Interphase Nuclei of Many Mammalian Cell Types Contain Deep, Dynamic, Tubular Membrane-bound Invaginations of the Nuclear Envelope

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Abstract. The nuclear envelope consists of a double-membraned extension of the rough endoplasmic reticulum. In this report we describe long, dynamic tubular channels, derived from the nuclear envelope, that extend deep into the nucleoplasm. These channels show cell-type specific morphologies ranging from single short stubs to multiple, complex, branched structures. Some channels transect the nucleus entirely, opening at two separate points on the nuclear surface, while others terminate at or close to nucleoli. These channels are distinct from other topological features of the nuclear envelope, such as lobes or folds.

The channel wall consists of two membranes continuous with the nuclear envelope, studded with features indistinguishable from nuclear pore complexes, and decorated on the nucleoplasmic surface with lamins. The enclosed core is continuous with the cytoplasm, and the luminal space between the membranes contains soluble ER-resident proteins (protein disulphide isomerase and glucose-6-phosphatase).

Nuclear channels are also found in live cells labeled with the lipophilic dye DiOC<sub>6</sub>. Time-lapse imaging of DiOC<sub>6</sub>-labeled cells shows that the channels undergo changes in morphology and spatial distribution within the interphase nucleus on a timescale of minutes.

The presence of a cytoplasmic core and nuclear pore complexes in the channel walls suggests a possible role for these structures in nucleo–cytoplasmic transport. The clear association of a subset of these structures with nucleoli would also be consistent with such a transport role.

In recent years, the view that the nucleoplasm is organized into a number of morphologically distinct and functionally significant domains has gained ground, supported by increasing evidence for subnuclear localization of processes such as replication (Banfalvi et al., 1989; Mills et al., 1989; Hozak and Cook, 1994; Hutchinson et al., 1994), repair (Jackson et al., 1994a, b), transcription (Jackson et al., 1993; Wansink et al., 1993), and RNA splicing and processing (Huang and Spector, 1991; Xing and Lawrence, 1993; Xing et al., 1993). In some cases these localized functions, or proteins associated with them, can be correlated with specific subnuclear structures, identified either morphologically or immunocytochemically at the light or EM level (Carmo-Fonseca et al., 1991, 1992; Spector et al., 1991; Lamond and Carmo-Fonseca, 1993; Huang and Spector, 1996; for review see Strouboulis and Wolfe, 1996). Although these intranuclear domains or structures are not bounded by membranes as cytoplasmic organelles are, they appear to offer a means of segregation of function in much the same way as organelles in the cytoplasm do; such an analogy has already been suggested (Spector, 1990; Moen et al., 1995).

Although the morphological and functional subdomains of the nucleus are not membrane bound, there is evidence that isolated nuclei contain large amounts of phospholipid and inositides and that activity of some nuclear enzymes depends on the products of intranuclear lipids metabolism for normal function (Maraldi et al., 1993). Isolated nuclei that lack the membranes of the bulk nuclear envelope still contain phospholipid as well as polyphosphoinositides. Indeed, it has been suggested that a separate inositol phosphorylation–dephosphorylation cycle might exist within the nucleus (Divecha et al., 1991; Banfic et al., 1993). Much of this nuclear lipid is presumed to exist as proteolipid complexes, most probably associated with a nucleoskeletal matrix, as it is not removed by detergent extraction with Triton X-100 (Divecha et al., 1993; Mazzotti et al., 1995).

Recent advances in immunocytochemical detection of phosphoinositols have permitted analysis of the intranuclear distribution of lipids that are not in the form of membranes (Mazzotti et al., 1995). However, there have also been suggestions that some of this nuclear lipid is in the form of recognizable bilayer membranes. Intranuclear
membrane extensions have been described in plant cells (Dickinson and Bell, 1972; Li and Dickinson, 1987), insect cells (polytene cells of the Drosophila melanogaster salivary gland; Hochstrasser and Sedat, 1987; Parke and de Boni, 1992), and mammalian cells (Bourgeois et al., 1979; Stevens and Trogadis, 1986). Single and double membrane invaginations have been identified, and an association with evaginations of the nuclear envelope into the cytoplasm has also been described (Parke and de Boni, 1992). Experimental demonstration of a statistically significant association of nuclear invaginations with nucleoli has led to the suggestion that such structures might play an important role in nucleocytoplastic transport (Bourgeois et al., 1979).

