asterisks represent *: <0.05, **: <0.01, ***: <0.001. Same as below.

Reporter activities near Met4 binding sites are enhanced over a ~40kb range

Besides *MET13*, other loci that colocalize with Met4 puncta, like *MET6* (Fig. 3E-F), should also function as transcriptional hotspots. To test this idea, we first verified that *MET6* remains associated with Met4 puncta after the reporter insertion. We used the *MET6* yeast strain in Fig. 3E and integrated a *S.kud MET3pr* reporter nearby (Fig. 6A). This reporter contains a frame-shifted GFP (fsGFP) in order not to interfere with the Met4-GFP signal. As controls we integrated the same reporter near two genomic regions not bound by Met TFs (*PUT1* and *ATG36*). We imaged these strains in -met and analyzed them with the same method used in Fig. 3F. Interestingly, despite the fact that *S.kud MET3pr* is bound and activated by Met4, insertion of this promoter does not increase the local Met4 intensity at *MET6* and *PUT1* (Fig. 6B & C, $\pm$ rep). In comparison with the reporter-containing *PUT1* and *ATG36* loci, *MET6* again shows stronger co-localization with the Met4 puncta (Fig. 6B & C). This finding indicates that *S.kud MET3pr* alone is insufficient in nucleating a new Met4 condensate or recruiting the integrated locus to an existing condensate.

We then integrated the original *S.kud MET3pr*-GFP reporter into *MET6*, *ATG36*, and *LDS2* (the latter two are controls far from any endogenous Met4 binding sites) (Fig. 6D) and compared their GFP expressions. GFP shows significantly higher intensity at *MET6* in comparison with the two basal loci (P -value $< 1e-4$) (Fig. 6E). To test the generality of this observation, we inserted the reporter into two additional genomic loci associated with Met TFs, *MET17* and *RMD6*, and the reporter shows higher GFP expression in both loci (Fig. 6E). This data supports a model where Met4 condensates associated with the endogenous *MET* genes enhance the activity of a nearby reporter. This model also predicts that such enhancement effect should be specific to the met pathway, i.e., inducible genes activated by other TFs should not benefit from higher local concentrations of *MET* activators. We tested this idea by inserting into the same loci a GFP reporter driven by either *MET17pr*, a promoter in the same pathway, or by *GAL1Spr*, a promoter controlled by a different set of TFs in another pathway (Rohde et al. 2000; Slot and Rokas 2010). Indeed, *MET17pr* shows the same trend as *MET3pr*, while the strengths of *GAL1Spr* remain constant across all these loci (Fig. 6F). These results demonstrate the pathway-specificity of the transcriptional hotspots.

We next conducted experiments to determine the genetic range of the transcriptional hotspot. We inserted the GFP reporter driven by *MET3pr* or *GAL1Spr* at various distances from the Met4 binding site in the *MET6* promoter (Fig. 6G). *MET3pr*-GFP shows higher expression level when inserted up to 30kb upstream and 10kb downstream of the Met4 site. Beyond this region, the *MET3pr* activity returns to the basal level (Fig. 6H). In contrast, *GAL1Spr* shows similar activities among these insertion sites (Fig. 6H). These data indicate that, at least near the *MET6* gene, the elevated expression of the *MET3pr* can occur over a ~40kb region.

The deletion of a disordered region in Met4 reduces puncta formation and reporter expression at transcriptional hotspots

To determine the relation between Met4 condensates and its gene regulatory function more directly, we introduced mutations to Met4 to reduce its propensity to form condensates. Since LLPS is often promoted by IDRs, we first performed PONDR analysis of Met4 (Xue et al. 2010). This analysis reveals three distinct stretches of IDRs interspersed with two previously annotated functional domains: the activation domain (aa95-144), which interacts with mediator, and the auxiliary domain (aa312-375), which interacts with Met31/32 (Fig. 7A) (Kuras and Thomas 1995). AlphaFold also predicts that Met4 is largely devoid of folded structures (Fig. S2A) (Jumper et al. 2021). To probe the effect of these IDRs on Met4 puncta formation, we first truncated individual IDR stretches: Δ IDR1 (aa1-69), Δ IDR2 (aa117-359), or Δ IDR3 (aa397-651). As Met4 is essential for cell viability in the -met media, we constructed strains containing the endogenous unlabeled Met4 with an additional copy of truncated Met4-GFP driven by the native *MET4*

