Chromatin Dynamics in Interphase Nuclei and Its Implications for Nuclear Structure

James R. Abney,‡ Bryan Cutler,* Misty L. Fillbach,§ Daniel Axelrod,† and Bethe A. Scalettar*  
*Department of Physics, ‡Northwestern School of Law, and §Department of Mathematics, Lewis & Clark College, Portland, Oregon 97219; and †Biophysics Research Division and Department of Physics, University of Michigan, Ann Arbor, Michigan 48109

Abstract. Translational dynamics of chromatin in interphase nuclei of living Swiss 3T3 and HeLa cells was studied using fluorescence microscopy and fluorescence recovery after photobleaching. Chromatin was fluorescently labeled using dihydroethidium, a membrane-permeant derivative of ethidium bromide. After labeling, a laser was used to bleach small (~0.4 μm radius) spots in the heterochromatin and euchromatin of cells of both types. These spots were observed to persist for >1 h, implying that interphase chromatin is immobile over distance scales ≥0.4 μm. Over very short times (<1 s), a partial fluorescence recovery within the spots was observed. This partial recovery is attributed to independent dye motion, based on comparison with results obtained using ethidium homodimer-1, which binds essentially irreversibly to nucleic acids. The immobility observed here is consistent with chromosome confinement to domains in interphase nuclei. This immobility may reflect motion-impeding steric interactions that arise in the highly concentrated nuclear milieu or outright attachment of the chromatin to underlying nuclear substructures, such as nucleoli, the nuclear lamina, or the nuclear matrix.

In recent years, models of the structure and organization of the interphase nucleus have changed dramatically. The interphase nucleus was once believed to be a largely homogeneous organelle, containing little structural specialization beyond the nucleoli (Manuelidis, 1990; Berezney et al., 1995). In contrast, the interphase nucleus is now widely believed to be inhomogeneous, with many nuclear substructures and functions localized to specific nuclear subdomains (for review see van Driel et al., 1995). In particular, a considerable body of evidence suggests that the most prominent nuclear constituent, chromatin, is organized so that each chromosome occupies its own discrete domain (Cremer et al., 1982; Lichter et al., 1988; Leitch et al., 1990; for review see van Driel et al., 1995; Eils et al., 1996).

Considerably less is known about the mobility of chromatin and other constituents of interphase nuclei than is known about their organization. Despite being poorly understood, mobility has been the subject of interest for at least two reasons. First, the mobility of nuclear constituents should reflect and influence important aspects of their organization. Indeed, in the case of interphase chromatin, mobility has been most commonly inferred indirectly from observed changes in organization and distribution (Manuelidis, 1985; Bartholdi, 1991; Ferguson and Ward, 1992; Janevski et al., 1995). Second, the mobility of nuclear constituents should reflect their function. In the case of interphase chromatin, a functional significance for mobility is strongly suggested by recent observations of changes in centromere and chromosome distribution in response to cell differentiation, transcription signals, and stage of the cell cycle (Manuelidis, 1984; Bartholdi, 1991; Ferguson and Ward, 1992; Funabiki et al., 1993; Janevski et al., 1995; Pluta et al., 1995; LaSalle and Lalande, 1996).

To date, data bearing on the mobility of chromatin in interphase nuclei are not only limited, they are also somewhat contradictory. For example, some studies suggest that chromatin is relatively immobile during interphase. Notable examples include observations of nonrandom organization of chromosomal substructures and of chromosome confinement to domains in fixed interphase nuclei (Moroi et al., 1981; Lichter et al., 1988; Manuelidis and Borden, 1988; Dyer et al., 1989; Leitch et al., 1990; Popp et al., 1990; for review see van Driel et al., 1995). Moreover, in living HeLa cells, centromeres are generally motionless during interphase (Shelby et al., 1996), and in living Drosophila embryos, chromosomes undergo late decondensation and subsequent condensation at the same sites on the nuclear envelope (Hiraoka et al., 1989). Finally, photobleaching studies of chromatin in isolated interphase nuclei indicate that chromatin reorientational mobility is highly restricted (Selvin et al., 1990). In contrast, some studies have shown that chromatin can reposition during interphase; notable
examples include the three-dimensional movement of heterochromatin in nuclei of living interphase neurons (De Boni and Mintz, 1986), the coalescing and dispersing of centromeres in cultured cells in late G, and early G, respectively (Manuelidis, 1985), and the occasional slow movement of a centromere in living HeLa cells (Shelby et al., 1996). Data display on the mobility of nonchromosomal molecules and other objects in interphase nuclei are also somewhat limited and contradictory. In situ hybridization studies have shown that RNA molecules may move nonrandomly along “tracks” in the nucleus, perhaps because their motion is constrained by an underlying nuclear matrix (Lawrence et al., 1989). Studies of the trajectories of large, naturally occurring cytoplasmic inclusions have shown that the mobility of inclusion-sized objects in the nuclei of newt pneumocytes is several hundredfold slower than in dilute solution (Alexander and Rieder, 1991). In contrast, translational FRAP experiments on fluorescently labeled dextrans (3–150 kD) have shown that translation of dextran in the nuclei of Hepatoma cells is only about sevenfold slower than in dilute solution (Lang et al., 1986). These rather disparate results leave considerable uncertainty surrounding the rates and determinants of motion in the nucleus.