Nucleocytoplastic transport is bidirectional and energy requiring; almost certainly the majority of this flow passes through the nuclear pore complexes (NPC)1 that stud the nuclear envelope (Dingwall and Laskey, 1992; Hinshaw et al., 1992; Rout and Wente, 1994). Despite the identification of nuclear targeting signals, little is known of the routes by which imported polypeptides move from the NPC to their specific subnuclear compartment within the nucleoplasm. Recent results have suggested that proteins capable of repeatedly shuttling between the nucleus and the cytoplasm may play an important role in these processes (Laskey and Dingwall, 1993; Schmidt-Zachmann et al., 1993). EM immunocytochemistry hints at the possibility that a nuclear skeleton may provide tracks for the shuttling movement of these proteins (Meier and Blobel, 1992).

Similarly, newly made ribosomes and mRNA probably leave the nucleus through NPCs, but their routes from the sites of intranuclear synthesis to the NPC remain obscure (Rosbash and Singer, 1993). Again, linear tracks have been observed in some experimental systems by in situ hybridization, suggesting the possible involvement of a nucleoskeleton and reinforcing a linear production line model for postsynthetic processing (Lawrence et al., 1989; Xing and Lawrence, 1993; Xing et al., 1993; Moen et al., 1995). Other experimental systems have not found evidence for a tracked route of RNA transport through the nucleoplasm, but rather, suggest a diffusion through a network of channels preferentially accessible to nascent RNA (Zachar et al., 1993). Interestingly, even in the cases where nascent transcripts formed extended intranuclear tracks, the majority of these tracks was not observed to reach the nuclear envelope (Rosbash and Singer, 1993; Xing et al., 1993, 1995). The presence of a draining network of intranuclear membrane bound channels could account for this unexpected observation.

In this study we report a combination of serial section transmission electron microscopy (TEM) and confocal laser scanning fluorescence microscopy (CLSM) experiments designed to characterize intranuclear membranes in more detail and to provide a baseline for considering the functional significance of membrane bound structures in the nucleus. Our results suggest that long, branching, intranuclear membrane channels are derived from the ER as deep, narrow invaginations of the nuclear envelope. These structures, which traverse deeply into the nucleoplasm and may pass completely through the nucleus as the nucleoplasm forms an annulus in the process, are found in all tissue culture cell types examined. Primary cells in culture also contain similar structures. The number of channels and their morphological complexity vary widely but remain characteristic for a given cell type. Examination of the channels in living cells shows that these structures are dynamic, changing position and morphology within the interphase nucleus. Some of the results described here have been presented in abstract (Vaux, D., M. Hollinshead, and M. Fricker, 1994. The Eukaryotic Nucleus. Keystone Symposium)

Materials and Methods

Chemicals

Vectashield antioxidant mountant and fluorochrome-conjugated horse anti-rabbit secondary antibodies were obtained from Vector Labs, Inc. (Burlingame, CA). Sterile trypsin solution, DME, and Liebowitz 15 media were obtained from Gibco BRL (Gaithersburg, MD). Fetal calf serum was obtained from HyClone Labs (Logan, UT), and heat inactivated for 30 min at 56°C before use. Other chemicals were analytical grade and obtained from Sigma Chemical Co. (St. Louis, MO), Merck Chemical Div. (Rahway, NJ), or BDH Chemicals Ltd. (Dagenham, Essex, England).

Fluorescent Probes

Biotinylated or directly fluorochrome conjugated Con A was obtained from Vector Labs, Inc. Fluorescein isothiocyanate-labeled immunoglobulin (FITC-IG) fraction from normal goat serum, DAPI, and propidium iodide (PI) were obtained from Sigma Chemical Co. 3,3-dihexyloxacarbocyanine iodide (DiOC6), lissamine-rhodamine-conjugated streptavidin, To-Pro-3 iodide, and dihydroethidium (DiHE) were obtained from Molecular Probes Inc. (Eugene, OR).

