Physical tethering and volume exclusion determine higher-order genome organization in budding yeast

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In this paper we show that tethering of heterochromatin regions to nuclear landmarks and random encounters of chromosomes in the confined nuclear volume are sufficient to explain the higher-order organization of the budding yeast genome. We have quantitatively characterized the contact patterns and nuclear territories that emerge when chromosomes are allowed to behave as constrained but otherwise randomly configured flexible polymer chains in the nucleus.

Remarking this constrained random encounter model explains in a statistical manner the experimental hallmarks of the *S. cerevisiae* genome organization, including (1) the folding patterns of individual chromosomes; (2) the highly enriched interactions between specific chromatin regions and chromosomes; (3) the emergence, shape, and position of gene territories; (4) the mean distances between pairs of telomeres; and (5) even the co-location of functionally related gene loci, including early replication start sites and tRNA genes. Therefore, most aspects of the yeast genome organization can be explained without calling on biochemically mediated chromatin interactions. Such interactions may modulate the pre-existing propensity for co-localization but seem not to be the cause for the observed higher-order organization. The fact that geometrical constraints alone yield a highly organized genome structure, on which different functional elements are specifically distributed, has strong implications for the folding principles of the genome and the evolution of its function.

[Supplemental material is available for this article.]

The structural organization of the genome in its nuclear environment is a key factor in the correct execution of nuclear functions (Misteli 2007; Takizawa et al. 2008; Taddei et al. 2010). For instance, in budding yeast, heterochromatin regions such as telomeres and silent mating-type loci are silenced by anchoring them to the nuclear envelope (NE), presumably through heterochromatin protein factors (Gotta et al. 1996; Hediger et al. 2002; Taddei et al. 2004, 2009; Mekhail and Moazed 2010; Horigome et al. 2011). For some other genes, the location at the NE has also been proposed to play a major role in their transcriptional repression (Csík and Henikoff 1996; Dernburg et al. 1996; Maillet et al. 1996; Brown et al. 1997; Cockell and Gasser 1999; Towbin et al. 2009). However, other genes relocate to the NE upon transcriptional activation (Casolari et al. 2004; Cabal et al. 2006), presumably instigated by forming interactions with nuclear pore complexes, facilitating mRNA export to maximize cellular transcription levels.

The spatial clustering of functionally related loci is also a key characteristic of genome organization. In budding yeast, all heterochromatic centromeres are located in a distinct region of the nucleus. This occurs because throughout interphase they remain attached through microtubules to the spindle pole body (SPB) (O’Toole et al. 1999; Jin et al. 2000). On the other hand, ribosomal DNA (rDNA) repeats appear to be clustered at the NE, opposite to the SPB in the nucleus (Yang et al. 1989; Dvorkin et al. 1991; Bystricky et al. 2005). There they form the core of a distinct subnuclear compartment named the nucleolus, which is the site of RNA pol-I–mediated rDNA transcription and ribosome biogenesis (Yang et al. 1989; Bystricky et al. 2005; Berger et al. 2008; Mekhail et al. 2008; Mekhail and Moazed 2010; Taddei et al. 2010).

There is also growing evidence for a territorial organization of the chromosomes in yeast (Bystricky et al. 2004, 2005; Schober et al. 2008). Large-scale fluorescence imaging experiments on budding yeast have revealed that several individual gene loci are strongly confined into distinct “gene territories” (Berger et al. 2008; Therizols et al. 2010). Also, several genome-wide conformation capture experiments have revealed highly structured chromatin contact patterns: Some chromosome pairs were found to interact rarely, while others interact more often than expected (Rodley et al. 2009; Duan et al. 2010). The contact patterns of chromosomes 3 and 6 in budding yeast agree with a Rabl-like configuration: Both chromosomes appear to be folded backward from their centromeres, so that their telomeres are juxtaposed (Jin et al. 2000; Dekker et al. 2002; Bystricky et al. 2005; Schober et al. 2008). Such a configuration and the resulting territorial chromosome organizations have been previously observed in live fluorescence imaging experiments (Bystricky et al. 2004, 2005; Schober et al. 2008; Taddei et al. 2010).

