Systematic kinetic analysis of mitotic dis- and reassembly of the nuclear pore in living cells

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During mitosis in higher eukaryotes, nuclear pore complexes (NPCs) disassemble in prophase and are rebuilt in anaphase and telophase. NPC formation is hypothesized to occur by the interaction of mitotically stable subcomplexes that form defined structural intermediates. To determine the sequence of events that lead to breakdown and reformation of functional NPCs during mitosis, we present here our quantitative assay based on confocal time-lapse microscopy of single dividing cells. We use this assay to systematically investigate the kinetics of dis- and reassembly for eight nucleoporin subcomplexes relative to nuclear transport in NRK cells, linking the assembly state of the NPC with its function. Our data establish that NPC assembly is an ordered stepwise process that leads to import function already in a partially assembled state. We furthermore find that nucleoporin dissociation does not occur in the reverse order from binding during assembly, which may indicate a distinct mechanism.

Introduction

Nuclear pore complexes (NPCs) mediate all traffic of macromolecules across the nuclear envelope (NE). They are large protein assemblies composed of multiple copies of ~30 different proteins, the nucleoporins (Nups), which are organized in about 10 subcomplexes and arranged with eightfold symmetry. In metazoans, NPCs are stable throughout interphase (Daigle et al., 2001) but disassemble into their subcomplexes during mitosis. When the NE breaks down in pro/metaphase, most Nups become cytoplasmic and transmembrane Nups relocalize to the ER together with other nuclear membrane proteins (Ellenberg et al., 1997; Yang et al., 1997; Daigle et al., 2001; Beaudouin et al., 2002). Reassembly occurs during anaphase and telophase when the NE is rebuilt around chromatin.

In live cells, NE disassembly has been shown to start by partial disassembly of NPCs, with Nup98 leaving the NE early followed by dissociation of Nup153 and Nup214 before the NE is completely permeabilized. The membrane Nup POM121 dissociates from NE fragments only after permeabilization (Beaudouin et al., 2002; Lenart et al., 2003). In fixed cells, the nuclear basket Nup Tpr dissociates from the NE before Nup107 but later than Nup98 and Nup50 (Hase and Cordes, 2003).

More is known about the mechanism of postmitotic NPC assembly. In vitro studies of nuclear assembly in Xenopus laevis egg extracts have shed light on the essential role of the Ran-importin system, which regulates the release of several Nups from importin in proximity to chromatin, enabling them to reassociate and form NPCs (Harel et al., 2003a; Walther et al., 2003b). Several Nups bind to chromatin in early anaphase before membrane association (Belgareh et al., 2001; Walther et al., 2003a), where they have been postulated to form a prepore (Suntharalingam and Wente, 2003; Wozniak and Clarke, 2003; Rabut et al., 2004b). The mechanism of subsequent insertion into the membrane and full assembly of the NPC remains to be understood.

For some Nups, the order of reassociation with the reforming NE was investigated in various experimental systems, fixed cells of different mammalian species, or nuclei assembled in X. laevis egg extracts. Together, these data predict that the Nup107–160 complex, Nup153, Nup98, and POM121 bind during anaphase, followed by the Nup62 and Nup93 complexes, Nup358, and Nup214 in telophase, whereas Tpr and gp210 reassemble only in early G1 (for review see Burke and Ellenberg, 2002).

Evidence for structural disassembly and reassembly intermediates has been provided by field emission scanning electron microscopy. Porelike structures of different levels of complexity could be visualized in egg extract nuclei (Goldberg et al., 1997; Wiese et al., 1997; Wiese et al., 1997; Kiseleva et al., 2001) and a rough time course of the formation of these structures could be established in Drosophila melanogaster embryos (Kiseleva et al., 2001). Their protein composition remained, however, unclear.

Our current knowledge predicts that NPC disassembly and reassembly are ordered processes that proceed via a defined set of intermediates formed by sequential interactions of NPC.
subcomplexes. However, the precise order in which the different subcomplexes bind, the kinetics of the assembly events, and the functional state of the different intermediates are unknown. To address this, we systematically investigated the kinetics of mitotic NPC disassembly and reassembly by time lapse confocal microscopy in single dividing cells. Simultaneously, we monitored import competence of the nucleus. We analyzed a set of GFP-tagged Nups (Rabut et al., 2004a) representing eight different NPC subcomplexes. Our results show that NPC assembly is indeed a highly ordered process that proceeds in a stepwise fashion. Partially assembled NPCs were already import competent, which indicates that several Nups may not be required to reestablish import function. Regarding NPC disassembly, we found it to occur more rapidly than assembly and not simply in the reverse order, which could indicate a distinct mechanism. Based on our data, we present the first comprehensive model for the order, composition, and functional state of NPC disassembly and reassembly intermediates in living cells.

