Nuclear Lamins A and B1: Different Pathways of Assembly during Nuclear Envelope Formation in Living Cells

Robert D. Moir, Miri Yoon, Satya Khuon, and Robert D. Goldman
Department of Cell and Molecular Biology, Northwestern University Medical School, Chicago, Illinois 60611

Abstract. At the end of mitosis, the nuclear lamins assemble to form the nuclear lamina during nuclear envelope formation in daughter cells. We have fused A- and B-type nuclear lamins to the green fluorescent protein to study this process in living cells. The results reveal that the A- and B-type lamins exhibit different pathways of assembly. In the early stages of mitosis, both lamins are distributed throughout the cytoplasm in a diffusible (nonpolymerized) state, as demonstrated by fluorescence recovery after photobleaching (FRAP). During the anaphase-telophase transition, lamin B1 begins to become concentrated at the surface of the chromosomes. As the chromosomes reach the spindle poles, virtually all of the detectable lamin B1 has accumulated at their surfaces. Subsequently, this lamin rapidly encloses the entire perimeter of the region containing decondensing chromosomes in each daughter cell. By this time, lamin B1 has assembled into a relatively stable polymer, as indicated by FRAP analyses and insolubility in detergent/high ionic strength solutions. In contrast, the association of lamin A with the nucleus begins only after the major components of the nuclear envelope including pore complexes are assembled in daughter cells. Initially, lamin A is found in an unpolymerized state throughout the nucleoplasm of daughter cell nuclei in early G1 and only gradually becomes incorporated into the peripheral lamina during the first few hours of this stage of the cell cycle. In later stages of G1, FRAP analyses suggest that both green fluorescent protein lamins A and B1 form higher order polymers throughout interphase nuclei.

Key words: nuclear envelope • mitosis • chromatin • intermediate filaments • green fluorescent protein

Introduction

The nuclear lamins are closely related to cytoplasmic intermediate filaments (IFs) and are classified as Type V IF proteins (Moir et al., 1995; Stuurman et al., 1998). The lamins are the major components of the nuclear lamina, a fibrous structure adjacent to the nucleoplasmic face of the nuclear membrane. The lamina is required for maintaining nuclear size and shape, and may also play a role in the organization of chromatin. In addition, the lamins form structures within the nucleoplasm that are distinct from the peripheral lamina. These include lamin foci that are associated with sites of DNA replication (Goldman et al., 1992; Bridger et al., 1993; Moir et al., 1994).

The lamin family of proteins is classified into Types A and B based primarily on their amino acid sequence and their expression patterns. In vivo, B-type lamins are expressed in all cells, while A-type lamins are expressed only in differentiated cells (Benavente et al., 1985; Stewart and Burke, 1987; Rober et al., 1989). There are two B-type lamins in somatic cells, B1 and B2, that are products of different genes (Hoger et al., 1990; Zewe et al., 1991). In most mammalian somatic cells, the A-type lamins include lamins A and C, produced by differential splicing of the same gene (McKeon et al., 1986; Fisher et al., 1986). During mitosis, there is evidence that lamin B isoforms remain associated with membranes when the nuclear envelope is broken down and therefore have different biochemical properties when compared with A-type lamins (Gerace and Blobel, 1980).

As is the case for cytoplasmic IFs, the nuclear lamins have a central rod domain distinguished by heptad repeats characteristic of an alpha-helical structure. This rod domain is responsible for the dimerization of the lamin polypeptide chains and is also required for higher order interactions of the lamin dimers into the polymerized structures that form the nuclear lamina (Stuurman et al., 1998). The lamins have a short NH₂-terminal non-α-helical domain that is also essential for assembly (Moir et al., 1991; Heitlinger et al., 1992). The COOH-terminal domain includes the nuclear localization sequence (NLS) as well as...
a recognition sequence for isoprenylation (CAAX sequence), thought to be required for proper targeting of lamins A and B into the nucleus and the nuclear envelope, respectively (McKeon et al., 1986; Fisher et al., 1986; Loewinger and McKeon, 1988; Vorburger et al., 1989; Holtz et al., 1989). In addition, there is evidence that lamins A and C have chromatin-binding domains based upon in vitro assays (Hoger et al., 1991; Glass et al., 1993).

During interphase, the lamina plays an essential role in maintaining the shape and stability of the nuclear envelope (Newport et al., 1990; Spann et al., 1997). In support of this, nuclei are very fragile when assembled in Xenopus laevis egg extracts from which the majority of lamins have been immunodepleted or in which the lamina has been disrupted with a dominant-negative mutant (Newport et al., 1990; Spann et al., 1997). Furthermore, DNA replication is inhibited in lamin-depleted or -disrupted nuclei (Newport et al., 1990; Meier et al., 1991; Ellis et al., 1997; Spann et al., 1997; Moir et al., 2000). When the assembly of the lamina is disrupted with a dominant-negative lamin mutant, the endogenous lamins form aggregates within nuclei and the replication factors PCNA and RFC colocalize with these aggregates (Spann et al., 1997). This result suggests that lamins are required for the elongation phase of DNA replication and is in agreement with the observation that lamin B and PCNA colocalize in nucleoplasmic foci during S phase in cultured mammalian cells (Moir et al., 1994).

After mitosis, the assembly of the lamins into a lamina takes place during the formation of the nuclear envelope in daughter cells. However, the steps involved in this assembly process and the molecular interactions of the lamins with other envelope components have yet to be defined. The role of the lamins in nuclear assembly has been tested using in vitro nuclear assembly systems and the results are conflicting (Gant and Wilson, 1997). For example, if a lamin antibody is added to Xenopus laevis interphase extracts to block lamin function, nuclear assembly does not take place in the presence of chromatin (Dabauvalle et al., 1991). Instead, large aggregates, resembling annulate lamellae and containing nuclear pore intermediates, assemble in the extract. Similar results have been reported after the addition of lamin antibodies to nuclear assembly extracts derived from Drosophila melanogaster embryos or when lamins are immunodepleted from these extracts (Ulitzur et al., 1992, 1997). Furthermore, when lamins are immunodepleted from nuclear assembly extracts from mammalian cells, nuclear envelopes do not assemble properly around chromatin templates (Burke and Gerace, 1986). In addition, the reduced expression of lamins due to partial insertional inactivation of a D. melanogaster lamin gene results in defects of nuclear envelope structure in cells of affected flies, including the formation of annulate lamellae-like structures in the cytoplasm (Lenz-Bohme et al., 1997). Finally, in some systems, a fraction of the nuclear lamins appears to bind to chromatin very early in the process of nuclear formation, as seen in immunofluorescence assays (Yang et al., 1997). These results suggest that the lamins are required for successful envelope assembly (Foisner, 1997).

