Dynamic as well as stable protein interactions contribute to genome function and maintenance

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Abstract The cell nucleus is responsible for the storage, expression, propagation, and maintenance of the genetic material it contains. Highly organized macromolecular complexes are required for these processes to occur faithfully in an extremely crowded nuclear environment. In addition to chromosome territories, the nucleus is characterized by the presence of nuclear substructures, such as the nuclear envelope, the nucleolus, and other nuclear bodies. Other smaller structural entities assemble on chromatin in response to required functions including RNA transcription, DNA replication, and DNA repair. Experiments in living cells over the last decade have revealed that many DNA binding proteins have very short residence times on chromatin. These observations have led to a model in which the assembly of nuclear macromolecular complexes is based on the transient binding of their components. While indeed most nuclear proteins are highly dynamic, we found after an extensive survey of the FRAP literature that an important subset of nuclear proteins shows either very slow turnover or complete immobility. These examples provide compelling evidence for the establishment of stable protein complexes in the nucleus over significant fractions of the cell cycle. Stable interactions in the nucleus may, therefore, contribute to the maintenance of genome integrity. Based on our compilation of FRAP data, we propose an extension of the existing model for nuclear organization which now incorporates stable interactions. Our new “induced stability” model suggests that self-organization, self-assembly, and assisted assembly contribute to nuclear architecture and function.

Keywords Nucleus · Fluorescence microscopy · FRAP · Multi-protein complex · Chromatin binding · Dynamics · Residence time · Induced stability · Assisted assembly · Self-assembly · Self-organization

Abbreviations
CB Cajal body
CBP C-AMP response element binding protein (CREB) binding protein
CT Chromosome territory
CP Continuous fluorescence photobleaching
FFM Fluorescence fluctuation microscopy
FLIM Fluorescence lifetime imaging
FLIP Fluorescence loss in photobleaching
FRAP Fluorescence recovery after photobleaching

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Structure and function in the nucleus

The storage, propagation, maintenance, and expression of the genetic material is executed by biochemical activities, namely DNA compaction/decompaction, DNA replication and segregation, DNA repair, and RNA transcription/processing, respectively (Diekmann and Hemmerich 2005). The corresponding machineries are highly structured, yet dynamic macromolecular assemblies (Misteli 2007) which must work on chromatin with high fidelity in a crowded nuclear environment (Richter et al. 2007). In addition, the mammalian cell nucleus contains a variety of subnuclear domains, nuclear bodies, or subnuclear compartments (Figs. 1 and 2). DNA is wrapped around nucleosomes and forms individual chromosomes which are compacted in mitosis (Fig. 2g). In interphase cells, chromosomes decondense into so-called chromosome territories (CTs), which occupy distinct volume regions (Figs. 1 and 2h) (Cremer et al. 2006; Heard and Bickmore 2007; Solovei et al. 2009; Finan

Fig. 1 Compartmentalization of the mammalian cell nucleus. The mammalian cell nucleus contains chromatin in the form of chromosome territories (CTs). CTs may overlap at their touching borders (intermingling) or create the so-called interchromatin space (white). Constitutive heterochromatin is mainly found as pericentricromeric chromatin in patches throughout the nuclear volume, at the nuclear periphery, as well as around nucleoli. Nuclear pore complexes, the double-layered nuclear membrane (dark green) and the meshwork-like nuclear lamina are structural hallmarks in the periphery of the nucleus. Chromatin loops with associated transcription factories may extrude out of chromosome territories within the nucleolus as well as throughout the nucleoplasm. Transcription (orange), replication (yellow), and DNA repair processes (light blue) usually occur in small domains with a diameter below 100 nm. A diverse set of nuclear bodies, such as speckles, paraspeckles, the perinucleolar compartment, Cajal bodies, or PML bodies are found in the interchromatin space.
et al. 2010). Staining of interphase chromatin using DNA dyes does not reveal CT structures but allows discrimination between transcriptionally active euchromatin and transcriptionally silent heterochromatin (Fig. 2e). Constitutive heterochromatin is mainly composed of pericentric DNA, and in this case, the chromosome’s centromere/kinetochore complex can be found embedded within or adjacent to this chromatin region (Fig. 2j) (Probst and Almouzni 2008). The nucleus obtains structural support through the nuclear lamina, which is attached to the nuclear double-membrane, together forming the nuclear envelope (Fig. 1). The nuclear envelope controls traffic of molecules between the cytoplasm and the nucleoplasm but has also emerged as a critical determinant in genome architecture (Starr 2009).

The most prominent subnuclear domains include the nucleolus, the perinucleolar compartment, speckles, paraspeckles, Cajal bodies, and promyelocytic leukemia (PML) bodies (Figs. 1 and 2). In addition, a variety of other nuclear bodies have been identified such as PcG bodies, Gemini bodies (Gems), the OPT domain, cleavage bodies, and the SAM68 nuclear body (Spector 2001; Handwerger and Gall 2006). Subnuclear structures are macromolecular complexes that consist of membrane-less accumulations of specific sets of functionally related molecules. For example, components of the ribosome biogenesis pathway are predominantly confined to the nucleolus. First thought to be exclusively devoted to the synthesis of ribosomal RNA and assembly of ribosomal subunits, it has become clear that the nucleolus serves a variety of additional functions, including regulation of mitosis, cell-cycle progression, proliferation, and various stress responses (Raska et al. 2006; Sirri et al. 2008).

The biochemical function(s) of the other subnuclear domains are less clear or unknown. PML bodies attract a selected set of nuclear proteins which are functionally quiet promiscuous. Therefore, PML bodies have been implicated in the regulation of diverse cellular functions, such as the induction of apoptosis and cellular senescence, inhibition of proliferation, maintenance of genomic stability, and antiviral responses (Bernardi and Pandolfi 2007). PML bodies are positionally stable structures at which controlled molecular traffic and posttranslational modifications may regulate the activity of specific proteins throughout the genome and the epigenome in response to various cellular stresses (Bernardi and Pandolfi 2007; Torok et al. 2009). Speckles, also referred to as interchromatin granule clusters, are enriched in pre-mRNA splicing factors. At the microscopic level, speckles appear as irregular, punctate domains varying in size and shape (Fig. 2). They are considered to be the main sites for storage, assembly,
and/or recycling of the essential spliceosome components (Lamond and Spector 2003). Because highly transcribed genes are found in the periphery of speckles and also other subnuclear domains, they may also serve to efficiently integrate and regulate mRNA transcription and mRNA processing machineries (Zhao et al. 2009). Cajal bodies (CBs) are involved in the biogenesis of several classes of small ribonucleoprotein particles (snRNPs) as well as their modification (Gall 2000; Matera et al. 2009). Resembling the speckles/gene association mentioned above, CBs associate specifically with histone and snRNA genes. This colocalization is transcription-dependent, requires expression of snRNA coding regions, and is probably based on an energy-driven motor activity in the nucleus (Frey and Matera 2001; Dundr et al. 2007).