In this study, we have directly monitored the translational motion of fluorescently labeled chromatin in the nuclei of living Swiss 3T3 and HeLa cells using FRAP. These cell lines were chosen because microscopy studies of nuclear organization have frequently employed cultured cells, including Swiss 3T3 and HeLa, generating a considerable body of germane literature (e.g., Fey et al., 1986; He et al., 1990; Bereznay et al., 1995; Shelby et al., 1996). Our results show that interphase euchromatin and heterochromatin are substantially immobile in these cell lines. This immobility is consistent with chromatin attachment to nuclear substructures and with chromosome confinement to discrete domains during interphase. Moreover, this immobility provides a dynamics-based foundation for the many observations of nonrandom organization of chromosomal substructures in fixed interphase cells.

Materials and Methods

Cell Culture and Chromatin Labeling

Swiss 3T3 fibroblasts and HeLa cells were grown on 25-mm-diam round glass No. 1 coverslips (Fisher Scientific Co., Pittsburgh, PA) in Dulbecco’s modified Eagle medium (low glucose) (GIBCO BRL, Gaithersburg, MD) supplemented with 10% fetal calf serum. Cells were maintained at 37°C in a 10% CO/90% air incubator and used before cells had reached confluence. Chromatin in living cells was labeled with ethidium by a 10–45 min incubation at 37°C with 2.5–10 M calcein acetoxymethyl (AM) ester (Molecular Probes, Inc.), a “premier indicator of cell viability” (Haugland, 1996). The latter molecule is nonfluorescent and membrane permeant but is converted into the green fluorescent molecule calcein by intracellular esterases in viable cells. After conversion, calcein is retained by cells only so long as their plasma membranes are not compromised. Thus, green cells are viable cells, and cell death can be assayed by monitoring for a loss of cytoplasmic green fluorescence. This was done visually and electronically to demonstrate short-term viability during photobleaching experiments. In addition, to demonstrate that calcein is not retained after cells are compromised, calcein-labeled cells were placed in buffer containing 0.1% Triton X-100 (Bio Rad Laboratories, Hercules, CA) and loss of cytoplasmic green fluorescence was then monitored.

Second, long-term viability and absence of DNA damage were demonstrated by verifying that cells were able to divide and proliferate after they were photobleached. To facilitate identification of cells over long periods of time, cells were grown on coverslips that had been scratched with a diamond pen. Cells were labeled with dihydroethidium and calcein AM well before they had reached confluence and were then transferred to a sterile Sykes-Moore chamber for photobleaching experiments. Cells in an easily identified region of the scratch (such as the end) were bleached and returned to the incubator after ~1 h. Cells were then examined frequently under a dissecting microscope; individual bleached cells could thus be tracked at low cell density even if they moved slowly. Division and proliferation were monitored for ~72 h after bleaching.

Antibody Samples

As a FRAP control, samples containing a fluorescently labeled macromolecule undergoing Brownian diffusion were prepared by diluting a rhodamine-labeled goat anti-rabbit antibody into Hanks’ buffer.

FRAP Experiments

Chromatin dynamics was monitored using FRAP (Axelrod et al., 1976). In a FRAP experiment, a brief, intense pulse of laser light is used to bleach (render nonfluorescent) many of the fluorescent molecules in a small subregion of a fluorescent sample. The return of fluorescence to this subregion is then monitored by shining much attenuated laser light onto the bleached spot and monitoring the temporal dependence of the post-bleach (probe) fluorescence. The time-scale over which the fluorescence recovers is determined by the rate at which the fluorescently labeled molecules diffuse and can be used in simple systems to determine a diffusion coefficient. Failure of the fluorescence to recover indicates that the fluorescently labeled molecules are immobile.

The FRAP apparatus used in these experiments has been described previously (Yuan and Axelrod, 1995). Briefly, its major components are (a) an argon-ion laser, which is used as a high-intensity source of light; (b) an acousto–optic modulator, which is used to modulate the intensity of the light incident on the sample; (c) an inverted microscope, which is used to focus the light onto the sample and to collect the fluorescence emitted by the sample; and (d) an avalanche photodiode (“single photon counting module” [SPCM-100]; EG&G Optoelectronics, Vaudrevil, Quebec, Canada), which is used to detect the fluorescence. An aperture diaphragm placed in the image plane of the microscope minimizes detection of background and out-of-focus light. Data were collected on a 486-based PC using custom FORTRAN software and a custom-programmed counter/timer board (Gibbs & Keithley Metrabyte Instruments Inc., Cleveland, OH).