In contrast, other experiments show that intact nuclear envelopes assemble around chromatin after the immunodepletion of lamins from X. laevis interphase extracts (Newport et al., 1990; Meier et al., 1991). Furthermore, scanning electron microscopic studies of assembling nuclei in normal Xenopus extracts suggest that the lamins accumulate after membrane/pore complexes form (Wiese et al., 1997). Finally, other immunofluorescence observations of fixed mammalian cells undergoing nuclear assembly have suggested that the lamina is assembled after the nuclear membrane and nuclear pores have been established (Chaudhary and Courvalin, 1993). These results suggest that the lamins are imported into the nucleus and assemble only after the nuclear membrane and pore complexes have formed and, therefore, do not play a role in the initiation of envelope assembly (Newport et al., 1990). The apparent contradictions of the immunodepletion results obtained from different laboratories appear to be related to the inefficiency of this method for removing lamins from nuclear assembly extracts. In support of this contention, it has been shown that small amounts of lamins remain after immunoprecipitation, which appear to be adequate for the initiation of nuclear envelope assembly (Lourim and Krohne, 1993a,b).

To address the role of the lamins in nuclear assembly after mitosis and entry into the G1 phase of the cell cycle, we have expressed green fluorescent protein (GFP)–lamin fusion proteins in cultured cells. The utility of this approach for studying the lamins has already been demonstrated for the A-type lamins (lamins A, AΔ10, and C) by Broers et al. (1999). All three of these constructs are correctly processed and incorporated into the lamina of living interphase CHO-K1 cells. This work primarily describes several interphase structures that are revealed by GFP-lamin proteins. These include an extensive system of intra- and transnuclear tubular structures throughout the interphase nucleus. From fluorescence recovery after photobleaching experiments, the authors concluded that the tubules as well as the lamina exhibit relatively slow subunit exchange. In contrast, a second interphase population has an even distribution throughout the nucleoplasm, but appears to be much more diffusible, as demonstrated by fluorescence recovery after photobleaching (FRAP) experiments. The authors also showed that lamin A is dispersed throughout the cytoplasm in metaphase and enters daughter cell nuclei in late cytokinesis.

In this study, we describe in detail the dynamic properties of both A- and B-type lamins from metaphase through nuclear assembly and into G1. We chose to focus on this period of the cell cycle, because the role of the lamins in nuclear assembly remains controversial, as described above. The use of GFP-lamins allows us to determine the temporal sequence of changes in the distribution and the assembly state of the lamins in living cells during a time when they undergo profound changes in their structure and assembly states. To this end, the fates of lamins A, B1 and C during mitosis and nuclear envelope assembly in daughter cells were examined in vivo. The results demonstrate that there are significant differences between the A- and B-type lamins during nuclear assembly after mitosis. Lamin B1 begins to associate with chromatin in anaphase/telophase as chromosomes reach the spindle poles in daughter cells and before some nuclear pore components can be detected. Therefore, it appears that lamin B plays an early role in the assembly of the nuclear envelope. In contrast, lamin A does not obviously become associated...
with chromosomes until after they have initiated the decondensation process in late telophase/mid-late cytokinesis. Furthermore, lamin B1 rapidly assembles into a relatively stable polymer as soon as it encloses the decondensing chromosomes in late telophase, as shown by FRAP, while lamin A is not incorporated into a stable structure in the peripheral lamina until much later in G1.

Materials and Methods

GFP-Lamin Constructs

The cDNAs for full-length human lamins A, B1, and C were cloned into the commercial GFP vector pEFGP-C1 (CLONTECH Laboratories, Inc.) (McKeon et al., 1986; Pollard et al., 1990) so that the GFP sequence was at the NH2 terminus of the lamin sequence. For lamin A, a BamHI-EcoRI fragment containing the entire coding sequence of prelamin A (Weber et al., 1989), as well as the 11 amino acid myc 9E10 epitope at the NH2 terminus (Evans et al., 1985) and a four amino acid linker, was cloned into the BglII-EcoRI site of pEFGP-C1. The human lamin C cDNA was cloned in an identical manner as lamin A. For lamin B1, a BamHI-EcoRI containing the entire coding region with a four amino acid linker was cloned into the BglII-EcoRI site of pEFGP-C1. In addition, lamin B1 was cloned into the cyan version of GFP (pECFP-C1; CLONTECH Laboratories, Inc.), and lamin A was cloned into the yellow version (pYFP-C1; CLONTECH Laboratories, Inc.) using the same strategy as was used for pEFGP-C1.

DNA sequences encoding only the COOH-terminal non-s-helical domain of lamin B1 were made by PCR using the full-length cDNA as a template. This construct encompassed residues 378–586 of the B1 sequence (LB1; Pollard et al., 1990). BamHI restriction enzyme sites were placed in the oligonucleotides used for the PCR to allow cloning of the fragments into pEFGP-C1. All mutant constructs were sequenced in their entirety using an ABI373 automated sequencer. Plasmids were purified by chromatography on QIAGEN columns or by CsCl gradient centrifugation.

Cell Culture and Transfection

An embryonic mouse epidermal cell line, PAM, was grown in DME with 10% fetal calf serum and antibiotics (100 μg/ml penicillin and streptomycin; Jones et al., 1982). The hamster kidney fibroblast line, BHK-21, was grown in DME containing 10% calf serum, antibiotics, and 10% tryptose phosphate broth, as previously described (Yoon et al., 1998).

The GFP-lamin fusion constructs were introduced into cells by electroporation. Cells at 50–80% confluency (~2–3 × 105 cells) were detached by trypsin treatment and resuspended in medium with 10 mM Hepes, pH 7.0. The cells were mixed with 7 μg of the GFP plasmid and 13 μg of carrier DNA (sonicated salmon sperm DNA; a gift of Dr. Sui Huang, Northwestern University, Evanston, IL) in an electroporation cuvette (BTX Inc.). An electroporator (Bio-Rad Laboratories) was used to deliver a pulse at 0.26 kV and 960 Ω. The cells were plated onto 22-mm No. 1 coverslips or onto dishes designed for use with the microscope stage temperature control system (Bioptechs). The transfection efficiency varied from 20 to 70%, depending on the construct used.