Primary Antibodies

A monoclonal antibody against a synthetic peptide corresponding to the COOH terminus of protein disulphide isomerase was obtained from culture supernatant of the ID3 hybridoma cell line. A rabbit polyclonal antiserum against an ER membrane fraction was the generous gift of Professor Daniel Louvard (Institut Pasteur, Paris, France). A rabbit polyclonal antiserum against a peptide corresponding to the first 32 NH2-terminal amino acids of human lamin A was the generous gift of Drs. George Sinios and Spyros Georgatos (European Molecular Biology Laboratory, Heidelberg, Germany). This antiserum recognizes lamin A, B, and C from a wide range of mammalian and avian species.

Cells

HeLa, Vero, NRK, 3T3, CHO, A431, J774, RAW, G8, C2, and NOR10 cell lines were maintained in culture at 37°C in 5% CO2 in DME supplemented with 10% heat inactivated fetal calf serum. Cells were removed from stock flasks with trypsin and plated onto sterile 25-mm diameter cover slips at least 1 d before use. Primary cells were obtained from the peritoneal cavity of BALB/c mice by sterile lavage and placed in culture for 1–3 d before use.

Tissue Sections

Rats were perfusion fixed using a periodate/lysine/paraformaldehyde fixative, and 10 μm frozen sections from blocks of a range of tissues were collected on gelatinized glass slides, as described (the slides were provided by P. Tree and L. Darley (Sir William Dunn School of Pathology, UK), following the procedure described for mice by Hughes et al., 1995). Before use, the tissue sections were permeabilized with 0.2% (vol/vol) Triton X-100 in PBS for 15 min and quenched in fresh 50 mM ammonium chloride solution for 15 min, both at room temperature. Sections were labeled for 45

1. Abbreviations used in this paper: CLSM, confocal laser scanning fluorescent microscopy; G-6-Pase, glucose-6-phosphatase; NPC, nuclear pore complex; PDI, protein disulphide isomerase; TEM, transmission electron microscopy; 3-D, three-dimensional.
min with a mixture of 4 μM To-Pro-3 iodide and 10 μg/ml TRITC Con A in PBS at room temperature before washing four times in PBS, mounting in Vectashield, and examining by confocal microscopy.

**Serial Section Electron Microscopy**

For conventional microscopy, cells were washed with ice cold PBS and fixed in 0.5% glutaraldehyde in 200 mM sodium cacodylate, pH 7.4, for 30 min, and then washed in buffer and postfixed in 1% osmium tetroxide/1.5% potassium ferrocyanide for 60 min at room temperature. After washing in water and then staining in 0.5% mg uranyl acetate overnight at 4°C, the cells were left attached to the culture dish and flat embedded in Epon. Sections 70–90 nm thick were cut and collected either onto 50 mesh or slot grids, contrasted with lead citrate, and examined using two microscopes (EM10 at 60 kV; Philips Technologies, Cheshire, CT; EM912 at 80 kV; Zeiss, Inc., Thornwood, NY).

For the cytochemical staining of G-6-Pase, we used a minor modification (Griffiths et al., 1983) of the procedure developed by Wachstein and Meisel (1956). To optimize staining of AR-T20 cells for G-6-Pase we found it necessary to fix the cells in <1% glutaraldehyde and to incubate the fixed cells with the substrate solution for 2 h at 37°C.

**Cryoinmuno-electron Microscopy**

HeLa cells were fixed for 10 min at 4°C in 4% paraformaldehyde in 250 mM Hepes (pH 7.4) and then transferred into 8% paraformaldehyde in Hepes at room temperature for at least 20 min, removed from the plastic culture dish with a rubber policeman, and spun to a pellet. Fresh 8% paraformaldehyde was added to cross-link the cell pellet for another 30 min at room temperature. Coverslips were then washed in PBS and stored at 4°C overnight. Coverslips were then washed in PBS and stored at 4°C for up to 3 d before labeling. Before labeling, PFA fixed samples were permeabilized with 0.2% NP-40 in PBS for 5 min at room temperature. For indirect lectin labeling, coverslips were first incubated with 10 μg/ml of biotinylated lectin in PBS for 30–60 min at room temperature, washed four times in PBS, and then incubated in 1 μg/ml fluorochrome conjugated streptavidin in PBS for 30 min at room temperature. Indirect immunofluorescence was carried out as described previously. Most samples were also counterstained with 0.1 μg/ml DAPI for 5 min during the final wash before mounting. Samples were examined using either a microscope equipped for epifluorescence illumination and a low light level CCD camera (Axiopt; Zeiss Inc.) or a CLSM.