Light was focused onto the sample using a 63× (NA = 1.4) Zeiss objective (Carl Zeiss Inc., Thornwood, NY), producing an in-focus spot of radius ~0.4 μm in the sample plane. The 514-nm (green) line of the argon-ion laser was used to excite fluorescence; excitation of ethidium with visible light, as contrasted with ultraviolet radiation, reduces the probability of radiation-induced damage to the cells. The bleaching power at the sample was typically in the range ~4–60 mW. The ratio of the intensities of the bleach and probe beams was ~10,000:1, ensuring insignificant bleaching during the probe phase of the experiment.

Photography

Photographs of cells were taken at defined times before and after photobleaching using Kodak TMX 400 film. Cells were illuminated for photo...
Data Analysis

Diffusion coefficients and mobile fractions were obtained by fitting photobleaching data to an equation describing photobleaching recovery in a system containing one immobile fraction and one mobile fraction diffusing in two dimensions (Axelrod et al., 1976). Complex generalizations of this equation describing diffusion in three dimensions have recently been derived (Blonk et al., 1993). These generalized equations show that the recovery time along each dimension is proportional to the square of the characteristic distance along that dimension. This implies that the fastest recovery time will be dominated by diffusion along the smallest dimension. For a focused spot, the smallest dimension is the characteristic e^-2 bleached radius of the spot (0.4 μm here), rather than the corresponding e^-2 axial distance over which substantial bleaching occurs (2.5 μm here). Thus, for the focused spot used in these experiments, the simpler photobleaching recovery equations describing diffusion in two dimensions can be used. In addition, our conclusion that a large fraction of interphase chromatin is immobile (diffusion coefficient D < 10^{-13} cm^2/s) is a model-independent result, which is also demonstrated visually (see Results).

Results

Fig. 1 shows a series of photographs of two dihydroethidium-labeled Swiss 3T3 cells taken (a) before; (b) a few min after; and (c) ~60 min after the nucleus in the upper cell was exposed to a brief pulse of intense laser light. A several hundred millisecond bleaching pulse was sufficient to bleach the ethidium molecules in the illuminated region and create a dark spot in the euchromatin (Fig. 1, b and c). The radius of the spot was calculated to be ~0.6 μm, using known magnification parameters and the size of the photograph. This spot persisted for at least 1 h (Fig. 1 c), indicating that a significant fraction of the interphase euchromatin in Swiss 3T3 cells is translationally immobile on the 0.6 μm distance scale. (This distance scale can be lowered to ~0.4 μm; see below.) Moreover, the spot did not move within the nucleus, providing further evidence for chromatin immobilization during interphase. Similar results were observed after photobleaching HeLa cells and cells maintained at 37°C (data not shown).

More quantitative translational mobility data were obtained by monitoring the temporal dependence of the post-bleach fluorescence using FRAP curves. Fig. 2, a and b, shows representative curves for Swiss 3T3 and HeLa cells, respectively. Both euchromatin and heterochromatin regions of the nucleus were studied. When the light used in the FRAP experiments was focused down to a small ~0.4 μm radius spot, the recovery curves were biphasic, reflecting the presence of two fractions. One fraction was “mobile,” leading to partial recovery on a time-scale of several hundred milliseconds; the other fraction was immobile, leading to an incomplete recovery in the FRAP curves.

FRAP curves were fitted to the theoretical expression for photobleaching recovery in two dimensions, assuming one mobile and one immobile fraction. The results of the fits for dihydroethidium-labeled Swiss 3T3 cells are 27 ± 4% immobile, 73 ± 4% mobile, and D = 1.1 (± 0.5) × 10^{-9} cm^2/s, where the quoted errors are standard deviations and typically data from ≥20 different nuclei were averaged. The analogous results for dihydroethidium-labeled HeLa cells are 36 ± 6% immobile, 64 ± 6% mobile, and D = 2.9 (± 0.2) × 10^{-9} cm^2/s. These results were independent of the depth of bleach. The mobile fractions obtained from the fits are fairly large but will be shown to represent a population of less tightly bound ethidium molecules, and not mobile chromatin. No more than a few percent of interphase chromatin is mobile on the ≥0.4 μm distance scale (see Discussion).

There are two important differences between the results shown in the photographs and in the photobleaching recovery curves. First, fluorescence from out-of-focus planes inevitably appears in the photographs (Scalettar et al., 1996) but is preferentially excluded from the photobleaching recovery curves by the image-plane diaphragm. Because laser beam divergence causes the spot radius to increase in out-of-focus planes, out-of-focus pick-up may cause the spots shown in the photographs to appear larger than their in-focus size. Nevertheless, the spot radius obtained by direct measurement from the photographs closely agrees with the in-focus spot size produced by the 63× Zeiss objective.