Microscopic Observations of Live Cells

Cells that had been transfected for 16–48 h were examined directly, or in some cases the cells were trypsinized and replated on standard or locator (Bellico Inc.) coverslips before examination 16–48 h later. For microscopic observations, the cells were transfected into Leibovitz’s L15 medium containing 10% fetal calf serum and 100 μg/ml penicillin and streptomycin. The coverslips were mounted onto glass slides using chips of cover glass as feet to prevent compression and sealed with a mixture of lanolin, beeswax, and vaseline. The cells were maintained at 37°C while on the microscope stage using an airstream stage incubator (ASI400; Nevtek). In some cases, cells were examined in Bioptechs dishes using the Bioptechs stage/objective temperature controller.

Most time-lapse observations of GFP-transfected cells were made with an LSM 410 confocal microscope (Carl Zeiss Inc.) equipped with a 100 × 1.3 NA oil immersion objective and an FITC filter set (excitation 488 nm, emission 515 nm). The observations relating to extraction of samples on microscopes used in the line scanning of the samples on the microscope were done using a 40x 1.0 NA oil immersion lens. Images were collected at time intervals using LSM software (Carl Zeiss Inc.). To minimize phototoxicity, the laser was set at 10–25% of full power and at

tenated at 10 or 30%. Phase contrast was used to identify mitotic stages. In addition, an LSM 510 was used to image pCFP-lamin B1 and pYFP-lamin A simultaneously in doubly transfected cells. The 458- and 514-nm lines of the argon laser were used at 25–50% power for CFP and YFP respectively. The appropriate emission filters were obtained from Chroma. FRAP was carried out as previously described (Yoon et al., 1998). In brief, bar-shaped regions were bleached using the line-scan function of the LSM 410 for 3–4 s at 488 nm with the laser set at 100% power with 1% attenuation. Fluorescence recovery was monitored by taking images at time intervals using 15% laser power with 10% attenuation. The time required for 50% recovery of the bleach zone (τ1/2) fluorescence was calculated using Metamorph imaging software as described previously (Yoon et al., 1998). In brief, the average fluorescence intensity (pixel value) of each postbleach image was equalized to that of the prebleach image to compensate for overall fading during image collection. A line along the area of the lamina rim was drawn for each image using the line tool function of Metamorph and the profile of fluorescence intensity along the line determined using the line-scan function.

Fluorescence intensity was also measured using the Metamorph software. The average intensity of the background was subtracted from optical sections from the middle of the nucleus. Specific areas (lamina rim or nucleoplasm) of the image were selected with the Regions tool. The mean of the pixel values in these regions was used to determine total fluorescence intensity.

Fluorescence energy transfer experiments (FRET) were carried out on an Axiovert microscope (Carl Zeiss, Inc.) equipped with the appropriate filter sets. Cells doubly transfected with pCFP-LB1 and pYFP-LA were excited with light in the CFP excitation range (420–440 nm) and the emission was sequentially collected with CFP (455–485 nm) and YFP (520–550 nm) filter sets (Chroma) mounted on an external filter wheel. Subsequently, an image of YFP-LA was collected after excitation in the YFP excitation range. Images were collected with a 63 × 1.4 NA oil immersion lens.

In some experiments, cells were extracted during microscopic observation. Transfected cells were plated into Bioptechs dishes as described above. Once a transfected cell was located, the culture medium was removed and IF extraction buffer (PBS containing 0.6 M KCl, 10 mM MgCl2, and 1% Triton X-100) was added to the dish, taking care not to alter the position of the cell within the field of view. Image capturing was initiated immediately before extraction and within 10 s after the addition of the IF extraction buffer.

Indirect Immunofluorescence and Western Blotting

Cells on coverslips were washed with PBS, and then fixed with 4% formaldehyde in PBS for 10 min at room temperature. After fixation, the cells were permeabilized with 0.5% Triton X-100 in PBS for 5 min at room temperature, and then rinsed with PBS. The rabbit polyclonal antibodies for lamins A/C and B have been described previously (Moir et al., 1994). In addition, monoclonal antibodies directed against nucleoporins, mAb14 (Babco) (Davis and Blobel, 1986), the nuclear pore protein NUP153 (a gift of Dr. Brian Burke, University of Calgary, Calgary, Alberta, Canada) (Doody et al., 1999), and lamin-associated protein 2 (LAP2) (a gift of Dr. Roland Foissner, University of Vienna, Vienna, Austria) (Dechat et al., 1998). A rabbit polyclonal antibody directed against LAP2β was also used in some studies (a gift of Dr. Kathy Wilson, Johns Hopkins University Medical School, Baltimore, MD) (Gant et al., 1999). Secondary antibodies consisted of lissamine rhodamine conjugated to donkey anti–rabbit IgG (Jackson ImmunolResearch Laboratories), fluorescein-conjugated goat anti–rabbit IgG and fluorescein-conjugated goat anti–mouse IgG (Molecular Probes). Fixed and stained cells were observed with either the LSM 410, LSM 510, or an Axioptoph (Carl Zeiss, Inc.) equipped for epifluorescence. Images were recorded as previously described (Yoon et al., 1998).

Western blotting was performed as described in Yoon et al. (1998), except that chemiluminescence (Amersham Pharmacia Biotech) was used to detect the reactive products. The antibodies used were rabbit polyclonal antibodies to lamins A/C and lamins B1/B2 (Moir et al., 1994) or a monoclonal antibody to GFP (CLONTECH Laboratories, Inc.).

Results

The Assembly and Organizational States of Lamins during Cell Division

GFP human lamins A, B1, and C (see Materials and Methods) were transfected into either PAM or BHK-21 cells.

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The expressed GFP fusion proteins were examined 16–48 h post-transfection by confocal or standard epifluorescence microscopy. Mitotic PAM cells retain the relatively flat appearance typical of dividing epithelial cells, making it easier to recognize each stage of mitosis. To determine the changes in lamin assembly states during these different stages, individual cells expressing GFP-lamins were followed from the metaphase/anaphase transition through cytokinesis and into G1 for several hours.