In mammalian cell nuclei, DNA replication, RNA transcription, and repair of damaged DNA occurs in dot-like structures with a mean diameter of ~100 nm (Fig. 3). With respect to transcription and replication, these focal sites have been coined “factories” as each site contains all of the enzymatic activity required. A general model was recently suggested for the organization of all genomes in which the transcription factories play a central role (Cook 1999; Cook 2010). Notably, the model proposes that active RNA polymerases do not move along their templates during elongation but are bound to a factory acting both as motors that reel in their templates and as fixed structural entities that hold active chromatin loops in place (Cook 2010). DNA replication also occurs at similarly specialized subnuclear sites where the factors directly or indirectly involved in replication are concentrated (Fig. 3b) (Leonhardt et al. 2000). Finally, repair of damaged DNA at focal sites throughout the genome is also a dynamic process that requires careful orchestration of a multitude of enzymes, adaptor proteins, and chromatin constituents (Fig. 3c) (Lukas et al. 2005).

**Fluorescence fluctuation microscopy**

The development of in vivo microscopy techniques employing the green fluorescent protein (GFP) has

![Fig. 3](image-url) DNA and RNA metabolism occurs in small foci. a Transcription can be visualized after brief exposure of living cells to the nucleotide analog Fluoro-Uridine (FI-U), followed by immunodetection of the FI-U epitope. b Bromo-deoxy-Uridine (BrdU) can be used in a similar incorporation assay to visualize nascent DNA during replication. c Sites of DNA double-strand break repair are detected using antibodies against a phosphorylated form of histone H2AX (γ-H2AX). Common to all of these active foci is the accumulation of many factors required to perform the biochemical activities. Size bar, 10 μm.
opened the door to probe cellular architecture and function in living cells (Lippincott-Schwartz et al. 2003). By analyzing macroscopic relaxation after disturbing the equilibrium state, fluorescence intensity images can be used to assess diffusion times, interactions, and binding constants of molecules. Fluorescence fluctuation microscopy (FFM) approaches have been developed to investigate a few molecules in small regions of a cell. These approaches supply dynamic spatiotemporal information by creating cellular diffusion and concentration maps (Fig. 4). A major consideration is the accessible resolution as nuclear processes can take place on a time scale ranging from microseconds to hours, and single molecules or huge macromolecular assemblies in well-defined stoichiometries can be involved.

FFM allowed for the first time to not only visualize protein dynamics and chromatin interactions but also to quantitatively determine biophysical properties of proteins in the nucleus of intact cells (Erdel et al. 2010, this volume). FFM approaches include time-lapse microscopy (Heun et al. 2001), fluorescence redistribution after photobleaching (FRAP, FLIP, iFRAP, etc.) (Bancaud et al. 2009; van Royen and

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**Fig. 4** Fluorescence fluctuation microscopy. **a** Upper panel FRAP experiment in a nucleus expressing GFP-tagged HP1. Fluorescence was bleached within two circled areas (green) and fluorescence redistribution was monitored over time. Bottom panel Quantification of a FRAP experiment. Graphs typically show mean values from 10 to 20 FRAP experiments as relative fluorescence intensity (RFI) after normalization to prebleach levels. The FRAP curve immediately delivers information on mobile and immobile or slow populations. Ideally, the FRAP experiment is extended to allow discrimination between immobile or slowly exchanging populations. Mathematical modeling can be performed to extract biophysical parameters such as the diffusion coefficient and binding constants from fit functions (red dotted line). **b** RICS. GFP-HP1 was expressed in HEp-2 cells. For RICS, a time series of GFP fluorescence images (512×512 pixels) was acquired in a subregion of the nucleus by confocal microscopy (left panel). Subregions (64×64 pixels) within this time series are than extracted and correlation spectra assessed from these subregions. A diffusion coefficient map can be generated by fitting with appropriate diffusion models (middle panel) from which diffusion coefficients of GFP-HP1 can be determined at different subnuclear positions. **c** In FCS, a confocal volume is generated by focusing of a laser beam through an appropriate objective (upper panel). Fluorescent molecules emit photons during their movement through the confocal volume leading to a fluorescence fluctuation over time (middle panel). The fluctuation data are then subjected to autocorrelation (bottom panel) from which fluorescent molecule concentration and diffusion coefficients can be extracted after fitting of the data with appropriate mathematical diffusion models. Size bar, 5 μm
Houtsmuller 2010; this volume), fluorescence correlation spectroscopy (FCS) (Haustein and Schwille 2003), CP (Wachsmuth et al. 2003), raster image correlation spectroscopy (RICS) (Digman and Gratton 2009), SPT (Levi and Gratton 2008), FRET and FLIM (Wallrabe and Periasamy 2005), and MFIS (Weidtkamp-Peters et al. 2009). Figure 4 highlights some of these techniques demonstrating their potential to study nuclear protein dynamics.

**Dynamic and stable interactions contribute to the assembly and maintenance of subnuclear compartments**

FRAP analyses of subnuclear domains such as nucleoli, speckles, and Cajal bodies have revealed that their component parts rapidly exchange with nucleoplasmic pools (Misteli 2001a, 2008). Typical residence times of proteins in these compartments are in the seconds range. Similarly, most (but not all, see below) factors acting at transcription, replication, and repair foci show rapid exchange at chromatin (Houtsmuller et al. 1999; Dundr et al. 2002; Sporbert et al. 2002). These observations have led to the conclusion that nuclear proteins undergo repeated and rapid cycles of association and dissociation between the nucleoplasm and the compartment they are mainly working in. As a consequence, nuclear bodies and factories are in perpetual flux. Their structure is determined by the ratio of on-rate versus off-rate of its components, clearly suggesting self-assembly and/or self-organization as the mechanism of their assembly (Misteli 2001b; Matera et al. 2009). To further test this hypothesis, we have compiled FRAP data of 264 nuclear proteins collected over the last 16 years. Data are presented as individual sheets of a single table (ESM 1) containing: chromatin-binding proteins, transcription factors, DNA metabolism (covering replication, repair, recombination), RNA metabolism (covering speckles, splicing, RNA modification), centromeres, nucleolus, nuclear envelope, Cajal bodies, PML bodies, and “others.” Surprisingly, while indeed most nuclear proteins are highly dynamic, we found a small but nevertheless important set of proteins showing either very slow turnover or complete immobility (ESM 1). Figure 5 provides an overview of the FRAP data with respect to the various subnuclear structures. In many cases, very slow FRAP recoveries probably reflect relatively long residence times on chromatin. However, it is also possible in principle that slow recoveries can arise from short residence times if the average fluorescent molecule undergoes many binding events as it progresses to the center of the bleach spot (Sprague et al. 2004). Accurate residence time estimates require a test for the role of diffusion in the FRAP and a proper mathematical model for fitting (Sprague and McNally 2005). This approach should be considered for the slowly recovering proteins discussed below.

