

Form and function of topologically associating genomic domains in budding yeast

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The genome of metazoan cells is organized into topologically associating domains (TADs) that have similar histone modifications, transcription level, and DNA replication timing. Although similar structures appear to be conserved in fission yeast, computational modeling and analysis of high-throughput chromosome conformation capture (Hi-C) data have been used to argue that the small, highly constrained budding yeast chromosomes could not have these structures. In contrast, herein we analyze Hi-C data for budding yeast and identify 200-kb scale TADs, whose boundaries are enriched for transcriptional activity. Furthermore, these boundaries separate regions of similarly timed replication origins connecting the long-known effect of genomic context on replication timing to genome architecture. To investigate the molecular basis of TAD formation, we performed Hi-C experiments on cells depleted for the Forkhead transcription factors, Fkh1 and Fkh2, previously associated with replication timing. Forkhead factors do not regulate TAD formation, but do promote longer-range genomic interactions and control interactions between origins near the centromere. Thus, our work defines spatial organization within the budding yeast nucleus, demonstrates the conserved role of genome architecture in regulating DNA replication, and identifies a molecular mechanism specifically regulating interactions between pericentric origins.

genome organization | budding yeast | DNA replication | cell cycle | systems biology

An important distinction between eukaryotic and prokaryotic cells is the presence of the eukaryotic nucleus, which compartmentalizes the cell. It is becoming increasingly clear that the eukaryotic nuclear compartment contains additional layers of spatial organization, including the nucleolus, splicing bodies, transcriptional foci, and the peripheral localization of telomeres (1, 2). In addition, high-throughput chromosome conformation capture (Hi-C) technologies have recently revealed the spatial organization of chromatin into topologically associating domains (TADs) on the 100-kb to 1-Mb scale for mammals (3, 4), as well as the fly *Drosophila melanogaster* (5), the worm *Caenorhabditis elegans* (6), and the fission yeast *Schizosaccharomyces pombe* (7). Loci within a TAD are much more likely to interact with one another than with loci outside the domain (5, 8, 9).

In metazoans, topological domains play important roles in coordinating the DNA-templated processes of replication and transcription (10–12). Chromatin within a TAD tends to have similar histone modifications, and consequently euchromatic or heterochromatic state, so that the genome is organized into self-associated globules that are either permissive or repressive of transcription. Repressive TADs are likely to be associated with the nuclear periphery (8). In addition to coordinating transcription, TADs also coordinate replication so that replication origins within a domain activate synchronously.

That TAD nuclear organization is important for transcription and replication has motivated much recent work on the molecular mechanisms underlying TAD formation. The regions separating

one TAD from another are referred to as boundaries and are essential for TAD organization. Removing a boundary region results in the merging of two adjacent TADs (9). Boundaries are enriched with insulator elements, such as CTCF, the loss of which disrupts TAD boundaries (5, 8–10, 13, 14). In addition, both fission yeast and mammalian TADs depend on cohesin (7, 15).

Previous work suggests that TADs are conserved across diverse phyla, but are not present in *Saccharomyces cerevisiae* (16). TAD organization has been associated with a fractal globule model of polymer folding, whose scaling relationship between genomic distance and contact frequency fits metazoan but not yeast data (17, 18). Furthermore, in silico modeling using polymer models and known constraints of the budding yeast nucleus showed that many features of yeast Hi-C data, including chromosome territories and self-association of centromeres, telomeres, and chromosome arms, could be explained without TADs (19–21).

Although previous work showed no evidence of TADs in budding yeast, one of the key features of topological domains, spatially coordinated DNA replication, was previously reported (22). More specifically, origins located near budding yeast centromeres are known to fire early and those near telomeres to fire late. Genetic manipulation to place early firing centromere-proximal origins near telomeres results in late firing, whereas placing centromeres near late-firing origins results in early firing (22). Furthermore, origins close to each other along a chromosome fire more synchronously

Significance

In metazoans, topological domains are regions in the genome that more frequently associate with themselves than with neighboring regions. These domains are important for regulating transcription and replication. However, topological domains were thought to be absent in budding yeast. Thus, we did not know the degree of conservation of topological organization and its associated functions. Herein, we describe the existence of topologically associating domains in budding yeast and show that these domains regulate replication timing so that origins within a domain fire synchronously. Our work showing conservation in budding yeast sets the stage to use yeast genetics to interrogate the molecular basis of the topological domains defining genome architecture.

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than distant origins, suggesting that nearby replication timing is coordinated (23, 24). In addition to 1D proximity on a chromosome, 3D proximity is also correlated with replication timing (19, 25). Taken together, this body of work suggests a role for the spatial organization of the nucleus in coordinating replication timing in budding yeast.

Although nuclear organization may be important for coordinating budding yeast replication, the genome architecture and molecular basis of this organization is poorly understood. A number of factors affect replication timing, including chromosomal location and proximity to binding sites of the Forkhead proteins, Fkh1 and Fkh2 (25). It has been suggested that not only do Fkh1/2 determine replication timing, but they may also be required for the increased frequency of contacts among early origins (25). This theory led to a model in which the 3D organization of origin–origin contacts regulates replication timing.

In contrast to previous work, herein we report the existence of TAD-like structures in budding yeast, in which chromosomal regions have more contacts within domains than across domain boundaries. Budding yeast TADs are ~200 kb in size, which distinguishes them from recently reported self-associated domains, which are less than 10 kb in size (26). We find that our TAD-like domains do not seem to play a significant role in transcription, but correlate strongly with replication timing. Origins within a TAD are much more likely to fire synchronously than origins in different TADs even when they are a similar distance apart on a chromosome. We find that the replication regulators Fkh1/2 control contacts among origins in TADs containing centromeres (referred to as pericentric domains). This finding indicates that there are likely distinct molecular mechanisms controlling pericentric and nonpericentric chromosome contacts. Taken together, our data suggest a model where TAD organization within chromosomes, in conjunction with