Results and discussion

A functional and quantitative assay for the kinetics of NPC disassembly and reassembly

The kinetics of Nup dissociation from and reassociation with the NE during mitosis was monitored in live NRK cells expressing 11 GFP-tagged Nups representative of eight different subcomplexes (Rabut et al., 2004a): Nup133, Nup107, Seh1, and Nup43 (all from the Nup107–160 complex); the cytoplasmic Nup Nup214, Nup98, Nup58 (Nup62 complex), Nup93 (Nup93 complex); the nucleoplasmic Nups Nup50 and Nup153; and the transmembrane Nup POM121. In triple color time-lapse sequences of individual dividing cells, we recorded each GFP-Nup together with a red fluorescent nuclear import marker (importin α [IBB]; Gorlich et al., 1996) and vital DNA staining (Fig. 1, A and B). DNA was used as spatial reference to quantify nuclear (envelope) intensities (for details see Materials and methods) and to monitor mitotic progression. The import marker IBB was efficiently imported into the nucleus during interphase, released into the cytoplasm at NEBD, and reimported in telophase, providing a functional reference for the import competence of the NPCs. In addition, we used the reimport/release of IBB to temporally align the assembly time series of the different Nups (Fig. 1, C and D). DNA was used as spatial reference to quantify nuclear (envelope) intensities (for details see Materials and methods) and to monitor mitotic progression. The import marker IBB was efficiently imported into the nucleus during interphase, released into the cytoplasm at NEBD, and reimported in telophase, providing a functional reference for the import competence of the NPCs. In addition, we used the reimport/release of IBB to temporally align the assembly time series of the different Nups (Fig. 1, C and D). In summary, this assay allowed us to analyze the kinetics of NPC disassembly and reassembly in detail and to determine the import competence of the nucleus in different states of NPC assembly in living cells.
and Nup43 are less stably associated with the complex and, indeed, this has been reported for Seh1 although not for Nup43 (Loiodice et al., 2004).

To test whether the binding of members of the Nup107–160 complex to chromatin represented formation of NPCs rather than a general “coating” of chromatin, we analyzed early assembly stages by high resolution microscopy of living cells. Binding of GFP-tagged members of the Nup107–160 complex to chromatin occurred in discrete patches and small dots of the appearance of single pores (Fig. 3 A). If these structures truly represent partially assembled NPCs, they should also contain Nups from other subcomplexes. We tested this by simultaneously imaging GFP-tagged Nup107–160 complex members and mCherry-tagged POM121. Indeed, POM121 first accumulated in patches around chromatin that also showed a strong localization of Nup107–160 complex members (Fig. 3 B). To rule out that this reflected the inability of the ER to contact other regions of chromatin, we also analyzed the localization of mCherry-tagged lamin B receptor (LBR), a protein of the inner nuclear membrane known to bind to chromatin (Ye and Worman, 1994). In contrast to POM121, the localization of LBR was relatively smooth and did not show a bias for sites of Nup107–160 labeling (Fig. 3 C). Our data therefore suggest that Nup binding previously (Belgareh et al., 2001). General association of Nup133 with chromatin was detected shortly after the metaphase–anaphase transition or 8.5 ± 0.5 min (n = 5) before the time point of half maximal IBB intensity in the nucleus (t1/2[import]; Figs. 2 and S1 A; and Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1). Nup133 had already reached its maximal concentration at t1/2[import]. These observations are in line with the essential function of the Nup107–160 complex in NPC assembly observed in vitro (Boehler et al., 2003; Harel et al., 2003b; Walther et al., 2003a; D’Angelo et al., 2006).

We analyzed the assembly of three additional proteins of this subcomplex (Nup107, Seh1, and Nup43). NPC subcomplexes are thought to be stable throughout the cell cycle (Matsuoka et al., 1999; Belgareh et al., 2001; Loiodice et al., 2004) and should thus bind to the reforming NE as a unit with identical kinetics. Indeed, we found Nup107 to faithfully recapitulate the assembly kinetics of Nup133 (Fig. S2 B, available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1). This suggests that stable subcomplexes are well represented by one member in our assay. Although the assembly of Seh1 and Nup43 also started early and was completed before t1/2[import], their kinetics were slightly but consistently delayed relative to Nup107 and Nup133 during early anaphase. This could indicate that Seh1 and Nup43 are less stably associated with the complex and, indeed, this has been reported for Seh1 although not for Nup43 (Loiodice et al., 2004).