During metaphase, GFP-tagged lamins A, B1, and C are distributed diffusely throughout the cytoplasm, and are excluded only from the chromosome region (for example, Fig. 1). If photobleach zones are introduced into the cytoplasm at this stage, full fluorescence recovery takes place within a few seconds, before an image of the bleach zone can be obtained in a subsequent confocal laser scan (data not shown). This suggests that the lamins can move freely throughout the extrachromosomal cytoplasm.

Interestingly, there are temporal and organizational differences between B- and A-type lamins during the metaphase/anaphase transition through cytokinesis and into G1 (Fig. 2–4). In the case of lamin B1 in PAM cells (Fig. 2), the GFP signal intensifies in the peripheral region of chromosomes, mainly in the area closest to the spindle pole during mid to late telophase (Fig. 2, a and e). A few minutes later, as cleavage furrowing begins, the diffuse cytoplasmic fluorescence decreases, while the fluorescence intensity increases at the surface of the chromosomes. Within a short time, intense fluorescence is seen to completely enclose the decondensing chromosomes (Fig. 2, b and f). Lamin B1 always defines the perimeter of the decondensing chromatin as the nucleus grows during the transition from late cytokinesis into G1 (Fig. 2, c and g). At a slightly later stage, the cytoplasmic fluorescence is barely perceptible (Fig. 2, d and h). Similar patterns are seen when BHK-21 cells expressing GFP-lamin B1 are followed through mitosis (Fig. 3). However, in these cells, lamin B1 appears to accumulate at the surface of chromosomes in late anaphase or early telophase, which is earlier than seen in PAM cells. This accumulation, as indicated by

![Figure 1](image1.png)
![Figure 2](image2.png)
fluorescence intensity, is most obvious at the chromosome surface closest to the spindle poles (Fig. 3, a and d, and b and e). A few minutes later, as the chromosomes reach the poles and a cleavage furrow is initiated, the fluorescence intensity associated with the surface of the chromosomes increases dramatically (Fig. 3, c and f). In addition, some bright lamin B1 foci can be seen in the cytoplasm of these cells throughout mitosis (Fig. 3, a–c). Similar cytoplasmic foci have been previously reported in dividing cells by indirect immunofluorescence (Chaudhary and Courvalin, 1993; Georgatos et al., 1997).

The organizational changes exhibited by GFP-lamin A are different from GFP-lamin B1 in PAM cells. Most obviously, lamin A initially accumulates throughout the region occupied by decondensing chromosomes in daughter cells (Fig. 4, a–f), rather than initially concentrating in the peripheral regions, as is the case for the B-type lamin. The lamin A fluorescence has a granular appearance during this time. Similar patterns of lamin A are seen in nontransfected PAM cells undergoing mitosis when examined by indirect immunofluorescence (compare Fig. 4, c with g). Cells expressing GFP-lamin C are indistinguishable from of G1 cells shows primarily nucleoplasmic staining with relatively little lamin A/C staining in the cytoplasm or in the region of the nuclear lamina (g). GFP-lamin C shows a similar pattern as GFP-lamin A in early G1 (h, compare with c). The differences between B- and A-type lamins in early G1 distribution are seen in doubly transfected cells expressing CFP-laminB1 (blue) and YFP-laminA (green, i). Lamin A is found throughout the nucleoplasm and lamin B1 exclusively at the rim, but there is some overlap at the rim of the two proteins. Bars, 10 μm.
GFP-lamin A with respect to the lamin fluorescence after its accumulation in nuclei (compare Fig. 4, h with c). The different distributions of A- and B-type lamins immediately after nuclear assembly in early G1 are most clearly illustrated in cells expressing both pCFP-LB1 and pYFP-LA (Fig. 4 i). In this case, the lamin A signal is found throughout the nucleoplasm without an obvious concentration at the rim, and lamin B1 is found only at the periphery. However, the two proteins do overlap at the periphery (Fig. 4 i).

In addition, lamin A shows an association with chromatin at a later stage of cell division than does lamin B1. For example, lamin A fluorescence does not appear to become concentrated in the nuclear region until late cytokinesis as indicated by a well-developed cleavage furrow (11 min after metaphase/anaphase transition), after lamin B1 becomes associated with chromosomes (5–7 min after anaphase begins; compare Fig. 4, a and d, with 2, b and f, and 3, c and f).

The timing of the association of B- and A-type lamins with chromosomes was also compared with other components of the nuclear envelope. Nontransfected PAM cells were fixed and prepared for double indirect immunofluorescence with both a lamin B antibody that reacts with both lamins B1 and B2 (Moir et al., 1994) and an antibody that reacts with a set of five related nuclear pore components (mb414; Davis and Blobel, 1986). In telophase cells, when lamin B just begins to accumulate in the poleward region of chromosomes, no significant increase in nucleoporin staining is seen in association with these chromosomes (Fig. 5, a and b). At this stage, the nucleoporins appear to remain distributed throughout the cytoplasm. This suggests that lamin B becomes concentrated at the surface of chromosomes before some of the major nuclear pore components. At slightly later times, both lamin B and nucleoporin staining are concentrated in the peripheral region of the decondensing chromosomes (Fig. 5, c and d). Cells in the same stages of the cell division process were also double labeled with lamin A/C and the 414 antibody.

In this case, the nucleoporin antibody became obviously concentrated at the perimeter of decondensing chromosomes before the lamin A/C antibody stained the chromosomal region (Fig. 5, e and f). At later times in G1, both lamins A/C and nucleoporins become concentrated in the nucleus (Fig. 5, g and h). Cells could not be found that showed increased lamin A/C staining in the nuclear region in the absence of nucleoporin staining during cell division and the early phases of G1.

These observations were extended using other nuclear pore and nuclear membrane markers. PAM cells were stained with both an antibody to the nuclear pore protein NUP153 (Bodoor et al., 1999) and the lamin B antibody. In contrast to the staining seen with the 414 antibody, NUP153 appears to become concentrated at the spindle pole side of the chromosomes before lamin B (Fig. 5, i and j). From these results, we infer the order of initial chromatin association to be NUP153, lamin B, 414-reactive nucleoporins, and lamin A/C. This is consistent with previous results suggesting sequential targeting of nuclear pore proteins (Bodoor et al., 1999; but see Haraguchi et al., 2000).