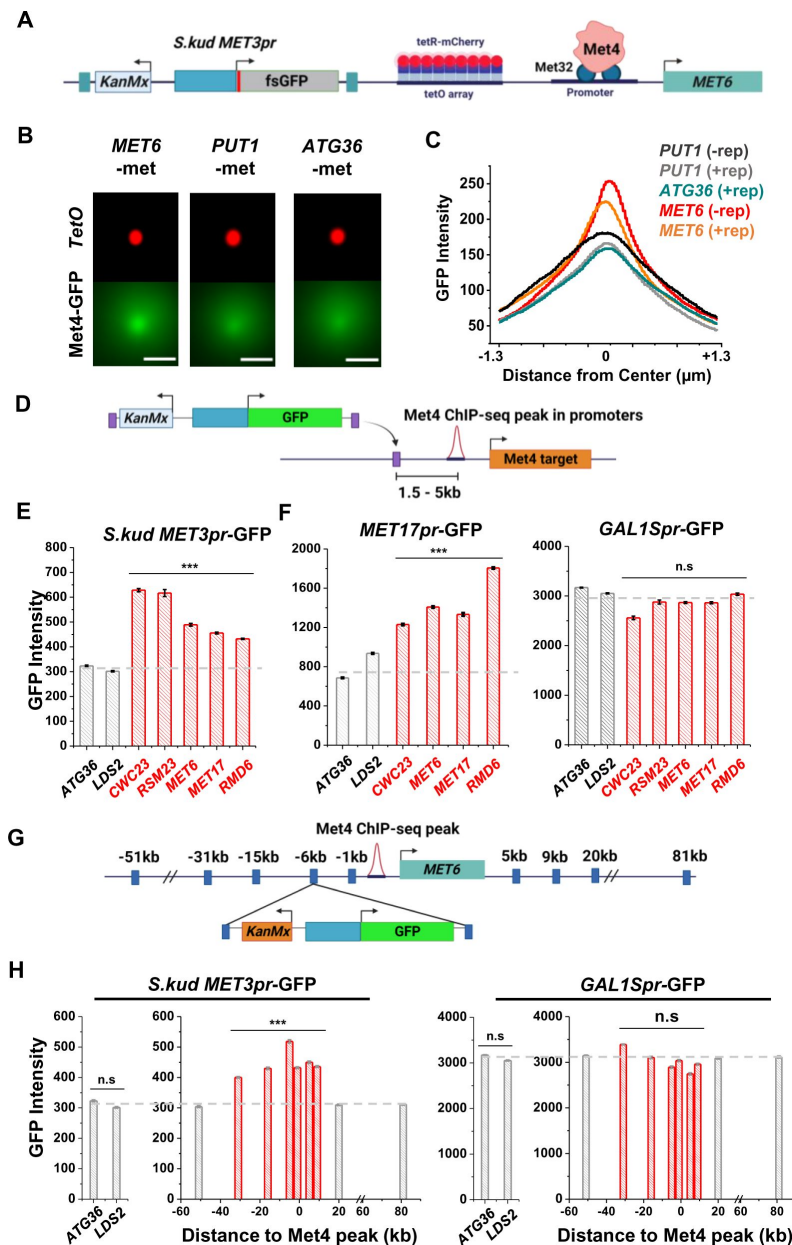


Figure 6.

Reporter activities near Met4 binding sites are enhanced over a ~40kb range.

A) Schematics of measuring the co-localization of the GFP reporter with Met4 puncta. *S.kud* MET3pr-fsGFP (frameshifted GFP) reporter gene and a tetO array (196x) are inserted side by side into the genome, in this case near the *MET6* gene. **B**) Average mCherry and GFP intensities centered at the mCherry labeled tetO arrays near *MET6* (Met4 target) and *PUT1* / *ATG36* (not Met4 targets) in the presence of nearby reporter. Scale bars represent 1 μ m. **C**) The GFP intensity profile across the dot center in panel B. For *MET6* and *PUT1*, the same type of data without the GFP reporter (-rep) are also included. **D**) Schematic of GFP reporter insertion near three additional Met4-bound loci, *MET17*, *RMD6*, and *MET6*. The distance and orientation of the insertion are labeled in the diagram. **E**) Mean GFP fluorescent intensities when *S.kud* MET3pr-GFP are inserted near indicated genes. The genes labeled in red have adjacent Met4-bound sites, while the ones labeled in grey do not. **F**) Same as in panel E except with *MET17pr*-GFP and *GAL1Spr*-GFP reporter. Strains with *MET17pr*-GFP were grown in -met, and the ones with *GAL1Spr*-GFP were pre-grown in raffinose and induced by galactose for 6 hours. **G**) Schematic of the *MET3pr*-GFP reporter inserted at various distances from the Met4 binding site near the *MET6* gene. The same orientation was used for all the loci as indicated. **H**) Mean GFP fluorescent intensities with *MET3pr*-GFP and *GAL1Spr*-GFP reporters inserted into locations indicated in panel G, in comparison to the same reporter inserted into two control loci far from Met4 binding sites.

promoter. This allows us to visualize Met4 mutants without hindering cell growth (**Fig. 7B** [↗](#)). Deletions of IDR2 and IDR3 both lead to significant reductions in Met4-GFP Fano number, with Δ IDR2 having a slightly more pronounced effect (**Fig. 7B & C** [↗](#), **S5A**). To further specify the part of IDR2 that affects Met4 condensates, we generated shorter truncations within IDR2, Δ IDR2.1 (aa117-135), Δ IDR2.2 (aa136-270), and Δ IDR2.3 (aa271-359). All three truncations result in intermediate GFP Fano number between the full length Met4 and Met4- Δ IDR2 (**Fig. 7C** [↗](#), **S5B & C**). These findings suggest that multiple IDRs contribute to Met4 condensation, and deleting parts of the Met4 IDR can diminish, but not completely eliminate, Met4 puncta formation.

When the full length Met4 is replaced by Met4 Δ IDR2.3, the strain is viable in the absence of met, allowing us to evaluate the functional consequence of this mutant on Met4 binding and transcriptional activation. Given our model and supporting evidence that Met4 condensates increase local TF concentration at Met4 targeted genes and enhance their expression, we expect a Met4 mutant less prone to condensate will result in diminished Met4 binding and reduced induction. We therefore compared the ChIP-seq and RNA-seq data of the full-length vs truncated Met4. Consistent with the idea above, the Met4 Δ IDR2.3 shows decreased binding across all the previously annotated Met TF binding sites, without generating new ChIP-seq peaks (**Fig. 7D** [↗](#) & **E**). Interestingly, the three Met4 binding sites near *MUP1*, *STR3*, and *YKG9* that cluster with *MET6* show significantly more reduction in Met4 ChIP-seq peaks (reduced to $42 \pm 4\%$) than other Met4 sites ($61 \pm 8\%$, P -value = 0.02) in the Met4 Δ IDR2.3 strain. Out of the 40 potential target genes, 22 show significant decreases in their mRNA level (**Fig. 7F** [↗](#), **S5D**, **Table S4**). Importantly, our model also predicts that decreased Met4 condensation should have more impact on target genes that are located near Met4 puncta than those farther away. To test this idea, we again took advantage of the *MET3pr*-GFP reporter inserted into hotspots and basal loci. We expect the reduced Met4 condensation to have a larger impact on the reporter activity in the former case because the reporter is only associated with Met4 puncta at hotspots (**Fig. 6B** [↗](#)). Indeed, after substituting Met4 with Met4- Δ IDR2.3 in these strains, we observed strong decreases in the reporter gene activity in hotspots but minimal effect on those in basal regions (**Fig. 7G** [↗](#)). These findings collectively support a model wherein Met4 condensates cluster target genes, enhance their activity, and give rise to the formation of transcriptional hotspots.

Discussion

In this study, we investigated TF distribution, 3D genome organization, and gene activation in the met response pathway using a combination of genetics, genomics, and imaging approaches. Our findings connect three emerging phenomena in gene regulation: TF condensates, co-regulated gene cluster, and transcriptional hotspot. More specifically, we show that the activator Met4 and its co-factor Met32 form puncta-like structures that co-localize with at least some Met4 target genes. Multiple *MET* genes on different chromosomes cluster in the 3D nuclei space, and such clustering requires the presence of Met4. Chromosomal loci near endogenous Met4 binding sites, in one case within a 40kb range, are hotspots for *MET3pr* reporter expression. Mutations that decrease Met4 puncta formation lead to selective reduction of the reporter transcriptional activity at the hotspots.

Condensate formation of Met4 and Met32

Using live-cell imaging, we found that fluorescently labeled Met4 and Met32 form puncta-like structures in yeast nuclei under the -met condition. This is reflected by the higher Fano numbers in Met4 / Met32 pixel intensities in comparison to free GFP, and in the Met4 case, other chromatin-associated factors. High resolution microscopy of yeast TFs is challenging due to the small size of the nuclei. The low intensities of the Met4 / Met32 signals require high excitation for imaging, which also makes them prone to photo-bleaching. Thus, we were unable to measure Met4 and Met32 puncta dynamics *in vivo*. When purified *in vitro*, Met32-mCherry readily forms droplets

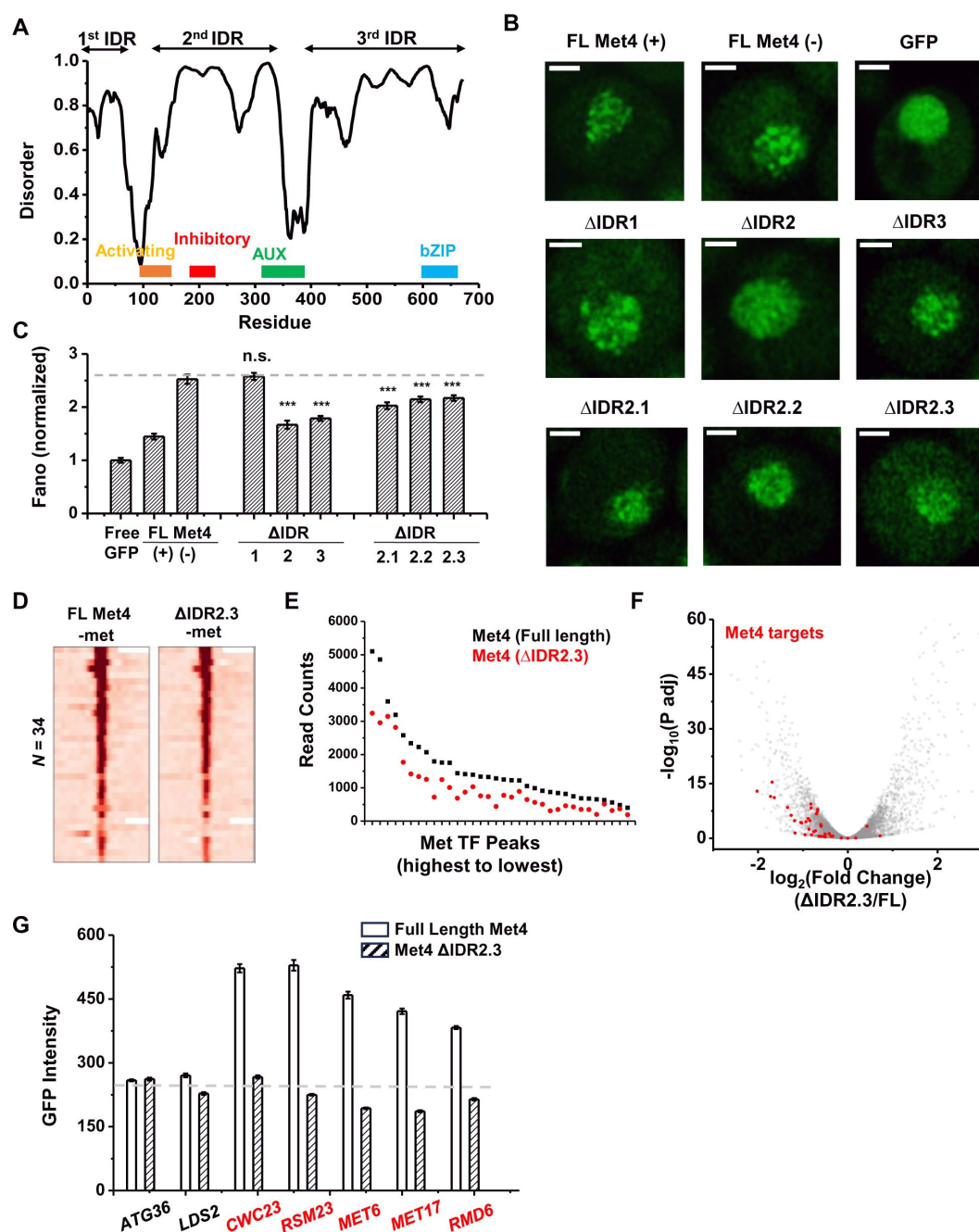


Figure 7.

The deletion of a disordered region in Met4 reduces puncta formation and reporter expression at transcriptional hotspots.

A) PONDR disorder plot of Met4 with previously annotated functional domains, including activation domain (aa95-144) that interacts with mediator, inhibitory domain (aa188-235), auxiliary domain (aa312-375) that interacts with Met31/32, and bZIP domain (aa595-660) that interacts with Cbf1. **B**) Single nuclei images of cells expressing full length Met4 ($\pm$ met) or Met4 with various truncations (-met) fused with GFP. **C**) The Fano numbers of nuclear pixel intensities for different versions of GFPs in panel B. **D**) Heatmaps of full length Met4 and Met4 Δ IDR2.3 ChIP-seq in -met over previously identified Met TF binding sites. **E**) Alignment counts underneath the Met TF ChIP-seq peaks over the same sites for TAP tagged full length Met4 and Met4 Δ IDR2.3 (sorted from high to low). **F**) Volcano plot from RNA-seq data from cells containing full length Met4 or Met4 Δ IDR2.3 in -met. Genes associated with Met TFs (Fig. 3C) are indicated in red. **G**) Mean GFP intensities from cells expressing full length Met4 and Met4 Δ IDR2.3 with *MET3pr*-GFP reporter inserted into loci near (red) or far away from (black) Met4-target genes. The dashed lines represent basal reporter expression.

with typical LLPS properties without molecular crowder, while His- and MBP-tagged Met4 remains diffusive at physiological salt concentration. We also found that Met4 partitions into the Met32 droplets when they are mixed, and inside cells, the two factors show significant co-localization. These observations raise the intriguing possibility that Met32 serves as a scaffold to nucleate the co-condensates of Met4 and other Met TFs. This may explain why the Met4 distribution becomes more puncta-like when switched from +met to - met condition, when Met32 becomes available inside the nuclei. However, we cannot exclude the possibility that the absence of Met4 LLPS is due to our *in vitro* condition (tag, impurity, lack of post-translational modification, etc.). Met4 is also reported to recruit the Mediator and SAGA complex (Leroy et al. 2006 [↗](#)), and these factors, together with Pol II and RNA, may also affect Met4 condensate formation *in vivo* (Henninger et al. 2021 [↗](#)).

TF condensates and the 3D genome topology

TF condensation *in vivo* occurs in a chromatin context, but the relation between TF condensates and 3D genome topology has not been extensively studied. By recruiting multiple TFs, mediator, and transcription machineries into proximity, TF binding loci can potentially serve as nucleation sites for transcription condensates. Looped enhancers and promoters, as well as clustered genes, may also function as scaffold and stabilize these condensates. This is best illustrated at super-enhancers (SEs), where clustered enhancer sites tend to associate with transcriptional condensates and lead to very high activities of their target genes (Hnisz et al. 2013 [↗](#); Sabari et al. 2018 [↗](#)). Synthetic LacO array and endogenous microsatellite repeats were also shown to promote local condensation of associated TFs (Chong et al. 2018 [↗](#)). Reversely, TF condensates can mediate long-distance chromosomal interactions by contacting multiple genomic sites and therefore inducing their 3D proximity. This idea is supported by the findings here: 1) multiple Met4-targeted genes are co-localized with Met4 condensates, 2) at least a subset of Met4-targeted genes form clusters with long-distance intra- and inter-chromosomal interactions, and 3) such clustering only occurs in - met activating condition in the presence of Met4. Similar observations were made during yeast heat shock response, where the activator Hsf1 coalesces and mediates transient clustering of heat shock genes (Chowdhary et al. 2019 [↗](#)). A few other mammalian TFs were also shown to contribute to 3D genome conformation. For example, YAP/TAZ organizes their target sites into clusters that co-localize with the YAP/TAZ/TEAD1 condensates (Lu et al. 2020 [↗](#)). NUP98-HOXA9, a chimeric TF related to the pathogenesis of AML, was also shown to mediate long-distance chromosomal loops between NUP98-HOXA9 bound enhancers and oncogenes into SE-like clusters (Ahn et al. 2021 [↗](#)).

Interestingly, a *MET3pr* reporter that is associated and activated by Met4 only co-localizes with Met4 puncta when inserted near native Met4 binding sites. Also, adding *MET3pr* reporter gene into these sites does not further increase the local Met4 concentration (Fig. 6B & C [↗](#)). These observations indicate that Met4 binding is not *sufficient* for a gene to nucleate or be recruited into a Met4 condensate. The underlying mechanism of this observation is unclear, but we speculate that some properties of the endogenous *MET* genes may facilitate their clustering with Met4 puncta. For example, some *MET* genes, like *MET13* and *MET6*, are associated with nuclear pore complex (NPC) even in the presence of met (Forey et al. 2021 [↗](#)). The tethering by NPCs may increase the chance for these genes to come into proximity, enhancing the local density of Met4 target sites to nucleate the Met4 condensates. In contrast, a reporter gene inserted into random genomic locations may not be optimally positioned to contact NPC and/or other Met4 target genes. Further experiments are required to test these ideas. Notably, interaction with NPC was found to be necessary and sufficient to promote clustering of the *GAL1-10* alleles (Brickner et al. 2016 [↗](#)).

TF condensates and gene expression

The Met4 condensates represent a local environment with elevated Met4 concentrations, which can promote Met4 binding and lead to enhanced gene expression. Consistent with this idea, *MET3pr* reporter inserted near these Met4 targets, which also co-localizes with the Met4 puncta, shows increased transcriptional activity. Near *MET6*, such hyper-activity of the reporter occurs

over a 40kb range. According to previous measurement (Bystricky et al. 2004 [↗](#)), 40kb linear chromatin explores a 3D space with ~200 nm diameter, which may reflect the size of the Met4 condensates. The elevation of transcription level is pathway-specific as *GAL1Spr*-GFP inserted into the same loci does not show the same effect. This finding agrees with the “specialized transcription factory” previously proposed (Bartlett et al. 2006 [↗](#)) and argues against a model where the transcription elevation solely relies on general transcription factors like mediator or transcription machinery.

The relation between Met4 condensation and transcriptional activity is further supported by Met4 Δ IDR2.3. This Met4 mutant, in comparison to full length Met4, reduces the Met4 puncta formation and the activities of most of its target genes in the genome. Importantly, Δ IDR2.3 does not have strong effect on the transcriptional activity of *MET3pr* reporter inserted into basal loci, suggesting that this truncation does not directly impact Met4 integrity and its activation function. Instead, it selectively reduces reporter activity at transcriptional hotspots, consistent with the idea that such reduction is due to the disruption of Met4 condensates.

In terms of mechanism, we speculate that 1) high Met4 local concentration inside the condensates may increase the binding rate of Met4, and/or 2) multivalent interactions among Met4, Met32, and possibly other co-activators may slow down the dissociation of Met4 from its target genes. Both ideas are consistent with our observation that Met4 Δ IDR2.3 shows less binding than the full length Met4. The same mechanism has been proposed for other TF condensates, and in some cases, supported by direct kinetic evidence (Trojanowski et al. 2022 [↗](#)). It should be noted that more TF binding does not necessarily lead to an increase in gene expression level, for example, transcription may be limited by a step downstream of TF binding. Consistent with this idea, a recent study reports that Gal4 condensation facilitates its recruitment to target genes but does not contribute to gene activation (Meeussen et al. 2023 [↗](#)). In human cells, artificially enhanced condensation of a TF Ews::Fli1 causes sequestering of this TF into the nucleolus, resulting in decreased expression of their target genes (Chong et al. 2022 [↗](#)). This indicates that the exact relation between TF condensates and gene expression relies on the nature of the multivalent interaction, location of condensate formation, transcription kinetics, etc., and it needs to be analyzed on a case-by-case basis.

TF condensates and stress response

The sulfur-containing amino acid methionine is a key metabolite for cell survival, and its depletion represents a major stress for budding yeast. Interestingly, the TFs that were found to form condensates in yeast, include Met4, Gal4, and Hsf1, all respond to certain types of environmental stress. Similar observations are made in higher eukaryotes. For example, YAP forms condensates in response to osmotic stress (Cai et al. 2019 [↗](#); Lu et al. 2020 [↗](#)), and phyB in plants uses altered LLPS properties to sense light and temperature cues (Chen et al. 2022 [↗](#)). In contrast, constitutive TFs in yeast, like Reb1, Cbf1, and Sth1, are more evenly distributed in the nuclei (Fig. 1 [↗](#)). The condensation of stress-responding TFs may reflect a functional need to rapidly concentrate related resources into a subset of the nuclei space for more efficient usage. In the case of Met4, its condensates indeed allow yeast cells to mount a stronger response to met depletion. In addition, TF condensates may accelerate TF target search and response rate, facilitate synchronized expression of target genes, and/or enhance the activation specificity. These potential condensate functions need to be further explored.

Materials and Methods

Plasmid and strain construction

Standard molecular methods were used to construct budding yeast strains and plasmids (**Table S1**). The haploid strain was derived from w303a background, and most genetic modifications here involve homologous recombination. More specifically, to create strains with GFP-tagged endogenous Met4, the *GFP-HIS3* fragment was PCR amplified from a plasmid (pJL1) with primers containing upstream and downstream homologous sequences of the endogenous *MET4* gene. The amplified PCR product was transformed into haploid yeast to insert into the *MET4* locus. A similar strategy was used to create strains with mCherry-tagged endogenous Met32, Met4-TAP, Met32-TAP, Met28-TAP and Cbf1-TAP strains. For strains with tetO-tetR-mCherry labeled loci, variations of plasmid pSR11 were created from containing 196X tetO repeats with flanking homologous sequences to targeted sites were linearized and transformed into Met4-GFP expressing haploid strains. For expression of tetR-mCherry, a plasmid (pMY39) containing the tetR-mCherry insert was linearized and transformed into the *ADE2* genomic locus. To create Met4 truncated strains, individual plasmids containing truncated versions of Met4 (pJL1) were created and were linearized with restriction digestion. The linearized plasmid was then integrated into haploid yeast directly upstream of the *MET4* locus. For endogenous replacement of *MET4* with *MET4* (Δ IDR2.3), a plasmid variant of pJL1 with *MET4* (Δ IDR2.3) was linearized and transformed to replace the endogenous *MET4* gene.

Microscopy and image analysis

For imaging of Met4-GFP and Met32-mCherry, yeast cells were grown in SCD + 10X Met liquid media at 30°C to OD660 ~ 0.2, washed, and then transferred onto 5 ml of SCD - Met media for 2 hrs for induction. Afterwards the cells were put on a SCD-Met agarose pad and put under the confocal microscope for imaging. Confocal microscopy was performed with Zeiss LSM880 scanning laser confocal (Penn State University Huck Life Sciences Institute) with 488 nm Argon laser (541 nm detection) and 594 nm HeNe laser (648 nm detection) using 63x/1.4 plan apochromat objective. For single cell fluorescent images, the images were taken with Airyscan detector and deconvoluted with Zen Black software.

The individual nuclei from single cells were segmented to extract the nuclear boundaries and pixel fluorescent information of single nuclei with NucleiSplicer (Matlab) and subsequent nuclei information was analyzed with NucleiAnalyzer (Matlab). Variance and standard deviation were calculated from the nuclear pixels and plotted with OriginPro (**Table S2**). The mean GFP intensities were calculated by subtracting the measure GFP intensities by the mean background intensities of unlabeled cells. Coefficient of Variation (CV) values were calculated by dividing the Std. Dev by the mean GFP values for individual cells. The Fano number was calculated by dividing the Variance by the mean GFP. To normalize the Fano number, the mean Fano number of Free GFP was set to a value of 1 and the rest of the conditions were divided by the mean GFP Fano number (**Table S2**).

Single cell images of yeast cells expressing tetR-mCherry and Met4-GFP were imaged with 10 z-stacks (0.4 μ m). The z-stack images of mCherry and GFP channels were analyzed by DotTracker (MATLAB). For individual cells, the program detects the z-stack with the highest mCherry dot intensity and extracts the GFP image from the same z position. These images were aligned with the mCherry dot at the center. The mean intensity projection of the GFP channel was analyzed with ImageJ.

To measure *S.kud* Met3pr-GFP expression, yeast cells were grown in SCD+10X Met liquid media at 30°C to OD₆₆₀ ~ 0.2, washed, and then transferred onto a SCD-Met agarose pad for induction. After 6 hrs, the agarose pad was put under a Leica DMI6000 B fluorescent microscope for imaging. The GFP fluorescent intensity within each cell boundary was quantified using Celltracker4 (MATLAB) (Zou and Bai 2019 [DOI](#)).

Met4 and Met32 purification

Constructs of Met32-mCherry, and Met4-MBP were inserted into pET28b(+) backbones with an N-terminal 6x-His tag. Met4 and Met32 genes were cloned from budding yeast genomic DNA. MBP-tag was cloned from Addgene CAT#98651 and fusion constructs were assembled using 2xHiFi mastermix (NEB #E2621L) and transformed into DH5a cells (NEB C2987I). A single cysteine sequence was added to the linker between Met4 and MBP to facilitate maleimide conjugation of fluorophores for imaging. For protein expression, all plasmids were transformed into BL21(DE3)-Sigma32 cells (courtesy of Xin Zhang) and induced according to reported protocol for the Sigma32 cell line (Zhang et al. 2014 [DOI](#)). Bacterial protein expression was carried out with cells grown in LB supplemented with 50 mg/L kanamycin and 100 mg/L ampicillin at 37°C to OD₆₀₀=0.3, induced with 2 g/L arabinose (TCI A0515). Cells were then grown at 30°C until OD₆₀₀=0.6, induced with 1% isopropyl b-D-1-thiogalactopyranoside (IPTG, Research Products International I56000) and grown overnight at 20°C. Cells were resuspended in wash buffer supplemented with 1 mM phenylmethylsulfonyl fluoride (PMSF), and frozen before lysis.

For protein purification, bacterial cells were lysed with sonication and treated with DNaseI and RNaseA according to published protocols for 15 mins (Carrillo et al. 2012 [DOI](#)). Met32-mCherry was purified using Ni-NTA chromatography (Thermo Scientific HisPur resin #88222) with proteins buffers A (20 mM Tris pH 8.0, 400 mM NaCl, 5 mM imidazole) and B (20 mM Tris pH 8.0, 400 mM NaCl, 250 mM imidazole) with a stepwise gradient of buffer B. Protein was buffer exchanged into buffer C (20 mM HEPES pH 7.5, 400 mM NaCl, 10% glycerol, 1 mM DTT) and concentrated to at least 80 mM using Amicon Pro centrifugal filters 10,000 NMWL (Milipore #ACS501024). The Met4 construct was purified using Ni-NTA chromatography with proteins buffers A and B with a stepwise gradient of buffer B. Ni-NTA elution was immediately loaded onto an amylose column (NEB E8021S) and purified using a stepwise gradient of buffers A and D (20 mM Tris pH 8.0, 400 mM NaCl, 10 mM maltose). Protein was labeled by incubating with 0.8 molar equivalents of Alexa Flour 488 maleimide (Invitrogen A10254). Proteins were buffer exchanged into buffer C and concentrated to less than 50 mM (to avoid aggregation).

Western blot analysis

Met4-MBP samples were run on a 4-20% SDS-PAGE gel (BioRad #4561094). Gels were wet transferred to a PVDF membrane (BioRad #1620177) at 80V for 3 hrs in Tris-Glycine buffer. Transfer was confirmed by Ponceau staining (Sigma P7170-1L). Membrane was blocked with 5% non-fat milk, washed with TBS with 1% Tween 20 (VWR 0777). Membrane was probed with anti-MBP antibody (Thermo Fisher MA527544). Membrane was stripped using PVDF stripping solution (DOT Scientific 65000-500) according to manufacturer instructions and reprobed with anti-6x-His primary antibody (Thermo Fisher MA1-21315) and conducted identically to the first probing. All incubation steps proceeded overnight. Blots were developed using Pierce ECL substrate (Thermo Scientific 32209) with a 2 mins exposure time.

Droplet formation

Proteins were diluted from 400 mM NaCl storage buffer so that the final concentration of the buffer was 150 mM NaCl, 20 mM HEPES pH 7.4, 2% glycerol. Samples were plated on a microscope slide with a hydrophobic spacer (Invitrogen #S24735) between the slide and untreated coverslip. Droplets were allowed to settle for 5 mins prior to imaging. All droplet formation and imaging was conducted at room temperature.

Fluorescent Recovery after Photobleaching (FRAP)

Bleaching was conducted after three frames using the 594 nm laser at full power for 45 iterations. Fluorescence intensity was monitored every 3 seconds for a total of 270 seconds per experiment. Fluorescence intensity levels were extracted using Zeiss Zen Black software. Fluorescence intensity was normalized to percent of pre-bleach intensity then averaged. Error bars show $\pm$ SD of the normalized fluorescence intensity. FRAP experiments were conducted at 20 μ M Met32-mCherry. Experiments were replicated with at least three preparations of each protein. Snapshots of the bleaching process were processed using FIJI.

Chromatin Immunoprecipitation (ChIP)

Yeast strains were grown to OD₆₆₀ of 0.4 in 50 mL of SCD-met medium and crosslinked with 1.39 mL of 37% formaldehyde for 20 mins at room temperature. Crosslinking was quenched by adding 2.7 mL of 2.5 M glycine and incubation for 5 mins. Crosslinked cells were centrifuged at $1882 \times g$ for 3 mins at 4°C, and then washed twice with cold 1x Tris-buffer saline (TBS). Cell pellet was then resuspended in 250 μ L of fresh FSPP buffer (50 mM HEPES-KOH, pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 1% protease inhibitor cocktail, and 1 mM phenylmethylsulfonyl fluoride (PMSF) and vortexed with $\sim$ 300 μ L glass beads for 2 cycles of 20 mins with an intervening 10 mins in a 4°C cold room. Another 250 μ L of fresh FSPP was added to each sample, and the cap and bottom of tubes were punched with a hot needle so that cell lysate could be collected by centrifuging at $836 \times g$ for 5 mins at 4°C. FSPP (0.5 mL) was added to the cell lysate to resuspend the pellet. The cell lysate was then sonicated using a 30 s on 30 s off cycle at 4°C for 7 cycles. Sonicated cell lysate was transferred to 1.5 mL tube and centrifuged at $17,136 \times g$ for 20 mins at 4°C. Two hundred microliters of supernatant was saved as input. Another 400 μ L of supernatant was mixed with 600 μ L of FSPP buffer and 20 μ L of pre-blocked Magnetic IgG beads and TAP antibody (Thermo CAB1001) or Rpb3 antibody (Biolegend 920204) for 10–12 hrs at 4°C. Chromatin was incubated with the beads overnight at 4°C, and then washed sequentially at 4°C with 1xFA-lysis buffer (50 mM HEPES-KOH, pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate), 1xFA-lysis buffer containing 150 mM NaCl, 1xFA-lysis buffer containing 500 mM NaCl, LiCl buffer (0.25 M LiCl, 1% NP-40, 1% sodium deoxycholate, 1 mM EDTA, and 10 mM Tris-HCl pH 8.0), and lastly TE buffer, pH 8.0. Chromatin was then eluted with 400 μ L ChIP elution buffer (50 mM NaCl, 50 mM Tris-HCl, pH 8.0, 10 mM EDTA, 1% SDS). Samples were mixed by rotation at 30°C for 30 mins. Beads were pelleted by centrifugation at $17,136 \times g$ and discarded. Five microliters of 20 mg/mL Proteinase K was added to supernatant; 180 μ L of 1xFA-lysis buffer, 20 μ L of 10% SDS and 5 μ L of 20 mg/mL Proteinase K were added into the input samples. ChIP and input samples were then incubated overnight at 65°C to reverse crosslinking. DNA was extracted by phenol:chloroform:isoamyl alcohol (25:24:1) followed by ethanol precipitation. Input samples were then treated with 200 μ g RNase A and precipitated with ethanol after adding 20 μ g glycogen. Enrichments of target TFs were quantified by quantitative PCR (qPCR). All ChIP experiments were done in biological duplicates.

RNA-seq

Cells were grown in 5 ml of SCD +10x met overnight. 5 ml of fresh SCD +10x was inoculated with 5 μ L of saturated culture until OD₆₆₀ $\sim$ 0.2. The culture is collected by centrifugation and washed 3x with sterile H₂O. The collected culture is then grown in SCD -met media for 2 hrs to reach OD₆₆₀ $\sim$ 0.4 and centrifuged at 300g for 4 mins at RT. The pellet was washed twice with sterile H₂O and resuspended in 250 μ L of RNA lysis buffer (10mM Tris-HCl pH8.5, 5mM EDTA, 2% SDS, 2% stock 2-mercaptoethanol) and placed on a heat block at 85°C for 20 mins mixing every 2 mins. The mixture was centrifuged at 12,000g for 5 mins and the supernatant was transferred to a new tube. The supernatant was then mixed with 1 ml of trizol and heated at 65°C for 20 mins with mixing in between. Standard trizol/chloroform RNA extraction protocol was followed. The extracted RNA was analyzed with TapeStation to check their integrity. RNA with RIN (>7.5) was used for

downstream analysis. Messenger RNA was captured using RNA purification beads (NEB). The eluted mRNA was fragmented and denatured for first and second strand synthesis for conversion into cDNA. The sequencing libraries were constructed using conventional protocols (NEB).

Methyltransferase Targeting-based chromosome Architecture Capture (MTAC)

Yeast cells were grown in 50 ml SCD +10x Met media at OD₆₆₀ 0.3. The cells were centrifuged and washed 3 times with autoclaved H₂O before transfer to 50 ml of SCD-Met and induced with 2 nM β -estradiol for 2 hrs. After induction the cells were centrifuged at 4000g for 5 mins and collected in 800 μ l of TE and 200 μ l lysis buffer (0.5 M EDTA pH 8.0, 1 M Tris pH 8.0, 10% SDS) and incubated at 65 °C for 30 mins. The cell lysate (1ml) was then sonicated using a 30 s on 30 s off cycle at 4°C for 11 cycles. The sonicated lysate was centrifuged at 12,000g at 4 °C for 20 mins and the supernatant was transferred to a new tube. The supernatant was mixed with 500 μ l for Phenol:Chloroform:Isoamyl Alcohol (25:24:1) nucleic acid extraction. The methylated genomic DNA (5 μ g) was incubated with 2 μ g of anti-methylcytosine antibody in 1X IP buffer on a rotating platform 4°C for 8-12 hrs. Input DNA (1%, 50 ng) was set aside for normalization. The lysate/antibody complex was incubated with protein A/G agarose beads for 2 hrs and washed 3x with 1x IP buffer. The beads were resuspended with 210 μ l digestion buffer and incubated on a rotating platform at 55 °C for 2 hrs. The eluted DNA was precipitated in 1 ml of ethanol and 3 μ l glycogen (20mg/ml) and eluted in 60 μ l of TE.

For strains with Met4 Auxin-induced degradation, we tagged the endogenous Met4 protein with V5-AID (Met4-V5-AID) and introduced OsTIR1 to the cells. Depletion of the Smc2 was achieved by adding auxin (Sigma, I2886) to a final concentration of 500 μ M. Strains were incubated with auxin for 30 mins prior to MTAC experiment. Depletion was confirmed by western blot.

Data availability

All sequencing data in this study can be accessed on GEO series GSE252386, which include 1) GSE252384: ChIP-seq raw data (fastq), bed files, and bigWig files, 2) GSE252385: RNA-seq raw data (fastq), counts data (featureCounts), and DESeq2 tables, and 3) GSE252985: MTAC raw data (fastq), bigWig files, counts data (featureCounts) and DESeq2 tables.

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Author Contributions

L.B., J.L., L.S. and Y.L. designed the experiments; J.L. performed most of the experiments and data analysis with help from S.B.; L.S. and Y.L. carried out most of the experiments in **Fig. 2** [↗](#) and **Fig. 4** [↗](#), respectively; F.Z. contributed to the image data analysis; X.Z. helped with protein purification; J.L. and L.B. wrote the manuscript.

Conflict of Interest Statement

The authors declare no conflict of interest.

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Editors

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Reviewer #1 (Public Review):

Summary:

In this study, James Lee, Lu Bai, and colleagues use a multifaceted approach to investigate the relationship between transcription factor condensate formation, transcription, and 3D gene clustering of the MET regulon in the model organism *S. cerevisiae*. This study represents a second clear example of inducible transcriptional condensates in budding yeast, as most evidence for transcriptional condensates arises from studies of mammalian systems. In addition, this study links the genomic location of transcriptional condensates to the potency of transcription of a reporter gene regulated by the master transcription factor contained in the condensate. The strength of evidence supporting these two conclusions is strong. Less strong is evidence supporting the claim that Met4-containing condensates mediate the clustering of genes in the MET regulon.

Strengths:

The manuscript is for the most part clearly written, with the overriding model and specific hypothesis being tested clearly explained. Figure legends are particularly well written. An additional strength of the manuscript is that most of the main conclusions are supported by the data. This includes the propensity of Met4 and Met32 to form puncta-like structures under inducing conditions, formation of Met32-containing LLPS-like droplets in vitro (within which Met4 can colocalize), colocalization of Met4-GFP with Met4-target genes under inducing conditions, enhanced transcription of a Met3pr-GFP reporter when targeted within 1.5 - 5 kb of select Met4 target genes, and most impressively, evidence that several MET genes appear to reposition under transcriptionally inducing conditions. The latter is based on a recently reported novel in vivo methylation assay, MTAC, developed by the Bai lab.

Weaknesses:

My principal concern is that the authors fail to show convincing evidence for a key conclusion, highlighted in the title, that nuclear condensates per se drive MET gene clustering. Figure 4E demonstrates that Met4 molecules, not condensates per se, are necessary for fostering distant cis and trans interactions between MET6 and three other Met4 targets under -met inducing conditions. In addition, the paper would be strengthened by discussing a recent study conducted in yeast that comes to many of the same conclusions reported here, including the role of inducible TF condensates in driving 3D genome reorganization (Chowdhary et al, Mol. Cell 2022).

Other concerns:

(1) A central premise of the study is that the inducible formation of condensates underpins the induction of MET gene transcription and MET gene clustering. Yet, Figure 1 suggests (and the authors acknowledge) that puncta-like Met4-containing structures pre-exist in the nuclei of non-induced cells. Thus, the transcription and gene reorganization observed is due to a

relatively modest increase in condensate-like structures. Are we dealing with two different types of Met4 condensates? (For example, different combinations of Met4 with its partners; Mediator- or Pol II-lacking vs. Mediator- or Pol II-containing; etc.?) At the very least, a comment to this effect is necessary.

(2) Using an in vitro assay, the authors demonstrate that Met4 colocalizes with Met32 LLPS droplets (Figure 2F). Is the same true in vivo - that is, is Met32 required for Met4 condensation? This could be readily tested using auxin-induced degradation of Met32. Along similar lines, the claim that Met32 is required for MET gene clustering (line 250) requires auxin-induced degradation of this protein.

(3) The authors use a single time point during -met induction (2 h) to evaluate TF clustering, transcription (mRNA abundance), and 3D restructuring. It would be informative to perform a kinetic analysis since such an analysis could reveal whether TF clustering precedes transcriptional induction or MET gene repositioning. Do the latter two phenomena occur concurrently or does one precede the other?

(4) Based on the MTAC assay, MET13 does not appear to engage in trans interactions with other Met4 targets, whereas MET6 does (Figures 4C and 4E). Does this difference stem from the greater occupancy of Met4 at MET6 vs. MET13, greater association of another Met co-factor with the chromatin of MET6 vs. MET13, or something else?

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Reviewer #2 (Public Review):

Summary:

This manuscript combines live yeast cell imaging and other genomic approaches to study how transcription factor (TF) condensates might help organize and enhance the transcription of the target genes in the methionine starvation response pathway. The authors show that the TFs in this response can form phase-separated condensates through their intrinsically disordered regions (IDRs), and mediate the spatial clustering of the related endogenous genes as well as reporter inserted near the endogenous target loci.

Strengths:

This work uses rigorous experimental approaches, such as imaging of endogenously labeled TFs, determining expression and clustering of endogenous target genes, and reporter integration near the endogenous target loci. The importance of TFs is shown by rapid degradation. Single-cell data are combined with genomic sequencing-based assays. Control loci engineered in the same way are usually included. Some of these controls are very helpful in showing the pathway-specific effect of the TF condensates in enhancing transcription.

Weaknesses:

Perhaps the biggest weakness of this work is that the role of IDR and phase separation in mediating the target gene clustering is unclear. This is an important question. TF IDRs may have many functions including mediating phase separation and binding to other transcriptional molecules (not limited to proteins and may even include RNAs). The effect of IDR deletion on reduced Fano number in cells could come from reduced binding with other molecules. This should be tested on phase separation of the purified protein after IDR deletion. Also, the authors have not shown IDR deletion affects the clustering of the target genes, so IDR deletion may affect the binding of other molecules (not the general transcription machinery) that are specifically important for target gene transcription. If the self-association of the IDR is the main driving force of the clustering and target gene

transcription enhancement, can one replace this IDR with totally unrelated IDRs that have been shown to mediate phase separation in non-transcription systems and still see the gene clustering and transcription enhancement effects? This work has all the setup to test this hypothesis.

The Met4 protein was tagged with MBP but Met 32 was not. MBP tag is well known to enhance protein solubility and prevent phase separation. This made the comparison of their in vitro phase behavior very different and led the authors to think that maybe Met32 is the scaffold in the co-condensates. If MBP was necessary to increase yield and solubility during expression and purification, it should be cleaved (a protease cleavage site should be engineered) to allow phase separation in vitro.

Are ATG36 and LDS2 also supposed to be induced by -met? This should be explained clearly. The signals are high at -met.

Figure 6B, the Met4-GFP seems to form condensates at all three loci without a very obvious difference, though 6C shows a difference. 6C is from only one picture each. The authors should probably quantify the signals from a large number of randomly selected pictures (cells) and do statistics.

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Reviewer #3 (Public Review):

Summary:

In this study, the authors probe the connections between clustering of the Met4/32 transcription factors (TFs), clustering of their regulatory targets, and transcriptional regulation. While there is an increasing number of studies on TF clustering in vitro and in vivo, there is an important need to probe whether clustering plays a functional role in gene expression. Another important question is whether TF clustering leads to the clustering of relevant gene targets in vivo. Here the authors provide several lines of evidence to make a compelling case that Met4/32 and their target genes cluster and that this leads to an increase in transcription of these genes in the induced state. First, they found that, in the induced state, Met4/32 forms co-localized puncta in vivo. This is supported by in vitro studies showing that these TFs can form condensates in vitro with Med32 being the driver of these condensates. They found that two target genes, MET6 and MET13 have a higher probability of being co-localized with Met4 puncta compared with non-target loci. Using a targeted DNA methylation assay, they found that MET13 and MET6 show Met4-dependent long-range interactions with other Met4-regulated loci, consistent with the clustering of at least some target genes under induced conditions. Finally, by inserting a Met4-regulated reporter gene at variable distances from MET6, they provide evidence that insertion near this gene is a modest hotspot for activity.

Weaknesses:

(1) Please provide more information on the assay for puncta formation (Figure 1). It's unclear to me from the description provided how this assay was able to quantitate the number of puncta in cells.

1. How does the number of puncta in cells correspond with the number of Met-regulated genes? What are the implications of this calculation?

2. A control for chromosomal insertion of the Met-regulated reporter was a GAL4 promoter derivative reporter. However, this control promoter seems 5-10 fold more active than the Met-regulated promoter (Figure 6). It's possible that the high activity from the control promoter overcomes some other limiting step such that chromosomal location isn't important. It would be ideal if the authors used a promoter with comparable activity to the Met-reporter as a control.

(4) It seems like transcription from a very large number of genes is altered in the Met4 IDR mutant (Figure 7F). Why is this and could this variability affect the conclusions from this experiment?

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