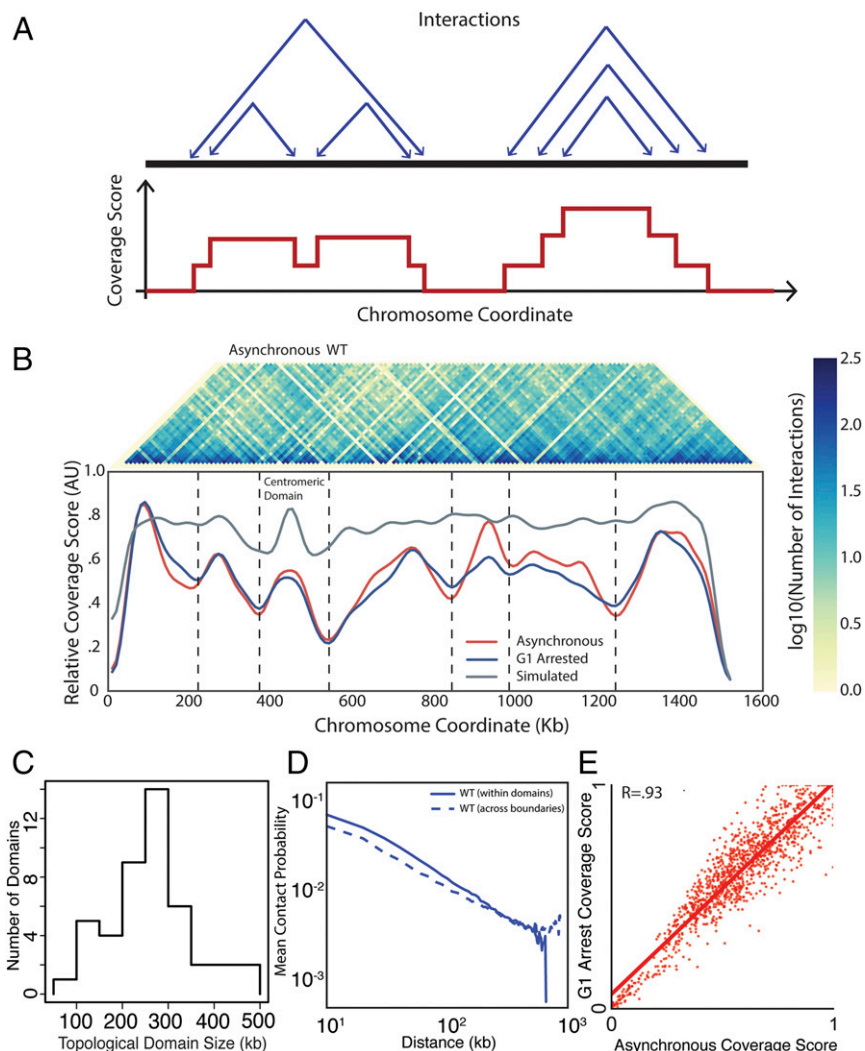


Fig. 1. Identification of TADs in the yeast genome. (A) Schematic of coverage score, which is the number of intrachromosomal interactions that span a specific locus (*Methods*). Highly covered loci are expected to be at the center of TADs, whereas poorly covered loci are expected to be in boundary regions between TADs. (Upper) Each set of two arrows points to loci denoting interactions. (Lower) Coverage score corresponding to the six interactions above. (B) Heatmap denoting the frequency of interactions in asynchronous WT Hi-C data between two loci on chromosome IV (Upper) and corresponding coverage score (Lower) for asynchronous (red) and G1-arrested (blue) cells. Vertical lines denote domain boundaries corresponding to minima in the coverage score (*Methods*). The null model (gray) that only identifies centromeric domains was generated by simulations following ref. 21, combining polymer physics and positional constraints. (C) Distribution of TAD sizes for all chromosomes in G1-arrested cells. (D) Normalized interaction frequency plotted as a function of distance for pairs of loci located within domains (solid) or across domain boundaries (dashed). (E) Correlation of coverage score for all chromosomes for pheromone arrested and asynchronous cells.

Fkh1/2-dependent associations across chromosomes, spatially organize the nucleus to determine replication timing.

Results

Analysis of *S. cerevisiae* Hi-C Data Reveals TADs. To test the hypothesis that yeast chromosomes adopt a domain-like structure, we developed a measure of association that we call “coverage score.” The coverage score for a specific locus is the sum of interactions that span that location in linear genomic coordinates (Fig. 1A). Thus, we expect coverage to drop at boundaries because few loci in one domain will interact with the neighboring domain: that is, few interactions will span the boundary. Significant oscillations in the coverage score indicate a domain structure, whereas a uniform coverage score indicates a lack of domain structure. We first applied our coverage analysis to Hi-C data for human fetal lung cells (8) and compared our coverage score with previously called domain boundaries. Nearly all previously identified domain boundaries are located near minima in the coverage score (Fig. S1A). Having validated our method on human Hi-C data, we next applied our coverage analysis to a previously generated yeast Hi-C dataset (19). Coverage varied along chromosomes, providing strong evidence for domains (Fig. 1B and Figs. S1B and S2A and B). When we applied our coverage score to a null model, based on steric interactions between chromosomes, centromeric and telomeric tethering, and DNA polymer physics (21), we only identified centromeric domains. We developed a procedure to identify TADs, which found 41 TADs with a median size of 260 kb, which is similar in size to TADs found in *D. melanogaster* and some of the smaller mammalian domains (Methods, Fig. 1C and Table S1) (4, 5, 8). The boundaries identified by minima in coverage score were similar to those found by identifying changes in directionality index, as used in Dixon et al. (8) (Fig. S1B). However, our coverage score method was more robust because the boundary number and position were less sensitive to changes in parameters than the directionality index method (Fig. S1C and D). Pairs of loci at a given distance within a predicted domain have a greater likelihood of interacting than pairs of loci in different domains calculated using the coverage score, but not the directionality index (Fig. 1D and Fig. S1F). Moreover, the interactions between loci of similar linear genomic distance were more similar if the loci were located within the same TAD (Fig. S2C).

Because our TADs are based on Hi-C data from an asynchronous culture, it was possible that our conclusions resulted from a small number of mitotic cells with a higher degree of chromosome condensation and therefore do not reflect interphase organization. To test this possibility, we performed Hi-C on cells arrested in G1 phase by exposure to mating pheromone. We observed that these G1-arrested cells exhibit a similar coverage score profile to the profiles of freely cycling cells (Fig. 1E and Fig. S3). Thus, TAD organization is not the result of mitotic condensation. In addition, the genome-wide changes in transcription associated with G1 pheromone arrest (27) do not influence domain organization. Similarly, mammalian TADs are already established in the G1 phase of the cell cycle (28) and remain nearly the same across cell types with very different transcriptional programs (29, 30). These observations suggest that stable chromosome conformation across cellular states is a conserved feature of genome organization.

Domain Boundaries Are Enriched for Markers of Transcriptional Activity. Metazoan topological domains correlate with histone modifications and transcriptional activity (5, 8, 10). Because correlated transcriptional activity within metazoan TADs is a hallmark of this type of chromatin organization, we decided to investigate the relationship between transcription and yeast TAD organization. To test this relationship, we examined the difference in active RNA polymerase II density at loci within and across domains. We did not find a significant difference in density, which is consistent with the notion that proximal elements generally determine tran-

scriptional regulation in yeast (Fig. 2A) ($P > 0.05$). We also found minimal differences in histone modifications within and across domains (Fig. 2B–D). Because metazoan TAD boundaries exhibit differential chromatin regulation, we tested for a similar