To test whether the binding of members of the Nup107–160 complex to chromatin represented formation of NPCs rather than a general “coating” of chromatin, we analyzed early assembly stages by high resolution microscopy of living cells. Binding of GFP-tagged members of the Nup107–160 complex to chromatin occurred in discrete patches and small dots of the appearance of single pores (Fig. 3 A). If these structures truly represent partially assembled NPCs, they should also contain Nups from other subcomplexes. We tested this by simultaneously imaging GFP-tagged Nup107–160 complex members and mCherry-tagged POM121. Indeed, POM121 first accumulated in patches around chromatin that also showed a strong localization of Nup107–160 complex members (Fig. 3 B). To rule out that this reflected the inability of the ER to contact other regions of chromatin in anaphase, we also analyzed the localization of mCherry-tagged lamin B receptor (LBR), a protein of the inner nuclear membrane known to bind to chromatin (Ye and Worman, 1994). In contrast to POM121, the localization of LBR was relatively smooth and did not show a bias for sites of Nup107–160 labeling (Fig. 3 C). Our data therefore suggest that Nup binding
times in interphase (Rabut et al., 2004a). In our assay, both Nup153 and Nup50 were detected at the periphery of the chromatin as early as 7.9 ± 1.4 (n = 4) and 6.6 ± 0.8 min (n = 6) before t_{1/2}(import), respectively (Figs. 2 and S2 A; and Video 2, available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1). However, this early pool accounted for <10% of the final nuclear intensity for Nup153 and only ~20% for Nup50 (Figs. 2 and S2 A, blue shading). The major pools of these Nups associated with the NE considerably later and reached their half maximal intensity at the NE only 1.0 ± 0.3 (Nup153) or 1.1 ± 0.5 min (Nup50) before t_{1/2}(import) (see Fig. 5 D).

The biphasic assembly behavior we observed is consistent with the interphase dynamics and reinforces the interpretation that both proteins have two distinct modes of binding at the pore. Because both proteins are bound on the nucleoplasmic side of the pore, the early association of a small pool to chromatin could be involved in the formation of functional pores. The second phase of assembly paralleled initiation of nuclear import and transport through the first functional NPC assembly intermediates may therefore add the full complement of Nup50 and Nup153 to the complex.

POM121 accumulates at the NE after several soluble Nups

In interphase cells, the vertebrate-specific membrane Nup POM121 localizes almost exclusively to the NE, whereas it disperses in the ER during mitosis (Daigle et al., 2001). In metaphase, the ER is largely excluded from chromatin and spindle regions. However, ER membranes come close to the poleward face of the separating chromosomes early in anaphase (Fig. 3, B and C). The resulting early increase of POM121 signal around chromatin does therefore not reflect a specific accumulation (Fig. 3 B and not depicted). Accumulation in the NE over ER background became visible at 5.9 ± 1.0 min (n = 5) before t_{1/2}(import) and then rapidly reached its maximal intensity at t_{1/2}(import) (Fig. 2).

Together with the colocalization with the Nup107–160 complex, our kinetic data suggest that POM121-binding sites on chromatin become available only in late anaphase. At this time point, ER membranes come into physical contact with the separated chromosome masses from all sides and POM121 associates with chromatin at sites where Nup107–160 components are already bound.

Nup93, Nup98, and Nup58 assemble after membrane association

The Nup93 as well as the Nup62 complex are thought to localize to central positions of the pore. In our assay, the Nup93 and Nup62 complexes (represented by Nup58) accumulated at the NE starting at 3.8 ± 0.4 (n = 5) and 3.3 ± 1.4 min (n = 11) before t_{1/2}(import), respectively. The more peripheral Nup98 was first detected 3.8 ± 0.6 min (n = 6) before t_{1/2}(import) (Figs. 2 and S2 A). All three Nups reached their maximal intensity at the NE shortly after t_{1/2}(import).

