High-throughput chromatin motion tracking in living yeast reveals the flexibility of the fiber throughout the genome

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Chromosome higher-order architecture has been increasingly studied over the last decade thanks to technological breakthroughs in imaging and in molecular biology ranging from painting single chromosomes in fixed cells (Cremer and Cremer 2001) to mapping physical interactions between genomic elements at the genome-wide level (Hi-C) (Dekker 2008). It is now established that chromosome large-scale organization plays important roles in all aspects of genomic transactions, allowing contact between distant chromatin loci (Göndör and Ohiłsson 2009) that trigger concomitant genomic processes in cis and in trans. Although several models have been proposed to describe the folding of chromosomes (Lieberman-Aiden et al. 2009; Bohn and Heermann 2010; Bancaud et al. 2012), the physical principles governing this organization are still largely debated.

Despite its small size, budding yeast has become a unique model that recapitulates some of the main features of higher eukaryotes, including heterochromatin-like regions, and an organization mediated by the nuclear envelope (NE), the nucleolus, and the spindle pole body (SPB). Structural data obtained by statistical positioning of a gene in a yeast cell population led to a surprisingly simple model to define yeast nuclear architecture (Zimmer and Fabre 2011): Chromosome position can be predicted by a few parameters such as genomic arm length, telomeres (TEL), and centromeres (CEN) tethered to the NE via nuclear-envelope-tethered proteins and to the SPB via microtubules, respectively (Bystricky et al. 2005; Therizols et al. 2010; Zimmer and Fabre 2011). This description was recently complemented by the first Hi-C comprehensive maps (Rodley et al. 2009; Duan et al. 2010), which confirmed an organization guided by nuclear landmarks, including TEL that congregate in foci (Gotta et al. 1996; Schober et al. 2008). Recent Brownian dynamics (BD) simulations confirmed this structural model by recapitulating Hi-C and imaging data, assuming that physical tethering at TEL and CEN and volume exclusion were driving chromosome conformations (Tjong et al. 2012; Wong et al. 2012).

The large spatial fluctuations of chromosome loci inferred from particle tracking in living yeast suggested a highly dynamic character of chromosome architecture at the molecular level (Marshall et al. 1997; Heun et al. 2001; Hihara et al. 2012). Chromatin dynamics appeared to be determined by nuclear constraints, in particular, the NE (Heun et al. 2001; Bystricky et al. 2004), and by the position along the chromosome, e.g., TEL (Bystricky et al. 2005). Furthermore, a large body of data provides information on spatio-temporal dynamics in wild-type versus mutant cells (see, e.g., Bystricky et al. 2009) and elucidates how chromatin properties are regulated by structural proteins. However, the quantitative...
analysis of these movements remains controversial, and essentially two models have been used: (1) Chromatin segments were suggested to undergo normal Brownian fluctuations at small time scales and to be confined in regions of $-0.3 \mu m^2$ at time scales longer than $-100$ sec (Marshall et al. 1997); or (2) the movement of GAL1-10 genes on chromosome II appeared to follow an anomalous behavior characterized by a diffusion coefficient of $-0.4$ (Cabal et al. 2006). This anomalous behavior was also detected in bacteria (Weber et al. 2010) and thoroughly studied using polymer models. In yeast, the narrow temporal range of measurements within less than two decades was insufficient for a truly quantitative analysis of chromatin dynamics.

Here we develop high-speed live cell imaging and high-throughput image analysis techniques to monitor the spatio-temporal fluctuations over an extended temporal range spanning more than four decades ($10^{-2} - 10^5$ sec) of nine chromosome loci on chromosomes III, IV, XII, and XIV of the yeast Saccharomyces cerevisiae. We show that chromatin segments move subdiffusively and that the characteristics of the anomalous response are largely conserved for every locus distant by more than $-50$ kb from TEL and CEN. Similar dynamic behavior was also observed for two telomeric loci whenever they adopt a central localization in the nucleus, thus leading to a consistent picture for chromosome movement within the entire nuclear volume. We then demonstrate that these dynamics are consistent with the Rouse model using BD simulations and statistical analysis of the trajectories and suggest that chromatin is a flexible polymer in yeast, characterized by a persistence length of $<30$ nm. Finally, we show that the Rouse model is also relevant in mutants for proteins involved in chromatin structure, and propose that this model describes a consistent framework to study chromosome motion. Our findings significantly advance the quantitative understanding of chromosome dynamics and their implications for yeast genome architecture.

Results and Discussion

The dynamics of chromosome loci are uniform throughout the nucleus and anomalous over a broad temporal range

We assayed the dynamics of chromatin in a collection of yeast strains each bearing a single fluorescently labeled chromatin locus (see Methods for details). The labels were inserted on chromosomes III, IV, XII, and XIV at loci distant from TEL and CEN (Figs. 1, 2A). The chromosomes tested ranged in size from 300 kb to 1500 kb, which is well suited to assess whether the dynamics are affected by chromosome length. Motion was analyzed over a temporal range spanning more than four decades (15 msec to 400 sec) (Fig. 2B). This broad temporal analysis is more than two orders of magnitude larger than in previous studies performed in living yeast (Heun et al. 2001; Bystricky et al. 2004; Cabal et al. 2006) or in bacteria (Weber et al. 2010), and investigations on similar time domains were only conducted in mammalian cells for telomere dynamics (Bronstein et al. 2009). The motion of chromosome loci was recorded using a bright-field microscope and analyzed using a customized high-throughput tracking software based on the Multiple-Target Tracing algorithm (Supplemental Fig. S1; Serge et al. 2008; Albert et al. 2013). Because physical tethering of chromosomes may occur at the nuclear periphery (Heun et al. 2001; Hediger et al. 2002), the nucleus was divided into two regions of equal surfaces, and every tracked locus was automatically assigned to a central or peripheral localization based on the segmentation of the first image of the acquisitions (see image in Fig. 1).

We first focused on loci located in the nuclear center, far from constraints imposed by the nuclear membrane, and representing $-60\%$ of the observed positions (Fig. 2A). We then characterized their motion by plotting the temporal evolution of the mean square displacement (MSD), which describes the average displacement after a given time lag $\tau$:

$$MSD(\tau) = \langle (\vec{r}(t + \tau) - \vec{r}(t))^2 \rangle,$$

with $\vec{r}$ the position vector, and $\langle \rangle$ denoting the temporal average. We observed similar responses for every locus (data points in Fig. 2B), which appeared to fold onto a single master curve (solid line). This curve follows an anomalous diffusive behavior over a broad temporal range characterized by a power-law scaling response in $\tau$ with $\alpha = 0.52 \pm 0.08$ ($\alpha = 1$ for normal diffusion). This result is consistent with an earlier study describing the motion of GAL1-10 genes, which exhibited a nonlinear MSD response characterized by an exponent of $-0.4$ in the temporal range of $4-80$ sec (Cabal et al. 2006), but it departs from other contributions showing normal diffusion and a plateau in the long time limit (see, e.g., Heun et al. 2001). The anomalous subdiffusion response is associated with a slow increase in MSD over time. Its persistence over an extended period indicates that chromatin loci explore a broad region of the nucleus of $\sqrt{0.2 \mu m^2} \sim 450$ nm in $\sim 400$ sec. This estimate is consistent with the extent of “gene territories” of $\sim 500$ nm, which are obtained by statistical mapping of the position of loci in cell populations of $\sim 2000$ individuals (Berger et al. 2008).
Next we studied the motion of chromosome loci located at the periphery and observed that they frequently oscillated between the nuclear center and periphery during the time course of the acquisition (Supplemental Fig. S1), suggesting that the localization classification based on a single image is somewhat artificial. We did not detect significant differences in the MSD response for peripheral or central localization for the three loci on chromosome XII (see, e.g., position 240 kb in Fig. 2C) and on chromosomes IV and XIV (data not shown), so the anomalous diffusive response appears to be largely conserved and appropriate to describe chromosome segmental dynamics for loci distant by more than ~50 kb from TEL and CEN.