**Scrape Loading of Cells**

HeLa or NRK cells were plated in 60-mm diam tissue culture dishes and allowed to grow to 90% confluence. The monolayers were washed twice with ice cold PBS, and the cells were scraped from the dish in 0.5 ml of ice cold PBS containing 10 mg/ml of FITC conjugated normal goat immunoglobulin (150) using a soft rubber policeman. The cells were allowed to stand on ice for 5 min and then diluted in 10 ml ice cold complete medium and pelleted by centrifugation at 400 g for 5 min. The cells were resuspended in complete culture medium and plated onto 25-mm diam coverslips. The samples were either examined live or after fixation in 4% PFA.

**Low Light Level Fluorescence Microscopy**

The microscope (Axiopt; Zeiss Inc.) was equipped with a 50 W mercury arc lamp for epifluorescence illumination, a 100× 1.3 NA Plan Neofluar oil-immersion objective, and an Optimar set at 1.25×. Images were captured using a cooled CCD camera (CDC 242 Peltier; Hamamatsu Photonics K.K., Hamamatsu City, Japan) under the control of software (BioVision; ImproVision Ltd., Warwick, UK) running on a Macintosh Quadra 900 computer. Optimal settings were obtained by adjustment of camera gain and offset and on-chip integration time, using the intensity histogram to ensure that the range of fluorescence intensities fell within an 8 bit recordable range. Settings were optimized individually for each fluorochrome and then held constant for all samples in a given experiment. Images were stored as 8 bit greyscale files. Serial Z axis images separated by 0.5-μm steps were collected using a motor driven focus system under computer control (BioVision Ltd.). The 8-bit images from each fluorochrome were median filtered with a 3 × 3 kernel and merged using the 24 bit merge capability of Adobe Photoshop. No other processing was used in the images displayed here. Vitally stained cells were also examined with the low light level CCD system attached to an inverted microscope (Axiovert; Zeiss Inc.) with a temperature controlled stage.

**Confocal Microscopy**

Two confocal microscopy systems were used. The first was based on a scan head (MRC600; BioRad Laboratories, Cambridge, MA) attached to a Nikon Diaphot TMD inverted microscope with modification of the laser excitation path to provide co-aligned beams from a number of different lasers including 488-nm line (25 mW Argon-ion; Ion Laser Technologies Ltd, Salt Lake City, UT) and 543-nm line (1.3 mW Gre-Nc; Melles Griot, Rochester, NY) introduced through single-mode fiber optics (Fricker and White, 1992; Fricker et al., 1994). Neutral density filters were used to roughly balance the intensities of the two beams. Fluorescence emissions were collected at 540 ± 15 nm and >600 nm using a Nikon 60× 1.4 NA Plan apochromat oil-immersion objective.

The second system comprised an MRC 1000 scan head attached to a Nikon Diaphot 300 inverted microscope with excitation at 488 nm, 568 nm, and 647 nm (15 mW Argon-Krypton; Ion Laser Technologies Ltd). Fluorescence emissions were collected at 522 ± 35 nm for fluorescein, 605 ± 32 nm for rhodamine, Texas red, and propidium iodide, and 680 ± 32 nm for To-Pro-3 using a Nikon 60× 1.4 NA Plan Apochromat oil-immersion objective.

Three dimensional (3-D) datasets were collected with an XY pixel spacing of 0.05–0.09 μm/pixel for fixed cells and 0.18 μm/pixel for live cell recordings and a focus motor (z axis) increment of 0.4 or 0.5 μm. Individual frames were averaged or Kalman filtered over four to eight frames. The gain required to visualize the intranuclear structures was often sufficient to saturate signal from the ER outside the nucleus.