In addition, the integral membrane protein LAP2β (Dechat et al., 1998) also appeared to aggregate around chromatin before lamin B (Fig. 5, k and l). From these results, we infer the order of initial chromatin association to be LAP2β (i), lamin B, 414-reactive nucleoporins, and lamin A/C. This is consistent with previous results suggesting sequential targeting of nuclear pore proteins (Bodoor et al., 1999; but see Haraguchi et al., 2000).

In this case, LAP2β has almost completely accumulated around chromatin before lamin B. The nonmembrane-bound form of LAP2, LAP2α, showed similar staining (data not shown). We also did these experiments using the intrinsic GFP-laminB1 fluorescence, without antibody staining, in conjunction with the 414, NUP153, and LAP2 antibodies (data not shown) and obtained identical results.

Properties of Lamin B1 during Nuclear Assembly

To assess the state of lamin B1 polymerization during nuclear assembly, FRAP studies were carried out in trans-
fected cells. Dividing cells in the metaphase-to-anaphase transition that had not yet shown an accumulation of GFP-lamin B1 at the surface of chromosomes (as indicated by an increase in fluorescence intensity) were identified and followed for up to 3 h into G1. At various times in the nuclear assembly process, bar-shaped bleach zones across entire nuclei were introduced in either one or both daughter cells. The rate of recovery for each bleach zone was monitored by capturing images at regular time intervals, and an average fluorescence recovery rate ($t_{1/2}$) was calculated for the bleach zone in the lamina rim. Typical results are illustrated in Fig. 6. For each FRAP experiment, three images are shown: images taken immediately before bleaching, immediately after bleaching, and an image taken during the recovery period. In Fig. 6, a–c, a bleach zone was introduced ~10 min after GFP-lamin B1 had first enclosed the decondensing chromosomes in daughter cells. The bleach zone appeared to recover completely within 10 min. This value is approximately the same as seen for normal vimentin IF in interphase (Yoon et al., 1998) and is also much greater than that seen for assembly-incompetent GFP-lamin B1 (see below). This suggests that lamin B1 starts to form a stable polymer almost immediately upon associating with the surface of the chromosomes. However, the FRAP rate was difficult to determine because nuclear shape, as defined by the peripheral lamin B1 fluorescent signal, was irregular and changed rapidly during the earliest stages of nuclear assembly. In general, nuclear shape appeared to stabilize within 20–30 min after lamin B1 enclosed chromatin, although the nuclear diameter continued to increase as cells progressed in G1. FRAP experiments during this time period gave a $t_{1/2}$ of 29 ± 15 min ($n = 8$; Fig. 6, d–f).

In FRAP studies carried out on daughter PAM cells 45–60 min after lamin B1 enclosure, recovery was very slow and bleach zones remained for more than 3 h into G1 (Fig. 6, g–i). We used the partial fluorescence recovery attained at the 3-h time point to estimate the $t_{1/2}$ to be ~120 ± 40 min ($n = 8$). This value suggests that lamin B1 has formed a polymer in the region of the lamina that undergoes relatively slow subunit exchange by this stage of G1. This slow recovery time of the peripheral lamina is similar to that seen in typical interphase cells chosen at random. In these cells, complete recovery of photobleach zones takes longer than 4 h (see Figs. 10, d–f, and 11), and the average $t_{1/2}$ appears to be >180 min ($n = 3$; Figs. 10 and 11). These FRAP experiments suggest that lamin B1 assembles into a relatively stable polymer rapidly after it associates with the surface of chromosomes in the late telophase/early G1 phases of the cell cycle.

As a further indicator of the state of polymerization of lamin B1 during nuclear assembly, and early G1, cells were extracted with a high salt/nonionic detergent buffer (IF buffer; see Materials and Methods) while being observed by confocal microscopy. This IF buffer has been used extensively to assess the polymeric state of IF proteins in cells (see Yoon et al., 1998). Individual transfected cells were identified in the metaphase/anaphase transition, followed...
through lamin B1 enclosure of decondensing chromosomes and into the G1 phase. At different stages, the culture medium was removed and rapidly replaced with IF buffer on the microscope stage such that the same live cell could also be examined within 10 s after extraction (Fig. 7). When cells were extracted 10 min after enclosure of decondensing chromosomes in daughter cells, no remaining GFP-lamin B1 signal could be detected (data not shown). In daughter cells extracted 15–30 min after enclosure, some lamin B1 is removed or solubilized and the remaining lamin B1 is seen as an array of bright foci, or occasionally short lines, located at the nuclear periphery (Fig. 7 b, arrow). In contrast, the vast majority of GFP-lamin B1 remained as a continuous rim around the nucleus after extraction within 45–60 min after enclosure is initiated.

**Lamin B1 Polymerization Is Required for Its Targeting and Stable Integration into the Lamina**

To assess further the requirement for lamin B1 polymerization in lamina assembly during the earliest stages of nuclear assembly, PAM cells were also transfected with a GFP construct consisting only of the COOH-terminal domain of lamin B1 (termed LBT; amino acids 378–586). This fragment retains the NLS and the CAIM sequence (amino acids 583–586) that undergoes isoprenylation, but lacks the alpha-helical rod domain required for lamin polymerization. The NLS and CAIM motifs are required for proper lamin B targeting to the nuclear envelope (Loewinger and McKeon, 1988; Holtz et al., 1989; Kitten and Nigg, 1991; Frangioni and Neel, 1993; Hennekes and Nigg, 1994). Thus, this mutant lamin cannot polymerize independently or interact with endogenous laminas through rod domain, but it retains other sequences required for nuclear localization. When cells were examined within 20 min after the time when lamin B1 enclosure of chromatin normally takes place, the mutant protein appears to be dispersed throughout the nucleoplasm (Fig. 8 a). In contrast, wild-type lamin B1 is located almost exclusively at the nuclear periphery (Fig. 8 b). When bleach zones were introduced in cells expressing LBT at any stage of cell division or interphase, the fluorescence recovered before an image of the bleached area could be obtained in a subsequent confocal laser scan (<5 s). This suggests that the mutant protein is in a diffusible, nonpolymerized state (data not shown). In addition, the mutant protein can be completely extracted with IF buffer from early G1 cells, as well as cells at all other stages of the cell cycle (data not shown). These results further show that lamin B1 polymerization is a critical, early step in the establishment and maintenance of a stable nuclear lamina during nuclear envelope assembly after mitosis.