The dynamics of telomeres are anomalous in a central localization

In yeast, the 32 TEL tend to cluster near the NE, and the motion of tagged chromosome sites in the vicinity of a TEL is constrained (Heun et al. 2001; Sage et al. 2005). Using our reference behavior for chromosome loci located halfway along chromosome arms, we set out to perform an MSD analysis on TEL 3R and 14L (Fig. 1; hereafter denoted Tel3R and Tel14L). Although TEL predominantly accumulate near the periphery (Fig. 3A, right panel), their positioning in the nuclear center can be detected with an occurrence of ~30%, and we analyzed their dynamics in this central region. The amplitude of spatial fluctuations in the short time regime appeared to be relatively similar compared with the reference response (Fig. 3A), although Tel3R appeared to be slightly less mobile after ~10 sec than Tel14L, presumably because telomeric interactions between the two ends of chromosome XIV are much less frequent than that of chromosome III (Bystricky et al. 2005; Duan et al. 2010).

In contrast to the bulk of chromosomes, telomere dynamics appeared to be reduced at the nuclear periphery (Fig. 3B), in agreement with the idea that telomere peripheral localization is mediated by protein interactions involving, among others, Sir4 and Yku70/Yku80 (Hediger et al. 2002; Taddei et al. 2004; Bupp et al. 2007; Schober et al. 2009). It has been shown that deletion of either YKU70 or SIR4 leads to delocalization of some TEL to the nuclear center (Laroche et al. 1998; Hediger et al. 2002), but it remains unclear whether this deletion affects TEL dynamics aside from chromosome III

Figure 2. Chromatin dynamics in living budding yeast. (A, upper panel) Fluorescence micrographs of different yeast strains for interframe intervals of 200 msec (for interframe intervals of 16–50 msec and mutants, see Supplemental Fig. S2); (lower plot) the spatial distribution of these loci in the nuclear volume. (B) The temporal evolution of the MSD for seven different chromatin loci with central localization is plotted over a broad temporal range covering more than four orders of magnitude. These data show a universal behavior characterized by an anomalous diffusive response associated to power low scaling comprised between $t^{0.44}$ and $t^{0.60}$ (thin dashed lines). (Black solid line) The Rouse regime, which is associated with an exponent of 0.5 (see details in Fig. 4). Note that each data set is the average over 30–200 trajectories and that we selected tracks with signal-to-noise ratios (SNR) >30 dB. (C) The MSD response for loci with central or peripheral localization is compared (yellow and green data sets, respectively), showing similar dynamics. (Black solid line) The behavior measured in B. The departure of the two curves in the short time regime is associated with the lower SNR of peripheral loci (data not shown).
(Bystricky et al. 2009). We thus investigated the movements of Tel14L in wild-type and in SIR4 mutant strains. Deletion of SIR4 leads to a predominant localization of Tel14L in the center (62%), likely due to loss of anchoring to the nuclear membrane. Moreover, telomere mobility is increased in SIR4 mutant strains when the tagged locus resides at the periphery and even more drastically at the nuclear center (Fig. 3B), confirming that physical interactions of the extremities of chromosomes are an essential player of their spatio-temporal dynamics. Overall, this study tends to support the anchoring regions.

Chromosomal anchoring is provided that they are sufficiently distant from chromatin loci, giving rise to the "Rouse" regime (Rouse 1953), has been extensively documented for concentrated solutions of synthetic polymers (see, e.g., Ewen and Richter 1997) and was recently validated in vivo for the bacterial chromosome (Weber et al. 2010).

Chromosomes are modeled by a Rouse chain, which is composed of a series of N elastic segments of length b. The motion of each bond is determined by its elasticity defined by the stiffness $3k_BT/b^2$ with $k_BT$ the thermal energy, and its viscous drag $\zeta$. Assuming that chromosomes are tethered at CEN and TEL, the MSD of monomers can be solved analytically in the short time regime (see Model section):

$$\text{MSD}(\tau) \sim \left[ \frac{12k_BT}{\pi \zeta} \right]^{1/2} \tau^{1/2}.$$