**Image Reconstruction and Display**

3-D confocal images were median filtered using a 3 × 3 box and visualized using software (ThruView Plus; BioRad Laboratories) as a rotation series of height-coded projections (White, 1995). Typically the interior of the nucleus was viewed from the coverslip side, omitting the first two to three sections at the base of the cell that contained the thin layer of peripheral cytoplasm and nuclear envelope. An intensity threshold was manually selected to retain detail of the intranuclear channels. The first occurrence of a pixel above this threshold was recorded along the line of sight for each angle in the rotation series and displayed as a height-coded image, where intensity represents the Z position of the pixel, white nearest the viewer, black furthest away (White, 1995). Smooth animations for inspection were calculated with rotation steps of 4° and a constant tilt angle of 30°. The Z-stretch factor required to compensate for the asymmetric sampling in X, Y, and Z was calculated from the focus motor increment to account for the difference in refractive index between the oil-immersion fluid, coverslip, and mounting medium (White et al., 1996). This correction factor was calculated using a weighted average of rays across the full NA of the lens. An average factor of 0.829 was used for cells in Vectashield, and a factor of 0.7 was used for live cells. Voxel dimensions in these images are thus approximately scaled as a 1 × 1 × 1 aspect ratio.

**Vital Staining and Live Cell Confocal Microscopy**

For live cell analysis, small tissue culture dishes were prepared by gluing 1-cm diam Perspex cylinders (OD 25 mm, ID 19 mm) onto 25-mm diam coverslips using a silicone liquid adhesive and sterilizing them in 100% ethanol before use. An arrangement was designed to use the oil immersion objectives on an inverted microscope with a temperature controlled stage for either low light level CCD or laser scanning confocal microscopy. The cells were transferred to Liebowitz 15 medium containing 10% fetal calf serum (L10) to remove the requirement for a 5% CO2 atmosphere during recordings and a focus motor (z axis) increment of 0.4 or 0.5 μm. Individual frames were averaged or Kalman filtered over four to eight frames. The gain required to visualize the intranuclear structures was often sufficient to saturate signal from the ER outside the nucleus.
examination. Similar results were obtained in short term experiments by sealing a second coverslip to the top of the dish with vacuum grease after equilibration of the contents at 5% CO₂, but this system maintains pH for only 30 min. Preliminary experiments using <20 mM Hepes to buffer normal growth medium revealed substantial changes in cell morphology, as described previously (Bowers and Dahm, 1993), which were not seen when L10 was used.

Cells were labeled with dihydroethidium (diHE) at 1 μg/ml in complete culture medium for 90 min at 37°C. The cells were then perfused with PBS, followed by methanol at −20°C, which was left on the cells for 6 min (i.e., comparable to the fixation conditions used to prepare fixed cells for labeling). 3-D datasets were collected during fixation. The DiOC₆ signal was extracted completely from the cells by the methanol within 30 sec. After fixation, the dish was perfused with PBS to wash the monolayer. The cytoplasm is nucleoli that are not apparently associated with this channel. Bars: (a–j) 200 nm; (k) 1 μm.

**Results**

**TEM Identifies Tubular Membrane Profiles within the Nucleoplasm**

Examination of tissue culture cells sectioned parallel to the growth substrate after in situ embedding revealed frequent membrane profiles within nuclei. Fig. 1, a–j shows 10 consecutive 70-nm sections through an oriented 3T3 nucleus (a low magnification view is shown in k). The double membraned figure in the center can be followed through all sections. a grazes the nuclear envelope, giving en face views of several nuclear pore complexes; the dark structure to the left is an HRP-loaded endosome within the cytoplasm. The predominant feature running through the series is a double membraned channel surrounded by an electron dense coat and with an electron lucent core. Additional, finer strands of membrane are associated with this larger structure (for example, Fig. 1, g–j). Small electron dense particles may be found attached to the inner membrane of the lucent core (Fig. 1 e).