**Fig. 5 (Im)mobilities in the nucleus. Illustration of subnuclear compartments and overview of their component’s residence times ($t_{res}$) determined by FRAP**
Most nucleosomes are stable for long periods of time. FRAP analyses of GFP-tagged core histones have shown that far more than 50% of H2A, H2B, H3, and H4 are stably incorporated into chromatin and do not exchange with soluble pools (ESM 1; Kimura and Cook 2001; Gautier et al. 2004). These observations are consistent with the idea of a structural foundation of genome architecture and maintenance based on stable nucleosome arrays. The transiently incorporated sub-fraction of core histones is most likely attributable to chromatin remodeling events associated with transcription, replication or repair. All four core histones are present in the same nucleosome, yet their stabilities vary. H3 and H4 are loaded into nucleosomes during DNA replication, after which >80% remains permanently bound (Kimura and Cook 2001). In contrast, ~50% of H2A and H2B exhibit substantial turnover (Kimura and Cook 2001; Gautier et al. 2004), compatible with the assembly properties of nucleosomes in vitro and in vivo (Annuziato 2005). While the immobile fractions might be associated with heterochromatin, the stability of the central H3 and H4 coupled with the lability of the outer H2A and H2B suggest simple ways of maintaining epigenetic marks (Kimura and Cook 2001). This is consistent with many posttranslational modifications found at the N-terminal ends of H3 and H4 providing long-term memories (Margueron and Reinberg 2010).

Genome maintenance relies on hyper-stable protein complexes at the centromere

Cell division is a highly dynamic process in which the chromosomes are segregated in a coordinated fashion. The centromere is the genetic locus required for precise and accurate chromosome segregation and provides a platform on which the kinetochore multiprotein complex assembles (Przewloka and Glover 2009). Accurate chromosome segregation is essential for cell survival and genome maintenance. Aberrant mitotic segregation can result in aneuploidy, cell death, or cancer (Kops et al. 2005). Intuitively and confirmed by FRAP data, stable chromatin–protein and protein–protein interactions are required during mitosis when sister chromatids are pulled along microtubules into daughter cells (ESM 1). However, the kinetochore/centromere complex could in principle disassemble completely during interphase, since a stable epigenetic mark such as a unique histone modification would be sufficient to transmit centromere identity through interphase. Shortly before mitosis starts, the epigenetic mark would be used as a landing platform for cell cycle-regulated factors that initiate de novo kinetochore assembly. This, however, is not the case as FRAP of centromeric proteins performed at any time during cell cycle show that at least Cenp-A and Cenp-I (ESM 1) stably associate with centromeres for the complete cell cycle (Hemmerich et al. 2008; Hellwig et al. 2008).

Because there is absolutely no dynamic exchange of Cenp-A between centromeres and freely diffusible nucleoplasmic pools, the question arises how the binding sites created by centromere replication are occupied. This issue has been addressed by FRAP experiments which clearly showed that Cenp-A is assembled during the G1 phase of the cell cycle via a “loading only” mechanism (Hemmerich et al. 2008): while the first FRAP experiment showed fluorescence recovery, a subsequent FRAP showed no recovery of CENP-A indicating an undetectably small off-rate. This shows that CENP-A is “loaded only,” without exchange at its binding sites. Interestingly, Cenp-A is incorporated via the “loading only” mechanism exclusively in early G1 (Hemmerich et al. 2008) corroborating earlier SNAP-tagging results (Jansen et al. 2007). The epigenetic marking of the centromere is believed to be conveyed by Cenp-A, because it is required for the association of probably all other kinetochore proteins (Buscaino et al. 2010) and because of its sustained presence at centromeres without dynamic exchange (Hemmerich et al. 2008; ESM 1). Although Cenp-A and Cenp-I are stably bound, many rapidly exchanging centromere components have also been identified (ESM 1). Taken together, the identification by FRAP of proteins which bind to the centromeres with dynamic exchange but also without dynamic exchange for longer than one cell cycle clearly indicates that in addition to organization and self-assembly, also heredity mechanisms contribute to centromere structure and function and hence genome maintenance.

A fraction of stably bound cohesin glues sister chromatids together

Sister chromatid cohesion after replication is mediated by the multiprotein complex cohesin which acts as a topological linker (Wong 2010). Gerlich et al. (2006)
performed FRAP analyses identifying two distinct binding modes of cohesin to chromatin during the cell cycle. One pool of cohesin dynamically exchanged on and off chromatin throughout the entire interphase, moving through the nucleus by diffusion. This pool was absent in metaphase. About one third of the subunits SA1 and Scc1 are stably fixed to chromosomes after replication with a residence time of 6 to 7 h (ESM1). This pool persisted until just before chromosome segregation (Gerlich et al. 2006). These analyses provided the first experimental evidence in living cells for a permanent and stable link between sister-chromatids. This stable link may functionally contribute to homologous recombination repair of DNA double-strand breaks during the G2 phase of the cell cycle.

HA95 and lamin A immobility in the nucleoplasm is consistent with the idea of a stable nuclear matrix