Second, the FRAP curves show an initial partial recovery of the fluorescence in the spots (mobile fractions ~64–73%) that is not evident from the photographs. The recovery time associated with this component was studied as a function of the size of the illuminated spot to identify its origin. If molecular motion is the source of the initial partial recovery, the recovery time should increase systematically as the size of the illuminated spot is increased. For example, for a recovery driven by Brownian diffusion, the recovery half-time is expected to increase as the square of the spot size (Axelrod et al., 1976). Fig. 3 illustrates this effect in data obtained from a fluorescein-labeled antibody solution. Fig. 2, a and b, shows FRAP curves and associated, fitted theoretical recovery curves obtained from two different nuclei samples for two different spot sizes. As with the antibody samples, the initial recovery time for the nuclei samples increases as spot size is increased, implying that molecular motion is the source of the initial recovery.

A motion-derived origin for the partial recovery is also suggested by results from two additional experiments; see Fig. 4, a and b. First, when samples are dried down and immobilized, the recovery largely disappears. Second, when hydrated samples are deoxygenated, the recovery time is essentially unaltered, which indicates that the initial recovery does not arise from reversible photobleaching (see Discussion). It is important to eliminate this latter possibility because the effects of reversible photobleaching have previously been observed in FRAP experiments on DNA and chromatin (Scalettar et al., 1990; Selvin et al., 1990).

The mobile fraction could represent chromatin, RNA, or a subpopulation of the ethidium bromide that transiently comes off the chromatin. To distinguish these possibilities, FRAP control experiments were performed on cells fixed and labeled with either ethidium bromide or ethidium homodimer-1, which binds essentially irreversibly to DNA and RNA (Gaugain et al., 1978; Markovits et al., 1985; Rye and Glazer, 1995). Fig. 5 shows that the mobile fraction is present in the fixed ethidium bromide–labeled samples but is absent from the fixed ethidium homodimer-
Figure 1. Pre- (a) and post-bleach (b and c) photographs of two dihydroethidium-labeled Swiss 3T3 cells in interphase. The ethidium stains nucleic acid in both the nucleus and cytoplasm, and reveals the distribution of euchromatin and heterochromatin within the nucleus. a shows the cells immediately before bleaching; b and c show the cells a few minutes and 1 h after bleaching the upper cell, respectively. The long-lived dark spot in the upper cell in b and c was created in an initially fluorescent region of euchromatin by illuminating the region with a 300 ms pulse of 514 nm (green) light from an argon-ion laser (Power = 50 mW at the sample). The failure of this spot to fill back up with fluorescently labeled chromatin demonstrates that a large fraction of the chromatin is immobile. The focus may have shifted slightly or the cell may have moved slightly in c after the sample spent 60 min on the microscope stage. Some nonuniformity in fluorescence in the photographs may arise from interference in the illuminating light. Bar, 10 μm.
1-labeled samples. Specifically, the results of curve fits for the fixed ethidium bromide-labeled samples are 29.6% immobile, 71.6% mobile, and $D = 7.1 \times 10^{-10}$ cm$^2$/s. The results for the fixed ethidium homodimer-1-labeled samples are essentially 0% mobile, 100% immobile, and $D < 0$ cm$^2$/s. Fixation should largely inhibit motion of protein-containing macromolecules on the FRAP distance scale. This, together with the fact that only the nucleic acid label differs between the two control samples, strongly suggests that the mobile fraction is a subpopulation of the ethidium bromide that transiently comes off the chromatin and diffuses freely (see Discussion). The mobile chromatin fraction is thus small (or zero).

A potential artifact in interpretation could arise if the lack of full fluorescence recovery is due to depletion of the finite reservoir of fluorescence in the nucleus, and not to an immobile chromatin fraction. However, the photographs show that the volume of the bleached region is small, indicating that the bleach is unlikely to significantly deplete the total nuclear fluorescence. Moreover, if the “immobile fraction” simply reflected depletion of that fraction of the total nuclear fluorescence, a few bleaches would render the nucleus essentially nonfluorescent. This was not found to be the case. Finally, the long-lived spots shown in the photographs demonstrate that the nucleus does not become uniformly less fluorescent with time after the bleach.

Artifacts in interpretation could also arise if the cells were not viable or were damaged during the FRAP experiments. For this reason, cell viability during and after photobleaching experiments was tested using the two methods described under Materials and Methods. No leakage of the viability dye calcein from bleached or unbleached cells was observed during the ~1-h period that the cells remained on the microscope stage, demonstrating viability during the experiments. In contrast, substantial leakage of calcein occurred <2 min after cells were placed in buffer.