Binding of these three complexes occurred only after several other Nups were already present on chromatin. Their addition may be the last step for the formation of an import competent NPC assembly intermediate because IBB import initiated concomitant...
NPC disassembly in prophase occurs rapidly and synchronously

The same set of eight representative Nups was followed during dissociation from the NE in prophase (Figs. 4, S1 B, and S3 A, available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1). Disassembly proceeded more rapidly than assembly and more synchronously for the different Nups so that distinct steps in the disassembly process could not be clearly resolved (compare Fig. 5, A and B). This could be caused by insufficient time resolution of the assay or simply the fact that disassembly occurs in fewer steps than assembly. Disintegration of a large part of the pore could be triggered in a single step. Also, recent EM data suggest that the disassembly of individual pores within one nucleus in X. laevis egg extract is asynchronous, leading to pore intermediates in different states of disassembly at the same time (Cotter et al., 2007). If this occurs in live mammalian cells, it would compromise our ability to detect the order of the process because we measure the mean of many pores simultaneously.

Figure 4. Time series representing the dissociation of four Nups from the NE during prophase. The contrast of the image series was normalized to a common maximal mean intensity on the nuclear rim at the first time point of each series. Plots show the data obtained from the series shown (red) and the mean of n series (black). As a reference, mean intensities of Nup98 (cyan) and POM121 (green) are shown in all plots. Time stamps give min:s relative to \( t_{1/2} \) (import). Videos 3 and 4 (available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1) show representative full-image sequences for Nup98 and POM121. Error bars indicate SD.

with their assembly (see Fig. 5, B and D). At this time, the Nup107–160 complex and POM121 were assembled already to ~80%, whereas only the minor early fractions of Nup50 and Nup153 were present.

Nup214 association with the NE lasts well into G1

Nup214 is a peripheral cytoplasmic Nup with a residence time of several hours at interphase NPCs (Rabut et al., 2004a). We first detected Nup214 at the NE 0.8 ± 0.2 min (\( n=4 \)) before \( t_{1/2} \) (import) (Fig. S2). It was thus the last Nup to associate with the newly forming NPC investigated in this study. Its first appearance was concomitant with the regaining of nuclear import activity but its concentration continued to increase over cytoplasmic background long after the maximal IBB intensity in the nucleus was reached. High import rates were reached already when Nup214 had only reached 50% of its maximal intensity at the NE (see Fig. 5 D). These kinetics suggest that Nup214 may not be required for IBB import, which is consistent with previous findings that show no role of Nup214 in protein import via classical import routes but rather suggest an activity in protein export (Walther et al., 2002; Hutten and Kehlenbach, 2006). A newly assembled nucleus will likely have to establish import function first and export only later when nuclear biosynthesis has restarted. This would explain the late assembly time of factors not required for import such as Nup214.
Figure 5. Summary of NPC disassembly and reassembly kinetics. (A and B) Overview over all means of disassembly (A) and assembly (B) kinetics. (C) Time points of first visible nuclear accumulation over background for all analyzed Nups. (D) Time points of 50% assembly of Nups relative to the first derivative of IBB intensity as a measure for import rate. Because of the change in concentration distribution of IBB between cytoplasm and nucleus during the import phase, the first derivative of IBB intensity systematically underestimates true instantaneous import rates. The maximum reached at time point 0 therefore does not reflect the true maximal import rates, which may be reached later. (E and F) Models for mitotic NPC disassembly and reassembly. Filament structures are included in the model in gray on the basis of previous data. The precise positions of the Nups in the NPC are unknown and thus drawn schematically. Because the different Nup-expressing cell lines showed some variability in the timing of mitotic progression (10.6 ± 1.5 min from anaphase onset to t₁/₂(import); not depicted), the time between anaphase onset and t₁/₂(import) was normalized to 10 min in B to D. Error bars indicate SD.
Electron microscopy of *D. melanogaster* embryos has revealed disassembly intermediates similar to assembly; however, one intermediate dominated all prophase nuclei, indicating that other intermediates may be very transient (Kiseleva et al., 2001). This fits well with our observation in living mammalian cells that disassembly is very rapid. The similar ultrastructural appearance of NPC intermediates lead to the hypothesis that disassembly could be the reversal of assembly. Despite the limitations of our assay, our data indicate that this may not be the case. For example, the Nups that assembled earliest and latest during anaphase, i.e., Nup133 and Nup214, dissociated from the NE in the middle of the disassembly process. Nup98, which assembles at an intermediate time point in anaphase, was clearly the first Nup to dissociate from the nuclear periphery in prometaphase, which is in agreement with data from starfish oocytes (Lenart et al., 2003). Finally, Pom121, which is assembled after the Nup107–160 complex in anaphase, also dissociated clearly after the Nup107–160 complex during disassembly.