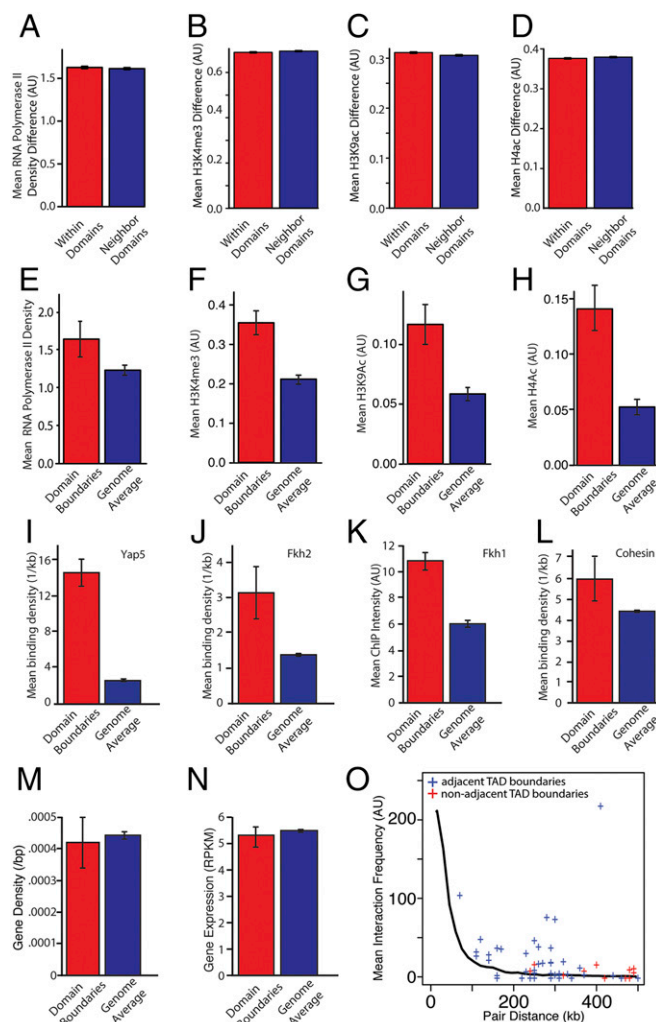


Fig. 2. Transcription is enriched in TAD boundaries. (A–D) For each gene we calculate the mean RNA polymerase II density and level of histone modification. We then determine the difference in these values for all gene pairs in two neighboring TADs. For gene pairs within a TAD and for gene pairs spanning a TAD boundary, the differences in polymerase II density and histone modifications are not statistically different ($P > 0.05$). (E–H) A gene was defined as being within a TAD boundary region if it is within 5 kb of the boundary. Compared with the average for the whole genome, RNA polymerase and associated histone modifications are significantly enriched at TAD boundary regions. (I and J) Binding site density relative to TAD boundaries for Yap5 (I) and Fkh2 (J). ChIP-chip data were used to estimate the mean signal for (K) Fkh1 and (L) cohesin. The transcription factors Fkh2, Yap5, and Fkh1 are enriched at TAD boundaries ($P < 0.01$ for all comparisons). P values were calculated using Bonferroni correction for multiple hypothesis testing. (M) Gene density at domain boundaries compared overall gene density. (N) Transcript abundance from genes at domain boundaries compared with all other genes. (O) Interaction frequency for TAD boundaries on the same chromosome as a function of their genomic distance apart. Each + indicates intrachromosomal interaction frequency for a pair of boundaries. The solid line denotes the average distance-dependent interaction frequency for all fragments. TAD boundaries interact more frequently with one another than expected given their genomic distance ($P < 0.01$ compared with datasets having the same number of pairs the same distance apart whose interaction frequency was determined by sampling from the distance-dependent interaction distribution).

effect in yeast. We consistently found significant differences in RNA polymerase II density and histone modifications within 5 kb of boundaries compared with the rest of the genome (Fig. 2 *E–H*). Next, we sought to examine if boundary elements were enriched for specific transcription factor binding sites. We analyzed the location of binding sites, as determined by MacIsaac et al. (31), relative to TAD boundaries. We found enrichment for Fkh2, Yap5, and other transcription factor binding sites (Fig. 2 *I* and *J*, Fig. S44 and Table S2). Examination of ChIP-chip data from refs. 25, 32, and 33 revealed that Fkh1 was enriched at boundaries, whereas topoisomerase II was depleted (Fig. 2 *K* and Fig. S4B). Cohesin shows slight enrichment at boundaries, but the effect is not statistically significant (Fig. 2 *L*) ($P = 0.07$). We ensured that these observations were not the consequence of different gene density at boundaries, which were similar to the rest of the genome (Fig. 2 *M*). Next, we tested whether the difference in transcriptional activity led to a steady-state difference in transcript abundance as measured by RNA-seq. However, we found that there was no significant difference in overall expression, which might be because of mRNA processing and degradation (Fig. 2 *N*). In addition, domain boundaries interact much more with one another than would be expected for loci of similar genomic distance apart (Fig. 2 *O* and Fig. S2D) ($P < 0.01$). We speculate that these interboundary contacts may play some role in

organizing yeast TADs. Taken together, these data indicate that boundary elements might contribute to gene expression but, unlike in metazoans, transcriptional activity differs little, if at all, between TADs.

Timing of Cell Cycle-Dependent Gene Expression Is Independent of Genome Position. Although mRNA abundance and markers of transcriptional activity are most often examined, the ability to dynamically activate or repress transcription is also affected by chromatin and is associated with nuclear position (34). To test if transcription dynamics are affected by genome architecture, we examined the timing of transcriptional activation during the cell cycle. We restricted our attention to 291 genes activated by the related heterodimeric transcription factors SBF (Swi4–Swi6) and MBF (Mbp1–Swi6), which were previously defined as the G1/S regulon (35, 36). To test if topological domains contribute to transcriptional activation, we examined differences between the activation time of genes located within a domain compared with those located across different domains. Timing of activation within domains was only slightly (about 1 minute) more similar than across domains ($P < 10^{-4}$), suggesting that topological domains could explain a little of the 30-min variation in timing across the G1/S regulon (Fig. 3A). To test if domain location affects the timing of transcriptional activation, we integrated 1 kb of an early