The existence of a nuclear matrix was proposed by Zbarskii and Debov (1948) based on their observation that high-salt extractions of purified nuclei produced microscopically visible lattice-like residual structures. These observations were confirmed in the Berezney lab using similar extraction protocols (Berezney and Coffey 1974). The lack of direct, particularly in vivo evidence, however, raised considerable skepticism about the existence of a stable nuclear structure (Pederson 2000). Homologous to A-kinase anchoring protein 95 (HA95) is a nuclear protein harboring two zinc fingers and a putative nuclear localization signal (Ørstavik et al. 2000). HA95 binds to chromatin and the nuclear lamina network where it has a role in anchoring nuclear membranes and lamins to chromatin in interphase and in releasing membranes from chromatin at mitosis. Astonishingly, bleaching experiments revealed no detectable recovery of the diffusely localized GFP-tagged HA95 in the nucleus of living 293 T fibroblasts (Martins et al. 2000). Roughly one quarter of the nuclear pool of lamin A resides outside the nuclear lamina and distributes as a diffuse veil throughout the nucleoplasm (Moir et al. 2000). FRAP analysis of the nucleoplasmic GFP-lamin A pool revealed a recovery halftime of more than 3 h, clearly indicating a stable lamin-containing structure in living cells (Moir et al. 2000). Although overexpression artifacts of the GFP fusion proteins cannot be ruled out, the immobile properties of HA95 and lamin A in the cell nucleus provide compelling evidence for the existence of a nuclear matrix in living cells. It has been suggested that HA95 provides a stable platform at the chromatin/interchromatin interface that could regulate nuclear envelope–chromatin interactions during the cell cycle (Martins et al. 2003). Recent evidence suggests that HA95 is also functionally involved in pre-mRNA splicing (Kvissel et al. 2007). Thus, HA95 could also serve as a stable platform on chromatin, linking RNA transcription with RNA processing. It will be essential to analyze if nuclear stability and particular nuclear functions are affected in HA95-depleted cells.

Constitutive heterochromatin: more stable than initially thought?

The main function of constitutive heterochromatin is to serve as a kinetochore attachment site in mitosis, where its particular chromatin structure is thought to ensure chromosome segregation. Pericentric heterochromatin also plays important roles in organizing transcription-repressive compartments of the nucleus and to ensure regulation of particular genes (Probst and Almouzni 2008). Constitutive heterochromatin is characterized by three repressive epigenetic marks: tri-methylation of H3-K9, mono-methylation of H3-K27, and tri-methylation of H4-K20. The histone methyltransferases Suv39H1/2 and Suv420H1/2 play crucial roles in the initial steps of constitutive heterochromatin formation in mammals. Mice deficient for the Suv enzyme family members display impaired pericentric heterochromatin function leading to chromosome mis-segregation and increase of sister-chromatid exchange. H3-K9me3 marks placed by Suv39H activities stabilize heterochromatin protein 1 (HP1) binding to heterochromatin. HP1 proteins self-oligomerize and interact with both Suv39H and Suv420H (Fodor et al. 2010).

According to its structural function, heterochromatin had been perceived as a relatively densely packed, rigid, and inert region. Surprisingly, however, FRAP had demonstrated that GFP-tagged HP1 isoforms display dynamic binding properties, suggesting that heterochromatin is a dynamic and plastic domain, in which access to the underlying DNA would not necessarily be prevented (Cheutin et al. 2003; Festenstein et al. 2003; Schmiedeberg et al. 2004). In contrast to HP1, Suv39H1 consists of both a mobile and an immobile
population in heterochromatin where ~30% of molecules do not exchange over a time period of 7 min (ESM 1; Krouwe et al. 2005). The tight association of Suv39H1 with chromatin is obviously a more common feature because a large fraction of Suv420H2 is also tightly bound at pericentric heterochromatin (ESM 1). FRAP analyses has revealed that more than 90% of GFP-tagged Suv420H2 is immobile at these domains over several minutes (Souza et al. 2009). It is, therefore, possible that the residence times of Suv39H1 and Suv420H2 in heterochromatin are even in the hours range, indicating that these enzymes may, in addition to their catalytic activity, also serve a structural role in chromatin and act as a stable interaction platform for other heterochromatin proteins. Further structural support for heterochromatin architecture may also be provided by the heterochromatin-binding protein Ki-67 which, by FRAP, also shows a cell cycle dependent turnover in heterochromatin (ESM 1; Saiwaki et al. 2005). While Ki-67 dynamically binds to mitotic chromatin, it becomes very tightly bound to peri-nucleolar heterochromatin during G1 phase. Approximately 60% of GFP-Ki-67 is immobile over 10 min during FRAP analyses in interphase cells (Saiwaki et al. 2005). These data suggest that Ki-67 may have very long residence times on chromatin.

Some transcription factors can become stably bound to chromatin: pRB, E2F, Mdm2, VHL, TFIIH, HSF, and Gal4

Most of the time, transcription factors diffuse throughout the cell nucleus, encounter gene promoters in a random fashion, and bind to them for a very short time (Misteli 2001b). This, however, does not hold for all situations.

**pRB and E2F** The retinoblastoma protein (pRB) is a key cell-cycle protein that regulates the critical G1-S phase transition through interaction with E2F family members. The latter are cell-cycle transcription factors that repress transcription of genes required for G1-S checkpoint transition. pRB activity is regulated through networks sensing intracellular and extracellular signals which block or permit phosphorylation (inactivation) of the Rb protein (Poznic 2009). Quite unexpectedly, in bleaching experiments, fluorescence recovery of active (hypophosphorylated) GFP-RB and E2F is minimal (Angus et al. 2003). This behavior is in stark contrast to the mobility of most other transcription factors which are usually highly mobile (ESM 1). Only after inactivation by phosphorylation, pRB and E2F become highly mobile in chromatin. The regulation of the affinity of pRB to chromatin may constitute a new mechanism of transcriptional control.

**Mdm2 and VHL** Complete immobilization, in this case within the nucleolus, was also observed for the ubiquitin ligases Mdm2 and VHL in FRAP experiments (ESM 1; Mekhail et al. 2005). Upon an activation stimulus, the nucleolus rapidly releases these enzymes from static detention, thereby restoring their high mobility profiles. These observations provide strong evidence that cells have evolved mechanisms to regulate molecular networks by reversibly switching proteins between mobile and static states (Mekhail et al. 2005).

**TFIIH** In tissue-derived primary cells such as post-mitotic neurons, hepatocytes, and cardiac myocytes, the transcription factor TFIIH is effectively immobilized on the chromatin for hours during transcription, whereas in proliferative cells, it has the same highly dynamic behavior as in cultured cells (Giglia-Mari et al. 2009). It was proposed that static chromatin binding of TFIIH may be established during differentiation- and cell lineage-specific transcriptional programs. Interestingly, stable chromatin binding was not irreversible in the post-mitotic cells because induction of local DNA lesions remobilized TFIIH, probably based on its requirement in DNA repair (Giglia-Mari et al. 2009).