Figure 2. Typical FRAP curves and associated theoretical recovery curves obtained from dihydroethidium-labeled Swiss 3T3 cells (a) and HeLa cells (b). The FRAP curves were smoothed using an eleven-point fit to a fourth-order polynomial (Savitzky and Golay, 1964) to facilitate distinguishing the different recovery curves. The bleach pulse was 10 ms in duration, and data were collected in 10-ms increments. The upper (diamonds) and lower (squares) curves in each graph were obtained from relatively smaller (focused) and larger (defocused) spots, respectively. The defocused spot was obtained by translating a lens along the optical path so that the light was not focused exactly on the sample plane. Appropriate neutral density filters were inserted into the path of the bleach beam so that the bleach depths for the two curves in each graph were comparable; results did not depend on bleach depth (data not shown), because relatively shallow bleach depths were employed. Two important features are qualitatively apparent from the FRAP curves. First, a large fraction of the fluorescence fails to recover, indicating that a large fraction of the chromatin is immobile. Second, the recovery time associated with the fraction of the fluorescence that does recover increases with increasing spot size, indicating that molecular motion is the source of the initial recovery.

Figure 3. Typical FRAP curves and associated theoretical recovery curves obtained from a rhodamine-labeled antibody undergoing Brownian diffusion in dilute solution ($\sim 0.5$ mg/ml antibody) for smaller (diamonds) and larger (squares) spots. The bleach pulse was 200 $\mu$s in duration, and data were collected in 100-$\mu$s increments. The upper curve represents an average of 1,000 experiments and the lower an average of 3,000 experiments, and each was smoothed using an 11-point fit to a fourth-order polynomial. The fluorescence recovers almost completely, indicating that all molecules are mobile. In addition, the recovery time increases with increasing spot size.
containing 0.1% Triton X-100. Moreover, bleached cells in an easily identified (scratched) region of the coverslip were observed to divide and proliferate after being returned to the incubator. Both of these results strongly suggest that the FRAP data reflect interphase chromatin dynamics in living and unperturbed Swiss 3T3 and HeLa cells.

Several other results, observations, and aspects of the experimental protocol argue against light-induced artifacts in the FRAP experiments. First, the mobility results were independent of light intensity and bleaching time. Second, only a very small fraction of the nuclear volume was typically exposed to light. Finally, ethidium was excited through its visible absorption band, using green light; this is likely to result in significantly less nuclear damage than would excitation of ethidium through its UV absorption band.

**Discussion**

The experiments described here were directed at (a) measuring the translational mobility of chromatin, especially euchromatin, in living interphase cells using FRAP; and (b) relating the results to current ideas about chromatin function and organization during interphase.

**Visually Constrained Motion**

Even in the absence of photobleaching data, it is evident that there are some constraints on chromatin motion in living interphase Swiss 3T3 cells. For example, the photographs in Fig. 1 show that heterochromatin is largely immobile, since the bright heterochromatic spots remain in approximately the same place in the nucleus during long periods of observation.

The mobility of euchromatin in living Swiss 3T3 cells, and of chromatin in general in small HeLa cells, is more difficult to assay visually. Hence, we turned to photobleaching techniques to give us more complete information on chromatin mobility in interphase cells. When we bleached the fluorescence in euchromatic regions of Swiss 3T3 nuclei, the subsequent temporal dependence of the fluorescence could be monitored both photographically (Fig. 1) and through photobleaching recovery curves (Fig. 2). Both types of data show that it is possible to bleach a long-lived spot in Swiss 3T3 euchromatin, indicating that a substantial fraction of Swiss 3T3 chromatin is immobile over distance scales comparable to, or larger than, the size (~0.25 μm) of a chromatin looped domain (Nelson et al., 1986). Similar results were obtained after photobleaching a spectrum of regions in HeLa nuclei, indicating that chromatin is similarly immobile in this human cell line.
immobility of interphase chromatin is in marked contrast to the known mobility of ethidium bromide and ethidium bromide–stained naked DNA in isolated samples (Icenogle and Elson, 1983a,b; Scalettar et al., 1989).

**Origin of Initial Recovery**

Although long-lived, the bleached spots quickly refill partially, as manifest in the partial recoveries in Fig. 2. We have studied the properties of this initial recovery in detail to verify that it is attributable to molecular motion. This is essential because there are mechanisms that can lead to recovery in FRAP experiments, such as reversible photobleaching, which are not mobility derived (Velez and Axelrod, 1988; Scalettar et al., 1990; Selvin et al., 1990).