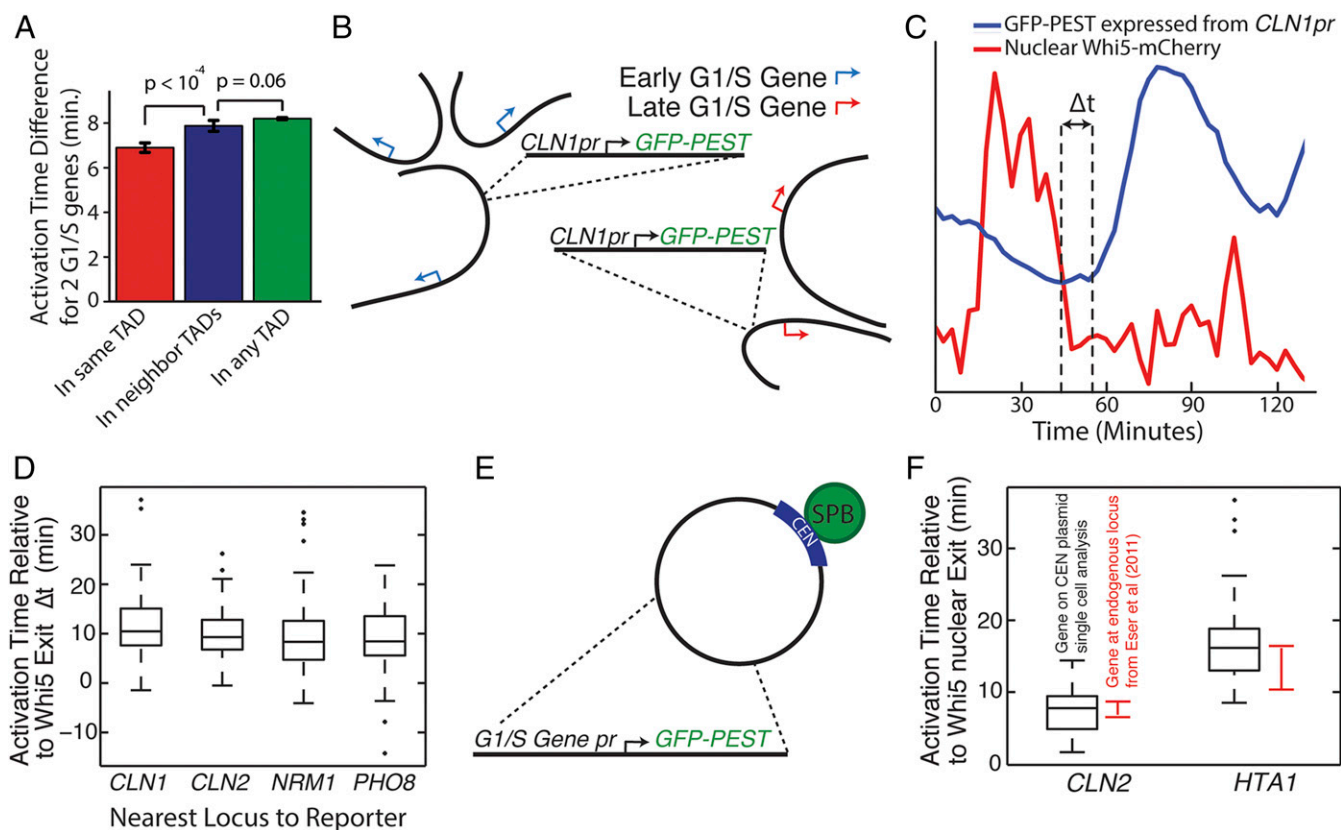


Fig. 3. Cell cycle-dependent transcriptional activation is independent of gene nuclear location. (A) The timing of transcriptional activation in the cell cycle for all genes regulated by the SBF and MBF G1/S transcription factors was calculated in Eser et al. (35). Genes within the same TAD are activated at similar times compared with genes in neighboring domains. (B) Schematic of G1/S gene translocation experiment in which a construct containing an early gene promoter, *CLN1pr*, regulating a destabilized GFP, is integrated next to either an early activated gene or a late activated gene. (C) Transcriptional activation time in the cell cycle was calculated relative to the exit of the transcriptional inhibitor Whi5 from the nucleus, which marks the point of commitment to the cell cycle in late-G1 phase. Cells express a Whi5-mCherry fusion protein from the endogenous locus. (D) *CLN1pr-GFPpest* was integrated at the *CLN1*, *CLN2*, *NRM1*, and *PHO8* loci, but this did not change the distribution of promoter activation times ($P > 0.05$ all comparisons). (E) Schematic for experiment comparing gene activation times for promoter GFP constructs expressed from CEN (centromere) plasmids, which localize near the spindle pole body (SPB). (F) The relative timing of transcriptional activation is similar for an early activated (*CLN2*) and a late activated (*HTA1*) gene expressed from either CEN plasmids or the genome. Data for transcriptional activation from the genomic loci are from Eser et al. (35).

gene promoter (*CLN1*) driving the expression of a destabilized GFP reporter at loci near early or late transcribed SBF- or MBF-regulated genes (Fig. 3*B*). Distributions of transcriptional activation timing relative to Whi5–mCherry nuclear export, which marks the G1/S transition known as *Start* (37), was the same for all loci, suggesting that there is little if any spatial effect on transcriptional activation timing ($P > 0.05$) (Fig. 3*C* and *D*). Consistent with this conclusion, the timing of the SBF-regulated genes *HTA1* and *CLN2* expressed from CEN (centromere) plasmids that maintain a constant nuclear position near the spindle pole body, was similar to their timing when expressed from their endogenous loci (Fig. 3*E* and *F*). Thus, timing of cell cycle-dependent transcriptional activation was independent of gene location.

Topological Domains Predict Replication Timing. Having examined the relationship between TADs and transcriptional regulation, we turned our attention to DNA replication, which is influenced by topological domains in metazoans (12, 38). DNA replication is activated at different times at different origins, but neighboring origins on the same chromosome tend to replicate synchronously (39, 40) (Fig. 4*A*). Replication starts earlier near centromeres and later near telomeres, and is likely affected by nuclear organization (22, 25). Origins replicating at similar times preferentially associate in three dimensions, which is reflected in two broad genome-wide clusters comprised mostly of early or late replicating origins (19, 25).

Two hypotheses might explain spatial correlations in replication timing. Either there is a length scale over which origins lose temporal correlation or there are discrete boundaries separating groups of similarly timed origins. To distinguish between these two possibilities, we set out to determine if the boundaries of topological domains separate clusters of similarly timed origins. When comparing origins of similar distance apart in linear chromosome coordinates, we found that the difference in replication timing between origins within topological domains was less than that be-

tween origins separated by a domain boundary (Fig. 4*B* and *C*) ($P < 10^{-3}$ all comparisons for distance bins in Fig. 4*C*). We note that the analysis in Fig. 4 is based on replication timing data from Raghuraman et al. 41. Our analysis suggests that topological domains in yeast may represent regions of coregulated DNA replication separated by boundary elements.

We emphasize that our 41 yeast TADs define intrachromosomal replication domains on linear coordinates, which is very different from previous analysis that defined two interchromosomal clusters of similarly timed origins (19, 25). However, it was also noted that the replication timing of groups of neighboring origins were similarly affected in cells lacking *FKH1* and expressing a mutant allele of *FKH2* (25). Intriguingly, these *FKH* mutant cells synchronously replicated their genome, resulting in an overall narrowing of the distribution of replication times (Fig. 4*D*), and exhibited a modified distribution of interactions for ARS305 (25). Taken with our observation that Fkh1 and Fkh2 are enriched at domain boundaries (Fig. 2*J* and *K*), these data suggest a potential role for Fkh factors in organizing genome architecture.

Fkh1 and Fkh2 Are Not Required for TAD Formation, but Promote the Association of Pericentric Origins. To test the hypothesis that Forkhead transcription factors regulate genome architecture, we sought to measure interaction frequencies across the genome in cells lacking the two Forkhead transcription factors, Fkh1 and Fkh2. Because of the severe phenotype of *fkh1Δfkh2Δ* cells (42), we generated a strain conditionally expressing *FKH1* from a galactose-regulated *GAL1* promoter in a *fkh2Δ* background (*GAL1pr-FKH1 fkh2Δ*). We grew these cells overnight on media containing galactose before switching cells to media containing glucose to shut off exogenously controlled *FKH1* expression. Cells were grown for another 8 h on glucose media so that there was no detectable Fkh1 protein at the time we performed DNase Hi-C (Fig. S5). The resultant data were used to calculate coverage scores for all chromosomes (Fig. S4 and Fig. S6). We note here that we now performed