**HSF** In Drosophila salivary glands, the heat shock factor (HSF; the transcription activator of hsp70) trimerizes and translocates from the nucleoplasm to chromosomal loci after heat shock (Yao et al. 2006). FRAP experiments showed a rapid exchange of HSF at chromosomal loci under non-heat shock conditions ($t_{1/2}=15$ s) but a very slow exchange after heat shock ($t_{1/2}>6$ min). Five minutes after the FRAP bleach pulse, ~50% of HSF was still immobilized on chromatin. The unbound pool of HSF was found by FCS to diffuse freely, indicating that this pool was not able to compete for chromatin binding (Yao et al. 2006).
Gal4 Nalley and colleagues developed a new variant of the chromatin immunoprecipitation assay that allowed a direct time-resolved observation of exchange between the transcription factor Gal4 and its DNA binding site on native promoters in yeast (Nalley et al. 2006). The assay revealed that after transcriptional activation, the Gal4–promoter complex was highly stable with a half-life of approximately 1 h. Again, this stability was established although unbound, competition–competent Gal4 molecules were present in excess amounts. During immobilization on its promoter, expression of the reporter gene occurred indicating that Gal4 functions via long-lived complexes with promoters during transcriptional induction (Nalley et al. 2006).

The nuclear envelope contains large pools of immobile proteins

Proteins of the nuclear lamina interconnect chromosomes with the nuclear envelope. The lamina harbors specific integral proteins including the lamin B receptor (LBR), lamina-associated polypeptides (LAPs), emerin, nurim, and MAN1. LBR, LAPs, and emerin bind to lamins A/C and B in vitro while LBR and LAP2b also bind chromatin via interactions with HP1 and the small DNA-binding protein BAF (Mekhail and Moazed 2010). Thus, a highly interconnected protein network provides multiple anchoring sites between chromatin and the nuclear envelope. Many of these proteins have meanwhile been subjected to FRAP analyses revealing stable incorporation of most of them into the nuclear lamina (ESM 1). Although mathematical modeling has not been employed yet, the extremely long recovery half-times (i.e., more than 3 h for lamin B, Moir et al. 2000) of nuclear lamina constituents suggest residence times in the hours range. This is particularly true for the nuclear pore complexes, some components of which persist for long periods of time (residence times up to 70 h) within well-defined spatial regions without dynamic exchange (ESM 1; Rabut et al. 2004). Because the nucleus also undergoes dramatic morphological changes, a completely static nuclear lamina is not beneficial. Transitions between a stable and a flexible lamina might be regulated by small pools of the dynamically exchanging lamina constituents (ESM 1). Posttranslational modifications may serve as an additional mechanism to alter nuclear lamina stiffness.

Nuclear bodies contain stable components

FRAP analysis of subnuclear domains has revealed that most of their component parts rapidly exchange with nucleoplasmic pools with residence times in the seconds range (Misteli 2007; Matera et al. 2009). As a consequence, it was thought that these structures are determined by the ratio of on-rate versus off-rate of their constituents (Misteli 2008). This assembly mechanism is certainly true for speckles and nucleoli (ESM 1), which usually form in response to particular functions (splicing factor assembly and rRNA synthesis, respectively). However, FRAP experiments and mathematical modeling have revealed residence times of up to 1 h for Cajal and PML body components (ESM 1).

There are six nuclear isoforms of the PML protein in human cells from which PML V showed an extremely long residence time (48 min) (ESM 1). All other isoforms show substantially faster exchange, although they share the same nuclear body-binding domains with PML V (Weidtkamp-Peters et al. 2008). When analyzed in nuclei devoid of any endogenous PML protein (PML−/− MEFs), fractions of PML II becomes completely immobile within nuclear bodies and the residence time of PML IV is extended to 1.5 h (Brand et al. 2010). These observations indicate that probably all PML isoforms have the potential to build a stable scaffold within nuclear bodies, and that during evolution PML V was selected to execute this function (Weidtkamp-Peters et al. 2008). In Cajal bodies, p80 coiling was identified as the component with the longest residence time both, in human cells (Dundr et al. 2004) and in Xenopus germinal vesicles (Handwerger et al. 2003; Deryusheva and Gall 2004). These data indicate that the architecture of PML bodies and Cajal bodies may be primarily dictated by protein interactions forming stable scaffolds (Handwerger et al. 2003; Deryusheva and Gall 2004; Brand et al. 2010). On the other hand, their functionality certainly involves stochastic encounter of specific, fast exchanging components (Dundr et al. 2004; Weidtkamp-Peters et al. 2008).

Replication and repair factories contain a few stable components

RNA transcription sites and DNA repair foci have been visualized in mammalian cells, and initial FRAP
studies indicated that they are stochastically assembled de novo in each round of transcription or repair from freely diffusible components (ESM 1; Houtsmuller et al. 1999; Dundr et al. 2002). DNA repair factor recruitment dynamics point towards diffusion-based DNA lesion recognition by individual components or small subcomplexes as the primary mode of DNA repair, instead of the recruitment of large preassembled holoenzymes (Luijsterburg et al. 2007; Dinant et al. 2009). Nevertheless, FRAP analyses also indicate that some replication and repair factors can reside on chromatin for extended periods of time (ESM 1). Data for ERCC1 for example suggest that this component is assembled at sites of DNA damage with other constituents of the nucleotide excision repair (NER) machinery into immobile holocomplexes during one repair event with a mean residence time of 4 min which is the estimated time of a single repair event (Luijsterburg et al. 2010). Furthermore, strip-FRAP on globally irradiated cells showed incomplete recovery of DDB2 for several hours after 16 J/m² UV exposure, indicative of a significant immobile fraction over a period of hours at damaged chromatin (Luijsterburg et al. 2007). Twenty-four hours after exposure, most UV-induced lesions have been removed by NER, suggesting full release of bound molecules when repair is completed. The UV-induced immobilization was also shown for the NER proteins TFIH, XPA, and ERCC1-XPF but was not observed for RAD52 group proteins (ESM 1; Houtsmuller et al. 1999; Essers et al. 2002; Politi et al. 2005). In UV-treated cells, the fraction of lesion-bound XPA molecules increases to maximally 35%, and the increase is directly proportional to the DNA damage load. The immobile pool of XPA at damaged chromatin remains over a period of 2 to 4 h post-UV exposure (Rademakers et al. 2003).

While cells in living tissues may never experience DNA damage loads comparable to the cell culture studies, the immobilization of DNA repair factors ERCC1, XPA, and DDB1 on chromatin suggests that stable protein scaffolds may functionally contribute to DNA repair pathways.

Proliferating cell nuclear antigen (PCNA) is a further example of stable chromatin attachment. FRAP revealed elevated residence times for this factor at sites of DNA replication and repair (Sporbert et al. 2002; Solomon et al. 2004a, b; Mortusewicz et al. 2006; ESM 1). As a consequence, it was proposed that PCNA does not “bring” other factors to sites of replication or repair. Instead, PCNA appears to act as a stationary loading platform that is reused over multiple DNA synthesis or repair cycles with PCNA-binding proteins associating transiently and subsequently dissociating rather than being part of one stable, multifunctional, processive replication machinery (Mortusewicz et al. 2008).