In our case, several lines of evidence suggest that the initial recovery arises from motion and not effects such as reversible bleaching. First, the relaxation time associated with mobility-derived recoveries is expected to increase with increasing size of the illuminated spot, an effect which can be seen experimentally in the data in Fig. 2. Also, the relaxation time associated with reversible recoveries is expected to be independent of spot size but dependent on oxygen concentration, effects that we do not observe experimentally (Figs. 2 and 4). (The dependence on oxygen concentration is expected because reversible recovery reflects a long-lived residence in a triplet state, and triplet lifetimes are very sensitive to oxygen concentration; Scalettar et al., 1990; Selvin et al., 1990.) Second, the relaxation time associated with reversible photobleaching of ethidium bromide when bound to DNA or chromatin is <2 ms (Scalettar et al., 1990), about two orders of magnitude shorter than that of the initial recovery observed here. Finally, the initial recovery disappears when the molecules are dried down; drying leads to molecular immobilization but does not eliminate reversible bleaching (Scalettar et al., 1988b).

**Effects of High and Low Affinity Binding of Ethidium Bromide**

To identify the molecular species giving rise to the partial recovery, control experiments were performed on fixed samples that were labeled with either ethidium bromide or ethidium homodimer-1, which binds to naked DNA about 1,000 times more tightly than does ethidium bromide (Gaugain et al., 1978). Fixed samples were used because neither dye is membrane-permeant; however, fixation in formaldehyde was not expected to immobilize the dye, because formaldehyde cross-links amino groups (Baker, 1958) and because the dye was added after fixation. This expectation was borne out by the fact that the mobile fraction was still present in fixed, ethidium bromide–labeled samples and in dihydroethidium-labeled samples that were fixed after labeling.

In contrast, the mobile fraction was absent from fixed, ethidium homodimer-1–labeled samples. Because fixing and labeling with ethidium bromide leaves the mobile fraction unaltered, whereas fixing and labeling with the more tightly binding ethidium homodimer-1 eliminates the mobile fraction, we have attributed the initial recovery to independent motion of ethidium bromide arising from its weaker binding to DNA. In addition, the effects of on/off binding of ethidium bromide have been previously observed in FRAP data (Icenogle and Elson, 1983a,b).

Over the time-scale we have probed, only a subpopulation of the ethidium molecules is transiently moving independently of the chromatin. If all ethidium molecules were moving independently, the FRAP curves could be described by a single recovery time reflecting the fraction of ethidiums bound and the diffusion coefficient of ethidium alone (Icenogle and Elson, 1983a,b; Kao et al., 1993). Our data show two recovery times (one approximately several hundred milliseconds and one around infinity), indicating that not all ethidiums are moving independently.

Previous studies (Lawrence and Daune, 1976) of the interaction of ethidium bromide with chromatin suggest a mechanism that could give rise to two ethidium populations in our samples. These studies identified high and low affinity ethidium binding sites, which differ by close to three orders of magnitude in binding constant (Lawrence and Daune, 1976). Moreover, the low affinity sites constitute the dominant fraction (~82%) of the ethidium sites in native chromatin and are present in formaldehyde-fixed chromatin. It thus seems likely that the mobile species observed here is ethidium that diffuses after release from low affinity sites. This interpretation is strengthened by the fact that our mobile fractions (~64–73%) are in reasonable agreement with the fraction of sites in native chromatin that have a low affinity for ethidium bromide.

This interpretation is further strengthened by the quantitative agreement between mobile fractions in living dihydroethidium-labeled samples and fixed ethidium bromide–labeled samples. The mobile fractions in the living dihydroethidium labeled Swiss 3T3 samples were 73 ± 4% and in the fixed ethidium bromide–labeled Swiss 3T3 samples were 71 ± 6%. Thus, the entire mobile fraction can be attributed to transiently unbound ethidium bromide, and the FRAP data show that interphase chromatin is immobile on the ≈0.4 μm distance scale.

**Comparison with Previous Studies of Chromosome Organization and Dynamics**

Results obtained in a number of studies suggest that chromosome substructures can be nonrandomly organized and that chromosomes are confined to domains within interphase nuclei. From a dynamic perspective, these results suggest that there are restrictions on the translational mobility of interphase chromatin, as observed here.

Examples of nonrandom organization and confinement to domains are numerous. For example, a nuclear membrane–apposed/peripheral distribution has been observed for inactivated X chromosomes in fibroblasts, centromeres in some types of cells, and centromeres and telomeres of polytene chromosomes in Drosophila salivary gland nuclei (Moroi et al., 1981; Mathog et al., 1984; Manuelidis and Borden, 1988; Dyer et al., 1989). Similarly, chromosomal confinement to domains has been observed or deduced from in situ hybridization studies and DNA irradiation studies (Cremer et al., 1982; Hens et al., 1983; Schardt et al., 1985; Cremer et al., 1988; Lichter et al., 1988; Manuelidis and Borden, 1988; Pinkel et al., 1988; Leitch et al., 1990; Popp et al., 1990). Finally, active genes can exhibit nonrandom organization, concentrating in the nuclear periphery.
of mouse L and P19 embryonal carcinoma cells and near
the borders of condensed chromatin in nucleated newt
erthrocytes (Hutchison and Weintraub, 1985; de Graaf
et al., 1990). These examples all suggest constraints on
chromosome lateral mobility during interphase.