The Rouse segmental dynamics are characterized by an anomalous diffusive response uniform along the chain, in good agreement with our data. We wished to test the predictions of the Rouse model further by extracting the histogram of the displacement after a time lag of 0.18, 1.0, and 1.6 sec from the trajectories of locus 240 on chromosome XII (Fig. 4A). The histograms were first fitted with a normal diffusion model (Fig. 4A, left panel and equation in inset), which relies on one fitting parameter, the diffusion coefficient $D$. The diffusion coefficient of 4.8, 2.2, 1.8 $\times 10^{-3} \mu m^2/sec$ was very different for $\tau = 0.18, 1.0,$ and 1.6 sec, respectively, showing the inconsistency of this model. We then used the formalism described in Guerin et al. (2012) to compute the histogram of the displacement for a Rouse chain tethered at its extremities (see expression in Model section). One single parameter was sufficient to reproduce the three distributions (Fig. 4A, right panel), and the quality of the fitting was significantly improved (note the departure of the red fit from the conventional model), hence strongly supporting the relevance of the Rouse model to describe the segmental dynamics of yeast chromosomes.

We then performed BD simulations of yeast chromosome dynamics in a volume the geometry of which is consistent with yeast nuclear size, and using the structure derived from Hi-C experiments as the initial state (Duan et al. 2010). We assumed that chromosomes were attached to the SPB at their CEN (i.e., neglecting the contribution of microtubules connecting SPB to kinetochores), and freely diffusing at their TEL (chromosome III is represented in the left panel of Fig. 4B). The computed MSD for loci distant from CEN (>400 nm; blue data set in the right panel of Fig. 4B) reproduced the anomalous diffusive response associated with a power-law scaling of $-0.54$ over a temporal domain of four orders of magnitude that we observed experimentally (dashed line in Fig. 4B, right panel). The motion is more restricted for segments located at 10–100 nm from the CEN (pink data set) versus at a distance comprised between 200 and 400 nm (red data set). In addition, su-
that broken ends of chromosomes are free to search the entire genome for appropriate partners (Haber and Leung 1996), or partial as telomere colocalization and coordinated movement would indicate (Bystricky et al. 2005). Future BD simulation studies should therefore evaluate whether and how compact exploration is related to chromosome territoriality in the yeast genome.

**MSD fluctuations suggest that chromatin is highly flexible**

The level of flexibility of a chromatin fiber is a matter of debate. In vitro single-molecule experiments have shown that the persistence length \( \ell_p \) of nucleosome arrays is 20–30 nm (Cui and Bustamante 2000; Bancaud et al. 2006; Celledon et al. 2009; Kruithof et al. 2009). Conversely, analyses based on polymer models of in vivo distance measurements or Hi-C experiments (Bystricky et al. 2004; Dekker 2008; Tjong et al. 2012) provided estimates of \(~200\) nm and 66–134 nm, respectively. Physical modeling of the mechanical properties of nucleosome arrays predicts a persistence length smaller than 30 nm for chromatin (Ben-Haim et al. 2001; Schiessel et al. 2001), except in the case of highly ordered arrays with strong nucleosome–nucleosome interactions (Wedemann and Langowski 2002), in which case \( \ell_p \) increases to \(~250\) nm.

The amplitude of spatial fluctuations can be qualitatively linked to the flexibility of chromatin using the Rouse model, given that the bond length \( b \) and the viscous drag coefficient \( \zeta \) are equal to \( 2b \) and \( 6m_0 \eta b \), respectively (de Gennes 1979). Injecting this term in Equation 2 implies that the amplitude of the MSD increases with the square root of the polymer persistence length. The extent of spatio-temporal fluctuations therefore increases as the rigidity of the fiber increases, in qualitative opposition to free diffusion of an isolated tracer that increases as the diameter of the particle decreases. The MSD response can then be predicted for a persistence length comprised between 50 and 200 nm, given that the nuclear viscosity is \( \eta \sim 7 \times 10^{-3} \) Pa.s based on GFP diffusion coefficient measurements (Kawai-Noma et al. 2006; Slaughter et al. 2007). This conjecture exceeds our measurements by a factor of 10–20 (dashed lines in Fig. 5A). In contrast, fitting of our data with the Rouse model leads to an amplitude of the MSD of \( 1.0 \pm 0.2 \times 10^{-2} \) \( \mu m^2/sec^{1/2} \) (solid line in Fig. 5A), suggesting that the persistence length of chromatin in living yeast is \(~5\) nm. This length scale is coherent with the dimensions of a single nucleosome, which is a \(~5\)-nm by \(~10\)-nm disk-like structure. Hence, the analysis of chromosome motion suggests that chromatin is a flexible polymer in vivo.

Because the Rouse model is a simplified model that does not take volume exclusion between monomers into account,
we wished to strengthen our conclusions by running BD of a single chain composed of self-avoiding segments (see Model section). The extracted MSD shows an anomalous diffusive response, as well as the dependence on the persistence length and the viscous drag (Eq. 2) expected from the Rouse model (Fig. 5B). This analysis showed that the numerical prefactor of Equation 2 is underestimated by 1.4, suggesting that volume exclusion appears as a flexible polymer, and its persistence length can be qualitatively estimated to be less than 30 nm.

Chromatin structural proteins alter chromosome mobility

Finally, we investigated whether the dynamics of chromosomes was altered in the absence of chromatin structural proteins or chromatin assembly proteins. We first considered the HMG-like protein Spt2 in S. cerevisiae, which interacts with the histones H3 and H4 (Horn et al. 2002). The statics and the dynamics of one locus on chromosome XIV were assayed in wild-type (WT) and mutant spt2 cells, showing an accumulation at the periphery for the mutant and similar MSD responses in the short time regime (Fig. 6, right and left panels, respectively). We then performed the same analysis with Hho1p (Levy et al. 2008), which is considered as the linker histone analog in S. cerevisiae, although its structure differs from the linker histone in other organisms because it bears two globular domains. So far it remains unclear whether and how this protein participates in chromatin condensation. The deletion of HHO1 increases the proportion of spots detected in the nuclear center compared with wild type, and increases ~1.5-fold the amplitude of chromatin loci displacements (Fig. 6). Given that the ratio Hho1p versus nucleosome remains poorly estimated (compared between 1:37 and 1:4) (Freidkin and Katcoff 2001; Downs et al. 2003), these enhanced dynamics may be associated with the reduced frequency of chromosome interactions mediated by the two domains of Hho1p and/or with an
onset in flexibility due to conformational changes at the nucleosome level. Finally, we probed the role of the nucleosome assembly chaperone Asf1p, which is involved in replication-dependent and replication-independent chromatin assembly (Loyola and Almouzni 2004). As for hho1 deletion mutant, the labeled locus appeared to be more frequently central than in wild-type nuclei, and its mobility was 2.2-fold increased. The Rouse model therefore seems to account for the dynamics of chromosome loci in WT cells, as well as in hho1, asf1, and spt2 mutants, suggesting that this approach is sufficiently generic to perform systematic motion analyses.

Conclusion
We demonstrate that the motion of chromosome loci in living yeast is characterized by anomalous subdiffusion with an exponent of 0.52 over a temporal domain spanning more than four orders of magnitude. We then show that this response, which is relevant for the analysis of WT and mutant cells, is consistent with the Rouse dynamics. Because the Rouse regime is characteristic of polymers in crowded environments, our result strengthens the relevance of polymer models for the description of chromosome interactions in cis and trans, and envision that this hypothesis can be tested using BD simulations in combination with Hi-C experiments aiming to evaluate the dynamics of loop formation may interfere with the segmental motion of human chromosomes, a hypothesis that remains to be tested quantitatively in vivo.

The folding of interphase chromosomes in metazoans involves loops of variable sizes defining topological domains of 0.2–5 Mb (Nora et al. 2012; Sexton et al. 2012). Because the dimension of these domains, which can be described as end-tethered chromosome fragments, is consistent with the length of yeast chromosome arms, we speculate that the Rouse model is relevant to higher-order eukaryotes. This model has recently been implemented in numerical simulations (Bohn and Heermann 2010), showing that the dynamics of loop formation may interfere with the segmental motion of human chromosomes, a hypothesis that remains to be tested quantitatively in vivo.

Model
Dynamics
The motion of an end-tethered Rouse chain can be studied analytically using normal mode decomposition (Vandoolaeghe and Terentjev 2007), and the MSD of the nth segment (MSD$_n$) reads:

$$\text{MSD}_n(t) = \frac{4N\xi^2}{\pi^2} \sum_{p=1}^{\infty} \frac{1}{p^2} \left(1 - e^{-t/\tau_p}\right) \sin^2\left(\frac{p\pi}{N}\right),$$

with $\tau_p$ the relaxation time of each mode $\tau_p = \tau_0/p^2$ and $\tau_0 = L^2\xi/3\pi^2k_BT$ the Rouse time identified as the longest relaxation time of the chain. Note that this equation is valid in three dimensions (3D), and its two-dimensional (2D) version is readily obtained with a multiplication by 2/3, assuming that the motion is isotropic. The temporal evolution of the MSD is plotted in Supplemental Figure S4As. In the short time limit (t $\ll$ $\tau_0$), Equation 3 is dominated by the terms with large $p$, and it can be simplified to obtain Equation 2. In addition, the formalism described in Guérin et al. (2012) (see details in the Supplemental Material) enables us to extract the distribution of the displacement $R$ for the nth segment after a time lag $\tau$:

$$P_n(R, \tau) = \frac{R}{\Psi_0(\tau)} \exp\left(-\frac{R^2}{2\Psi_0(\tau)}\right)$$

$$\Psi_0(\tau) = \frac{2k_BT}{N^2} \sum_{p=1}^{N} \sin^2\left(\frac{p\pi}{N} \right) \left[1 - \exp\left(-\frac{2\xi^2}{k_BT} \left(1 - \cos\left(\frac{p\pi}{N+1}\right)\right)\right)\right] \left[1 - \cos\left(\frac{\pi}{N+1}\right)\right].$$

The movements of chromosomes, which were modeled by a series of bead linked by springs or ball-in-socket joints, was also analyzed by BD. The size of each bead was set to 15 nm in diameter, and it was assumed to contain approximately three nucleosomes, so that the full genome consists of 27,000 beads. Chromosomes were constrained in a sphere of 2 $\mu$m in diameter,
and the CEN were fixed to the SPB using harmonic constraints. We used the Langevin dynamics algorithm implemented in the NAMD software (Phillips et al. 2005) with an appropriate in-house force field. The Langevin equation for the nth monomer has the form:

\[ m_n \frac{d^2 x_n}{dt^2}(t) = -\text{grad}_x E + f_n(t) - m_n b_n \frac{dx_n}{dt}(t), \]

where the random force \( f_n(t) \) is derived from a Gaussian distribution with the properties:

\[ \langle f_n(t) \rangle = 0 \]

\[ \langle f_n(t)f_n(0) \rangle = 2k_b T m_n \delta(t), \]

with \( \zeta \) the friction coefficient. The energy \( E \) of each bead is the sum of three contributions associated with the confinement in the nucleus, the elasticity of the chain, and the repulsive interactions between monomers, which are expressed as:

\[ E = E_{\text{conf}} + E_{\text{spring}} + E_{\text{diss}} \]

\[ E_{\text{conf}}(r < r_1) = 0 \quad \text{and} \quad E_{\text{conf}}(r > r_1) = E_1(r-r_1)^4, \]

with \( r \) the distance of the bead to the center of the nucleus, and \( r_1 \) the radius of the nucleus.

\[ E_{\text{spring}}(n, n + 1) = k_s (r_n - r_{n+1})^2 \]

\[ E_{\text{diss}}(i, j) = U \left\{ \left( \frac{R_{\text{min}}}{|r_i - r_j|} \right)^{12} - 2 \left( \frac{R_{\text{min}}}{|r_i - r_j|} \right)^6 \right\} \]

The parameters of this model can be found in Table 1. BD were run during \( 3 \times 10^7 \) time steps, starting from the conformation obtained by Hi-C (Duan et al. 2010) equilibrated during \( 10^6 \) time steps. We also performed BD to investigate whether spatio-temporal fluctuations were different for 1, 4, or 10 segments, or for one segment of large diameter along the chain. We did not detect differences on the MSD response (data not shown), in agreement with our observation of similar dynamics with Lac or Tet labeling, which measure \(-10\) and \(7\) kb in length, respectively.

**Methods**

**Plasmids and strains**

All the strains we used are described in Supplemental Table 1.

**Table 1.** Set of parameters to run Brownian dynamics simulations

| \( T \) | \( 300 \text{ K} \) |
|---|---|
| \( k_s \) | 1 kcal mol\(^{-1}\) nm\(^{-2}\) |
| \( R_{\text{min}} \) | 7 nm |
| \( U \) | \(-0.01\) kcal mol\(^{-1}\) |
| \( m_n \) | 10 a.u. |
| \( b_n \) | 10 a.u. |
| \( E_1 \) | 0.01 kcal mol\(^{-1}\) nm\(^{-4}\) |

**Cell culture**

Cells were grown in YPD or YNB at 30°C starting from a fresh overnight culture. They were diluted at \(10^6\) cells/mL, and harvested when OD\(_{600}\) reached \(4 \times 10^6\) cells/mL and rinsed twice with the corresponding SC media. Cells were then spread on a SC media patch containing a 2% agarose and 2% glucose coated slide. A coverslip was deposited over the cells, and the construction was finally sealed with “VaLaP” (one-third vaseline, one-third lanoline, one-third paraffin). Live microscopy was limited to 20 min after mounting the coverslip. Extreme care was taken to ensure that cells are at approximately equal stages of growth based on their shapes, as inferred from a single image of transmission light microscopy, and based on the nucleus shape using the roundness parameter \( R \), which was defined as:

\[ R = 4\pi \times \text{Surface/Perimeter}^2, \]

with \( \text{Surface} \) the number of pixels after segmentation of the nucleus, and \( \text{Perimeter} \) the outline in pixels (see Supplemental Fig. S2). We considered nuclei with \( R > 0.8 \) to avoid mitotic phenotypes.

**Live cell imaging**

Yeast cells were imaged at room temperature with a BX-51 upright microscope equipped with a laser diode (Lumencor), a 100× oil immersion objective (NA = 1.4), and an EMCCD camera (Andor DU-897), as described in Hajjoul et al. (2009) (see Supplemental Video 1). The excitation emission at 470 ± 10 nm was set to 7.53 W/μm\(^2\), about 100 times less than the value that is necessary for normal cell growth (Carlton et al. 2010), and we observed consecutive rounds of divisions during ~2 h (data not shown). For slow acquisitions of interframe intervals of 2 sec or more, we used a 100× oil immersion objective with a variable NA set to ~1.1–1.2. The acquisition was performed with a shutter synchronized with the camera to minimize photobleaching, and we checked for drifts by monitoring the position of the center of the nucleus in the first and last image of the recording. Optimal interframe intervals of ~20 msec were reached by cropping regions of interest and using a 2 × 2 binning with the Andor iQ imaging software. The loss in pixel size resolution was compensated with a 2× lens placed in the microscope light path.

We focused on 2D single-particle tracking and checked that similar information was retrieved with 3D tracking using microscopes (Supplemental Fig. S3; Hajjoul et al. 2009, 2011). We note that 2D acquisitions may bias MSD responses for loci preferentially localized at the periphery, because in some cases central loci in the projected view may be peripheral in the 3D nucleus.

**Data analysis**

Video sequences consisted of 300 consecutive images, and we displayed MSD traces on 150 time intervals at most in order to ensure the statistical relevance of mean displacements. The movements of chromosome loci were systematically analyzed using a custom software that was developed in Matlab (Mathworks) (Serge et al. 2008; Albert et al. 2013 [source code at http://jcb.rupress.org/content/202/2/201/suppl/DC1, and the executable file at ftp://intermt:MTTinterface@ftp.laas.fr/]). This software enabled us to extract \((x, y)\) coordinates by Gaussian fitting, to reconstruct the trajectories, and to compute the MSD and the step distribution function (see Supplemental Video 1). Note that the MSD in fixed cells was 20- to 30-fold lower than in living cells (Supplemental Fig. S3B).
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