An extension of models for the assembly of macromolecular structures in the nucleus: the “induced stability” model

Current models of the functional organization of the cell nucleus mainly consider short-lived interactions between freely diffusing components (Misteli 2001b, 2007). Undoubtedly, these events play crucial roles in gene regulation and genome maintenance (Darzacq et al. 2005; Lukas et al. 2005; Dinant et al. 2009; Hübner and Spector 2010). However, the observation of many stable interactions in the nucleus (ESM 1), even at transcription start sites on chromatin (TFIIH, HSF, Gal4, see above), leads us to propose an extension of the model for the formation of macromolecular complexes in the cell nucleus (Fig. 6).

The revised model is based on initial rapid binding and unbinding of proteins on DNA. Subsequent formation of dimers or multimers or interactions with additional DNA-binding proteins can induce an increase in the overall residence time of the complex on chromatin. A similar effect may be realized by cooperative binding of several DNA-binding domains within the same protein, as recently demonstrated for linker histones (Stasevich et al. 2010). The more stable protein/chromatin complex may now serve as a platform for further binding partners. As the complex “grows,” the binding partners may increase their residence time on chromatin through multiplication of binding contacts. Chaperones may support the addition/incorporation of specific components as has been shown for the centromere complex (Foltz et al. 2009; Shuaib et al. 2010) or nucleosome (dis)assembly (De Koning et al. 2007; Park and Luger 2008a, b). Complex-induced alterations in chromatin structure may also contribute to increased stability (not shown in the model). All components, including the initial proteins, may exchange with short residence times at the stable core (Fig. 6a), as in chemical self-organizing systems.
Fig. 6 Model for the assembly of macromolecular complexes in the cell nucleus. a Formation of a chromatin-binding protein complex. Protein A binds transiently to DNA with very short residence times (1). By dimer formation or interaction with a second DNA-binding protein the overall residence time increases (2). Chromatin-association may be further increased by homo- or hetero-multimerization leading to various degrees of stability of the complex (3 and 4). The assembly of these complexes may be supported by chaperones. At these more or less stable protein/chromatin complexes, further binding partners (X, Y, Z) but also sub-fractions of the endogenous components (A, B, C) may exchange with short residence times (5). b Formation of a macromolecular protein complex. Protein A with homo-oligomerization properties assembles into small scaffolds (1, 2) which may serve as seeds for attachment of additional binding partners (3). The resulting intermediates may consist of the initial scaffolding protein only or may also contain additional components with scaffolding properties (B, C). Stable scaffolds may arise through accumulation of additional binding partners (4). The assembly of intermediates and stable scaffolds may be supported by chaperones. Stable scaffolds acquire functional properties by transient retention of a specific set of nuclear proteins (X, Y, Z) (5). Even a subpopulation of the scaffold components may exchange rapidly at the macromolecular structure depending on the number and strength of binding events upon contact (5).
(Aniansson and Wall 1974; Diekmann 1979). This model describes active transcription, replication, and repair complexes but also heterochromatin, cohesin, telomere, and centromere assembly.

An analogous model may also hold for the assembly of macromolecular protein complexes in the interchromatin space. A protein with homooligomerization properties, or interactions between different proteins, can lead to the assembly into small scaffolds onto which additional binding partners or further scaffolding proteins may attach (Fig. 6). The accumulation of additional binding partners may lead to larger, more stable scaffolds. These larger structures could acquire functional properties by transient retention of a specific set of nuclear proteins. As above, a subpopulation of scaffold components may exchange rapidly at the macromolecular structure depending on number and strength of binding events upon contact (Fig. 6). This model likely holds for the assembly of PML nuclear bodies, Cajal bodies, snRNPs, speckles, pre-nucleolar bodies, and the nuclear envelope including nuclear pore complexes. Chaperones may also be involved in the assembly process. For example, the survival of motor neurons complex can act as a chaperone for the assembly of snRNPs (Battle et al. 2006), although in vitro U1-snRNPs self-assemble (Hamm et al. 1987). Unlike Cajal bodies where most, if not all components are part of a large interaction network capable of Cajal body induction (Kaiser et al. 2008), formation of PML nuclear bodies require PML protein since typical PML body components are disperse in PML-negative cells (Ishov et al. 1999). Even quantitative expression of Sp100 failed to recruit PML body components to Sp100 nuclear bodies in PML-negative cells (Schwanitz and Hemmerich, unpublished observations).

The formation of stable complexes also raises problems. Immobilization requires tight control because it can lead to “pathological immobility” as shown for nuclear inclusions seeded by the polyQ proteins ataxins or huntingtin, which immobilize not only themselves but also important nuclear proteins, such as CBP (Chai et al. 2002; von Mikecz 2009). Therefore, the models described above raise the fundamental question how the stability of a complex is controlled. The regulation of the stability of a complex in the nucleus may be conducted through posttranslational modifications or a “stabilizing” factor (Fig. 7). Such alterations of the complex may be regulated in a cell cycle-dependent manner as proposed for many components of the centromere/kinetochore complex (Hemmerich et al. 2008). We, therefore, propose an “induced stability” mechanism which allows controlled switching between dynamic and stable complex formation (Fig. 7).

**Self-assembly, self-organization, and assisted assembly in the nucleus**

Are the slowly recovering FRAPs for some nuclear components evidence for self-assembled nuclear structures? Self-assembly and self-organization are the main driving forces to create living systems from single molecules (Kirschner et al. 2000; Fraden and Kamien 2000; Gabora 2006; Pelesko 2007; Karsenti 2008). With respect to nuclear organization and function, self-organization was proposed to be the major underlying mechanism (Misteli 2001a, b, 2009; Matera et al. 2009; Rajapakse et al. 2010). Self-assembly and self-organization are often used interchangeably and sometimes in association. In current literature, the boundary between these two mechanisms is rather elusive and elastic (Bensaude-Vincent 2009).

Self-assembly implies spatial structuring as a result of free energy minimization in a closed system (John and Bär 2005; also called “static self-assembly,” Pelesko 2007). Thus, self-assembly drives the system towards the thermodynamic equilibrium, as for example in phase separation of lipids or the stable assembly of virus particles from coat protein monomers (Makowski 1980). Tobacco mosaic virus can self-assemble in a test tube starting only with its basic protein and nucleic acid components (Fraenkel-Conrat and Williams 1955; Butler and Klug 1978). Also, the bacterial ribosome self-assembles. However, the large and the small subunit must be allowed to assemble separately first. Once both subunits have assembled in separate test tubes, they can be combined forming a fully functional ribosome (Nomura 1973).