More direct evidence for constraints on chromosome
lateral mobility during interphase comes from particularly
pertinent recent studies of the mobility of centromeres in
living HeLa cells (Shelby et al., 1996). Motion of cen-
tromeres was tracked by labeling centromeres with a fu-
sion protein consisting of green fluorescent protein and
CENP-B, a centromeric satellite DNA–binding protein.
Most centromeres were found to remain motionless for a
time period of up to 2 h.

In contrast, some studies have indicated that chromatin
can be mobile during interphase. For example, in living in-
terphase neurons the position of heterochromatin has
been shown to change in tandem with changes in nucleoli
position (DeBonii and Mintz, 1986). Similarly, centromeres
have been observed to relocate in a variety of interphase
cells (Manuelidis, 1985; Bartholdi, 1991; Ferguson and Ward,
1992; Janevski et al., 1995; Shelby et al., 1996), as have X
chromosome centromeres in certain pathological states
(Borden and Manuelidis, 1988). Finally, chromosome arms
in polytene nuclei (Hochstrasser and Sedat, 1987) and
neuronal cells (Manuelidis and Borden, 1988) can exhibit
some randomness in configuration and position.

Our work is in basic agreement with the body of work
suggesting the existence of constraints on interphase chro-
matin mobility. A unique feature here is that the results
reflect the mobility of interphase chromatin in general,
rather than the mobility of a particular chromosome or
chromosome substructure. The results thus give a global
view of chromatin mobility while also defining a distance
scale for chromatin immobilization. In addition, the results
were obtained from living cells, unlike many of those dis-
cussed above.

Chromatin in Transformed and Cultured Cells
Swiss 3T3 and HeLa cells are commonly used in studies of
chromatin organization and nuclear structure. Neverthe-
less, there are potential differences between nuclear struc-
ture in these cell lines and nuclear structure in untrans-
formed, uncultured cells. For example, transformed cells
may have less heterochromatin and a somewhat altered
nuclear matrix protein composition when compared with
untransformed cells (for review see Manuelidis, 1990; Nick-
erson et al., 1995). Similarly, chromatin in cultured cells
may partially expand to a more euchromatic state (Man-
uelidis, 1990). However, because these potential changes
in chromatin structure would probably tend to “loosen
up” chromatin and increase its mobility, our results should
represent a lower bound on the immobility of interphase
chromatin in cells in general.

Origin of Chromatin Immobilization on the $\geq 0.4 \mu m$
Distance Scale
There are two fundamentally different types of interac-
tions that could act alone or in combination to produce the
chromatin immobilization observed here. These are (a)
outright attachment of chromatin to nuclear substructures,
chromosomes appear not to be entwined (for reviews see Manuelidis, 1990; Jackson, 1991; van Driel et al., 1995), suggesting that entanglement does not have a major effect on chromatin motion. Steric interactions can also produce such major inhibition of mobility if the system contains a very high concentration of immobile obstacles that block all potential paths for motion (Luby-Phelps et al., 1987, 1988; Saxton, 1993). However, if such a situation exists in interphase nuclei, it would once again suggest that immobile nuclear substructures extensively bind nuclear constituents, thereby producing a high concentration of immobile obstacles.

Conclusions

We have shown that interphase euchromatin and heterochromatin in living Swiss 3T3 fibroblasts and HeLa cells are translationally immobile over distance scales \( \geq 0.4 \mu m \). This immobilization may reflect motion-impeding steric interactions that arise in the highly concentrated nuclear milieu or outright attachment of the chromatin to underlying nuclear substructures, such as nucleoli, the nuclear lamina, or the nuclear matrix. However, independent of its origin, this immobilization ensures that spatial organization of chromosomes within interphase nuclei is maintained, as postulated in current models of a highly ordered interphase nucleus.

We thank Dr. John Langmore of the University of Michigan for a critical reading of the manuscript.

This work was supported by grants CC3819 from Research Corporation (to B.A. Scalettar), BIR-9510226 from the National Science Foundation (to B.A. Scalettar), and MCB-9405298 from the National Science Foundation (to D. Axelrod).

Received for publication 27 December 1996 and in revised form 25 March 1997.

References

Abney, J.R., B.A. Scalettar, and J.C. Owicky. 1989. Self diffusion of interacting membrane proteins. Biophys. J. 55:817–833.

Alexander, S.P., and C.L. Rieder. 1991. Chromosome motion during attachment to the vertebrate spindle: initial saltatory-like behavior of chromo- somes and quantitative analysis of force production by nascent kinetochore fibers. J. Cell Biol. 113:805–815.