Interestingly, the Nup107–160, Nup93, and Nup214 complexes, which are the most stable NPC subcomplexes during interphase (Rabut et al., 2004a), dissociated early and rapidly, whereas Nup50 and Nup58 (Nup62 complex) together with POM121 remained longest in fragments of the NE (Figs. 4 and S3 A; and Videos 3 and 4, available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1). Thus, the NE identity of POM121-containing membranes appears to be lost only gradually in prometaphase, which is in agreement with previous observations (Beaudouin et al., 2002).

The persistence of Nup50 at the NE might be caused by chromatin rather than NPC association because we found Nup50 to coat chromatin throughout mitosis from prophase until anaphase (Fig. S3 B). It formed a dynamic coat, which rapidly exchanged with the cytoplasmic pool as assayed by photobleaching (unpublished data). This localization is consistent with the presence of the *Aspergillus nidulans* homologue of Nup50 on mitotic chromatin (Osmani et al., 2006) and could indicate a conserved mitotic function. However, it could also be caused by an inherent chromatin affinity of Nup50 because the yeast Nup50 homologue has been implicated in NPC associated gene regulation (Schmid et al., 2006).

**Conclusion**

In summary, our systematic study allows us to propose the first comprehensive model for mitotic NPC disassembly and reassembly (Fig. 5, E and F). Disassembly occurs in mammalian cells in a similar manner to starfish oocytes (Lenart et al., 2003) but with faster kinetics (Fig. 5 A). The composition of disassembly intermediates appears to differ from assembly intermediates, which suggests a distinct mechanism.

Our data provide detailed insight into the kinetics of pore assembly with high time resolution. Consistent with previous studies, we find NPC assembly to be a highly ordered process (Fig. 5 C). For the first time, we can relate the composition of the different assembly intermediates to import function. Our data supports the model that assembly starts with formation of a prepore on chromatin and indicates that such a structure contains the Nup107–160 complex as well as substoichiometric amounts of Nup153 and Nup50 (Fig. 5 F). These may provide the binding platform for additional components like the transmembrane Nup POM121.

In our live cell assay, we measure the mean concentration of Nups over all NPCs in the imaging plane to determine their assembly kinetics. We therefore cannot formally decide whether the fact that the association kinetics of individual Nups stretch over several minutes reflects asynchronous assembly of different NPCs in the nucleus, the sequential addition of multiple copies of the same Nup to NPCs in the same state of assembly, or a mixture of the two processes. However, our high-resolution imaging data showed similar concentration of Nups in adjacent pores at single time points during assembly (Fig. 3). Furthermore, electron microscopic data from *D. melanogaster* indicate that specific assembly intermediates dominate at any stage of mitosis (Kiseleva et al., 2001). We therefore assume that our kinetics reflect at least to a large extent the synchronous assembly process of many NPCs after mitosis.

What then is the first assembly intermediate that is competent for nuclear import? Comparing the time of half maximal concentration for each Nup with the rate of import (Fig. 5 D), our data show that the assembly intermediate containing mainly the Nup107–160 complex and POM121 does not support protein import (Fig. 5, B and D). Only upon association of Nup93, Nup58 (Nup62 complex), and Nup98 does IBB import initiate, which suggests that these complexes add transport activity to the new pore, possibly by providing many phenylalanine-glycine repeats. At this time point, at least a fraction of the pores in the nucleus contain all subunits necessary to support protein import function. In addition, the presence of a sealed or nearly sealed membrane around the nuclear compartment is likely required for IBB to accumulate in the nucleus. In contrast, the nucleoplasmic Nup50 and Nup153 as well as the cytoplasmic Nup214 are probably not required for import activity in stoichiometric amounts.

In the future, it will be very interesting to analyze the behavior of additional Nups, especially the membrane-bound Ndc1 and ELYS/Mel28, which have very recently been reported to play crucial roles in NPC assembly (Galy et al., 2006; Mansfeld et al., 2006; Rasala et al., 2006; Stavru et al., 2006; Franz et al., 2007). In addition, similar data obtained for interphase assembly will allow to test whether the insertion of NPCs into an intact interphase NE follows the same mechanism as postmitotic assembly.

Our assay using IBB as a functional and temporal marker should furthermore prove very useful to study additional aspects of NEBD and NE assembly. Besides a detailed kinetic understanding, the assay can also yield mechanistic insight when combined with molecular perturbations by RNAi or the expression of dominant-negative proteins.