Self-assembly is a process in which components, either separate or linked, spontaneously form ordered aggregates (Whitesides and Boncheva 2004; Bensaude-Vincent 2009), although every single step is reversible. Nuclear structures that can be generated spontaneously by self-assembly in vitro include complete nuclei, the nuclear envelope, nuclear pore complexes, subcomplexes of the replication machinery, kinetochore subcomplexes, U1-snRNPs, NuMa scaffolds, prenucleolar...
bodies, and PML bodies (Table 1). In addition to the classical four forces (H-bonds, van der Waals forces, hydrophobic, and charge interactions), nonspecific (entropic) forces also influence these self-assembly processes. Entropic forces can induce the formation of human chromosome territories and position them appropriately in the nucleus (Finan et al. 2010). Since self-assembled structures form spontaneously and are stable, FRAP of their components might be expected to yield slow recoveries.

Self-organization requires a situation far away from thermodynamic equilibrium in a steady state and is possible only in open systems with an external energy source (Gerhart and Kirschner 1997; Toussaint and Schneider 1998; John and Bär 2005; Halley and Winkler 2008). Self-organization may also be regarded as “dynamic self-assembly” (Pelesko 2007). Self-organizing complexes and self-assembly processes can also be controlled efficiently either by posttranslational modifications or by chaperone action. The

| Table 1 Evidence for self-assembly of nuclear structures |
|---------------------------------------------------------|
| Self-assembled nuclear structure | Biochemical assay | Reference |
|---------------------------------|-------------------|-----------|
| Nucleus-like structures         | Xenopus cell free extract | Forbes et al. 1983; Zhao et al. 2000 |
| Nucleosome                      | Biochemically purified components | Ruiz-Carrillo et al. 1979 |
| Nucleosome                       | Xenopus cell free extract | Noll et al. 1980; Daban and Cantor 1982 |
| Chromatin fibers                | Biochemical purification | Newport 1987; Sheehan et al. 1988 |
| Mitotic chromosomes             | Xenopus cell free extract | Leforestier et al. 2001 |
| Nuclear lamina                   | Biochemical purification | Lohka and Masui 1983; Aebi et al. 1986; Glass and Gerace 1990 |
| Nuclear envelope                 | Xenopus cell free extract | Newport 1987; Sheehan et al. 1988 |
| Nuclear envelope                 | CHO cell free extracts | Burke and Gerace 1986 |
| Nuclear pore complex             | Xenopus cell free extract | Newport 1987; Sheehan et al. 1988 |
| Nuclear pore complex             | Bacterially expressed components | Lutzmann et al. 2002 |
| Replication machinery           | Xenopus cell free extract | Blow and Laskey 1986; Newport 1987 |
| Replication factor C            | Bacterially expressed | Uhlmann et al. 1996 |
| Kinetochore subcomplexes        | Bacterially expressed | Gestaut et al. 2010 |
| Dam1 complex                     | Purified components | Westermann et al. 2005 |
| Mis12 complex                    | Bacterially expressed components | Kline et al. 2006 |
| U1-snRNP                         | Xenopus cell free extract | Hamm et al. 1987 |
| NuMa scaffolds                   | Bacterially expressed | Harborth et al. 1999 |
| Prenucleolar bodies              | Xenopus cell free extract | Bell et al. 1992 |
| PML-like bodies                  | Bacterially expressed | Kentsis et al. 2002 |
assembly of NuMa scaffolds in the nucleoplasm for example is tightly controlled by phosphorylation (Saredi et al. 1997). Similarly, nuclear lamina self-assembly is regulated by phosphorylation and methylation of lamins (Gerace and Blobel 1980; Chelsky et al. 1989). SUMO-attachment to PML (which harbors three SUMOylation sites) is a requirement for PML nuclear body assembly (Ishov et al. 1999; Duprez et al. 1999; Shen et al. 2006), and the number of SUMO modifications regulates the residence time of PML molecules at nuclear bodies, ranging from seconds to hours (Weidtkamp-Peters et al. 2008; Brand et al. 2010). Gradients are used in a number of biological systems to transmit spatial information over a range of distances. Intracellular gradients utilize intrinsic auto-regulatory feedback loops and diffusion to establish regions of activity, for example within the mitotic cytosol (reviewed in Fuller 2010). These intracellular gradients bear several hallmarks of self-organizing biological systems that designate spatial information during pattern formation. Since self-organized systems are dynamic, FRAP of their components might be expected to yield fast recoveries. However, our model for nuclear assembly discussed in the preceding section proposes that initially dynamic assemblies can become more stable over time or during specific phases of the cell cycle, yielding eventually slower FRAP recoveries for some components. Thus, slow FRAP recoveries could also arise from self-organization or a mixture of self-organization and self-assembly.

In biological systems, it is obviously very difficult to clearly separate self-assembly from self-organization. In vitro, self-assembly and self-organization can act in concert to create functional large molecule aggregates (Cuccia et al. 2002; Drain 2002). For example, purified DNA can be assembled in the test tube into structures that closely resemble cell nuclei. By combining naked DNA, cytosol, and light membrane fractions from Xenopus egg extracts, complete nuclei can be formed, including nuclear membranes with pore complexes, and these reconstituted nuclei are capable of normal nuclear processes (Laskey and Leno 1990; Cross and Powers 2008). The assembled nuclei are morphologically indistinguishable from normal eukaryotic nuclei. Nuclear assembly involves discrete intermediate steps, including nucleosome assembly, scaffold assembly, and nuclear envelope assembly, indicating that during reconstitution nuclear organization is assembled one level at a time (Newport 1987). The nuclear envelope contains functional, self-assembled nuclear pore complexes (Sheehan et al. 1988). After envelope assembly, these artificial nuclei are even capable of DNA replication (Newport 1987). Notably, nuclear lamina and envelope assembly in vitro do not require energy (Burke and Gerace 1986), a hallmark of the self-assembly process. The inherent ability of a primitive genome to induce its own inclusion in a membrane has even led to a model of the origin of eukaryotic cells solely based on self-assembly mechanisms (de Roos 2006).

We suggest that both self-assembly and self-organization contribute to the formation and function of nuclear networks (Table 2). The cellular thermodynamic ground state is dictated by minimizing the free energy of intracellular matrices based on self-assembly mechanisms (Fraden and Kamien 2000). Depletion attraction may support self-assembly. Depletion attraction occurs when aggregation of large complexes increases the entropy of the system through an increase of small crowding molecules within the system (Marenduzzo et al. 2006a). It has been demonstrated in vitro, that depletion attraction can be used to control the self-assembly of colloidal particles (Yodh et al. 2001). Theoretical approaches and experimental evidence suggest that the same mechanism helps to organize chromatin architecture in living cells (Marenduzzo et al. 2006b; Misteli 2009; Finan et al. 2010, this issue). Our proposal is supported by in silico analyses: When the formation of protein patterns near membranes of living cells is analyzed by mathematical modeling, self-assembly and self-organization eventually lead to

| Table 2 | Potential assembly mechanisms in living cells |
|---------|---------------------------------------------|
| **Self-assembly** | **Self-organization** | **Assisted assembly** |
| Nuclear lamina | Chromosome territory | Nucleosome |
| Nuclear pore complex | DNA Replication sites | Centromere |
| Nuclear envelope | DNA Repair sites | |
| PML nuclear bodies | Transcription sites | |
| Prenucleolar bodies | Heterochromatin | |
| snRNP pre-complexes | Telomere | |
| Centromere pre-complexes | Nucleolus | |
| NuMa | Cajal bodies | Speckles |
similar patterns (John and Bär 2005). However, their evolution occurs on different length and time scales. Self-assembly produced periodic protein accumulations below 0.1 μm in a few seconds followed by extremely slow coarsening, whereas self-organization resulted in a pattern of 100 μm within a few minutes (John and Bär 2005). These observations support a model in which both processes contribute to particular nuclear functions depending on size and required activity. Self-assembly is required for the formation of kinetochore subcomplexes (Kline et al. 2006; Gestaut et al. 2010), which are then incorporated into the centromere by self-organization (Westermann et al. 2005). Similarly, nucleoporins self-assemble to pre-complexes before stable incorporation into the nuclear pore (Lutzmann et al. 2002). Self-assembly in living cells may also be crucial to create localized sites of specific enzymatic activity. For example, enzyme self-assembly is observed in vitro (Gao et al. 2010) as well as in vivo (Noree et al. 2010). Self-assembly may, therefore, control enzymatic activity depending on the metabolic status. This may hold for PML nuclear bodies at which SUMO and ubiquitin pathways locally and, therefore, probably efficiently merge in the nucleus (Häkli et al. 2005; Lallemand-Breitenbach et al. 2008; Sharma et al. 2010).

For essential cellular processes which require precise timing at well-defined locations, stochastic self-assembly and self-organizing reactions might not be sufficiently effective. In these cases, assisting or helper molecules seem to be required to secure function. Nucleosomes spontaneously self-assemble in vitro from biochemically purified components (Ruiz-Carrillo et al. 1979). The in vivo assembly of nucleosomes (Corpet and Almouzni 2009) and the (re)positioning of nucleosomes at transcriptional start sites and gene boundaries (Mito et al. 2007; Dion et al. 2007) are regulated by chaperones, loading factors, chromatin remodeling enzymes, and histone modifiers (Fig. 8) (Park and Luger 2008a, b). Recent research dedicated to the question how and when histone variants are deposited into nucleosomes has for example revealed that Cenp-A incorporation into centromeric chromatin requires specific loading factors (Furuyama et al. 2006; Shuaib et al. 2010). In addition, some processes like the transport of larger proteins through nuclear pores will not happen stochastically but require molecular support by helper molecules like importins (Harel and Forbes 2004). Being mechanistically overlapping but clearly different from self-assembly and self-organization we would like to introduce the term “assisted assembly” for those complex formation processes in the nucleus that rely on helper molecules (Table 2). This model, of course, raises fundamental questions: When do helper molecules attach to their substrate during the cell cycle? Do they interact in the nucleoplasm or at the complex where the substrate will be incorporated in? Will depletion of the helper molecule destabilize the complex? The first two questions will be answered by fluorescence fluctuation

![Fig. 8 Nucleosome dynamics based on “assisted assembly.”](image-url)

During replication nascent DNA is loaded with histones into nucleosomes. The first step involves formation of an H3/H4 tetramer and two H2A/H2B dimers. In the second step ca. 80 bp of DNA stably associates with the H3/H4 tetramer together forming the tetrasome. Nucleosome assembly is completed when two H2A/H2B dimers combine with the tetrasome and ca. 40 bp of DNA (Ruiz-Carrillo et al. 1979). Adjacent nucleosomes then self-assemble to form chromatin (Noll et al. 1980). The rigid structure of the nucleosome needs to be dissociated when active biochemistry, such as transcription, repair or replication occurs, or if variant histones ($H_{\text{var}}$) need to be incorporated. These processes require remodeling enzymes, chaperones, modifiers, and energy (ATP).
microscopy of GFP-tagged chaperones in the future. The third question has been solved already for Cenp-A. At least one of its chaperones, HJURP, only appears at the centromere in early G1 when Cenp-A is incorporated into centromeric chromatin. Depletion of HJURP by siRNA abrogates Cenp-A incorporation, leading to kinetochore malfunction (Dunleavy et al. 2009; Foltz et al. 2009; Shuaib et al. 2010).

Conclusions

Chromatin-binding proteins are highly dynamic; they roam the nucleus in an energy-independent manner in search for high-affinity binding sites, and their residence times on chromatin are typically in the order of several seconds. This dynamic behavior is thought to play a major role in generating combinatorial protein complexes on chromatin which provide a mechanism to fine-regulate transcription, chromatin organization, and genomic plasticity (Misteli 2001a, b, 2008). By analyzing FRAP data of about 264 nuclear proteins, we identified nuclear factors that do not follow this rule of highly transient interactions, instead some of them exhibit residence times in the hours range (Fig. 5).

Thus, although dynamic transient interactions are a general and important property of nuclear proteins, they are not universal. Stable complexes may arise from dynamic ones through a mechanism we coined “induced stability.” The demonstration in living cells of two proteins (HA95 and lamin B) which are almost completely immobile throughout the nucleoplasm may even reanimate research into the nuclear matrix. In particular cases, helper proteins support correct complex assembly, a phenomenon we have termed “assisted assembly.” The idea of immobile nuclear entities in the nucleus was already put forward by Bazett-Jones and colleagues in 2001 (Hendzel et al. 2001). The combined action of self-assembly and self-organization processes may lead to nuclear structures which are stable for hours. It should be considered in the future that stable interactions also contribute to the outcome of physiological cues impacting on nuclear structure and function. Therefore, “immobile populations” inferred from FRAP experiments should be studied in more detail. More examples of stable nuclear protein networks may surface as the community continues to perform fluorescence fluctuation microscopy on hitherto unexplored nuclear components.

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