At the same time, there is ample evidence that the structure of the genome is highly dynamic (Marshall et al. 1997; Heun et al. 2001). Fluorescence imaging shows considerable cell-to-cell variation of gene and chromosome locations (Ferguson and Ward 1992; Csík and Henikoff 1998; Heun et al. 2001; Berger et al. 2008). Also, chromosome contacts are observed over a wide range of frequencies, indicating that not all contacts can be present simultaneously (Dekker et al. 2002; Lieberman-Aiden et al. 2009; Duan et al. 2010; Kalhor et al. 2012; Misteli 2012).

Some intrachromosomal contact probabilities are consistent with a diffusion-driven contact formation (Cook and Marenduzzo 2009; de Nooijer et al. 2009; Mateos-Langerak et al. 2009; Bohn and Heermann 2010; Dorier and Stasiak 2010). However, purely random chain behavior as studied in isolated model chromosomes...
cannot explain many of the specific patterns observed in experiments (Rosa et al. 2010).

To fairly assess the principles of chromosome folding and the possible role of molecular interactions in establishing nuclear order, we must first examine the genome structure that arises when chromosomes are tethered but otherwise randomly configured in the confinement of the nuclear environment. Previous work points toward an important role of nuclear constraints and relative chromosome arm lengths in genome organization (Berger et al. 2008; Taddei et al. 2010; Zimmer and Fabre 2011) as shown for the dynamic relationship of subtelomeric regions (Therizols et al. 2010). However, it remains to be seen if entirely random configurations of tethered chromosomes are sufficient to reproduce in a statistical manner all the available quantitative data about the yeast genome organization and gene loci interactions, including the high variability of chromosome configurations and the clustering of replication start sites as well as tRNA genes.

Our findings demonstrate that purely random configurations of tethered chromosomes do indeed reproduce in a statistical manner a wide range of data related to genome structure: genome-wide chromatin interaction frequencies; the emergence, shape, and location of specific gene territories; the relative distances between telomeres; and even the spatial clustering of functionally related chromosome regions such as early replication start sites and tRNA gene loci. Specific molecular interactions between chromatin regions, although possible, are not required to explain the available experimental data on the higher-order genome organization. Moreover, the large structural variability among individual cell’s genome configurations indicates that no single average genome structure can adequately reflect the wide range of structural features relevant to a population of cells.

Results

Population modeling for determining the three-dimensional organization of the genome

To address the challenge of representing highly variable genome structures, we construct a large population of three-dimensional (3D) genome structures, which represent a spectrum of all possible chromosome configurations, and interpret the result in terms of probabilities of a sample drawn from a population of heterogeneous structures (Methods).

All chromosomes are modeled as random configurations that are subject to the following constraints: (1) All chromosomes are confined in the nucleus; (2) all the centromeres are attached to the SPB through microtubules; (3) all the telomeres are located near the nuclear periphery; and (4) the nucleolus is inaccessible to chromosomes, except for those regions containing tRNA repeats (Methods) (Table 1; Fig. 1).

To generate a population of genome structures, we defined an optimization problem (Methods). In order to sample a representative set of all possible structures, we created a sample of 200,000 independently optimized genome structures, hereafter referred to as the structure population. We also generated a control population with an identical setup but without imposing any landmark constraints (Methods), referred to as the random control. We also calculated a structure population for a nucleus containing only a single chromosome, constrained in a manner identical to the full simulation. We refer to this population as the single chromosome population.

Probabilistic analysis of chromosome structural features

In the following sections, we analyze the spatial properties of the structure population in terms of several statistical quantities: (1) chromosome territory locations, (2) chromosome and gene loci interaction frequencies, (3) locus localization probabilities, (4) telomere distance distributions, (5) physical proximity of functionally associated genomic loci including early and late replication origins, and tRNA genes. Each property of the simulated structure population will be compared with available experimental data.