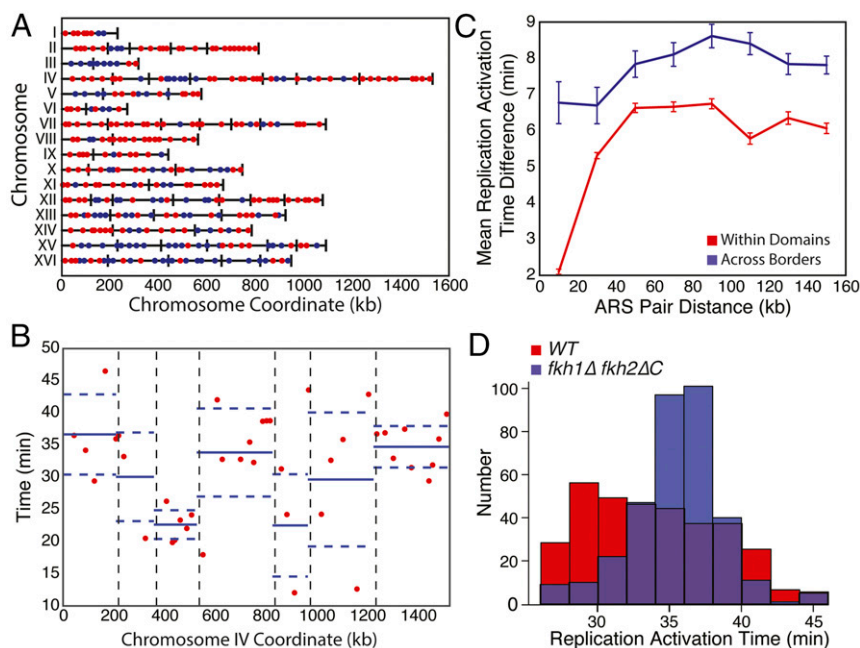


Fig. 4. Replication origins within TADs synchronously initiate DNA replication. (A) Origins were classified as early (blue) or late (red). Replication timing exhibits spatial correlation. Black lines denote TAD boundaries. (B) TAD boundaries (vertical dashed lines) separate clusters of similarly timed origins on chromosome IV. Each dot denotes the location and activation time of a single origin as determined by Raghuraman et al. (41). For all origins within a TAD, the blue solid and dashed horizontal lines represent the mean activation time and its SE. (C) The difference in replication timing between origins (ARS, autonomously replicating sequences) within TADs is smaller than for origins of similar distance apart in different TADs. (D) *fkh1Δ fkh2ΔC* mutants more synchronously replicate their DNA compared with WT. Data from Knott et al. (25) were analyzed.

Hi-C using DNase digestion (43), a method not previously applied to yeast. Coverage scores were similar for WT cells applying either restriction enzyme or DNase digestion, indicating that genome architecture is not strongly sensitive to Hi-C method (Fig. S7A–C). Coverage scores for WT and Fkh1/2-depleted cells were similar, suggesting that overall genome architecture is not strongly dependent on Fkh1/2 (Fig. 5A and B). Moreover, domain boundaries determined in Forkhead mutants did not differ dramatically from WT domain boundaries (Fig. 5C and D). Further supporting the preservation of domains in the absence of Forkhead, for loci a similar distance apart, interactions within domains remained more likely than interactions across domain boundaries (Fig. 5E). Taken together, our data strongly suggest that Fkh1 and Fkh2 are not required for the formation of topological domains.

Intriguingly, Fkh1/2-depleted cells had a lower interaction frequency at longer distances than WT, suggesting a role for Fkh1/2 in promoting long-range interactions (Fig. 5F). To test the idea that

Forkhead factors directly mediate long-range interactions, we examined the relationship between Fkh1/2 binding site density and changes in interaction frequency in WT and Fkh1/2-depleted cells for all bins. Consistent with this idea, Fkh1/2 binding site density correlates with changes in interaction frequency (Fig. S7D). However, so did the binding site density of the unrelated transcription factor Swi4 (Fig. S7E) and gene density (Fig. S7F). It is therefore unclear how specific Fkh1/2 binding affects interaction frequency.

Previous observations suggested that the proximity of replication origins correlates with the synchrony of firing (19, 25). This finding indicates that although Fkh1/2 do not influence TAD formation, they might be critical for origin–origin interactions, which could be important for replication timing. To test this model, we compared the frequency of these contacts in WT and Fkh1/2-depleted cells. Consistent with previous observations, some early-activated replication origins (cluster 1) interact much more frequently with each other than late-activated replicating origins (cluster 2), and the origin–origin contacts cluster broadly into two groups (Fig. 6A and B). Most of these interactions do not change in Fkh1/2-depleted cells. However, the subset of highly interacting early origins have an increased frequency of interaction with each other and a decreased frequency with the other origins (Fig. 6C and D). Next, we examined the location of the Fkh1/2-dependent interacting origins and found they were exclusively located in centromere-proximal regions (Fig. 6E). Moreover, the long-range interactions with these pericentric origins that disappear following Forkhead depletion could be directly mediated by Forkhead factors because Fkh1/2 binding sites appear frequently near these origins (Fig. 6F). This finding is consistent with our previous observation that Forkhead factors promote longer-range interactions in general (Fig. 5F). We note that Hi-C data do not measure absolute interaction frequency. Rather, Hi-C data reflect the relative frequency of interactions. Therefore, our analysis cannot distinguish between an increase in absolute contact frequency within cluster 1 origins or a decrease in interactions of other loci with origins in cluster 1.

We note that it is unlikely that contacts with the centromeric sequence promote origin clustering as similar origin–origin clusters are observed when bins containing the centromeres are excluded from analysis (Fig. S8A). Previous work showed that centromeric regions are among the earliest replicating regions in budding yeast (44, 45). This raises the possibility that the chief effect of Fkh1/2 on replication timing is through modulation of replication and chromatin organization at the centromeres. To test this theory, we examined replication timing within domains and across domain boundaries after excluding all domains containing centromeres. As previously, we found that replication origins that are a given genomic distance apart fired more synchronously if the origins were in the same noncentromeric domain (Fig. S8B). Thus, we conclude that Fkh1/2 influence some origin–origin association and modulate replication timing, but do not appear to influence the overall genome architecture.

Discussion

Here, we have shown that budding yeast chromosomes are organized into topological domains that separate regions of early- and late-replicated DNA (Fig. 7). The strong correlation between topological domains and replication timing functionally validates our TADs. That TADs are present in G1 before DNA replication supports an active role in regulating replication timing. Thus, genome architecture may explain the long-standing observations in yeast that replication synchrony is spatially organized along chromosomes and that moving origins from one place to another on the genome affects replication timing (22). This role for genome organization in replication timing was also found in mammalian cells (12, 46, 47). An additional similarity with metazoan TADs is that yeast domain boundaries are enriched for transcriptional activity markers as determined from RNA polymerase II density and histone modifications. However, in contrast to metazoan TADs,

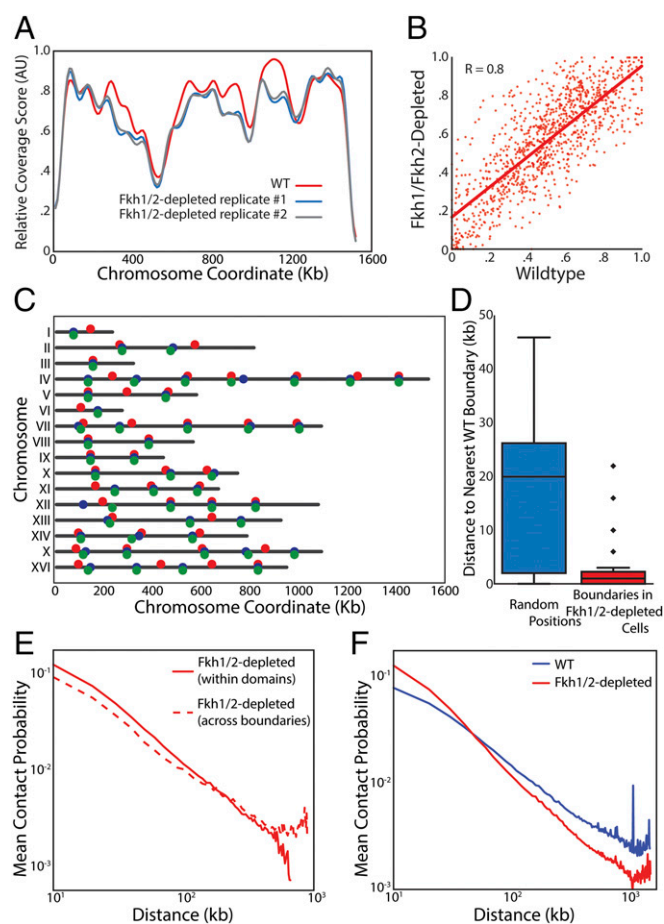


Fig. 5. TAD architecture does not depend on Fkh1/2. (A) Coverage score for chromosome IV for one WT and two Fkh1/2-depleted replicates showing no major changes in coverage in the absence of Fkh1/2. (B) Coverage in Fkh1/2-depleted cells is highly correlated with coverage in WT cells. Red line indicates the linear regression. (C) Diagram of chromosomes with boundaries marked as independently identified for WT (red) and Fkh1/2-depleted cells (blue and green denote biological replicates). (D) Comparison of distances from a boundary in Fkh1/2-depleted cells to the nearest boundary in WT cells (Right) compared with the distribution of distances from randomly chosen locations to the closest boundary in WT cells (Left). (E) Normalized interaction frequency plotted as a function of distance for pairs of loci located within a domain (solid) or across domain boundaries (dashed). (F) Normalized interaction frequency plotted as a function of distance for all pairs of loci in WT (blue) and Fkh1/2-depleted (red) cells.

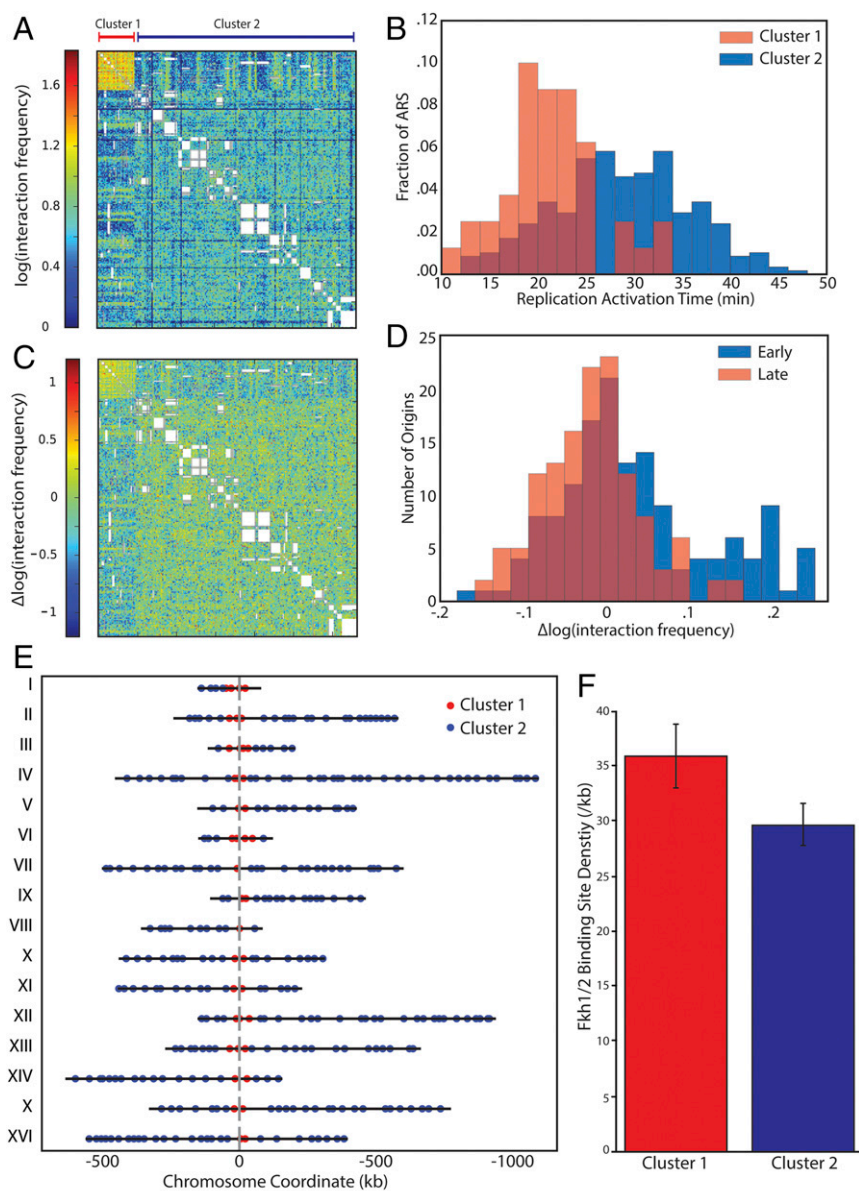


Fig. 6. Fkh1/2-depletion controls the interaction frequency of pericentric early-replicating origins. (A) K-means clustered heat map showing the interchromosomal interaction frequency between all origins for WT cells. Interactions are shown only for fragments containing an origin as determined from Raghuraman et al. (41). (B) Distribution of replication timing for origins in either of the two clusters identified in A indicates that the smaller cluster is highly enriched for early-replicating origins. (C) K-means clustered heat map showing differences in origin–origin interaction frequency between WT and Fkh1/2-depleted cells. The subset of clustered origins had increased relative interaction frequency with each other and decreased relative interaction frequency with the rest of the genome. (D) Distributions of the change in interaction frequencies for early- and late-replicating origins. (E) Early-replicating origins, whose interactions are increased by Fkh1/2-depletion, are located near centromeres. (F) Density of Fkh1 and Fkh2 binding sites for bins containing origins in cluster 1 or cluster 2.

average transcriptional activity and the ability to activate transcription within different TADs is similar. Taken together, the relationship between yeast TADs, replication timing, and boundary transcriptional activity motivate our use of TAD nomenclature and suggest a conserved role for TADs in regulating replication timing across eukaryotes.

Although spatial localization affects replication timing, it is not the only determinant and we do not know what factors account for the remainder of variation in replication timing both within and across domains. In budding yeast, previous work showed that replication synchrony is regulated by Forkhead transcription factors (25). The regulation of replication timing by Forkhead transcription factors could be through the regulation of genome architecture. Indeed, it was suggested that the C-terminal domain of Fkh2 was

required for origin–origin interactions, which led to a model of locally synchronized replication (25). However, this model was based on the examination of the contact frequencies of a single origin with the rest of the genome. In contrast, herein we used Hi-C data to perform a comprehensive analysis of all origin interaction frequencies to show that Forkhead transcription factors are not required for TAD formation. These factors do promote long-range interactions in general, including those with early-activated origins in pericentric domains. This finding suggests that Forkhead transcription factors regulate replication timing and specific genome contacts, but not overall genome architecture. Thus, our analysis supports the idea that replication timing likely requires both Forkhead transcription factors and topological domains. However,