A double membrane ring was the predominant morphology of membrane channel in 3T3, NRK, A431, and HeLa cells (Fig. 2). The annular double membranes frequently contain local fenestrations indistinguishable from nuclear pores sectioned transversely at the nuclear envelope (compare Fig. 2 a and the higher magnification inset in d with the nuclear envelope shown in e).

To test the possibility that these tubular membrane fig-

ures form extensions of ER membrane, we studied cells labeled histochemically for an ER marker, glucose-6-phosphatase (G-6-Pase). The lumen between the two membranes of the intranuclear tubules is decorated with the histochemical reaction product for G-6-Pase, confirming an ER origin (Fig. 3). Both short stubs of ER extending into the nucleoplasm and long channels that persist as double membrane extensions throughout the sections can be seen. The latter are distinct from the plate-like infoldings of the nuclear envelope that are characteristic of certain cell types, such as the polymorphonuclear leukocyte. The histochemical reaction product fills the lumen of the ER, the intermembrane space of the nuclear envelope, and the intranuclear channels, but is excluded from nuclear pores in both the nuclear envelope and the intranuclear channels. Shallow angle sections through the margins of nuclear channels (e.g., Fig. 3 a, top left) show that the number of nuclear pores per unit area in the channels is comparable to that in the bulk nuclear envelope (Fig. 3 d, top left).

The high electron density of the lumenal reaction product highlights the presence of fronds of membrane extending into the lumen of the channel as discrete clumps on the wall of a long channel (Fig. 3 d). Sometimes they appear to invade the core of a channel as a fine finger of ER from the cytoplasm, as can be well seen in the short channel at top left (Fig. 3, a–e).

**Intranuclear Con A Binding Sites Form Linear Arrays**

Fluorochrome conjugated Con A labels the endoplasmic reticulum and nuclear envelope. Confocal microscopy showed it also labels a limited number of nucleoplasmic sites in all cell lines examined (Fig. 4). This Con A labeling was completed via mannose-terminated oligosaccharides, because binding was abolished by competition with α-methyl mannoside (not shown).

The dataset illustrated in Fig. 4, consisting of 12 XY planes, was used to construct a height-encoded, 3-D view of nuclear volume seen from below with the basal cytoplasm removed. Different tilted views of the reconstruction confirmed that the channels were often orthogonal or near-orthogonal to the growth substrate. In addition, near horizontal Con A positive nucleoplasmic channels were also seen.

**Intranuclear Membranes Are Found in Many Cell Types**

To see how widespread intranuclear tubules were, we examined a range of cell types by labeling with Con A, followed by confocal microscopy and 3-D visualization. Labeled channels were found in all cell types tested, at varying frequencies. In non-synchronized populations not all cells contained detectable intranuclear Con A positive structures. However, by increasing the sensitivity (i.e., by...
using conditions that gave saturating levels of signal from the ER and nuclear envelope), we could detect more intranuclear structures. This suggests that some nuclear structures are either smaller than many cytoplasmic ER features or that they contain fewer accessible Con A binding sites. 3-D reconstruction of nucleoplasmic Con A positive channels from several cell types showed that some cells contained only a single tubule, and others contained a complex branching network. Fig. 5 illustrates the various degrees of complexity of the network seen in five different cell types.

NRK cells (Fig. 5 E) tend to have few intranuclear channels, almost invariably oriented vertically to the growth substrate and often offset within the nucleoplasm. HeLa cells (Fig. 4) tend to have a larger number of channels and more channels that are not vertically oriented to the growth substrate. Primary mouse peritoneal exudate cells in culture also show intranuclear membrane channels, often single and usually vertical to the growth substrate (Fig. 5 D). 3T3, CHO, and G8 cells (Fig. 5, A–C, respectively) have a more complex, frequently branched network.

We counted channels in complete 3-D datasets of >80 nuclei of each cell type (Fig. 6). We used a strict definition of a channel. It must extend at least 1.5 μm in X, Y, or Z to exclude short indentations found in some cell types (especially along the basal–nuclear margin) and residual “tails” of fluorescence along the Z axis from intense labeling of the nuclear envelope or cytoplasmic structures. More than

Figure 2. The intranuclear channels in a range of cell types contain nuclear pore complexes and are associated with an electron dense halo in the nucleoplasm. Representative profiles of channels from 3T3 cells (a), NRK cells (b), A431 cells (d and inset), and HeLa cells (f) are shown. Double membrane structures are clearly visible, together with local fenestrations, and comparable to the appearance of the nuclear pore complexes in the nuclear envelope of the 3T3 nucleus shown in e. The electron lucent core is also seen to contain small circular or crescent shaped features, best seen within the channel in b. Electron dense material in the nucleoplasm is associated with the channels (a, b and d, inset). In addition, channel association with nucleoli is shown for NRK cells (b). Thawed frozen thin section colloidal gold immunocytochemistry confirms that the channels are associated with strong immunoreactivity for lamins (c) and protein disulphide isomerase (f), as shown by the 9-nm gold particles decorating the intranuclear channels in these HeLa nuclei. Bars: 200 nm; d, 1 μm.
differentiated, postmitotic primary cell population (3 d ad-
Fricker, manuscript in preparation).

elsewhere (Vaux, D., M. Hollinshead, N. White, and M.
the nuclear channels as described here will be presented
shown; a complete analysis of the cell cycle dependence of
measured over the entire interphase population (data not
found in S phase nuclei was not lower than the abundance

gen, PCNA, confirmed that the abundance of channels
Con A and an antibody to the cell cycle–dependent anti-
tures could not be restricted to early postmitotic cells in G1.

60% of NRK cells do not have such channels, whereas
>90% of CHO cells contain at least one, with half having
three or more. We found no cell type with fewer channels
than NRK cells, nor any cell type with a more abundant
array than CHO cells.

The differences were reproducible over time, as shown
by the independent counts for NRK cells carried out 4 mo
apart (Fig. 6, NRK-1 and NRK-2).

The presence of channels in >90% of the cells in an un-
synchronized CHO population suggested that these struc-
tures could not be restricted to early postmitotic cells in G1.
Double labeling of unsynchronized cell populations with
Con A and an antibody to the cell cycle–dependent anti-
gen, PCNA, confirmed that the abundance of channels
found in S phase nuclei was not lower than the abundance
measured over the entire interphase population (data not
shown; a complete analysis of the cell cycle dependence of
the nuclear channels as described here will be presented
elsewhere (Vaux, D., M. Hollinshead, N. White, and M.
Fricker, manuscript in preparation).

Channels were found in the nuclei of a predominantly
differentiated, postmitotic primary cell population (3 d ad-

Composition of the Channels

The serial section TEM, G-6-Pase labeling, and presence
of nuclear pores together suggest that the nuclear enve-
lope invaginates to form these structures. If so, the invagi-
mating membranes should contain ER membrane proteins,
the annular space between the double membranes should
contain soluble ER proteins, and the central lumen should
respond to the cytoplasm. The nuclear channels could be
labeled with a monoclonal antibody directed against pro-	ein disulphide isomerase (PDI), a soluble resident enzyme
of the ER (Fig. 5 G); similar structures were labeled by an
antiserum recognizing ER membrane proteins (Fig. 5 G).

To test the possibility that the cores of the intranuclear
channels are not only topologically equivalent to the cyto-
plasm but are in continuity with it, we preloaded the cyto-
plasm with a high molecular mass (150 kD) fluorescent
tracer that was too large to pass through the nuclear pores.
The channels were filled with the tracer (Fig. 5 H), con-
fiming that they were extensions of the cytoplasm.

Double labeling experiments confirmed that PDI (Fig.
8 A) was contained within the structures that bound Con A
(Fig. 8 B; compare arrows in A and B). PDI was also found
within the intermembrane space of channels seen by EM
(Fig. 2 f), with immunolabeling gold particles decorating
the nucleoplasmic membrane–bound channels.

If the channels are invaginations of the whole nuclear
envelope, then the electron dense material on the nucleo-
plasmic side of the double membrane seen by EM (Figs. 1–3)
might be similar to peripheral heterochromatin and in-
clude a structure similar to the nuclear lamina. Confocal
microscopy confirmed that the channels defined by Con A
(Fig. 9 a) are associated with lamins (Fig. 9 b), often in ar-
eas poor in nucleic acids (Fig. 9 c). These results were con-
firmed by colloidal gold immunocytochemistry at the EM
level (Fig. 2 c).