**Materials and methods**

**DNA constructs and cell lines**

pIBB-DiHeRRed was generated by ligating the fragment of the IBB domain from the plasmid pQE60-IBB-GFP (Ribbeck and Gorlich, 2002) into pDIHeRRedN1 [Gerlich et al., 2003] with a 5'–amino acid linker (GPVAT) between the IBB domain and DiHeRRed.

pPOM121-mCherry was cloned by exchanging 3EGFP in pPOM121-3EGFP [Rabut et al., 2004a] with mCherry [Shorer et al., 2004]. pLBR1TM-mCherry contains the N terminus of LBR and its first transmembrane domain.
It was cloned by exchanging YFP in pLBR1TMYPF (Daigle et al., 2001) with mCherry.

NRK cells were grown in standard medium. NRK cell lines stably expressing Nups tagged with EGFP (Nup50, Nup58, Nup93, Nup98, Nup133, Nup153, Nup214, Pom121, Nup43, and Seh1) as described previously (Rabut et al., 2004a) were maintained at 0.5 mg/ml G418. Some experiments were performed by transient transfection with the same plasmids used for generation of the stable cell lines. Transient transfections with pBB-DHtRcd and Nup plasmids were performed with FuGene 6 (Roche) 24–72 h before imaging. For dual-color high-resolution imaging (Fig. 3), cells coexpressing GFP-tagged members of the Nup107–160 complex and IB- or POM121-mCherry were enriched by FACS.

Live cell microscopy

For live cell microscopy, cells were grown in Lab-Tek chambered coverglasses (Thermo Fisher Scientific). 30 min before imaging, the medium was exchanged for prewarmed CO₂-independent medium without phenol red supplemented with 2% FCS, 2 mM glutamine, 100 mg/ml penicillin and streptomycin, and 0.2 µg/ml Hoechst 33342. The chambers were sealed with silicone grease. Time lapse sequences of 2–4 µm thick confocal slices were recorded at 57°C on confocal microscopy systems (LSM 510) using a 63x 1.4 NA Plan Apochromat objective (Carl Zeiss, Inc.). Fluorescent chromatin was automatically tracked and focused during imaging using in-house developed macros (Rabut and Ellenberg, 2004). High-resolution imaging for Fig. 3 was performed with a 100x Plan Apochromat NA 1.4 objective (Carl Zeiss, Inc.).

Quantification and image analysis

Images were segmented on the chromatin channel in Image J (http://rsb.info.nih.gov/ij/) by successive application of a Gaussian and an anisotropic diffusion filter and thresholding of the filtered image with an in-house-developed macro. The segmentation was applied to the raw images of the IBB channel and the mean nuclear fluorescence intensities were quantified. For the assembly of most Nups, the same segmentation was used to quantify the mean intensity of the Nups on the chromatin region. During interphase, the soluble pools of both Nup50 and Nup153 localize to the nucleoplasm and a clear discrimination between nuclear rim association and nuclear import in later stages of mitosis could therefore not be achieved with the assay. However, the quantification on the nuclear rim region alone as compared with the complete chromatin region did not yield significantly different results for any of the two proteins, which suggests that the contribution of import to the measured kinetics is minor.

Manual rim segmentation was applied for all disassembly series to avoid folded regions of the NE. The apparent decrease in Nup133 fluorescence in the nuclear region after t1/2 (import) is caused by dilution of the signal avoid folded regions of the NE. The apparent decrease in Nup133 fluorescence distribution of import to the measured kinetics is minor.

Different results for any of the two proteins, which suggests that the contribution of import to the measured kinetics is minor. Different results for any of the two proteins, which suggests that the contribution of import to the measured kinetics is minor.

Online supplemental material

Fig. S1 shows all individual disassembly/reassembly curves used to derive the mean kinetics shown in Fig. 5 (A and B). Fig. S2 shows representative image series for the assembly of Nup50, Nup93, and Nup214 and mean assembly curves for all analyzed members of the Nup107–160 complex. Fig. S3 shows a representative image series for the disassembly of Nup133, Nup153, Nup93, and Nup214 and the localization of Nup50 on chromatin during mitosis. Videos 1 and 2 show representative assembly series for Nup133 and Nup93, respectively. Videos 3 and 4 show disassembly series for Pom121 and Nup98. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200707026/DC1.

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