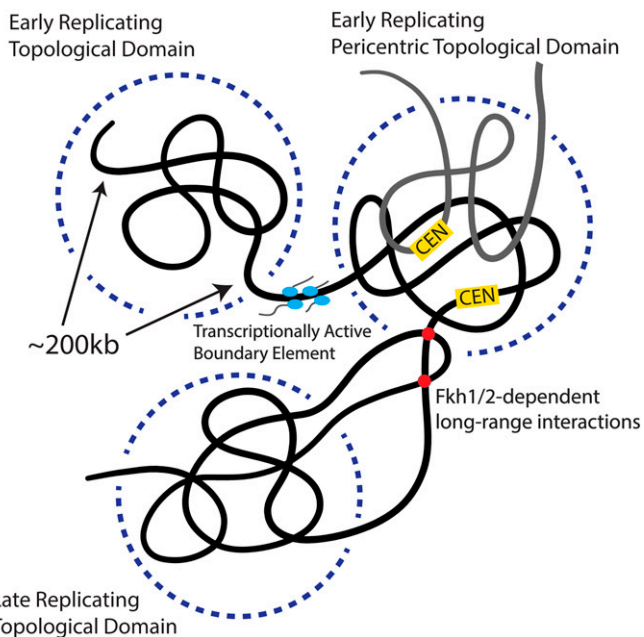


Fig. 7. Model for how the genome is organized into early- and late-replicating TADs separated by regions enriched for transcriptional activity.

definitive testing of this model will require identification of genetic determinants of topological domains in budding yeast.

That genome architecture was nearly unchanged in cells lacking the Forkhead transcription factors Fkh1 and Fkh2 leaves the question of the molecular determinants of budding yeast TAD formation unanswered. Recent progress in metazoans may shed light on the budding yeast mechanism. Metazoan TAD boundaries are enriched for CTCF, a pleiotropic transcriptional regulator, which likely plays a mechanistic role in TAD architecture (5, 8–10, 13, 14). Although CTCF is not present in yeast, other factors with which it interacts are present and may participate in TAD formation. Specifically, CTCF interacts with cohesin, which is a key molecule for the formation of TADs in fission yeast and mammalian cells (7, 15). A direct mechanistic role for cohesin in TAD formation is supported by its enrichment at fission yeast TAD boundaries (7). Although cohesin could play a similar role in budding yeast, we did not find strong cohesin enrichment at TAD boundaries, suggesting the possibility of a distinct mechanism for domain boundary determination. Nevertheless, cohesin-mediated domain formation should not yet be ruled out for budding yeast.

In metazoans, topological domains affect transcription in addition to replication timing (48, 49). In contrast, in fission yeast, cohesin mutations affecting TAD structure have only a modest effect on transcription as transcript abundance is unchanged, but the termination location is altered (7). Similarly, we show here that transcriptional activity and histone modifications in budding yeast are similar from domain to domain. Although budding yeast TAD boundary elements are enriched for transcriptional activity, we do not yet know the implication of this observation. Whereas it remains possible that transcription determines TAD boundaries, we found that altering transcription by treatment with pheromone does not change the TAD structure.

Taken together, our results firmly establish functional topological domain architecture in budding yeast, thereby demonstrating its conservation across eukaryotes. Our identification of TADs implies extensive spatial organization within the nucleus. However, this is based on Hi-C experiments performed on populations of cells and direct experiments measuring distributions of physical distances between specific loci are much needed to better interpret our data.

Nevertheless, our TADs likely represent a spatially organized cluster that impacts replication timing, which could be related to “replication factories,” the clusters of replication forks observed during DNA replication (50). Domain formation in chromatin might generate such a factory by placing several origins in close proximity (51). Thus, domain formation could ensure similar firing times because clustered origins would then share similar concentrations of replication factors. More broadly, colocalization of transcription and replication factors suggests that spatial clustering is a widely used mechanism to enrich local enzyme concentrations to increase the efficiency of multistep biochemical reactions (52–54).

Methods

Strains and Conditions. For Hi-C of pheromone-arrested cells, we used the strain BY4741 (*MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 bar1::KanMX*). Strains used for single cell transcription analysis were congenic to W303. *CLN1* promoter reporter strains were generated from laboratory stocks by standard methods. Reporters were integrated in the 5' end of *G1/S* gene promoters without disrupting nearby ORFs. For all activation timing experiments, cells were grown in synthetic complete media with 2% (wt/vol) glucose (SCD) to logarithmic growth phase, sonicated and plated on 1.5% (wt/vol) agarose SCD pads for imaging. Conditions for the Fkh1/2-depleted Hi-C experiment are described below. See also Table 1 for strains and conditions used.

Hi-C. To perform Hi-C on a population of G1 cells, BY4741 cells were cultured at 30 °C by shaking overnight in 50 mL of YEP media plus 2% (wt/vol) glucose and diluted the next morning to an $OD_{600} = 0.2$ in 1 L of YEP plus 2% (wt/vol) glucose, and 30 nM α -factor. After 120 min, cells were fixed with 1% formaldehyde for 10 min at room temperature and Hi-C was performed as described in Duan et al. (19).

Fkh1/2-depleted Hi-C was performed by growing BY4741 or *GAL1pr-FKH1 fkh2Δ* cells overnight in YEP with 1% galactose and 1% raffinose before diluting to 0.05 OD_{600} . Cells were then grown for 8 h to an OD_{600} of 1. Cross-linked nuclei were then prepared as described in Duan et al. (19). In situ DNase Hi-C was performed as described previously (43, 55). For each experiment, $\sim 2 \times 10^8$ isolated yeast nuclei were used.

Hi-C Data Processing and Normalization. It is important to normalize raw Hi-C data to avoid technical artifacts (56). To generate normalized interaction maps from the newly generated DNase Hi-C data, we processed the samples using a standard Hi-C pipeline. Briefly this pipeline: (i) maps the paired-end reads to the budding yeast reference genome; (ii) filters out the reads that have either a mapping quality score less than 30 or an edit distance greater than 2; (iii) eliminates PCR duplicates among the remaining uniquely mapped read pairs; (iv) uses 10-kb genomic windows to bin the data at this reso-

Table 1. Strains and conditions

Strain	Genotype
BY4741	<i>MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0</i>
BY4741 bar1Δ	BY4741 <i>bar1::KanMX</i>
DCY69	BY4741 <i>HIS3:GAL1pr:FKH1 fkh2Δ::KanMX</i>
DCY80	BY4741 <i>HIS3:GAL1pr:FKH1-Flag:LEU2 fkh2Δ::KanMX</i>
W303a	<i>MATa leu2-3,112 trp1-1 can1-100 ura3-1 ADE2 his3-11,15</i>
DCY85	W303a <i>pRS316:CLN1pr:yeGFP-PEST</i>
DCY86	W303a <i>pRS316:HTA1pr:yeGFP-PEST</i>
DCY61	W303a <i>CLN1:CLN1pr:yeGFP-PEST</i>
DCY38	W303a <i>CLN2:CLN1pr:yeGFP-PEST</i>
DCY40	W303a <i>NRM1:CLN1pr:yeGFP-PEST</i>
DCY39	W303a <i>PHO8:CLN1pr:yeGFP-PEST</i>

For Hi-C of pheromone-arrested cells, we used the strain BY4741 (*MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 bar1::KanMX*). Strains used for single-cell transcription analysis were congenic to W303. *CLN1* promoter reporter strains were generated from laboratory stocks by standard methods. Reporters were integrated in the 5' end of *G1/S* gene promoters without disrupting nearby ORFs. For all activation timing experiments, cells were grown in synthetic complete media with 2% (wt/vol) glucose (SCD) to logarithmic growth phase, sonicated, and plated on 1.5% (wt/vol) agarose SCD pads for imaging. Conditions for the Fkh1/2-depleted Hi-C experiment are described below.

lution; and (v) applies matrix balancing using a previously published Hi-C data normalization method (57).

Coverage Score. To analyze genome organization using Hi-C data, we binned the genome into 10-kb regions. For each fragment created by restriction enzyme or DNase digestion, we distributed its reads to bins based on the proportion of the fragment falling into each bin. Next, we calculated the interaction frequency for each bin pair. The coverage score for each 10-kb bin was calculated by finding the total number of interactions between fragments in a 100-kb window centered on the bin of interest. Our analysis identifying domain boundaries is insensitive to the window size because the coverage scores calculated with a wide range of window sizes were highly correlated (Fig. S2 A and B). Restricting our analysis to interactions less than 100 kb prevents fragments at the center of the chromosome from having larger coverage scores as a result of chromosome end effects. In addition, when the window is truncated by a chromosome end, we normalize the coverage score by multiplying the score with the ratio of the full-window size to the truncated window size. We normalized the coverage score for each chromosome by setting its minimum value to 0 and maximum value to 1.

To estimate the variation in coverage for an unstructured genome, we examined an interaction frequency matrix generated by simulating chromosome positions following the model developed in ref. 21. This simulation takes into account the physical properties of the genome as a semiflexible polymer, steric affects, and tethering at the centromere and telomere regions.

Defining TADs. TAD boundaries are defined as local minima in the coverage score using a python package from Notes on Scientific Computing for Biomechanics and Motor Control (GitHub repository). The coverage profile was smoothed with Scipy function *SignalSmooth*, using hamming window of size 50 (<https://github.com/scipy/scipy-cookbook/blob/master/python/SignalSmooth.ipynb>). The smoothing parameter was chosen to be 10 for all chromosomes, which suppresses fluctuations less than the estimated SD of the distribution of coverage scores.

Directionality Index Analysis. Directionality index was calculated similarly to that in Dixon et al. (8) and used to identify TAD boundaries in yeast (Fig. S1B). Briefly, directionality index is defined as the sum of squared differences between half the total number of interactions (E) and the upstream (A) and downstream (B) interactions, normalized on the expected interactions. That is, $[(B - A)/|B - A|] \times [(A - E)^2/E + (B - E)^2/E]$ where $E = (A + B)/2$. The bin and window sizes for estimating interactions were altered from the metazoan analysis because the yeast genome is much smaller. The bin size for our analysis was 10 kb as for coverage score and the upstream and downstream windows were 50 kb in size. The resultant directionality index was smoothed using the python Scipy function *SignalSmooth* with a smoothing parameter of 10. Boundaries were identified as locations where this smoothed signal changed sign from negative to positive. To perform sensitivity analysis, the smoothing parameter was varied from 1 to 20 and window size from 10 to 500 kb.

Additional Hi-C Analysis. For Figs. 1D and 5 E and F, the intrachromosomal interaction frequency as a function of interlocus difference was calculated by dividing the number of interactions between each pair of bins by the total number of interactions on that chromosome. To calculate the difference in interaction frequency between WT and Fkh1/2-depleted cells, the interaction frequency for each bin in WT cells was subtracted from the corresponding bin in Fkh1/2-depleted cells.

Replication Analysis. We analyzed autonomously replicating sequence (ARS) timing data from Raghuraman et al. (41) for Fig. 4. We analyzed spatial correlations in genome replication timing data from Knott et al. (25) for Fig. 4D.

We estimated replication time for each origin by first fitting a smoothing spline to the fraction of replicated DNA. Then, we took the activation time to correspond to when 50% of the DNA was replicated as described in Raghuraman et al. (41).

Polymerase Density and Histone Modifications Analysis. Histone modification data were obtained from ChIP-chip experiments reported in Pokholok et al. (58). To compare histone modifications across TADs and 10-kb boundary regions (± 5 kb from boundary), we first identified the genes within each TAD or boundary region. Next, we averaged the ChIP-chip signal for each gene. We then calculated the difference in signal for all gene pairs within and across neighboring TADs. For comparisons of boundary regions to the rest of the genome, we averaged the signal for all genes in each category. Polymerase II density on the genome was taken from ref. (59). Similarly, we determined the average density for each gene and performed the analysis just as we did for the histone modifications described above. Binding site density was estimated using the FIMO motif finding package using the default parameters (60). Binding site motifs for each transcription factor were taken from the Yeast database (61). Transcription factor binding site density was estimated using a kernel density estimate with a 0.01 bandwidth. The mean density in bins was calculated from this value.

Western Blotting. Protein abundance was measured by Western blotting. Cells were frozen in liquid nitrogen and then lysed in 150 μ L urea lysis buffer (20 mM Tris-Cl pH 7.5, 7 M urea, 2 M thiourea, 65 mM CHAPS, 65 mM DTT, 50 mM NaF, 100 mM β -glycerophosphate, 1 mM NaVO₃, 1 mM PMSF), and homogenized for 40 s at 4 °C in a FastPrep homogenizer (MP Biomedicals) using 300 μ L 0.5-mm ceramic beads. Lysates were transferred from the homogenizing tube to a clean tube by puncturing the tube and briefly centrifuging. Lysates were cleared by centrifugation for 10 min at 17,000 \times g. Total protein was normalized based on measurement by Bradford protein assay (Bio-Rad) to a maximum of 10 μ L to which 5 μ L 6 \times Laemmli buffer was added. This protein loaded on a 10% (wt/vol) polyacrylamide gel and run for 60 min at 15 Amps. Protein was transferred to nitrocellulose membrane using the iBlot system on program 8 for 7 min. Membranes were blocked in Licor Odyssey blocking buffer (TBS; 927-50010) for 60 min at room temperature. Membranes were incubated with 1:1,000 M2 mouse monoclonal anti-FLAG (Sigma F1804) and 1:500 goat polyclonal anti-CDC28 overnight diluted in 5% (wt/vol) nonfat dehydrated milk TBS + 0.1% Tween-20 at 4 °C. Membranes were washed 1 \times 15 min and 2 \times 5 min in TBS-T and then incubated with Li-Cor IRDye 800 donkey anti-goat (product # 926-32214) and Li-Cor IRDye 680LT donkey anti-mouse (product # 926-32214) antibodies. Membranes were scanned using a Licor Odyssey CLX-0670.

Microscopy and Analysis. Images were taken using a Zeiss Axio Observer. Z1 microscope with an automated stage using a plan-apo 63 \times /1.4 NA oil-immersion objective. Automatic focusing was performed using the Definite Focus system. Whi5-mCherry was visualized by exposing for 600 ms with excitation from the Colibri 540- to 580-nm LED system at 50% intensity. Transcriptional reporter eGFP-PEST was detected by exposing for 300 ms using excitation from a Colibri 505 nm LED at 25% intensity. Images were taken for at most 20 positions every 3 min. Image analysis was performed as previously described (62). Gene activation time was calculated as in Skotheim et al. (63) relative to the time Whi5 concentration in the nucleus is half-maximum, which corresponds to the point of commitment to cell division (37).

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