**Lamin A follows a Different Pathway during Nuclear Assembly**

Observations of lamin A during nuclear assembly in PAM cells indicate that it first accumulates throughout the region occupied by chromatin and is not immediately concentrated at the periphery (Fig. 4). This distribution is in contrast to that seen with lamin B1 (see above) and persists without evidence of a lamina rim pattern for ~60 min after GFP-lamin A begins to accumulate in assembling nuclei (Fig. 4 c). If live PAM cells showing this lamin A distribution are followed for an additional 90 min into G1, then the nucleoplasmic GFP-lamin A signal decreases as the lamina becomes more prominent at the nuclear periphery (Fig. 9, a and b). In addition, lamin A become more obvious within the nucleoplasm (Fig. 9 b). We also observed a similar pattern of lamin A organization in BHK-21 cells (data not shown). GFP-lamin C shows an identical organization, in contrast to that seen with lamin B1 (see above) and persists dispersed throughout the nucleoplasm (Fig. 4). This distribution is in contrast to that seen with lamin B1 (see above) and persists without evidence of a lamina rim pattern for ~60 min after GFP-lamin A begins to accumulate in assembling nuclei (Fig. 9 a). In contrast, wild-type lamin B1 is located almost exclusively at the nuclear periphery (Fig. 8 b). When bleach zones were introduced in cells expressing LBT at any stage of cell division or interphase, the fluorescence recovered before an image of the bleached area could be obtained in a subsequent confocal laser scan (<5 s). This suggests that the mutant protein is in a diffusible, nonpolymerized state (data not shown). In contrast, wild-type lamin B1 is located almost exclusively at the nuclear periphery (Fig. 8 b). When bleach zones were introduced in cells expressing LBT at any stage of cell division or interphase, the fluorescence recovered before an image of the bleached area could be obtained in a subsequent confocal laser scan (<5 s). This suggests that the mutant protein is in a diffusible, nonpolymerized state (data not shown). In addition, the mutant protein can be completely extracted with IF buffer from early G1 cells, as well as cells at all other stages of the cell cycle (data not shown). These results further show that lamin B1 polymerization is a critical, early step in the establishment and maintenance of a stable nuclear lamina during nuclear envelope assembly after mitosis.
from telophase to early cytokinesis and followed for an additional 60–120 min as they entered the G1 stage. Bar-shaped bleach zones were made across the nuclei of one or both daughter cells at various time points. In early G1 nuclei containing both nucleoplasmic fluorescence and a distinct, more intensely fluorescent lamina region (60–120 min after cytokinesis), the bleach zone recovers immediately in the nucleoplasm, but much more slowly in the rim region (Fig. 9 a, arrows). The same cell was immediately extracted in IF buffer while being viewed on the microscope stage and an image was captured 10-s later (e). The nucleoplasmic fluorescence is almost completely extracted. Bar, 5 μm. FRET analysis was also performed on cells in G1 (e–j). Cells were doubly transfected with pCFP-LB1 and pYFP-LA and examined using a FRET filter setup, as described in Materials and Methods. In cells early in G1 (120–180 min after telophase), the CFP-LB1 (e) was able to activate YFP-LA, resulting in a FRET signal (f), implying these molecules interact. Furthermore, cells very early in G1 (<90 min after telophase), when lamin A is largely nucleoplasmic (YFP-LA; j) also have a FRET signal, indicating an interaction at this time (h and i). Bar, 10 μm.

Figure 9. Mitotic PAM cells were identified and allowed to proceed until ~60–90 min after telophase. At this time, both the nucleoplasm and lamina fluoresce. When daughter cell nuclei are photobleached at this time, the recovery in the nucleoplasm is so fast that a bleach zone cannot be detected in a subsequent confocal scan (a). However, the bleached areas in the lamina region recover much more slowly and remain detectable for up to 70 min and later. Only one daughter cell nucleus is shown due to the size of the nucleus. The overall nucleoplasmic fluorescence becomes less intense during this period of observation and the number of nucleoplasmic foci increases. Many of these foci are not continuous with the nuclear surface, as shown through a focus series of images using the confocal microscope (not shown). Bar, 10 μm. A live PAM cell in early G1 expressing GFP-lamin A was photobleached across the entire cell and an image was captured in the subsequent confocal scan (d). As expected for an early G1 cell, the bleach zone is apparent only at the lamina rim (arrows). The overall nucleoplasmic fluorescence becomes less intense during this period of observation and the number of nucleoplasmic foci increases. Many of these foci are not continuous with the nuclear surface, as shown through a focus series of images using the confocal microscope (not shown). Bar, 10 μm. A live PAM cell in early G1 expressing GFP-lamin A was photobleached across the entire cell and an image was captured in the subsequent confocal scan (d). As expected for an early G1 cell, the bleach zone is apparent only at the lamina rim (arrows). This same cell was immediately extracted in IF buffer while being viewed on the microscope stage and an image was captured 10-s later (e). The nucleoplasmic fluorescence is almost completely extracted. Bar, 5 μm. FRET analysis was also performed on cells in G1 (e–j). Cells were doubly transfected with pCFP-LB1 and pYFP-LA and examined using a FRET filter setup, as described in Materials and Methods. In cells early in G1 (120–180 min after telophase), the CFP-LB1 (e) was able to activate YFP-LA, resulting in a FRET signal (f), implying these molecules interact. Furthermore, cells very early in G1 (<90 min after telophase), when lamin A is largely nucleoplasmic (YFP-LA; j) also have a FRET signal, indicating an interaction at this time (h and i). Bar, 10 μm.
Figure 10. GFP-lamin A and B show intranuclear fluorescence in the majority of interphase PAM cells. (a–c) A live cell expressing GFP-lamin B shows nucleoplasmic (the lamin “veil”) and lamina fluorescence (a) with confocal optics. In b, the cell was photobleached, and after 240 min (c) the bleach zone could still be detected. The t½ for both the nucleoplasm and lamina is similar (~180 min, see text and Fig. 11). This veil can be seen throughout the nucleoplasm in a confocal through focus series (not shown). (d–f) Live PAM cells expressing GFP-lamin A have a more prominent nucleoplasmic veil. An optical section from an interphase cell expressing GFP-lamin A is shown just before (d), and just after (e) photobleaching. Note that the prominent bleach zone is retained across the entire nucleus, suggesting that this is not an early G1 cell (see text). In f, the same cell was extracted immediately after photobleaching with IF buffer and an image was captured. The veil fluorescence is greatly reduced, but lamina fluorescence appears unaltered. Bars, 10 μm.

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Discussion

In the work described here, GFP fusion proteins have been used to observe the dynamic properties of nuclear lamins during nuclear assembly in daughter cells. A recent report also described the properties of GFP lamin A fusion proteins, primarily in interphase nuclei (Broers et al., 1999). The results reported here describe for the first time the different pathways of assembly and the organization of A- and B-type lamins in living cells during mitosis, nuclear formation in daughter cells, and the G1 phase of the cell cycle.

When we follow cells expressing GFP-lamins, we observe that lamin B1 is targeted directly to the periphery of chromosomes before significant decondensation begins, and remains at the nuclear boundary. In contrast, lamin A is targeted to the nucleoplasm of newly formed nuclei and is not initially concentrated at the periphery of chromosomes. In PAM cells, lamin B1 begins the process of enclosure in telophase, as the chromosomes reach the spindle poles, before the decondensation process begins. In contrast, lamin A does not begin to associate with chromosomes until the late stages of cytokinesis after the decondensation process has been initiated. In BHK-21 cells, increased lamin B1 fluorescence is seen associated with chromosomes in mid-to-late anaphase/early telophase, earlier than seen in PAM cells. In both cell types, lamin B1 first accumulates on the spindle pole side of the chromosomes, in the region of the kinetochore, suggesting lamina assembly may be initiated at this site. Therefore, this may
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to enter the nucleus. Our results on the timing of recruit-
ported into the nucleus after the nucleoporins become as-
completed. The results also show that lamin A is trans-
ous accumulation of some nuclear pore proteins, implying
the nuclear envelope. In support of this, lamin B1 target-
also represent an early nucleating site for the assembly of
the nuclear envelope. In support of this, lamin B1 targeting
to the surface of chromosomes occurs before an obvi-
ous accumulation of some nuclear pore proteins, implying
that lamin B1 assembly begins before pore assembly is
completed. The results also show that lamin A is trans-
ported into the nucleus after the nucleoporins become as-
associated with the surface of decondensing chromosomes,
suggesting that lamin A requires functional nuclear pores
to enter the nucleus. Our results on the timing of recruit-

Figure 11. The fluorescence recovery rates for bleach zones in
the lamina rim of interphase cells are shown as an X-Y plot of
fluorescence intensity (pixel value) versus pixel position. The av-
average fluorescence intensity (pixel value) of each postbleach image
was equalized to that of the prebleach image to compensate
for overall fading during image collection. A line along the
bleached area of the lamina rim was drawn for each image using
the line tool of the Metamorph image analysis program and the
profile of fluorescence intensity along the line determined using
the line-scan function. The profiles obtained from different time
points were plotted and the bleach zone appears as a trough rela-
tive to adjacent unbleached areas. Fluorescence recovery was de-
determined by measuring the relative fluorescence recovery in the
bleached to the unbleached areas over time. (Top) Recovery for
an interphase cell expressing GFP-lamin B1 in interphase. Ap-
proximately 50% recovery occurs in 142 min. (Bottom) A cell at
interphase expressing GFP-lamin A. This bleach zone has under-
gone <50% recovery in 3 h.

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suggesting that lamin A requires functional nuclear pores
to enter the nucleus. Our results on the timing of recruit-
ment of nuclear envelope components are in agreement
with results from fixed cells, which suggest sequential
binding of different nuclear pore proteins to chromatin
(Bodoor et al., 1999). However, a recent study in live cells
indicates that NUP153, p62 (a 414-reactive protein), lamin
B receptor (LBR), and emerin associate with chromo-
somes almost simultaneously in early telophase (Hara-
guchi et al., 2000). These results also show that LBR and
emerin are initially punctate in their distribution around
chromatin and do not become uniform until later in telo-
phase. Therefore, it is tempting to speculate that the bind-
ing of lamin B1 that we detect at this same time in the cell
division process is required to establish LBR and emerin
distribution, through direct interactions or indirectly through
the formation of a lamin polymer (see below).

Different patterns of A- and B-type lamins in telophase/
early G1 have also been described in other cell types using
immunofluorescence. For example, the same overall dis-
tributions have been reported for A- and B-type lamins in
normal rat kidney cells during early G1 (Dechat et al.,
1998). Nucleoplasmic staining with antibodies directed
against lamin A/C has also been reported during early G1
in other cells (Goldman et al., 1992; Bridger et al., 1993;
Hozak et al., 1995; Neri et al., 1999). In addition, the re-
cent results with GFP-tagged lamins A, LAΔ10, and lamin
C in CHO cells indicate that they exhibit a nucleoplasmic
distribution during G1 (Broers et al., 1999).

In this study, we show that as cells progress through G1,
lamins A and B1 exhibit significantly different assembly
properties and locations. In the case of lamin B1, a higher
order polymer is established rapidly at the nuclear periph-
ery based on the fluorescence recovery rates of cells at this
time in the cell cycle. At the earliest stages of lamin B1 en-
closure around chromatin, photobleach zones recover in
~10 min. This value is similar to that obtained for poly-
merized interphase cytoskeletal IF networks (Vickstrom
et al., 1992; Yoon et al., 1998) and suggests that lamin B1 is
beginning to polymerize on the surface of decondensing
chromosomes almost immediately after mitosis. FRAP
rates increase in the first 60 min of G1, indicating that
lamin B1 is rapidly assembled into a higher order structure
after mitosis.

The importance of lamin B1 polymerization in the early
stages of lamina assembly is also supported by the expres-
sion of the mutant protein, LBT. The absence of the alpha-
elical rod prevents this mutant lamin from assembling the
coiled-coil dimers required for the subsequent steps in
lamin assembly, but it retains the NLS and isoprenylation
sites required for nuclear targeting and nuclear membrane
association (Peter et al., 1991). LBT remains distributed
uniformly in the nucleoplasm throughout the cell cycle,
and it does not appear to be targeted specifically to the
lamina (see Fig. 8). Furthermore, FRAP analysis shows
that this mutant protein remains in a diffusible state
throughout the nucleoplasm, demonstrating that lamin B1
polymerization, in addition to the NLS and CAAX signals,
is required to establish normal lamin B1 organization.

Lamin A exhibits dramatically different dynamic prop-
erties relative to lamin B1 during G1. Immediately after
mitosis and into the early phase of G1, lamin A is distrib-
uted throughout the nucleoplasm of the assembling nu-
cleus of both PAM and BHK cells. The fast fluorescence
recovery rates recorded at this time suggest that lamin A is present in a form that is more mobile and hence represents a lower order structure. For ~90–120 min after its initial targeting to the nucleoplasm, lamin A is gradually incorporated into the lamina, as shown by the shift in the intensity of the GFP-lamin A signal from the nucleoplasm to the nuclear periphery. During this time, lamin A exists in two populations with different polymeric states as shown by the FRAP rates and extraction properties.

The differences in lamins A and B1 distributions may reflect their different interactions with other nuclear envelope components. For example, it is known that B-type lamins are associated with nuclear envelope-derived membrane vesicles in mitotic cells (Gerace and Blobel, 1980; Burke and Gerace, 1986; Vigers and Lohka, 1991; Lourim and Krohne, 1993a; Meier and Georgatos, 1994; Foisner, 1997; Maison et al., 1997; Drummond et al., 1999). Therefore, lamin B1 would be expected to be localized peripherally during the early stages of assembly as part of the forming nuclear membrane. Furthermore, LAPs probably mediate the interactions of lamins with membranes (Foiser, 1997; Yang et al., 1997). The LAP family includes LAP2α and β, emerin and MAN1 (Lin et al., 2000). In particular, LAP2 has been implicated in the regulation of the early stages of nuclear assembly and also in the growth of the nucleus during G1 (Yang et al., 1997; Dechat et al., 1998; Gant et al., 1999; Vleck et al., 1999). Different fragments of LAP2 can inhibit either nuclear assembly or nuclear growth, perhaps reflecting the binding of this protein to chromatin or lamins (Yang et al., 1997; Gant et al., 1999).

LAP2a, a non–membrane-bound isoform, colocalizes with A-type lamins during nuclear formation and may specifically regulate the assembly of lamin A (Dechat et al., 1998). Since LAP2β may interact primarily with lamin B isoforms, the distributions of lamins A and B1 that we observe may reflect interactions with different forms of LAP2 during nuclear assembly (Dechat et al., 1998). Furthermore, it has been proposed that the membrane-bound LBR interacts with lamin B only (Meier and Georgatos, 1994; Ellenberg et al., 1997; Drummond et al., 1999), although this putative lamin B-LBR interaction has been questioned (Mical and Monteiro, 1998; Gajewski and Krohne, 1999).

The different distributions of lamins A and B1 may also be due to posttranslational modifications. For example, lamins A and B are isoprenylated at a conserved COOH-terminal cysteine (Holtz et al., 1989; Kitten and Nigg, 1991). The isoprenyl group remains on lamin B throughout the cell cycle (Firmbach-Kraft and Stick, 1993, 1995), but it is rapidly removed from lamin A by an endoprotease (Weber et al., 1989; Kilic et al., 1999). The mutation of the cysteine residue prevents isoprenylation and results in an exclusively nucleoplasmic distribution of lamin A (Holtz et al., 1989). However, other experiments using inhibitors of isoprenylation suggest that lamin A incorporation into the lamina is not affected by the inhibition of this posttranslational event (Dalton et al., 1995; Sasseville and Raymond, 1995). In our studies, it is most likely that the majority of the lamin A that we observe localizing to the nucleus immediately after mitosis is synthesized in the previous cell cycle (Gerace et al., 1984) and therefore would not be isoprenylated as a consequence of the proteolytic cleavage step. It is possible, therefore, that as new, isoprenylated lamin A is synthesized during G1, it interacts, perhaps by forming dimers or tetramers, with the nucleoplasmic lamin A synthesized in the previous cell cycle, resulting in the targeting of both populations to the envelope.

We have also obtained evidence in this study that lamins A and B can exist in a polymerized state within the nucleoplasm in late G1 or other interphase stages, as a veil of fluorescence that is distinct from the peripheral lamina. Both the lamin A and B veils are distributed throughout the nucleus, as shown by through-focus series in the confocal microscope (data not shown). Furthermore, very slow FRAP rates were obtained for the A- and B-type veils, as well as the lamina. In the case of lamin A, this is contrasted with the extremely rapid fluorescence recovery obtained in early G1 cells. However, throughout interphase, the structure of the veil appears biochemically distinct from the lamina, as indicated by a significant reduction in fluorescence intensity after extraction with IF buffer. The recent report using GFP–A-type lamins in CHO cells also describes intranuclear fluorescence, although these workers report that fluorescence recovery takes only 1.5 s (Broers et al., 1999). This discrepancy with our results, especially in the later phases of G1, could be explained by the stage of the cell cycle during which FRAP experiments were carried out. In comparison with our data, the result of Broers et al. (1999) would be expected from early G1 cells. It will be an important priority, therefore, to determine the relationships of the different intranuclear lamin structures to each other during specific stages of the cell cycle.

There are now a number of reports of nucleoplasmic lamin structures in fixed, nontransfected cells (Hozak et al., 1995; Hozak, 1996; Neri et al., 1999). In support of this, immunoelectron microscopy of extracted, fixed cells has revealed putative lamin filaments within the nucleus (Hozak et al., 1995). We have also observed lamin staining within the nucleoplasm using immunoelectron microscopy (data not shown). Based on these considerations, it is interesting to consider the possibility that the nuclear lamins, as a member of the IF family of proteins, form the framework for an extended and pervasive nucleoskeletal system that is continuous with the lamina during interphase (Hozak et al., 1995). The detailed organization, state of polymerization, or structure of this lamin-based nucleoskeleton remains to be explored. However, its existence has profound implications for many nuclear functions (Pederson, 1998, 2000a,b). Among these are the determination of nuclear size, shape, mechanical properties, the organization of chromatin, and DNA replication. In support of the latter, it has been shown that lamin B colocalizes with DNA replication centers in situ (Moir et al., 1994), and there is a significant amount of evidence showing that an intact nuclear lamina is required for the completion of DNA replication (Newport et al., 1990; Spann et al., 1997; Ellis et al., 1997; Moir et al., 2000). Therefore, the existence of a dynamic nucleoplasmic lamin network may ultimately prove to represent the infrastructure required for many important nuclear functions.

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