Axelrod, D., E.D. Koppel, J. Schlessinger, E. Elson, and W.W. Webb. 1976. Mobility measurement by analysis of fluorescence photobleaching recovery kinetics. Biophys. J. 16:1055–1069.

Baker, J.R. 1958. Principles of Biological Microtechnique. John Wiley & Sons Inc., New York. 357 pp.

Bartholdi, M.F. 1991. Nuclear distribution of centromeres during the cell cycle of human diploid fibroblasts. J. Cell Sci. 99:255–263.

Berezney, R., M.J. Mortillaro, H. Ma, X. Wei, and K. Samarabandu. 1995. The nuclear matrix: a structural milieu for genomic function. Int. Rev. Cytol. 162A:1–65.

Blonk, J.C.G., A. Don, H. Van Aalst, and J.J. Birmingham. 1993. Fluorescence photobleaching recovery in the confocal scanning light microscope. J. Microsc. (Oxf). 169:363–374.

Borden, J., and L. Manuelidis. 1988. Movement of the X chromosome in epithelial mammalian cells and quantitative analysis of force production by nascent kinetochore fibers. J. Cell Sci. 99:255–263.

Langmore, J.M., and C.L. Rieder. 1991. Chromosome motion during attachment (to B.A. Scalettar), BIR-9510226 from the National Science Foundation (to D. Axelrod).

Manuelidis, 1990; Jackson, 1991; van Driel et al., 1995),

Haugland, R.P. 1996. Handbook of Fluorescent Probes and Research Chemicals. Molecular Probes, Inc., Eugene, OR. 679 pp.

Hiraoa, Y., J.S. Minden, J.R. Swedlow, J.W. Sedat, and D.A. Agard. 1989. Fo- cal points for chromosome condensation and decondensation revealed by three-dimensional in vivo time-lapse microscopy. Nature (Lond.) 342:293–296.

Hochstein, M., and J.W. Sedat. 1987. Three-dimensional organization of Drosophila melanogaster interphase nuclei. II. Chromosome spatial organi- zation and gene regulation. J. Cell Biol. 106:1471–1483.

Hou, L., F. Lanni, and K. Luby-Phelps. 1990. Tracer diffusion in F-actin and fi- colli mixtures: toward a model for cytoplasm. Biophys. J. 58:31–43.

Hutchison, N., and H. Weintraub. 1985. Localization of DNA isoelectric sequences to specific regions of interphase nuclei. Cell. 43:471–482.

Icenogle, R.D., and E.L. Elson. 1983. Fluorescence correlation spectroscopy and photobleaching recovery of multiple binding reactions. I. Theory and FCS measurements. Biopolymers. 22:1919–1948.

Jackson, D.A. 1991. Structure-function relationships in eukaryotic nuclei. Bioessays. 13:1–10.

Jacobson, K., A. Ishihara, and R. Inman. 1987. Lateral diffusion of proteins in membranes. Annu. Rev. Physiol. 49:163–175.

Janevska, J., P.C. Park, and U. De Boni. 1995. Organization of centromeric domains in hepatocyte nuclei: rearrangement associated with de novo activation of the vitellogenin gene family in Xenopus laevis. Exp. Cell Res. 217:227–239.

Kao, H.P., J.R. Abney, and A.S. Verkman. 1993. Determinants of the transla- tion mobility of a small solute in cytoplasm. J. Cell Biol. 120:175–184.

Lang, I., M. Scholz, and R. Peters. 1986. Molecular motility and nucleocyto- plasmic flux in hepatoma cells. J. Cell Biol. 102:1183–1190.

LaSalle, J.M., and M. Lalande. 1996. Homologous association of oppositely im- printed chromosome domains. Science (Wash. DC). 271:2872–2875.

Lawrence, J.J., and M. Daune. 1976. Ethidium bromide as a probe of conforma- tion. Sci. STKE. 169:363–374.

Leitch, A.R., W. Mosgoller, T. Schwarzzacher, M.D. Bennett, and J.S. Heslop- Harrison. 1990. Genomic in situ hybridization to sectioned nuclei shows chromosomal domains in grass hibrises. J. Cell Sci. 95:335–341.

Lichter, P., T. Cremer, J. Borden, L. Manuelidis, and D.C. Ward. 1988. Delini- cation of individual human chromosomes in metaphase and interphase cells by in situ hybridization using recombinant DNA libraries. Hum. Genet. 80:224–234.

Livolant, F., and M.F. Maestre. 1988. Circular dichroism microscopy of compact forms of DNA and chromatin in vivo and in vitro: cholesteric liquid- crystalline phases of DNA and single dinoflagellate nuclei. Biochemistry. 27:
