

Strategies to regulate transcription factor–mediated gene positioning and interchromosomal clustering at the nuclear periphery

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In budding yeast, targeting of active genes to the nuclear pore complex (NPC) and interchromosomal clustering is mediated by transcription factor (TF) binding sites in the gene promoters. For example, the binding sites for the TFs Put3, Ste12, and Gcn4 are necessary and sufficient to promote positioning at the nuclear periphery and interchromosomal clustering. However, in all three cases, gene positioning and interchromosomal clustering are regulated. Under uninducing conditions, local recruitment of the Rpd3(L) histone deacetylase by transcriptional repressors blocks Put3 DNA binding. This is a general function of yeast repressors: 16 of 21 repressors blocked Put3-mediated subnuclear positioning; 11 of these required Rpd3. In contrast, Ste12-mediated gene positioning is regulated independently of DNA binding by mitogen-activated protein kinase phosphorylation of the Dig2 inhibitor, and Gcn4-dependent targeting is up-regulated by increasing Gcn4 protein levels. These different regulatory strategies provide either qualitative switch-like control or quantitative control of gene positioning over different time scales.

Introduction

Within the nucleus of eukaryotic cells, genomes are spatially organized. Chromosomes loop, fold, and interact with subnuclear structures, occupying distinct “territories” (Cremer et al., 2006). Individual genes can assume different subnuclear positions, depending on their expression state (Pombo and Dillon, 2015). For example, in metazoan cells, developmentally regulated genes often move away from the nuclear lamina upon induction (Peric-Hupkes et al., 2010). Likewise, genes can reposition within chromosome territories and colocalize with RNA polymerase II foci called transcription factories (Brown et al., 2006; Xu and Cook, 2008; Schoenfelder et al., 2010). The spatial arrangement of the genome thus has the potential to create functionally distinct subdomains and contribute to transcriptional regulation.

The interaction of genes with stable nuclear structures can affect their regulation. Large transcriptionally repressed lamin-associated domains localize at the nuclear periphery and punctuate metazoan genomes (Guelen et al., 2008; Luperchio et al., 2014). In contrast, many active genes interact with nuclear pore proteins in yeast, flies, worms, and mammalian cells (Brickner and Walter, 2004; Casolari et al., 2004; Brown et al., 2008; Kalverda et al., 2010; Rohner et al., 2013). In budding

yeast, these interactions occur at the nuclear pore complex (NPC), whereas in metazoan cells, the interactions occur both at the NPC and with soluble nuclear pore proteins in the nucleoplasm (Ahmed et al., 2010; Capelson et al., 2010; Kalverda et al., 2010). In both yeast and metazoan cells, interaction with nuclear pore proteins correlates with transcription (Brickner and Walter, 2004; Taddei et al., 2006; Brickner et al., 2007; Brown et al., 2008; Ahmed et al., 2010; Capelson et al., 2010; Kalverda et al., 2010; Liang and Hetzer, 2011). Thus, the interaction of genes with distinct compartments at the nuclear periphery can lead to opposite transcriptional outcomes.

Coregulated regions of the genome often cluster together (Pombo et al., 2000; Brown et al., 2006; Schoenfelder et al., 2010). For example, transcriptionally silent subtelomeric regions in yeast (Aparicio et al., 1991), tRNA genes (Thompson et al., 2003), and Klf1-regulated genes all cluster (Schoenfelder et al., 2010). The spatial proximity of coregulated genes through interchromosomal clustering may create distinct subnuclear environments that affect gene regulation. Alternatively, changes in chromatin state and expression can lead to the creation of subnuclear domains (Meister et al., 2011).

As a model for these phenomena, we have studied the spatial repositioning of inducible genes from the nucleoplasm

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Abbreviations used in this paper: BS, binding site; ChIP, chromatin immunoprecipitation; GRS, gene recruitment sequence; NPC, nuclear pore complex; PRE, pheromone response element; TF, transcription factor; UAS_{INO}, inositol-sensitive upstream activating sequence; uORF, upstream open reading frame; URS, upstream repressing sequence.

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to the NPC upon activation in budding yeast. Recruitment to the NPC is controlled by cis-acting transcription factor (TF) binding sites in gene promoters (Ahmed et al., 2010; Light et al., 2010; Brickner et al., 2012). These DNA elements function as DNA zip codes: they are necessary to recruit genes from the nucleoplasm to the nuclear periphery and promote stronger transcription, and they are sufficient to target ectopic sites to the NPC (Ahmed et al., 2010; Light et al., 2010; Brickner et al., 2012). Targeting to the nuclear periphery can also lead to interchromosomal clustering of genes that share zip codes (Brickner et al., 2012). For example, a DNA zip code called gene recruitment sequence I (GRS I) from the promoter of the *INO1* gene (encoding inositol 1-phosphate synthase) interacts with the Put3 TF. Inserting GRS I beside the nucleoplasmic locus *URA3* leads to targeting of *URA3* to the nuclear periphery and clustering of *URA3:GRS I* with the endogenous *INO1* gene (Ahmed et al., 2010; Brickner et al., 2012). Loss of Put3 disrupts both GRS I-mediated targeting and interchromosomal clustering. This suggests that some TFs can promote interaction with the NPC and clustering. (Consistent with this notion, interaction of Nup98 with genes in *Drosophila* is mediated by the MBD-R2 DNA binding protein [Pascual-Garcia et al., 2014]). If so, then genomes could encode their spatial organization through TF binding sites.

Yeast genes such as *INO1*, *GALI*, *HSP104*, and *TSA2* localize in the nucleoplasm before induction and relocalize to the nuclear periphery upon activation (Brickner and Walter, 2004; Casolari et al., 2004; Dieppois et al., 2006; Ahmed et al., 2010). Targeting of these genes requires DNA zip codes in their promoters (Ahmed et al., 2010). However, the molecular mechanisms that regulate zip code activity have not been described. Here we identify three different regulatory strategies used by yeast cells to provide dynamic control of gene positioning and interchromosomal clustering. We define the regulation of TFs representing different families: Put3 (a Zn²⁺-binuclear cluster TF), Ste12 (a helix-turn-helix TF), and Gcn4 (a basic leucine zipper TF). Targeting of the *INO1* promoter to the nuclear periphery by Put3 is regulated through local recruitment of the Rpd3(L) histone deacetylase complex by transcriptional repressors. Many repressors are capable of blocking zip code function through Rpd3-dependent and -independent mechanisms. Ste12-mediated gene positioning is not affected by Rpd3 but is regulated downstream of DNA binding by MAPK phosphorylation of the inhibitor Dig2. Finally, Gcn4-mediated gene positioning is controlled by Gcn4 abundance. Each mechanism provides distinct advantages: repressor regulation leads to a slow switch, whereas MAPK signaling leads to a rapid switch. Changes in TF levels allow a quantitative increase in peripheral localization and interchromosomal clustering over an intermediate time scale.

Results

TF-dependent and stimulus-specific recruitment to the nuclear periphery

Our previous work suggests that targeting of inducible genes to the nuclear periphery is mediated by cis-acting TF binding sites that function as DNA zip codes (Ahmed et al., 2010). To test the generality of this model and to understand the regulation of gene positioning, we asked whether the inducible genes *PRM1* (a cell surface transmembrane protein induced by mating pheromone) and *HIS4* (a multifunctional histidine biosynthetic enzyme

induced by histidine starvation) are also recruited to the nuclear periphery in a TF-dependent fashion. *PRM1* is among a large set of genes previously shown to physically interact with the NPC in the presence of pheromone (Casolari et al., 2005), and *HIS4* was identified as a putative NPC-interacting gene by in silico comparison of TF binding to promoters that interact with the NPC (Fig. S1; Casolari et al., 2004; Venters et al., 2011).

An array of 128 Lac repressor binding sites was integrated downstream of each of these genes as well as the nucleoplasmic gene *URA3*. Into these strains, GFP-tagged Lac repressor (GFP-LacI) and an mCherry ER/nuclear envelope marker were introduced (Egecioglu et al., 2014). Live cells were imaged by confocal microscopy, and the fraction of the cells in which the GFP-LacI focus was unresolvable from the nuclear envelope was scored under uninducing or inducing conditions (Fig. 1 A). For all three genes, the uninducing condition is the same (synthetic complete medium). However, the inducing conditions are distinct: *INO1* (inositol starvation), *PRM1* (mating pheromone stimulation), and *HIS4* (amino acid starvation). In cells grown under uninducing conditions, *URA3*, *INO1*, and *PRM1* colocalized with the nuclear periphery in ~30% of the cells counted (Fig. 1 B), similar to the fraction of the nucleus that is unresolvable from the nuclear envelope by light microscopy (baseline, blue hatched line in Fig. 1 B and throughout; Brickner and Walter, 2004). In contrast, *HIS4*, which is modestly expressed in the presence of histidine (Mueller and Hinnebusch, 1986), colocalized with the nuclear envelope in ~45% of cells, significantly above the baseline (Fig. 1, B and C). Under all inducing conditions, *URA3* remained nucleoplasmic. In contrast, specifically under their respective inducing conditions, peripheral localization of *INO1*, *PRM1*, and *HIS4* increased significantly to ~60% of the cells (Fig. 1, B and C).

We next defined the TFs required for the targeting of *INO1*, *PRM1*, and *HIS4* to the nuclear periphery. *INO1* targeting to the nuclear periphery requires one of two cis-acting DNA elements, GRS I and GRS II (Ahmed et al., 2010). The Put3 TF binds to the GRS I zip code and is necessary for GRS I-dependent positioning but is not required for GRS II-mediated positioning (Brickner et al., 2012). Because the TF Cbf1 binds near the GRS II upon *INO1* induction (Shetty and Lopes, 2010), we tested whether Cbf1 is required for GRS II function. Indeed, whereas loss of Put3 alone (Ahmed et al., 2010; Brickner et al., 2012) or Cbf1 alone (Fig. S2) does not block targeting of *INO1* to the nuclear periphery, strains lacking both Put3 and Cbf1 fail to target *INO1* to the nuclear periphery (Fig. 1 B). This suggests that Cbf1 is required for GRS II-mediated gene positioning.

The positioning of *PRM1* and *HIS4* to the nuclear periphery requires the same TFs that regulate their expression. *PRM1* transcription is controlled by cooperative binding of the Ste12 TF to three pheromone response elements (PREs; 5'-TGAACA-3') in its promoter (Hagen et al., 1991). Loss of Ste12 blocks *PRM1* relocalization to the nuclear periphery in the presence of mating pheromone (Fig. 1 B). *HIS4* expression requires binding of the Gcn4 TF to binding sites (5'-TGACTC-3') in the promoter (Arndt and Fink, 1986), and loss of Gcn4 blocked positioning of *HIS4* to the nuclear periphery (Fig. 1 B). Therefore, the subnuclear positioning of *INO1*, *PRM1*, and *HIS4* requires TF binding to their promoters.

The SAGA histone acetyltransferase complex is required for recruitment of genes such as *GALI-10* and *INO1* to the NPC (Rodríguez-Navarro et al., 2004; Luthra et al., 2007; Ahmed et al., 2010; Strambio-De-Castillia et al., 2010). SAGA is also

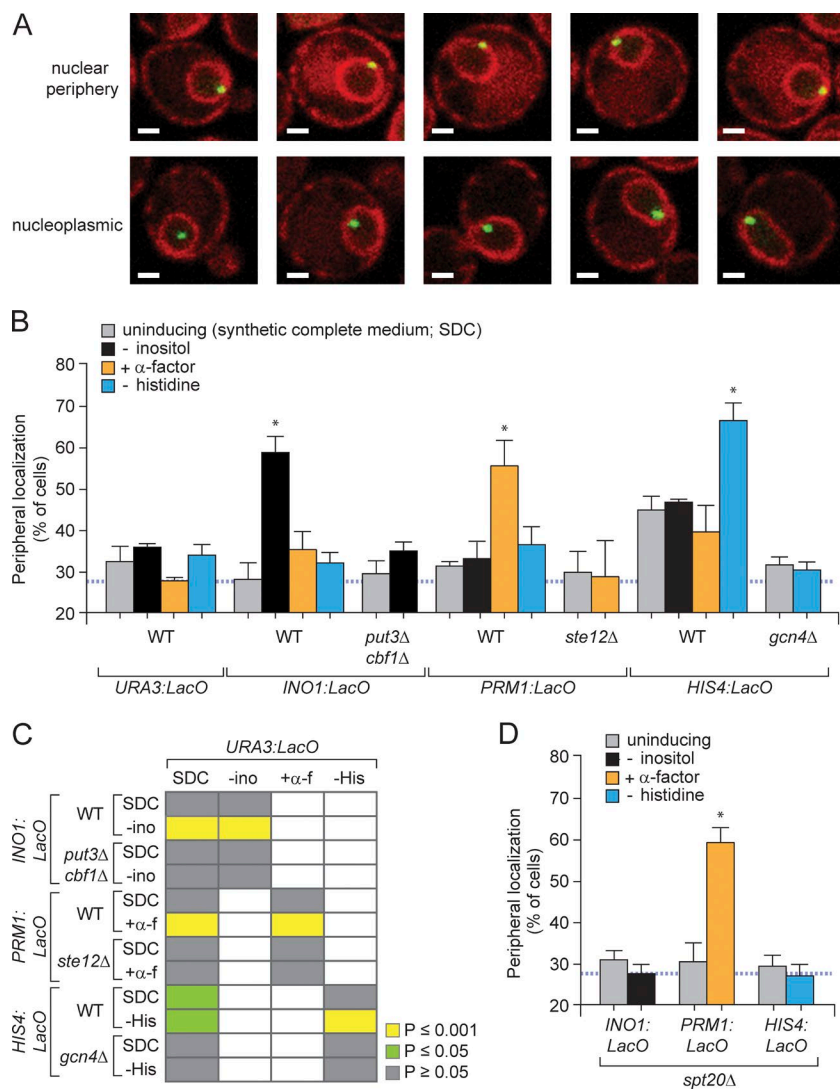


Figure 1. TF-dependent and stimulus-specific recruitment of *INO1*, *PRM1*, and *HIS4* to the nuclear periphery. (A) Confocal micrographs of cells having the Lac operator (LacO) array integrated at *INO1*, expressing GFP-LacI and an mCherry-ER marker (Egecioglu et al., 2014), scored as either peripheral (top) or nucleoplasmic (bottom). Bar, 1 μ m. (B) Wild-type or mutant strains having the LacO array inserted at *URA3*, *INO1*, *PRM1*, or *HIS4* were imaged by confocal microscopy under uninducing conditions (synthetic complete medium; gray bars), after inositol starvation overnight (black bars), after α -factor stimulation for 15–25 min (orange bars), or after histidine starvation for 45–75 min (cyan blue bars). (C) P-values (Fisher exact test) comparing the peripheral localization of the *URA3* locus with *INO1*, *PRM1*, or *HIS4* in the indicated conditions and strains from B. -ino, inositol starvation; + α -f, α -factor stimulation; -his, histidine starvation. (D) Peripheral localization of *INO1*, *PRM1*, and *HIS4* in *spt20Δ* strain. *, $P \leq 0.05$ (Fisher exact test) between SDC and inducing condition for specific strain. Mean and SEM from three or more biological replicates (30–60 cells per replicate).

required for the interaction of extrachromosomal circles with the NPC in yeast (Denoth-Lippuner et al., 2014). To test whether SAGA is required for recruitment of *HIS4* and *PRM1* to the nuclear periphery, we deleted *SPT20* (which is required for the structural integrity of SAGA; Roberts and Winston, 1997). Loss of Spt20 blocked recruitment of both *INO1* and *HIS4* to the nuclear periphery (Fig. 1 D). However, *PRM1* repositioning to the nuclear periphery was independent of Spt20 (Fig. 1 D). Thus, SAGA is necessary for recruitment of some, but not all, genes to the nuclear periphery.

TF binding sites function as DNA zip codes

We next asked whether the TFs that are required for peripheral localization mediate peripheral localization. Each binding site (BS) was inserted beside *URA3* to test its sufficiency to promote peripheral localization (Ahmed et al., 2010). As we have shown previously, both GRS I and GRS II are sufficient to reposition *URA3* to the nuclear periphery (Fig. 2 A). Put3 was required for GRS I-mediated gene positioning, Cbf1 was required for GRS II-mediated gene positioning, and SAGA was required for both (Fig. 2 A).

Insertion of three PREs (3 \times PRE) and the Gcn4 BS at *URA3* was also sufficient to promote peripheral localization, and these zip codes required Ste12 and Gcn4, respectively (Fig. 2 B).

Furthermore, the positioning to the nuclear periphery mediated by 3 \times PRE was SAGA independent, whereas the positioning mediated by the Gcn4 BS was SAGA dependent (Fig. 2 B).

Although repositioning of *INO1*, *PRM1*, and *HIS4* to the nuclear periphery was mediated by TFs, the regulation of these three TF-dependent repositioning events was different. Under uninducing conditions, recruitment of *INO1* and *PRM1* to the nuclear periphery is near baseline levels. In contrast, *HIS4* localization to the periphery under uninducing conditions is significantly higher than *URA3* recruitment or that in the *gcn4Δ* strain. This suggests that, under uninducing conditions, *HIS4* is targeted to the nuclear periphery by Gcn4 to an intermediate level and that it increases to a maximal level under inducing conditions. Also, whereas the zip codes from the *INO1* promoter functioned constitutively at the ectopic site, 3 \times PRE and Gcn4BS were both regulated at the ectopic site. This suggested that the regulation of GRS I and GRS II function in the *INO1* promoter is dependent on its local context and that the regulation of Ste12- and Gcn4-mediated positioning is independent of local context.

The Rpd3(L) histone deacetylase regulates targeting of *INO1* to the nuclear periphery

To understand how GRS I and GRS II are regulated by their promoter context, we identified the cis- and trans-acting factors

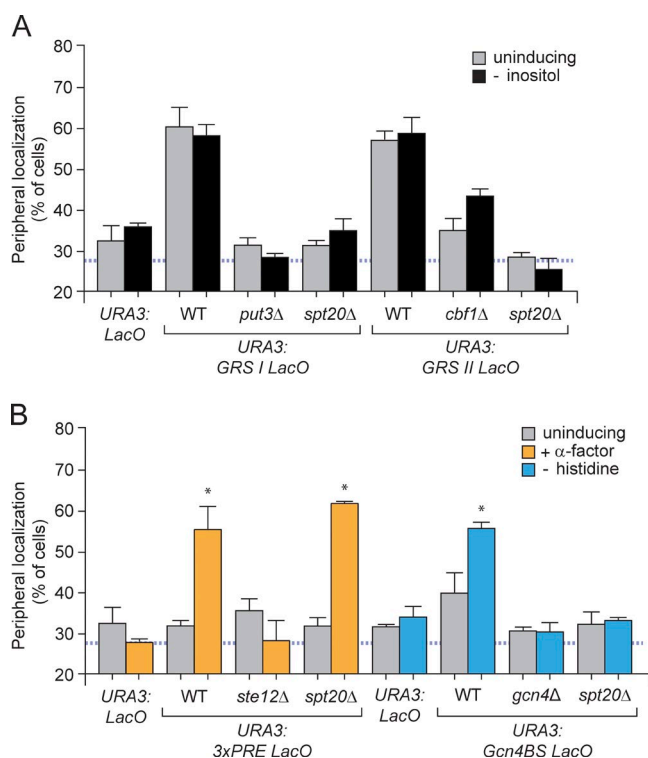


Figure 2. TF BSs function as DNA zip codes. (A and B) Peripheral localization of the *URA3* locus, \pm indicated DNA BSs, grown under uninducing and inducing conditions. GRS I or GRS II (A), 3xPREs (B), or Gcn4 BS (B) were inserted at *URA3* in wild-type (WT) and mutant strains. *, $P \leq 0.05$ (Fisher exact test) comparing uninducing and inducing condition. Mean and SEM from three of more biological replicates (30–50 cells per replicate).

that block their function in the context of the *INO1* promoter. Deletion of 100 bp of the *INO1* promoter ($\Delta 4$; Fig. 3 A) led to constitutive localization to the nuclear periphery (Ahmed et al., 2010). This part of the promoter contains two inositol-sensitive upstream activating sequence (UAS_{INO}) elements as well as an upstream repressing sequence (URS) that regulate the transcription of *INO1* (Fig. 3 A; Lopes et al., 1993; Nikoloff and Henry, 1994). Mutations that disrupt these UAS_{INO} elements prevent expression of *INO1*, and mutation of the URS element leads to constitutive expression of *INO1* (Lopes et al., 1993; Bachhawat et al., 1995). Therefore, we hypothesized that the URS element might both repress transcription and prevent peripheral targeting in the presence of inositol. Indeed, mutation of the URS element led to constitutive targeting of *INO1* to the nuclear periphery (Fig. 3 A). However, mutations that disrupted the UAS_{INO} elements also led to constitutive localization at the nuclear periphery (Fig. 3 A). Therefore, *INO1* targeting to the nuclear periphery under uninducing conditions is blocked by a mechanism that requires both the URS element and the UAS_{INO} elements.

Transcriptional repression of *INO1* is mediated by two mutually dependent repressors (Fig. 3 C). In the presence of inositol, the Ume6 repressor binds to the URS (Lopes et al., 1993; Kadosh and Struhl, 1997; Rundlett et al., 1998) and the Opi1 repressor interacts with the Ino2/Ino4 activator bound to the UAS_{INO} (Heyken et al., 2005). Neither mechanism is sufficient, because loss of either Ume6 or Opi1 leads to constitutive, high-level expression of *INO1* (Fig. S3 B). Loss of Ume6, Opi1, Ino2, or Ino4 led to constitutive targeting of *INO1*

to the nuclear periphery (Fig. 3 A). Furthermore, a mutation in Ino2 (L118A) that disrupts binding of Opi1 (Heyken et al., 2005) had the same effect (Fig. 3 A). This is not related to derepression of *INO1* transcription, because strains lacking Ino2 and Ino4 or the UAS_{INO} element show no expression of *INO1* (Graves and Henry, 2000). Furthermore, a strain lacking Isw2, a chromatin-remodeling factor required for *INO1* repression (Shetty and Lopes, 2010), showed normal, regulated peripheral targeting (Fig. 3 A). Therefore, this suggests that recruitment of Opi1 and Ume6 to the *INO1* promoter blocks GRS I and GRS II function. Consistent with this idea, the peripheral targeting of *INO1* in the *opi1Δ* mutant was lost when both GRS I and GRS II were mutated (Fig. 3 B).

Both Ume6 and Opi1 recruit the Sin3/Rpd3 histone deacetylase, which is essential for *INO1* repression (Kadosh and Struhl, 1997; Wagner et al., 2001), so we tested whether this complex prevents repositioning of *INO1* to the nuclear periphery under repressing conditions. Indeed, loss of Sin3 or Rpd3 or a catalytically inactive form of Rpd3 (His188A; Kadosh and Struhl, 1998) led to constitutive targeting of *INO1* (Fig. 3 A). The peripheral targeting of *INO1* in the *rp3Δ* mutant was lost when GRS I and GRS II were mutated (Fig. 3 B). Therefore, local recruitment of Rpd3 deacetylase activity blocks GRS I and GRS II zip code activity.

Rpd3 is the catalytic subunit of two distinct complexes, Rpd3(L) and Rpd3(S), that have distinct protein components, interact with distinct genomic sites, and have distinct effects on gene expression (Keogh et al., 2005). Whereas Rpd3(L) is associated with promoters and functions as a corepressor, Rpd3(S) is recruited cotranscriptionally to gene bodies to prevent cryptic transcriptional initiation (Kadosh and Struhl, 1997; Carrozza et al., 2005). We tested mutants lacking complex-specific subunits. Loss of Rpd3(L) components Pho23 or Sap30 led to unregulated targeting of *INO1* to the nuclear periphery, whereas loss of the Rpd3(S) component Rco1 did not (Fig. 3 A). Thus, Rpd3(L) blocks peripheral targeting of *INO1*.

To test whether Rpd3 affects local histone acetylation and Put3 binding to the GRS I, we performed chromatin immunoprecipitation (ChIP). Acetylation over the *INO1* promoter increased under inositol starvation and was constitutively high in the absence of Rpd3 (Fig. 3 D). Likewise, unlike the wild-type strain, in which Put3 binds to the GRS I only under inducing conditions (Brickner et al., 2012), in the *rp3Δ* mutant, Put3 binding to the *INO1* promoter was constitutive (Fig. 3 E). This suggests that Rpd3 deacetylase activity regulates Put3 binding to GRS I in the *INO1* promoter.

Rpd3(L) regulates interchromosomal clustering of *INO1*

Zip code-mediated targeting to the nuclear periphery also leads to interchromosomal clustering of genes that share the same zip codes (Brickner et al., 2012). For example, upon inositol starvation, the two alleles of *INO1* reposition to the nuclear periphery and cluster (Brickner et al., 2012). This requires the zip codes, the TFs that bind to the zip codes, and nuclear pore proteins (Brickner et al., 2012). In haploid cells, this can be observed by comparing the position of two loci that are targeted to the nuclear periphery by the same zip code. For example, active *INO1* clusters with both *URA3:GRS I* and another GRS I-containing gene, *TSA2* (Brickner et al., 2012).

To test whether Rpd3(L) also regulates clustering of *INO1* alleles in diploid cells, we measured the distribution of distances

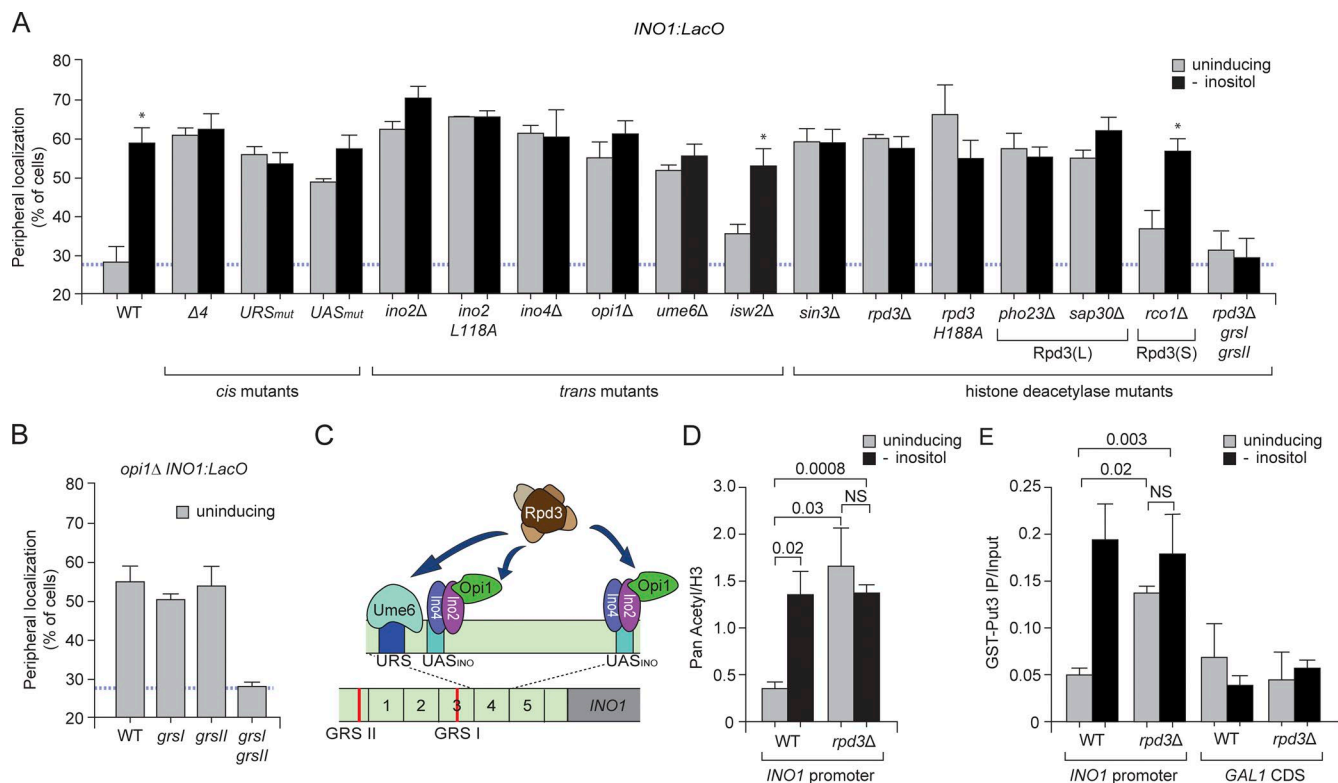


Figure 3. Rpd3(L) histone deacetylase regulates *INO1* gene recruitment to the nuclear periphery. (A and B) Peripheral localization of *INO1*. *, $P \leq 0.05$ (Fisher exact test) between SDC and inducing condition. Mean and SEM from three or more biological replicates (30–50 cells per replicate). (A) Strains having the indicated cis-acting, trans-acting, and histone deacetylase mutants were scored for *INO1* targeting to the nuclear periphery. A schematic of the *INO1* promoter showing the 100-bp “segment 4,” GRS I, GRS II, UAS_{INO}, and URS elements is shown in C. WT, wild-type. (B) Peripheral localization of wild type, *grsI*, *grsII*, or *grsI grsII* mutant *INO1* in the *opi1* Δ mutant strain grown under uninducing conditions. (C) Model for *INO1* transcriptional regulation. Ino2/Ino4 binds the UAS_{INO} elements and recruits Opi1 to the promoter under uninducing conditions. Both Ume6 and Opi1 recruit Rpd3(L). (D) ChIP with anti-pan-acetyl antibody, quantified relative to ChIP with anti-histone H3 antibody from *RPD3* or *rpd3* Δ strains. The recovery of the *INO1* promoter from each IP, normalized to input, was quantified by real-time quantitative PCR (qPCR). NS, not significant. (E) ChIP of Gst-Put3 (Ahmed et al., 2010; Brickner et al., 2012) expressed in either *RPD3* or *rpd3* Δ strains. The recovery of the *INO1* promoter or the *GAL1* coding sequence was quantified relative to input by qPCR. P-values determined by Student's *t* test.

between the alleles of *INO1* in a population of cells grown under both uninducing and inducing conditions (Fig. 4 A). Upon inositol starvation, the distribution of distances between alleles of *INO1* shifts to significantly shorter distances (Brickner et al., 2012; Fig. 4 B; $P = 2 \times 10^{-4}$, Wilcoxon rank sum test). The fraction of cells in which the two alleles are $\leq 0.55 \mu\text{m}$ apart, an alternative metric for clustering (Brickner et al., 2012), was 24% in uninducing conditions and 49% under inositol starvation (Fig. 4 E; $P = 6 \times 10^{-5}$, Fisher exact test).

In contrast to wild-type *INO1*, two alleles with mutated URS elements clustered constitutively, independent of inositol starvation (Fig. 4 C, $P = 0.173$). These alleles were $\leq 0.55 \mu\text{m}$ apart in 45.9% (uninduced) and 52.6% (induced) cells (Fig. 4 E, $P = 0.31$). Likewise, in the *rpd3* H188A mutant strain, the alleles of *INO1* were constitutively clustered, and this was dependent on GRS I and GRS II (Fig. 4, D and E). Thus, Rpd3(L) recruitment regulates both *INO1* positioning and interchromosomal clustering.

Opi1 and Ume6 are sufficient to block GRS I and GRS II function

To test whether Opi1 and Ume6 are sufficient to regulate GRS I- and GRS II-mediated gene positioning, we inserted a LexA BS into the endogenous *INO1* promoter 50 bp from the middle of GRS I and 395 bp from the middle of GRS II (Fig. 5 A).

This experiment used the Ino2 L118A mutant strain, which blocks binding of Opi1, leading to constitutive targeting to the nuclear periphery (Fig. 3 A). LexA-Ume6 or LexA-Opi1 were expressed in these strains, as confirmed by immunoblotting against LexA (Fig. S4 A), and both proteins repressed *INO1* transcription (Fig. S4 B). LexA-Ume6 and LexA-Opi1, but not LexA alone, blocked recruitment of *INO1* to the nuclear periphery (Fig. 5 A). Therefore, Opi1 and Ume6 are sufficient to block GRS I and GRS II function in the context of the *INO1* promoter.

We also reconstituted Opi1- and Ume6-mediated regulation of each zip code separately at the ectopic site by tethering LexA-Ume6 and LexA-Opi1 beside *URA3:GRS I*, *URA3:GRS II*, or *URA3:3×PRE* (Fig. 5 B). Tethering the repressors beside GRS I or GRS II blocked peripheral localization (Fig. 5 B). However, this effect was specific: neither repressor blocked Ste12-mediated targeting (Fig. 5 B). This suggests that Ste12 targeting must be regulated by a different mechanism and that repressors such as Ume6 and Opi1 can regulate peripheral targeting by some, but not all, TFs.

Finally, we asked whether tethered LexA-Opi1 was also sufficient to regulate clustering of *INO1* with *URA3:GRS I* LexA BS. In cells expressing LexA alone, *URA3:GRS I* LexA BS clustered with *INO1* under inositol starvation (Fig. 5, C and D). Expression of LexA-Opi1 disrupted this clustering (Fig. 5, C [$P = 0.002$] and D [$P = 0.003$]). Thus LexA-Opi1 was sufficient

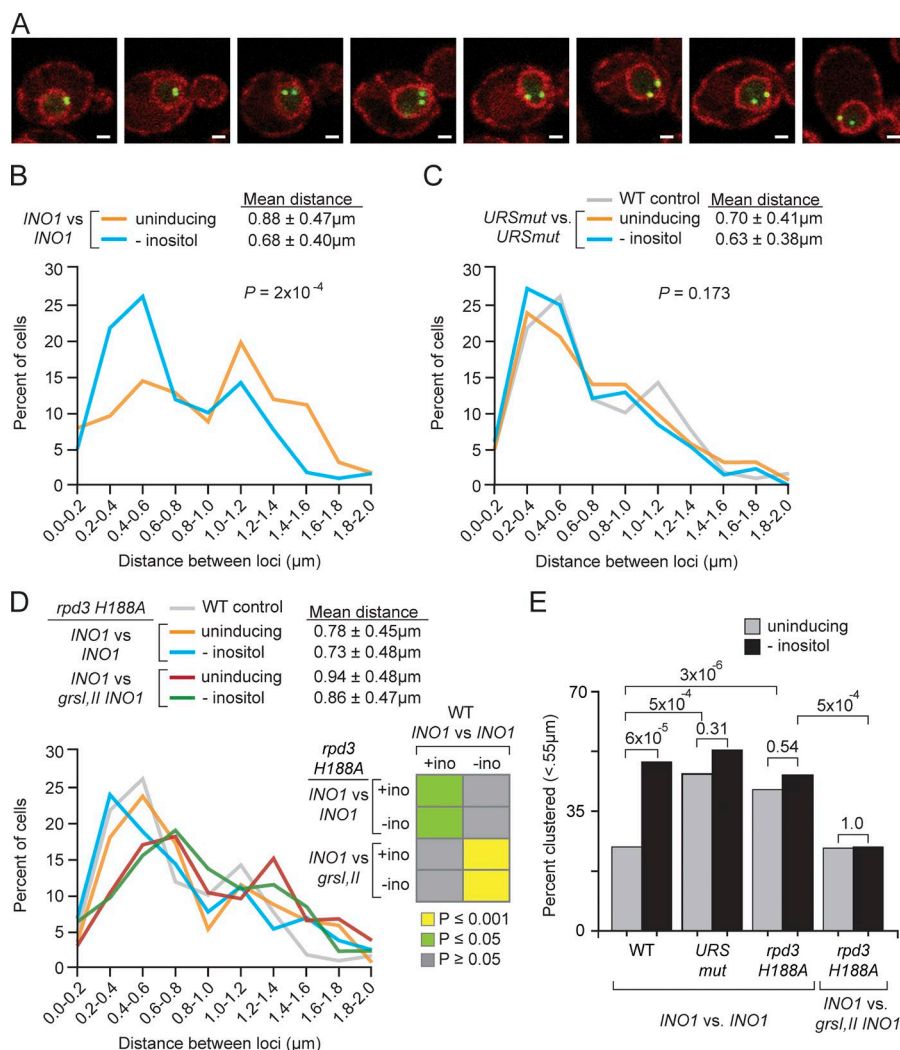


Figure 4. Interchromosomal clustering of *INO1* alleles is regulated by Rpd3(L). (A) Confocal micrographs of diploid cells having LacO arrays at each allele of *INO1*, expressing GFP-LacI and mCherry-ER marker. (B–D) Distances between *INO1* alleles were measured and binned into $0.2\text{-}\mu\text{m}$ bins, and the distribution of distances within the population was compared. In B and C, the p -value is from a Wilcoxon rank sum test comparing the distributions. In panels C and D, the gray line (wild-type [WT] control) is the distribution from B under the inducing condition. (D, inset) P -values from Wilcoxon rank sum test for the indicated comparisons. (E) From the same measurements used to generate the distributions in B–D, the percentage of cells in which the two alleles are within $0.55 \mu\text{m}$ of each other. P -values from Fisher exact test.

to regulate both peripheral localization and interchromosomal clustering mediated by GRS I.

General role for repressors in regulating zip code-mediated targeting to the nuclear periphery

We next asked whether the ability to repress zip code function is a general function of transcriptional repressors by creating LexA fusions to an additional 19 transcriptional repressors. These 24 fusion proteins were tested for expression (immunoblot), ability to repress *INO1* transcription (by tethering to the LexA BS used in Fig. 5 A), and ability to block recruitment of *URA3:GRS I* LexA BS to the nuclear periphery (Fig. S4). Three repressor fusions (Leu3, Rpd3, and Pho23) failed all three of these tests and were excluded (Fig. S4 C). By measuring the peripheral localization of *URA3:GRS I* LexA BS in strains expressing the remaining 21 LexA-repressor fusions, we found that 16 blocked GRS I-mediated targeting to the nuclear periphery in a LexA BS-dependent manner (Fig. 6, A and B). There was no correlation between the ability to block GRS I function and repressor size, expression level, or ability to repress *INO1* transcription (Fig. S4 C). Finally, 11 of the 16 repressors that blocked GRS I function required Rpd3; in the *rpd3 H188A* mutant, GRS I-mediated peripheral localization was restored. The remaining five repressors were Rpd3 independent.

Therefore, many repressors are capable of regulating DNA zip codes, both through Rpd3 recruitment and by other mechanisms.

Regulation of Ste12-mediated gene positioning by MAPK signaling

Ste12 binds to the promoters of genes such as *PRM1* constitutively (Zheng et al., 2010), but transcriptional activation (Roberts et al., 2000) and targeting to the nuclear periphery (Fig. 7 B) occur only in the presence of mating pheromone. The positioning mediated by $3 \times \text{PRE}$ is independent of context (Fig. 2), and tethering Opi1 and Ume6 beside $3 \times \text{PRE}$ failed to block targeting to the periphery (Fig. 3 B). Therefore, the regulation of Ste12 must be through a different mechanism.

In the absence of mating pheromone, Ste12 binds to two inhibitors, Dig1 and Dig2, that independently inhibit Ste12 by different mechanisms (Olson et al., 2000). Stimulation of the MAPK pathway by pheromone leads to phosphorylation of Dig1 and Dig2, causing them to dissociate from Ste12 (Fig. 7 A). Therefore, we tested whether Dig1 and/or Dig2 blocked Ste12-mediated positioning of *PRM1* at the nuclear periphery. Mutants lacking Dig1 showed normal, conditional targeting to the nuclear periphery (Fig. 7 B). However, mutants lacking Dig2 showed constitutive targeting of *PRM1* to the periphery (Fig. 7 B). Targeting of *PRM1* to the periphery in the *dig2Δ* mutant required Ste12 (Fig. 7 B). Therefore, although

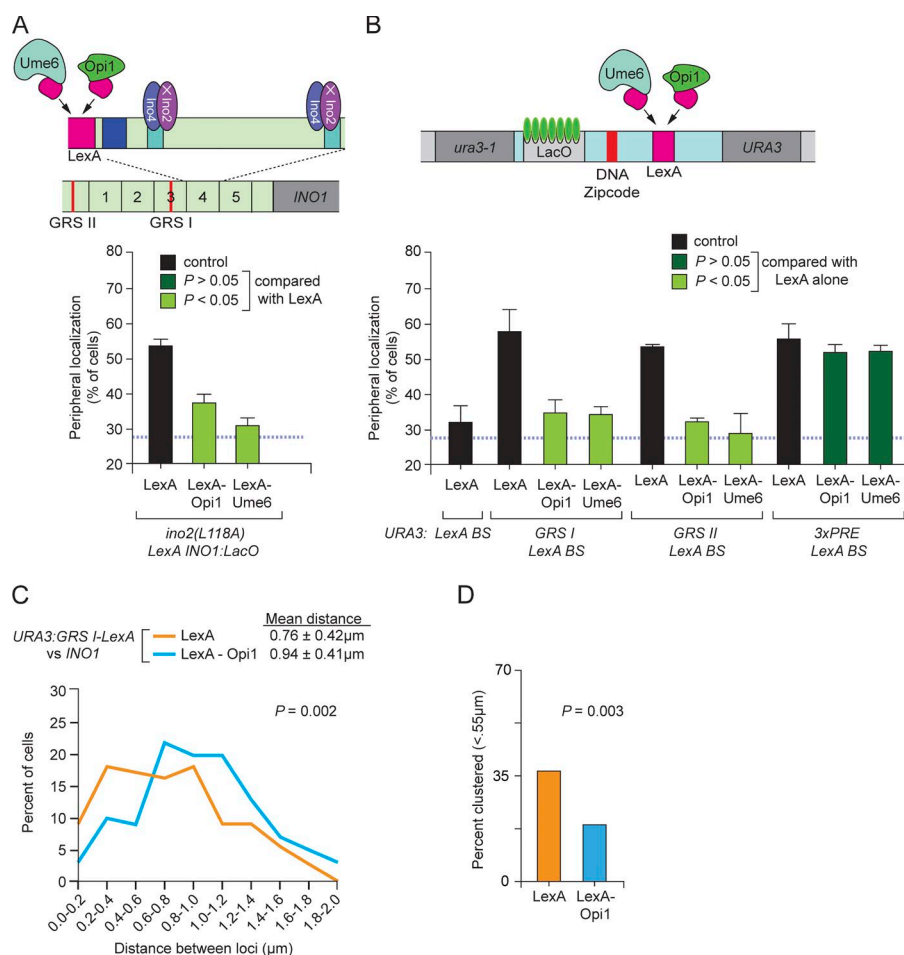


Figure 5. Opi1 and Ume6 are sufficient to block GRS I and GRS II zip code function, but not PRE zip code function. (A, top) Schematic of the *INO1* promoter having a LexA BS in the *ino2 L118A* strain (which blocks the interaction of Ino2 with Opi1; indicated with an X). (bottom) Peripheral localization of *ino2 L118A LexA BS-INO1* in strains expressing LexA, LexA-Opi1, or LexA-Ume6 grown under the uninducing condition. (B, top) Schematic of LacO128-LexA BS ± zip codes integrated at *URA3*. The spacing from the middle of the LexA BS to the middle of the zip codes was GRS I = 68 bp, GRS II = 58 bp, 3xPRE = 42 bp. (bottom) Peripheral localization of *URA3-LexA BS* ± the indicated zip codes in strains expressing LexA, LexA-Opi1, or LexA-Ume6. For A and B, the color indicates the p-value from the Fisher exact test, compared with the control. (C) Distribution of distances between *INO1* and *URA3-GRS I-LexA BS* in strains expressing LexA or LexA-Opi1, grown under inositol starvation. (top) Mean distances and standard deviations. P-value from Wilcoxon rank sum test comparing the distributions. (D) Fraction of cells in which *INO1* and *URA3-GRS I-LexA BS* were $\leq 0.55 \mu\text{m}$ apart (p-values from Fisher exact test).

both Dig1 and Dig2 regulate Ste12-mediated transcription, Dig2 alone blocks Ste12-mediated gene positioning.

We next tested whether MAPK signaling relieves Dig2 repression to allow *PRM1* targeting to the nuclear periphery. A phosphoproteomic study identified Ser34 of Dig2 as a pheromone-stimulated MAPK phosphorylation site (Gruhler et al., 2005). Ser34 was replaced with either an alanine (to block phosphorylation) or an aspartate (to mimic phosphorylation) in the chromosomal *DIG2* gene. The Ser34Ala mutation blocked *PRM1* targeting to the nuclear periphery (Fig. 7 B). Mimicking phosphorylation with the Ser34Asp mutation led to constitutive peripheral localization, like the *dig2Δ* mutant (Fig. 7 B). This suggests that MAPK phosphorylation of Dig2 Ser34 relieves inhibition of Ste12-mediated targeting to the nuclear periphery.

Ste12 is necessary and its BS is sufficient to control spatial positioning. To confirm that Ste12 is responsible for targeting, LexA-Ste12 was tethered to *URA3::LexA BS* (Fig. 7 C). Under uninducing conditions, *URA3::LexA BS* was nucleoplasmic (Fig. 7 C). However, in the presence of mating pheromone or in the *dig2Δ* mutant strain, LexA-Ste12 expression caused *URA3::LexA BS* localization at the nuclear periphery (Fig. 7 C). Thus, Ste12 is sufficient to induce peripheral positioning, and its regulation by Dig2 is independent of DNA binding.

To test whether Ste12-mediated targeting to the nuclear periphery leads to interchromosomal clustering, we created a *MATa* haploid yeast strain having LacO arrays at both *PRM1* and *URA3::3xPRE*. In the absence of mating pheromone, the distances between *PRM1* and *URA3::3xPRE* were broadly

distributed, with 23% of the cells having distances $\leq 0.55 \mu\text{m}$ (Fig. 7 D). In the presence of mating pheromone, *PRM1* and *URA3::3xPRE* clustered together, with a decrease in the mean distance ($P = 0.0017$; Fig. 7 D) and an increase to 43% of the cells with distances $\leq 0.55 \mu\text{m}$ ($P = 0.0006$; Fig. 7 E). Loss of Dig2 led to even higher levels of clustering in the absence of mating pheromone (Fig. 7, D and E). Therefore, Ste12-mediated positioning and interchromosomal clustering are regulated by MAPK signaling through Dig2 phosphorylation.

Increased peripheral gene positioning through regulated TF synthesis

Mutant strains lacking Gcn4 fail to target both *HIS4* (Fig. 1 B) and *URA3::Gcn4 BS* (Fig. 2 C) to the nuclear periphery. Therefore, we hypothesized that Gcn4 abundance, not DNA binding or activity, regulates Gcn4-mediated gene positioning. Several short upstream open reading frames (uORFs) in the 5' end of the *GCN4* mRNA compete with the *GCN4* coding sequence for translation (Mueller and Hinnebusch, 1986; Hinnebusch, 2005) and, in the presence of amino acids, Gcn4 is poorly translated. Amino acid starvation leads to an accumulation of uncharged tRNAs, stimulating the Gcn2 kinase to phosphorylate eIF2 α and leading to a global decrease in translation initiation rates (Cigan et al., 1993). This both reduces the global utilization of amino acids and, by decreasing translation initiation of the *GCN4* uORFs, leads to increased translation of Gcn4 protein.

To test the idea that Gcn4-mediated gene positioning is controlled by Gcn4 protein concentrations, we mutated the

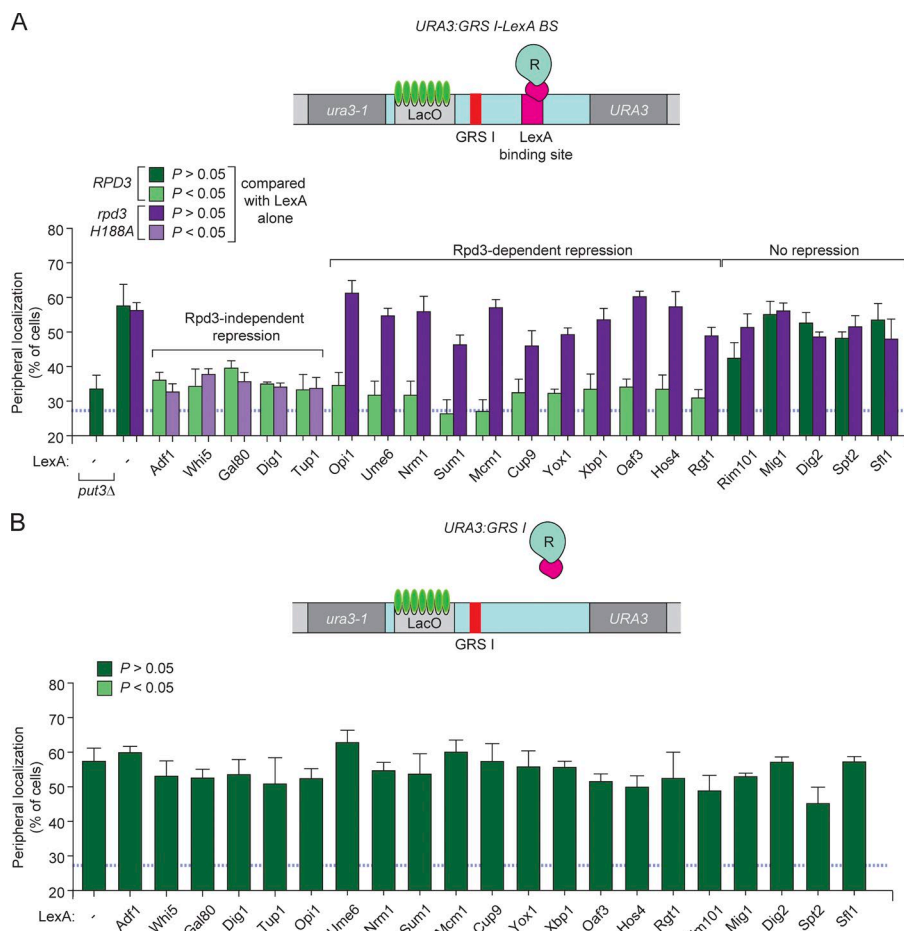


Figure 6. A general role for repressors in regulating zip code-mediated targeting to the nuclear periphery. (A and B) Peripheral localization of URA3-GRS I-LexA BS (A) or URA3-GRS I (B) in strains expressing LexA alone (–) or LexA fused to the indicated full-length repressors (see www.yeastgenome.org for descriptions of all gene names). These LexA fusions were validated as described in Methods and in Fig. S3. LexA fusions were tested for their ability to block GRS I-mediated targeting in both *RPD3* (A and B; light green or forest green) and *rpd3 H188A* (A; lavender or purple) strains. Repressors that resulted in a statistically significant decrease in peripheral localization compared with LexA alone ($P < 0.05$; Fisher exact test) are indicated in light green (*RPD3*) and lavender (*rpd3 H188A*); repressors that did not affect GRS I-mediated peripheral localization are indicated in forest green (*RPD3*) and purple (*rpd3 H188A*) bars. Mean and SEM from three or more biological replicates (30–50 cells per replicate).

initiation codon of the third and fourth uORF in the 5' end of the *GCN4* mRNA in the endogenous *GCN4* gene, which leads to constitutive translation of Gcn4 at levels comparable to those observed during histidine starvation (Mueller and Hinnebusch, 1986). These uORF mutations led to constitutive, high-level localization of both *HIS4* (~65%; Fig. 8 A) and *URA3:Gcn4 BS* (~60%; Fig. 8 B) at the nuclear periphery. LexA-Gcn4 is also sufficient to position *URA3:LexA BS* to the nuclear periphery, even under uninducing conditions (Fig. 8 C). These results suggest that the occupancy of Gcn4 on the DNA is regulated by the efficiency of Gcn4 translation and that the increase in occupancy leads to an increase in targeting to the nuclear periphery.

Gcn4-mediated targeting of *HIS4* to the nuclear periphery also leads to interchromosomal clustering of *HIS4* alleles in a diploid cell (Fig. 8 C). In the presence of histidine, the *HIS4* alleles were partially clustered and, upon histidine starvation, this clustering increased (Fig. 8 C). Clustering requires Gcn4; in strains lacking Gcn4, clustering was completely lost under both uninducing and inducing conditions (Fig. 8 C). In a strain having the uORF mutations, *HIS4* clustering was high and unregulated (Fig. 8 C). Thus, interchromosomal clustering of *HIS4* is quantitatively controlled by Gcn4 protein levels.

To compare the kinetics of spatial reorganization regulated by these three mechanisms, we analyzed the gene positioning and interchromosomal clustering of *INO1*, *HIS4*, or *PRM1* after shifting cells from uninducing to inducing conditions (Fig. 9). In each case, interchromosomal clustering occurred more slowly than targeting to the nuclear periphery, consistent with

the fact that interaction with the NPC is a prerequisite for clustering (Brickner et al., 2012). Dissociation of a histone deacetylase from the *INO1* promoter led to peripheral localization within 1 h and interchromosomal clustering after 3 h (Fig. 9 A). MAPK phosphorylation of Dig2 led to relocalization of *PRM1* to the nuclear periphery within 15 min and interchromosomal clustering within 30 min (Fig. 9 B). Relief of Gcn4 translational attenuation led to a statistically significant increase in peripheral localization within 30 min and interchromosomal clustering after 60 min (Fig. 9 C). Thus, these different mechanisms of regulation allow cells to change the spatial positioning of genes with respect to each other over different time scales.

Discussion

The work presented here argues that TFs play a critical role in controlling the spatial organization of the yeast genome and that their function can be regulated to allow this spatial organization to be dynamically altered. Put3, Cbf1, Ste12, and Gcn4 (and other TFs; unpublished data) mediate inducible repositioning of target genes from nucleoplasm to the nuclear periphery. Importantly, the BSs for these TFs function as DNA zip codes, encoding peripheral targeting and interchromosomal clustering (Brickner et al., 2012). These TFs represent four different families, suggesting that control of gene positioning is a common function of TFs. Not all TFs possess this activity; several TFs bind to the promoters of genes that interact with the NPC (Fig. S1). However, many of these BSs are not sufficient to confer

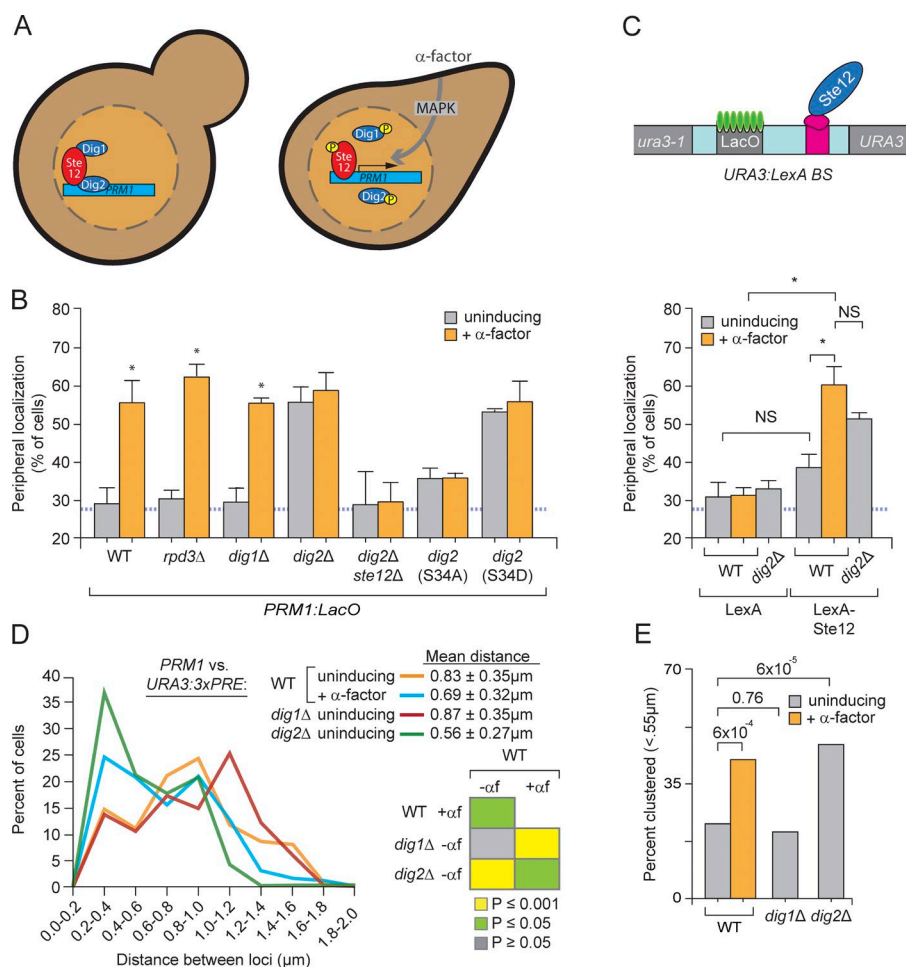


Figure 7. MAPK phosphorylation of Dig2 regulates Ste12-mediated peripheral targeting and interchromosomal clustering. (A) Schematic for regulation of Ste12-dependent transcriptional activation of *PRM1* by Dig1 and Dig2. (B) Peripheral localization of the *PRM1* gene in wild-type (WT) and mutant strains \pm α -factor. Point mutations in Dig2 were introduced into the chromosomal *DIG2* locus. (C, top) Schematic of *URA3-LexA BS*. (bottom) Peripheral localization of *URA3-LexA BS* in wild-type and *dig2 Δ* strains expressing LexA or LexA-Ste12, grown under uninducing and inducing conditions. Mean and SEM from three or more biological replicates (30–50 cells per replicate); *, $P \leq 0.05$ (Fisher exact test) between the uninducing and inducing condition. NS, not significant. (D) Distribution of distances in 0.2- μ m bins between *PRM1* and *URA3:3xPRE* in wild-type, *dig1 Δ* , or *dig2 Δ* *MATa* haploid strains \pm α -factor. (top) Mean distances and standard deviations. (inset) P-values from Wilcoxon rank sum test for each comparison. (E) Fraction of cells in which *PRM1* and *URA3:3xPRE* were ≤ 0.55 μ m apart (p-values from Fisher exact test).

targeting to the nuclear periphery (Fig. S1 and not depicted). Such TFs may be important transcriptional regulators of genes that are targeted to the NPC by different mechanisms. For example, although the Ino2/Ino4 BS is enriched among genes that interact with the NPC, these factors are neither necessary nor sufficient to promote peripheral localization. It will be important to understand what distinguishes the TFs that control spatial positioning from those that do not.

Our data suggest that TFs use multiple mechanisms to promote targeting to the nuclear periphery. Repositioning of *INO1* and *HIS4* or their respective zip codes to the nuclear periphery requires the SAGA complex, whereas repositioning of *PRM1* or *URA3:3xPRE* does not. It remains unclear what role the SAGA complex plays in recruiting chromatin. Because the TFs that require SAGA are regulated at the level of TF occupancy, whereas Ste12 is regulated downstream of DNA binding, perhaps SAGA functions to promote TF binding. For example, the ability of Put3 to bind to GRS I and mediate recruitment to the nuclear periphery is blocked by a histone deacetylase. In the absence of SAGA-mediated acetylation, Put3, Cbf1, and Gcn4 may fail to bind to the DNA, preventing peripheral targeting. Alternatively, SAGA may be recruited by the TFs and serve a more direct role in recruiting chromatin to the nuclear periphery (Cabal et al., 2006; Luthra et al., 2007; Schneider et al., 2015).

Gene positioning is regulated by either genomic context or direct modulation of TF function (Fig. 9 D). In the context of the *INO1* promoter, GRS I and GRS II zip code function is

blocked by local recruitment of the conserved Rpd3(L) histone deacetylase, which regulates hundreds of genes in yeast (Bernstein et al., 2000; Yang and Seto, 2008). Under uninducing conditions, Rpd3(L) recruitment by Opi1 and Ume6 blocks Put3 binding to GRS I. The requirement for both Ume6 and Opi1 suggests that these repressors associate poorly with the *INO1* promoter in the absence of one another. Upon induction, Opi1 dissociates from Ino2/Ino4 (Loewen et al., 2003; Brickner and Walter, 2004; Luthra et al., 2007; Schneider et al., 2015). This presumably leads to Ume6 dissociation as well, leading to loss of Rpd3(L) and increased histone acetylation, ultimately allowing binding of Put3. Consistent with this model, tethering either repressor to the *INO1* promoter was sufficient to block peripheral localization (Fig. 5 A). Thus, transcriptional repressors can regulate gene positioning by blocking the interaction of the TFs that mediate targeting.

Regulation of GRS I-mediated targeting represents a novel assay for Rpd3 function. Although loss of Rpd3 affects the expression of hundreds of genes, in most cases the mechanism of repression has not been defined. Our data provide new insight into Rpd3 function. For example, despite the fact Rpd3 shows physical interactions with Whi5 and Tup1 and genetic interaction with Gal80, none of these factors requires Rpd3 catalytic activity to block GRS I targeting. Furthermore, of the 11 that are Rpd3 dependent, only three (Cup9, Oaf3, and Xbp1) been shown to interact with Rpd3. Our results suggest that the rest may also repress transcription and gene recruitment to the nuclear periphery through local recruitment of Rpd3(L).

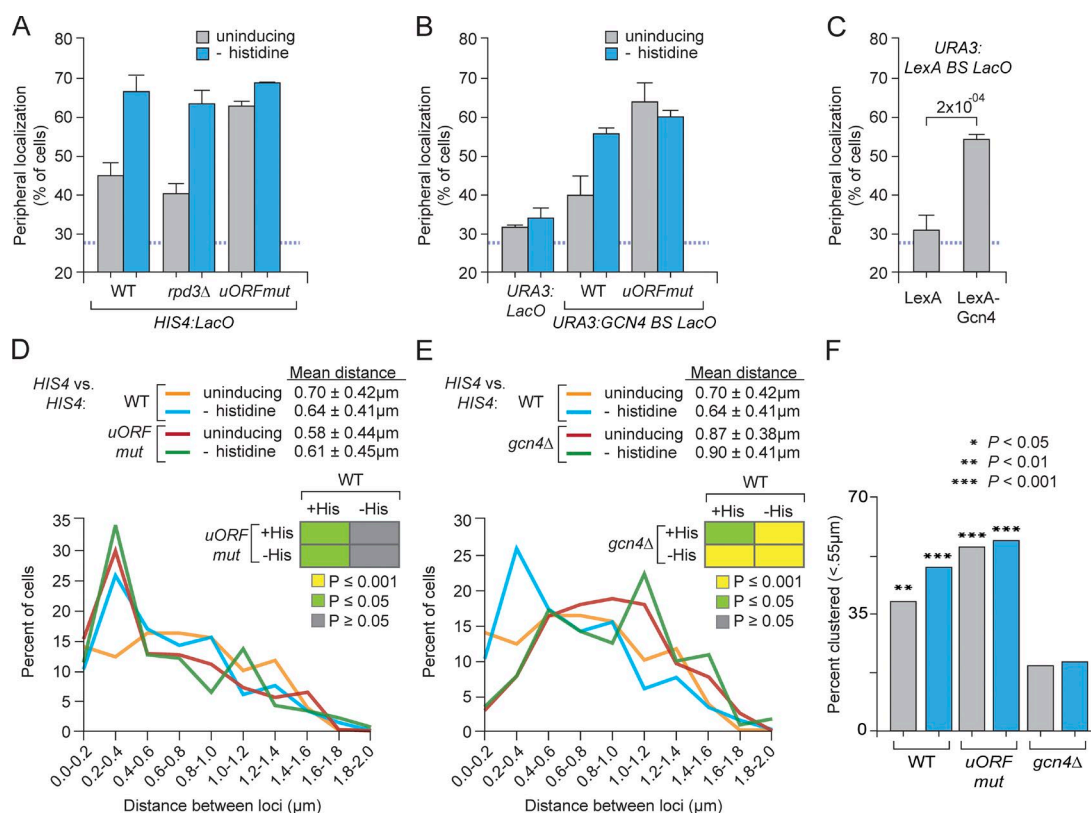


Figure 8. Translational control of Gcn4-mediated peripheral targeting and interchromosomal clustering. (A–C) Peripheral localization of *HIS4* (A), *URA3* (B), *URA3:Gcn4 BS* (B), and *URA3:LexA BS* (C) in the indicated strains grown under uninducing (gray bars) or inducing (cyan bars) conditions. Mutations in the *uORFs* were introduced into the chromosomal *Gcn4* locus (Methods). *, $P \leq 0.05$ (Fisher exact test) between uninducing and inducing condition. WT, wild-type. (C) Peripheral localization of *URA3:LexA BS* in strains expressing LexA or LexA-Gcn4. Mean and SEM from three of more biological replicates (30–50 cells per replicate; p-value from Fisher exact test). (D and E) Distribution of distances, binned into 0.2-μm bins, between alleles of *HIS4* in wild type, *Gcn4uORF* (D), or *gcn4Δ* (E) homozygous diploid strains grown ± histidine. (insets) P-values from Wilcoxon rank sum test for each comparison. (F) Fraction of cells in which the two alleles of *HIS4* were ≤ 0.55 μm apart (p-values from Fisher exact test compared with the *gcn4Δ* strain under uninducing conditions).

Another GRS I target, *TSA2*, is induced by protein folding stresses and is not regulated by Ume6 or Opi1 (Ahmed et al., 2010; Brickner et al., 2012). This gene is induced and targeted to the nuclear periphery much more rapidly than *INO1* (Ahmed et al., 2010), suggesting that Put3 binding can be regulated by other mechanisms. Consistent with this idea, we have found that many transcriptional repressors are capable of blocking Put3-dependent recruitment of an ectopic site to the nuclear periphery. These repressors regulate gene targeting by more than one mechanism: eleven repressors require Rpd3 deacetylase activity and five do not. Therefore, the same targeting mechanism can be regulated by different, context-specific regulatory strategies.

Ste12-dependent targeting is not dependent on its genomic context: when inserted at an ectopic site in the genome, 3xPRE mediated pheromone-responsive targeting to the nuclear periphery. Ste12 transcriptional activation is inhibited by Dig1 and Dig2. Previous work has suggested that Dig1 regulates the spatial arrangement of Ste12 target genes: in *dig1Δ* mutants, Ste12 forms foci in the nucleus, leading to increased intrachromosomal looping of Ste12 target genes (McCullagh et al., 2010). We found that Ste12-mediated targeting to the nuclear periphery and interchromosomal clustering was blocked by Dig2 but not Dig1. Blocking phosphorylation of serine 34 of Dig2 prevented Ste12-mediated targeting to the nuclear periphery in the presence of pheromone, and a phosphomimetic

substitution for serine 34 led to unregulated peripheral targeting and interchromosomal clustering. Dig2 regulates Ste12 function, not DNA binding; although Ste12 binding to some sites in the genome is increased by pheromone signaling, binding to the *PRM1* promoter is not (Zheng et al., 2010). Furthermore, tethering LexA-Ste12 to the *URA3* locus resulted in pheromone- and Dig2-regulated targeting to the nuclear periphery. Therefore, Dig2 must block events downstream of DNA binding that lead to targeting to the nuclear periphery.

Unlike *INO1* and *PRM1*, Gcn4 is capable of promoting significant peripheral targeting and interchromosomal clustering under uninducing conditions. When cells were starved for amino acids or when the translational regulation of Gcn4 was disrupted, targeting was much stronger. Thus, low-level production of Gcn4 leads to partial targeting to the nuclear periphery, and increased production of Gcn4 leads to increased targeting to the nuclear periphery, a quantitative, rather than a qualitative, change upon induction of its target genes (Fig. 9 D). Consistent with this idea, the number of Gcn4 BSs also impacts the efficiency of peripheral targeting. The peripheral localization of *HIS4* (which has five Gcn4 BSs in its promoter) was consistently higher than that of *URA3:Gcn4 BS*.

Cells use many strategies to regulate TF function. Here we have defined three different ways that TF-mediated gene positioning is regulated in yeast, although it is likely that this list is incomplete. Why should cells use different regulatory schemes

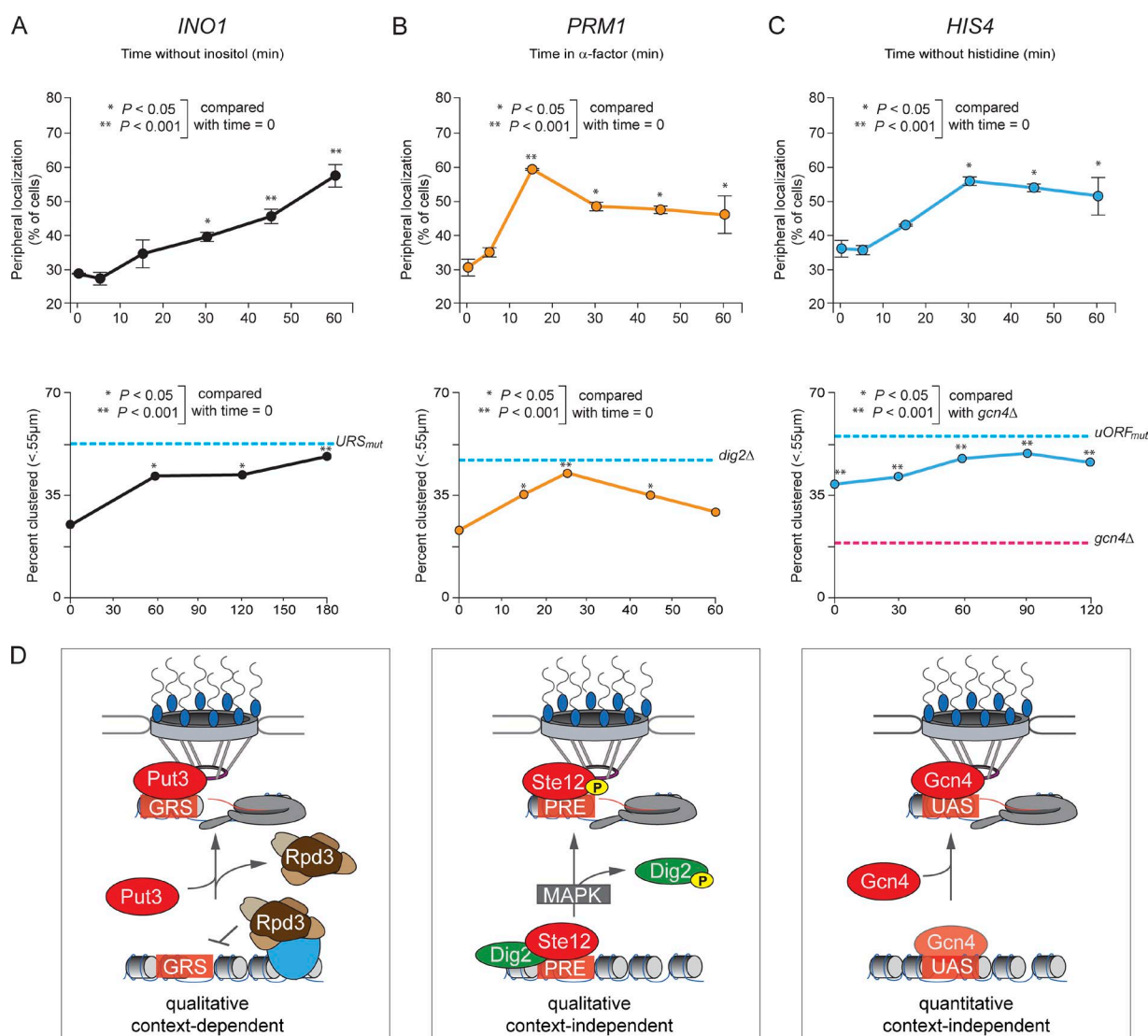


Figure 9. Different regulatory strategies lead to large-scale changes in nuclear organization over different time scales. (A–C) Time course after shifting to $-\text{inositol}$ (A), $+\alpha\text{-factor}$ (B), or $-\text{histidine}$ (C). (top) Peripheral localization of *INO1* (A), *PRM1* (B), and *HIS4* (C). (bottom) Percentage of cells in which the two loci were $\leq 55\mu\text{m}$ for *INO1* versus *INO1* in diploid cells (A), *PRM1* versus *URA3:3 \times P_{RE}* in haploid *MAT α* cells (B), and *HIS4* versus *HIS4* in diploid cells. (D) Schematic for three distinct mechanisms of regulation of gene positioning. *, $P \leq 0.05$; **, $P \leq 0.001$ (Fisher exact test) between SDC and inducing condition.

for different target genes? Because the rates of the three mechanisms we describe are very different, these regulatory strategies allow cells to dynamically alter the spatial reorganization of their genomes over very different time scales, which may have important adaptive value (Fig. 9).

Materials and Methods

Chemicals and media

All chemicals were purchased from Sigma-Aldrich, unless otherwise noted. Yeast media were obtained from Sunrise Science Products. Alpha factor was obtained from Zymo Research. Yeast and bacteria were grown with standard media as described previously (Maniatis et al., 1982). For experiments involving inositol starvation, cells were grown in SDC-inositol $\pm 100\mu\text{M}$ *myo*-inositol. For experiments involving α factor, $100\mu\text{M}$ (final concentration) was added to yeast suspended in $100\mu\text{l}$ of SDC media.

Yeast strains

Yeast were transformed with plasmids described in Eggecioglu et al. (2014). All yeast strains were derived from W303 (*ade2-1 URA3-1 trp1-1 his3-11,15 leu2-3,112 can1-100*) strains CRY1 (*MAT α*) or CRY2 (*MAT α*) and are listed in Table S1.

INO1 promoter mutants $\Delta 4$, *grs1 Δ* , *grsII Δ* , and *grsI Δ grsII Δ* were created by transforming *INO1*promoter Δ strains with PCR of the *INO1* promoter containing the desirable mutations and selecting on minimal medium without inositol. The following mutations were introduced at the endogenous loci by a different approach: *UASmut INO1pro*, *URSm_{ut} INO1pro*, *ino2-L188A*, *rdp3-H188A*, *GCN4-uORF*, *dig2-S34A*, and *DIG2-S34D* were created by integrating *URA3* and *SUP4-o* ochre suppressor (Goodman et al., 1977) to replace the endogenous locus surrounding the mutation. Mutated versions of the endogenous loci were then integrated in place of the *URA3-SUP4-o* cassette by counterselection with 5-fluoroorotic acid (against *URA3*) or canavanine (against *SUP4-o*). For *UASmut INO1pro*, two UAS_{INO} elements at -178 (5'-CAC ATG-3') to -172 and -243 (5'-CATGTG-3') to -237 were mutated to

(5'-CACTTC-3') and (5'-GAAGTG-3'), respectively; for *URSmut INO1pro*, -260 (5'-TCGGCGGCT-3') to -251 was mutated to (5'-GAT TATTAG-3'); for *GCN4*, 5' third uORF ATG was mutated to AGG and fourth uORF ATG was mutated to AUC; for *rdp3-H188A*, HIS188 5'-CAT-3' was mutated to Ala 5'-GCT-3'; and for *Dig2-S34A* and *DIG2-S34D*, 5'-TCT-3' was mutated to Ala 5'-GCT-3' and 5'-GAT-3', respectively.

Molecular biology

The plasmids p6LacO128 and p6LacO128-*INO1* have been described (Brickner and Walter, 2004). The plasmids pmCh-ER03, pmCh-ER04, and pmCh-ER05 were derived from pAC08-mCh-L-TM (Meinema et al., 2011). The *GALI-10* promoter of pAC08-mCh-L-TM was replaced with the GPD1 promoter as a *SacI*-*SpeI* fragment to produce pGPD-mCh-ER16. The promoter, mCherry fusion, and 3' UTR were then inserted into shuttle vectors pRS303 (*HIS3*), pRS304 (*TRP1*), and pRS305 (*LEU2*; Sikorski and Hieter, 1989) to generate pmCh-ER03, pmCh-ER04, and pmCh-ER05. These plasmids were digested with *Bst*XI and integrated at *HIS3*, *TRP1*, or *LEU2*, respectively.

Plasmid pADH-LexA was derived from p414-ADH1 (Mumberg et al., 1995). LexA was inserted into p414-ADH1 as a *NotI*-*PstI* fragment. Repressors were PCR amplified from W303 genomic DNA and cloned into a pADH-LexA plasmid as either a *Bam*HI C-terminal LexA fusion or a *XhoI* or *PstI* fragment creating an N-terminal LexA fusion (Fig. S3 C). *Ste12* and *Gcn4* were cloned into p414-ADH1 as *XhoI* and *Bam*HI fragments, respectively.

The following plasmids were derived from p6LacO128: p6LacO128-GRS I, p6LacO128-GRS I-LexA, p6LacO128-GRS II-LexA, p6LacO128-3XPRES-LexA, p6LacO128-*PRM1*, p6LacO128-*HIS4*, and p6LacO128-*GCN4*. They were created as follows. Plasmid p6LacO128-GRS I: -266 to -366 of *INO1* promoter was amplified and inserted into p6LacO128 as a *SpeI*-*SacI* fragment. For integration at *URA3*, p6LacO128-GRS I was linearized by digestion with *StuI*. Plasmid p6LacO128-GRS I-LexA: LexA BS was inserted into p6LacO128-GRS I as a *SacI* fragment with the sequence 5'-AAGGTTGGGAAGCCCTGCAAACTCATACTGTATATATACAGTATACAAAGCT-3'. Plasmid p6LacO128-GRS II-LexA and p6LacO128-3XPRES-LexA were created by inserting the zip code along with LexA BS as *Bam*HI fragments into the p6LacO128 plasmid. GRS II LexA fragment sequence is 5'-GATCCTTCTACTGTTATTCTTCCAGCAATCATTCACGCTTGCTACGTTGTATATGAAACGAGTAGTGATACTGTATATATATACAGTA-3'. 3XPRES LexA fragment sequence is 5'-GATCGAGTCCGGGTAATACATATGTTTCAATACTGTTTCAATACTGTTTTCAGAAGTGCGTCACATATTACTGTATATATATACAGTA-3'. Plasmid p6LacO128-*PRM1*: 1500 to 3067 of the *PRM1* CDS and 3' UTR was amplified and inserted as a *Bam*HI-*NotI* fragment into p6LacO128. This plasmid was linearized by digestion with *SnaBI* to integrate at *PRM1* locus. Plasmid p6LacO128-*HIS4*: 3171 to 3577 *HIS4* CDS and 3' UTR was amplified and inserted as a *Bam*HI-*SphI* fragment into p6LacO128. p6LacO128-*HIS4* was linearized by *HindIII* and integrated at *HIS4*. p6LacO128-*GCN4*: *GCN4* BS 5'-CATGCACAGTGACTACGTTTTTTT-3' from the *HIS4* promoter (228 to 253) was inserted into p6LacO128 as a *HindIII* fragment. Plasmid p6LacO128-*GCN4* was linearized by *StuI* and integrated at *URA3*. To create the *URA3:3XPRES LacO* strain, the sequence 5'-TACATATGTTTCAATACTGTTTCAATACTGTTTTCAGAAGTGCGTCACATATTAA-3' was cloned into an integration cassette within the plasmid pZipKan using flanking *StuI* sites. It was then integrated directly into the backbone of the p6LacO128 as described previously (Egecioglu et al., 2014).

Microscopy

For all experiments, cells were maintained at $OD_{600} < 0.8$. For inositol starvation experiments, strains were grown overnight in SDC-inositol

in the presence or absence of 100 μ M *myo*-inositol. For histidine starvation experiments, strains were grown overnight in SDC, harvested, resuspended in either SDC or SDC-His media, and incubated at 30°C for 45–75 min. For α -factor stimulation experiments, strains were grown overnight in SDC, and 100 μ l of cells were transferred to a 1.5-ml Eppendorf tube with 100 μ M final concentration of α -factor for 15–25 min. After treatment, cells were concentrated by brief centrifugation and imaged immediately. The images were captured as 0.34- μ m-thick *z*-stacks (the yeast nucleus is ~ 2 μ m in diameter) with an SP5 II Line Scanning Confocal Microscope (Leica Biosystems) with 100 \times 1.44-NA (oil immersion) objective using Argon 488-nm and Diode Pumped Solid State 561-nm lasers in the Northwestern Biological Imaging Facility as described (Egecioglu et al., 2014). Cells were scored using LAS AF or LAS AF Lite software. For each individual cell, the *z*-stack with the brightest and most focused LacO dot was scored. The slice used was not necessarily the same for every cell and, for cells in which the dot was at the top or bottom of the nucleus, localization was not scored. If the center of the dot overlapped with the membrane, the cell was scored as peripheral.

Clustering analysis

Samples were visualized on a Spinning Disc Advanced Fluorescence confocal microscope (Leica Biosystems) in the Northwestern University Biological Imaging Facility. Cells were deposited on a double-cavity microscope slide (VWR) prepared with 4% low-melt agar. Slide covers were sealed with nail polish to prevent drying. Images of the cells consisted of 16 *z*-slices spaced 0.4 μ m apart, with eight time points taken at 30-s intervals over 4 min. From the first time point of each image, the distance between the two loci was measured using ImageJ. When only a single locus was apparent in a cell but the entire cell was captured in the image, the additional time points were referenced to determine whether the spot subsequently later split into two loci. In those cases, a distance value of 0 μ m was assigned to these tightly clustered loci. For images of the *PRM1* versus *URA3:3XPRES* loci, only a single time point was taken in the interest of more rapid acquisition, and tightly clustered loci were excluded from our analysis. For each condition, >100 cells were measured. The distances were binned into 0.15- or 0.2- μ m classes to generate distributions of distance used in mountain plots and heat plots. For heat plots, the mean fraction of cells and standard deviation for all bins was used. An R script was used to generate the grayscale heat map of the number of standard deviations from the mean for each bin, scaled as shown in Figs. 3, 5, and 6. To control for the effect of GFP dimerization on interchromosomal clustering, we tested a mutant monomeric form of GFP (A206K)-LacI (Zacharias et al., 2002; Mirkin et al., 2014). This mutation had no effect on the degree of interchromosomal clustering between two alleles of *HIS4* (Fig. S5).

ChIP

ChIP experiments were performed as described in Egecioglu et al. (2014). For anti-pan-acetyl and anti-H3 ChIP, lysates were fixed for 15 min. For GST-Put3 ChIP, lysates were fixed for 1 h. 5 μ l anti-pan-acetyl (sc-8649-R; Santa Cruz Biotechnology), 3 μ l anti-H3 (ab180727; Abcam), 4 μ l α -GST antibody (G7781; Sigma-Aldrich), and 8 μ l α -rabbit IgG Dynabeads (11203D; Invitrogen) were incubated with 3.0 mg lysates. All immunoprecipitations were incubated overnight at 4°C.

Online supplemental material

Fig. S1 shows identifying TFs that mediate targeting to the nuclear periphery, related to Fig. 1. Fig. S2 shows that loss of Cbf1 does not block targeting of *INO1* to the nuclear periphery. Fig. S3 shows trans- and cis-mutant effects on *INO1* transcription, related to Fig. 2. Fig. S4 shows the phenotype of LexA-repressor fusions, related to Fig. 4. Fig. S5 shows the distribution of distances and fraction of cells between alleles

of HIS4. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201508068/DC1>. Additional data are available in the JCB DataViewer at <http://dx.doi.org/10.1083/jcb.201508068.dv>.

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