Chromosome territories as a result of constrained random encounters

We first ask to what extent the landmark constraints lead to preferred chromosome locations. We calculate the probability that each chromosome occupies any given region of the nucleus (i.e., the localization probability density [LPD] of a chromosome) (Supplemental Material). Based on the LPD, it is evident that all the chromosomes have preferred regions. Smaller chromosomes (e.g., chromosome 1 in Fig. 2) reside preferentially around the central axis, near the SPB. Interestingly medium-sized chromosomes are more likely to reside away from the central axis (e.g., chromosome 8 in Fig. 2A,B; Supplemental Fig. 1), while for large chromosomes

Table 1. Functional forms of the restraints in the scoring function

| Restrained type | Functional form | μ | d (nm) | Bead i | k |
|-----------------|----------------|---|--------|--------|---|
| Ucen | nuclear envelope restraint | $u^d$ | (0,0,0) | 1000 | All beads | 1 |
| Ulocus | nucleus localization restraint | $u^a$ | (-1200,0,0) | 1600 | $\forall i \in \delta$ | 1 |
| Uexcl | nucleus excluded volume restraint | $u^b$ | (-1200,0,0) | 1600 | $\forall i \notin \delta$ | 1 |
| Ucenl | centromere localization restraint | $u^a$ | (-700,0,0) | 300 | $\forall i \in \beta$ | 1 |
| Uloc | telomere localization restraint | $u^a$ | (0,0,0) | 950 | $\forall i \in \gamma$ | 1 |
| Uchb | chromatin chain bond restraint | $u^a$ | $r_{i+1}$ | 30 | $i = \{1...N-1\}$ | 1 |
| Uexchb | chromatin excluded volume restraint | $u^b$ | $r_{i+1}$ | 30 | $i = \{1...N-1\}$ | 1 |

All restraints are expressed as harmonic functions ($d^2$), as well as harmonic upper ($u^d$) and lower bounds ($u^a$), respectively: $d^2(r_i, \mu) = \frac{1}{2}k(r_i - \mu - d)^2$ for $r_i - \mu > d$, and $d^2(r_i, \mu) = \frac{1}{2}k(r_i - \mu - d)^2$ for $|r_i - \mu| < d$, where $r_i \in \mathbb{R}^3$ is the coordinate vector of a reference point, $d$ = reference distance, $k$ = harmonic constant, and $N$ = the total number of beads in a model. We also define several subsets of beads that share certain properties. More specifically, $\alpha$ is the set of beads assigned to the last bead of every chain, $\beta$ is the set of beads assigned to centromeric regions, $\gamma$ is the set of beads assigned to telomeric regions, and $\delta$ is the set of all beads flanking rDNA repeat regions.
(e.g., chromosome 4, whose size is 1.5 Mb), the LPD is highest in the central region of the nucleus again along the central axis.

We then ask what factors are responsible for the chromosomes’ preferred locations. For each chromosome, we calculate a new structure population for a nucleus containing only a single chromosome but otherwise constrained in a manner identical to the full simulation (i.e., the single chromosome population) (Fig. 2C). Comparing the two structure populations reveals great differences for each chromosome location (Fig. 2D). For example, in the full simulation, large chromosomes reside substantially farther from the SPB region toward the nucleolus than would be expected based on chromosome tethering alone. The differences are caused by a volume exclusion effect: Because of tethering, the chromosomes must compete for the limited space around the SPB. Smaller chromosomes are naturally more restricted to regions closer to the SPB, which in turn tends to exclude parts of larger chromosomes from these regions. For smaller chromosomes, the opposite effect is observed; in the full simulation, they exhibit an increased probability density around the SPB (Supplemental Fig. 1). Importantly, due to the volume exclusion effect, the preferred location of a chromosome is not defined by tethering alone but also depends on the total number and lengths of all other chromosomes in the nucleus.

**Genome-wide chromosome contact patterns**

Next, we measure how often any two chromosome chains come into contact with each other over the entire structure population. Interestingly, most chromosomes show distinct preferences for interacting with certain others. For instance, chromosome 1 has a significantly higher chance of interacting with chromosomes 3 and 6 than with any other chromosome. Its interactions with the large chromosomes 4, 7, and 12 are substantially depleted (Fig. 3A). Strikingly, almost identical chromosome interaction preferences are observed in an independent genome-wide chromosome conformation capture experiment (Fig. 3A; Supplemental Fig. 2A; Duan et al. 2010). Pearson’s correlation between the chromosome-pair contact frequencies in our structure population and those detected in the experiment is 0.94 ($P < 10^{-15}$). In the random control, the contact frequencies do not display any significant chromosome-pair contact preferences (Pearson’s correlation coefficient between experimental data and the random control is $-0.57$) (Supplemental Fig. 2B).

Next, we compare contact frequencies for all possible pairings of the 32 chromosome arms (Fig. 3B,C). It is evident that some pairs of chromosome arms have a greater propensity to interact than others. In particular, chromosome arms with <500 kb (chromosomes 1, 3, 5, 6, 8, and 9) are more likely to interact with each other than longer arms. For instance, the short arm of chromosome 1R is almost eight times more likely to interact with the short arm of chromosome 3L than with the long arm of 4R. Also these observations are in almost complete agreement with the conformation capture experiments (Pearson’s correlation coefficient of 0.93, $P < 10^{-15}$) (Fig. 3C,D; Duan et al. 2010).

Finally, when chromatin contacts are analyzed at a resolution of 32 kb, the contact frequency heat map of the structure population shows highly organized cross-shaped patterns (Fig. 3E).
tromere behave very differently. Contacts between subcentromeric regions are finite and cannot be exhaustive. Thus, the observed correlation coefficient of 0.54 represents a re-
magnitude that is expected for the experiment (Duan et al. 2010).

Interchromosomal locus-locus contacts

The interchromosomal contact frequencies in the structure population are correlated with those observed in experiments, with an average Pearson’s correlation of 0.54, which is highly significant ($P < 10^{-15}$) (Fig. 3E,F; Supplemental Fig. 2D). In contrast, the Pearson’s correlation between the random control and experiments is close to nil, and the distinctive contact patterns in the experimental data are completely absent in the random control (Supplemental Fig. 2C).

To examine the effect of limited sampling on the accuracy of chromosomal contact patterns, we compared our initial contact frequency map to maps generated from randomly sampling different proportions of these contacts (Supplemental Material; Supplemental Fig. 3D). In contrast, to interchromosomal contacts, the correlations between interchromosomal contact patterns are greatly affected by limited sampling. At a sampling rate of 0.1%, we find that the Pearson’s correlation between the two interchromosomal contact maps (even when assuming an ideal model) cannot exceed 0.5. Similar correlation values are observed in the Hi-C experiment when two interchromosomal contact maps are compared that are generated by using two different restriction enzymes (Yaffe and Tanay 2011). In our analysis, the observed correlation value of 0.54 corresponds to a sampling rate of ~0.2%, which is also the order of magnitude that is expected for the experiment (Duan et al. 2010). Thus, the observed correlation coefficient of 0.54 represents a remarkably good agreement between the interchromosomal contact patterns, given that the experimental and computational samplings are finite and cannot be exhaustive.

Gene localizations

We now focus on the nuclear locations of individual gene loci. The locations of eight genes have been determined by large-scale fluorescence imaging experiments (Berger et al. 2008). These locations...
are measured with respect to the two principal axes of the nucleus (Methods; Fig. 5A). We determined the two-dimensional (2D) density distributions of the same gene loci in our structure population, allowing for a direct comparison with fluorescence experiments (Berger et al. 2008) (Fig. 5A). The density distribution functions agree well with experiments, in that each locus occupies a well-defined territory. The volumes and shapes of these territories strongly resemble those observed in experiments (Berger et al. 2008). For instance, genes \textit{GAL2}, \textit{HMO1}, and \textit{SNR17A} are located near the nucleolus in the structure population, as seen in the experiment. Interestingly, the structure population places \textit{SNR17B} (no experiments available) and \textit{SNR17A} in similar positions near the nucleolus, despite the fact that these genes are located on different chromosomes. Both of these genes are involved in ribosome biogenesis and code the snoRNA U3. Also in agreement with experiments, the distribution patterns of the functionally related genes \textit{RPS5} and \textit{RPS20} are quite different. For instance, \textit{RPS5} positions are significantly more diffuse.

In order to compare quantitatively the relative positions of these eight genes, we measure their median distance along the central axis in the 2D density maps obtained from experimental data and in the structure population. These positions are in excellent agreement (Pearson's correlation is 0.95, \(P < 10^{-4}\)) (Fig. 5B).

**Pairwise telomere distances**

It is well known that telomeres are not positioned randomly on the nuclear periphery (Gotta et al. 1996; Bystricky et al. 2005; Berger et al. 2008; Therizols et al. 2010). Fluorescence imaging has revealed that the distance between any two subtelomeres increases gradually with the arm lengths of their chromosomes (Therizols et al. 2010). For a given subtelomere, this relationship is linear. In the structure population, we observe a very similar behavior. More specifically, after applying a change point analysis (Zeileis et al. 2003), we find that the distance between subtelomere pairs as a function of arm length is divided into two linear regimes (Fig. 6). For chromosome arms with lengths up to \(360\) kb, the distances observed in our structure population increase with a relatively steep slope. Above \(360\) kb, the slope decreases significantly. This behavior is entirely consistent with experiments, and the change in slope has been explained as follows (Therizols et al. 2010). For small arms, the accessible position of a subtelomere at the NE is entirely restricted by the arm length. Hence, the median distance between two subtelomeres increases rapidly with their accessible areas. However, at a certain arm...
length, the subtelomere is able to reach all points on the NE. Further increases in arm length do not dramatically increase the median subtelomere distance.

Interestingly, the change in slope occurs at a slightly different arm length in experiments (∼310 kb) (Therizols et al. 2010) compared to our structure population (356–440 kb) (Supplemental Fig. 5a). However, the incompleteness of the experimental data can explain some of this difference. If we only include those chromosomes that are also analyzed in the experiment (Supplemental Fig. 5b), the change in slope in our simulation shifts to 309–327 kb in remarkably good agreement with the experiment.

**Telomere clusters**

To identify subtelomere clusters in the structure population, for each subtelomere we calculated the fraction of structures with at least one other subtelomere within 250 nm. In agreement with another experiment (Therizols et al. 2010), such small subtelomere distances are infrequent (1%–3%) for most chromosomes (Supplemental Fig. 6A). However, a few co-localizations are observed more frequently between relatively short chromosome arms, namely, 1R:1L, 6R:6L, 1R:9R, and 3L:3R. These pairings occur in 12%–20% of the population, also in agreement with experiments (Bystricky et al. 2005; Schober et al. 2008; Therizols et al. 2010). For example, the pairs 3R:3L and 6R:6L were recently reported to form significant but transient interactions, leading to the formation of chromosome loops (Bystricky et al. 2005; Schober et al. 2008; Therizols et al. 2010). In general, we find that as the length difference between chromosome arms grows more pronounced, the probability of their telomeres being co-localized decreases. Interestingly, the volume exclusion effect has a pronounced effect on the co-location frequency of telomeres. For small and also large chromosomes, the volume exclusion effect increases significantly the co-location frequency, while for medium-sized chromosomes the opposite is observed by decreasing the co-location frequency (Supplemental Fig. 6B). For instance, the fraction of co-located telomeres increases by almost 20% for the small chromosome 6 upon the presence of all other chromosomes in the nucleus, while it decreases by 60% for the medium-sized chromosome 8.

**Co-localization of functionally related loci**

Next, we investigate whether functionally related gene loci are co-localized in the structure population. First, we compare the 3D spatial distributions of early and late replication start sites in the structure population. These sites are distributed across all chromosomes (Fig. 7, right panels). Experimental evidence exists that early replication sites are spatially clustered during interphase (Di Rienzi et al. 2009; Duan et al. 2010).

In each structure of the population, we calculate the mean pairwise distance between all early replication sites. The frequency distribution derived from these mean pairwise distances is compared to a distribution chosen from randomly selected sites in the genome. We observe significant spatial clustering of the early replication sites (Fig. 7A), in the sense that their mean pairwise distances are significantly less than would be expected from randomly selected sites (Stouffer’s Z-transform [Stouffer et al. 1949] tests z-scores < −160; Supplemental Material). This observation holds for all three sets of early replication origins identified in the literature (Feng et al. 2006; McCune et al. 2008; Sekedat et al. 2010). Remarkably, for late replication sites we see the opposite effect: a statistically significant increase in the mean pairwise distances between late replication sites compared with the background. It appears that, on average, early replication start sites are closer to the centromere on the chromosome sequence compared with the late start sites (all P-values < 10^{-5} for the three data sets) (Supplemental Material).

We also analyzed the spatial positions of all tRNA gene loci in the genome, which have been observed to cluster in experiments (Thompson et al. 2003; Duan et al. 2010). Again, we observe a statistically significant decrease in the pairwise distances between tRNA loci (Fig. 7B) compared with randomly picked loci.

Our observations clearly indicate that the chromosomal locations of these specific sites are not randomly distributed over the genome; they are positioned in such a way that early replication sites have a higher probability of being co-localized when the chromosome chains behave as random polymer chains that are subject to nuclear landmark constraints.

**Discussion**

In this paper, we demonstrate that purely random configurations of tethered chromosomes reproduce in a statistical manner all the experimental hallmarks of genome organization in Saccharomyces cerevisiae. Specifically, random configurations generate structural features that agree remarkably well with (1) the highly specific
interaction patterns between individual chromosomes, chromosome regions, and chromosome folding patterns obtained by genome-wide conformation capture experiments (Duan et al. 2010); (2) the emergence, shape, and position of individual gene territories revealed by probability maps from fluorescence experiments (Berger et al. 2008); (3) the distribution of median distances between telomeres; (4) the relative frequencies of telomere loci shown in imaging experiments (Bystricky et al. 2005; Schober et al. 2008; Therizols et al. 2010); and even (5) the physical proximity of functionally related gene loci, including early replication sites and rRNA gene loci.

In addition to chromosome tethering, the main organizing factor is a volume exclusion effect, as a result of the competition of all the chromosomes for the limited nuclear space. The fact that the chromosome arms have different lengths gives rise to important nuances of organization and implies that the locations of a gene or chromosome territory depends on all the other chromosomes. Therefore, the gene territory position and specific interaction patterns of a given gene locus is determined not only by its chromosome sequence position and the arm lengths of its own chromosome but also by the total number and the relative arm lengths of all other chromosomes. The volume exclusion effect can even create counterintuitive effects. For instance, for small and large chromosomes the volume exclusion effect leads to an increase in the frequency with which subtelomeres on the same chromosome are in proximity to each other, while for medium-sized chromosomes a decrease is observed.

Our findings have several important consequences. First, we show that a small number of purely geometrical constraints on otherwise randomly configured chromosomes can lead to a highly structured 3D genome organization. Second, the hallmarks of genome organization can be explained without calling on specific molecular interactions between chromatin regions or chromatin-bound proteins. For instance, random chromosome encounters can also statistically explain the spatial features often attributed to an apparent Rabl-like chromosome folding, which refers to the back-folding of subcentromeric chromosome regions so that chromosome arms appear juxtaposed. This pattern is mainly caused by the volume exclusion effect (Fig. 4). In response to the competition for the limited space around the SPB, chromosome regions on both sides of the centromere show a statistical preference for bending toward each other. When averaged over the entire cell population, this tendency gives rise to the distinctive cross-shaped intra-chromosomal contact patterns observed in experiments and in our structure population. However, most individual structures will not exhibit simultaneously all the features of such an apparent Rabl-like fold. We therefore emphasize that the data should be explained as a statistical preference for chromosome contacts but not necessarily be interpreted as a stable chromosome fold. An interesting predication of this model is that the Rabl-like subcentromeric contact pattern should not be expected in yeast species if the number of chromosomes was considerably smaller even if the chromosomes were all tethered to nuclear landmarks. Although S. pombe and S. cerevisiae have similar genome sizes, the former has only three chromosomes. The prediction is sustained: In genome-wide conformation capture experiments, Schizosaccharomyces pombe yeast does not show the cross-shaped intrachromosomal contact patterns characteristic of this type of folding (Tanizawa et al. 2010).

Another remarkable result is that the early replication sites in our structure population have a high probability of being in close proximity compared with the background distribution of pairwise separations. In contrast, late replication sites have a lower probability of being colocated compared with randomly selected sites. This difference may help regulate a naturally occurring order on replication timing. The existence of these and other co-location patterns may indicate that the relative positions of affected loci in the chromosome were selected by evolution. Due to excluded volume effects, the spatial position of a gene in the nucleus is not only modulated by its relative sequence position in its own chromosome, but also by the relative arm lengths and the total number of all other chromosomes in the nucleus.

We also note that our study provides additional evidence for the existence of a chromatin fiber in the yeast interphase nucleus with length and density properties similar to the 30-nm fiber. We created an alternative structure population consistent with a 10-nm chromatin fiber, and the statistical results do not agree with the described experimental evidence.

Finally, we believe that our results point toward a considerable structural variability of genome structures among individual cells. Each structure in our population not only differs considerably from the “average conformation” but also from all the other structures in the population (<0.3% of loci contacts are shared between any two structures; Supplemental Material). No single-genome structure or population-averaged structure is representative of the population. Although the true structural variability is unknown, our results indicate that a single structural model cannot adequately reflect all the spatial features of the genome. It is crucial to analyze genome structures from a statistical rather than an individual standpoint. Structural analysis should be performed by generating a population of 3D genomes, which represent the spectrum of all possible chromosome configurations consistent with the data. The structural organization of the genome can then be interpreted sta-
of every chain, the configurations of the chromosomes are optimized.

Nuclear architecture

The nuclear radius is set to 1 micron, as suggested by experiments (Gasser 2002; Chubb and Bickmore 2003; Berger et al. 2008; Meister et al. 2010). The relative position and size of the SPB and nucleolus are taken from imaging experiments (Berger et al. 2008). The SPB and nucleolus are located at opposite ends of the nucleus, while a central axis connects the centers of the SPB, nucleolus, and nucleolus (Fig. 1).

Scoring function

The scoring function is defined as a sum of spatial restraints and quantifies the degree of consistency between the structure and the imposed landmark constraints derived from experimental information. To optimize the structure, the scoring function is minimized to a score of zero. The scoring function is written as

\[ S(\mathbf{r}_1, \ldots, \mathbf{r}_N) = \sum_{i=1}^{N-1} U_{\text{ch}}(\mathbf{r}_i, \mathbf{r}_{i+1}) + \sum_{i=1}^{N} U_{\text{exc}}(\mathbf{r}_i) + \sum_{i=1}^{N} U_{\text{cen}}(\mathbf{r}_i) + \sum_{i,j} U_{\text{tel}}(\mathbf{r}_i, \mathbf{r}_j) + \sum_{i,b} U_{\text{nuc}}(\mathbf{r}_i) + \sum_{i,b} U_{\text{cen}}(\mathbf{r}_i) = 0, \]

where \( \mathbf{r}_i \in \mathbb{R}^3 \) is the coordinate vector of bead \( i \), and \( N \) is the total number of beads in a model. The restraints are expressed as pseudo potential energy terms as described in Table 1; \( \alpha, \beta, \gamma, \) and \( \delta \) are subsets of specific beads in the chromosome chains that share certain properties. More specifically, \( \alpha \) is the set of beads assigned to the last bead of every chain, \( \beta \) is the set of beads assigned to centromeres, \( \gamma \) is the set of beads assigned to telomeres, and \( \delta \) is the set of all beads flanking rDNA repeat regions.

**Chromatin chain restraint** \( U_{\text{ch}} \)

Two consecutive beads in a chromosome chain are restrained to be at a distance of 30 nm (Table 1).

**Chromatin chain excluded volume restraint** \( U_{\text{exc}} \)

Overlap between beads is prevented by imposing excluded volume restraints for all bead pairs (Table 1).

**NE restraint** \( U_{\text{nuc}} \)

All chromatin beads must remain within the nucleus, defined as a sphere with radius \( R_{\text{nuc}} = 1 \) micron (Table 1).

**Centromere localization restraint** \( U_{\text{cen}} \)

All the centromeres cluster near the SPB, which is the microtubule organization center in the yeast nucleus (Jin et al. 2000). The centromeric regions are attached to the SPB through microtubules up to 300 nm in length (O’Toole et al. 1999). Accordingly, all beads representing centromeric regions are restricted to a spherical volume with a radius 300 nm, centered on the SPB (Fig. 1). We follow ex-
rDNA repeat region (i.e., positions 458 kb from the left telomere and 620 kb from the right telomere in the sequence of chromosome 12) to the surface of the nucleolus (Table 1; Fig. 1).

**Nucleolus excluded volume restraint \( U_{\text{nuv}} \)**

All chromosomal regions except those containing rDNA repeats are excluded from the nucleolus. (Table 1).

**Chromatin persistence length**

During the optimization process, we imposed an angular restraint between each set of three consecutive beads to reproduce the desired chain stiffness. The constraint is expressed as a harmonic potential:

\[
U_{\text{angle}} = \frac{1}{2} k_{\text{angle}} \sum_{i=1}^{N/2} \left( \frac{\angle{r_{i+1}} - \angle{r_i}}{\varphi} \right)^2 \times \left( \frac{\angle{r_{i+2}} - \angle{r_{i+1}}}{\varphi} \right)^2
\]

for \( i, i+1 \), and \( i+2 \) on the same chain.

This restraint is considered only when calculating gradient forces during the optimization process. It makes no contribution to the total score of a model (below). With a force constant of \( k_{\text{angle}} = 0.2 \text{ kcal/mol} \), we obtain chromatin chains that behave like random polymers with a persistence length between 47 and 72 nm (the average is \( 61.7 \pm 7.7 \text{ nm} \)) (Supplementary Fig. 7), consistent with experiments. Estimated values for the persistence length from experiments fall between 30 and 220 nm (Cui and Bustamante 2000; Bystricky et al. 2004; Langowski 2006).

**Optimization**

The optimization is performed using a combination of simulated annealing molecular dynamics and the conjugate gradient methods implemented in the Integrated Modeling Platform (IMP; http://www.integrativemodeling.org) (Alber et al. 2007a,b, 2008; Russel et al. 2012). An individual optimization starts with an entirely random bead configuration, followed by an initial optimization of the structure. Next, we apply simulated annealing protocols to entirely equilibrate the genome configuration. Finally, conjugate gradient optimization ensures that all constraints are satisfied, leading to a structure with score zero. Many independent optimizations are carried out to generate a population of 200,000 genome structures with a total score of zero, hence consistent with all input data. A comparison between the frequency maps of two independently calculated populations, each with 100,000 structures, showed that our genome structure population is highly reproducible (Pearson’s correlation between the contact frequency maps of the two populations is 0.999).

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**Figure 7.** Spatial clustering of replication origins and tRNA gene loci. (Left) The histograms show the distribution of the mean pair distance ratio between a set of specific sites (e.g., early replication sites) and all sites in the structures of the population. The histograms are generated as follows: For a given structure in the population, the mean pair distance between a set of specific loci (e.g., all early replication origins) is calculated. This distance is divided by the mean pair distance of all sites in the same structure. The distribution of the distance ratio is then obtained from all structures in the population. If the distribution is centered on 1 (vertical dashed line), the selected sites behave similarly to a random sample of all sites. If the distribution is shifted toward smaller values, their pair distances are smaller than would be expected from the background control. If the distribution is shifted to larger values, the selected sites are more distant from each other than would be expected from the random control background. (Right) Distribution of distance ratios for all 275 tRNAs (loci extracted from SGD, http://www.yeastgenome.org). For all sets of sites in A and B, the shift of the mean pair distances is highly statistical significant (for details, see Supplemental Material and main text).
Control population
We also generated a control population of 25,000 structures without chromosome tethering constraints and nucleolus excluded volume constraints. Otherwise, the chromosomes are constrained in a manner identical to the full simulation.

Analysis
The analysis of the structure population and all statistical tests are described in great detail in the Supplemental Material.

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