Additional sites of intranuclear lamin staining, unassoci-
ated with membrane or Con A labeling, were also seen
(Fig. 9, a and b). These additional sites exhibited various
morphologies including punctate sites and extended linear
tracks, as is clearly demonstrated in the height-encoded re-
constructions of the entire datasets for Con A labeling
(Fig. 9, d–f) and lamin staining (Fig. 9, g–i).

Association of Some Channels with Nucleoli

This was confirmed by confocal microscopy, after staining
nucleic acid with propidium iodide and channels with Con
A. The channels often ran through holes in nucleic acid
staining (Fig. 10, A and B) and ran up to, or past, nucleoli
(Fig. 10 C). Electron microscopy suggested that some of the
channels were closely associated with nucleoli (Fig. 2 b).
Double labeling of tissue sections of normal liver also
demonstrated an association between hepatocyte nucleoli

Figure 3. The intranuclear channels are bounded by a double mem-
brane enclosing a G-6-Pase positive lumen. Consecutive 70 nm
serial sections of part of an AT20 nucleus sectioned parallel to
the growth substrate after histochemical visualization of G-6-
Pase activity. Dense reaction product fills the space between the
inner and outer nuclear membranes but is excluded from nuclear
pore complexes. Similar reaction product is seen within the chan-
nels in the nucleoplasm. Bar, 500 nm.

Figure 5. A. The channels often ran through holes in nucleic acid
with propidium iodide and channels with Con A (Fig. 5 A) and
PDI (Fig. 5 F); similar structures were labeled by an antiserum
recognizing ER membrane proteins (Fig. 5 F).

Figure 6. Additional sites of intranuclear lamin staining, unassoci-
ated with membrane or Con A labeling, were also seen (Fig. 9,
a and b). These additional sites exhibited various
morphologies including punctate sites and extended linear
tracks, as is clearly demonstrated in the height-encoded re-
constructions of the entire datasets for Con A labeling
(Fig. 9, d–f) and lamin staining (Fig. 9, g–i).
and Con A–labeled channels (Fig. 7, merged view of the left hand cell).

However, not all nucleoli show a closely associated channel (Fig. 10C, XY and XZ views), and some channels show no apparent association with nucleoli (by light microscopy in Fig. 10C, and by EM in Fig. 1k).

**Dynamic Channels are Present in Live Cells**

We next excluded the possibility that the channels were artifacts of cell preparation by visualizing them in living cells. The presence of channels in cells scrape-loaded with a 150-kD fluorescent tracer before fixation provided preliminary evidence that such structures are present in live cells (Fig. 5H). NRK, BHK, or HeLa cells in culture were labeled with dihydroethidium and with the lipophilic dye, DiOC₆, which selectively concentrated into cellular membranes (including the ER). The ER, the nuclear envelope (Fig. 11a), and intranuclear structures (Fig. 11, a and b) can then be seen in confocal optical sections cut through the center of the nucleus in living cells. The labeling of nucleic acids with dihydroethidium confirms that the confocal optical sections are indeed through the center of the nuclei (Fig. 11, e and f).

We next confirmed that the intranuclear membranes that label with DiOC₆ are the same structures that were visualized after fixation using Con A, by a sequential labeling experiment carried out on the stage of the confocal microscope. Fig. 11: (a) shows a single XY plane from the dataset for two DiOC₆ labeled cells; (e) shows the dihydroethidium label in the same plane. Fig. 11 (b and f) are from data collected 20 min later. The upper nucleus has changed considerably in shape. After methanol fixation, the DiOC₆ signal is lost (c), but the cells remain intact; and some dihydroethidium label persists (g). After Con A labeling, the intranuclear channels originally labeled by DiOC₆ are seen to be labeled with Con A. The result shown for a single plane in Fig. 11 (a–g) is confirmed by 3-D reconstruction in the lower part of Fig. 11. The DiOC₆ labeled structures found in the nuclei of live cells (Fig. 11, h–j) correspond to the Con A positive structures found in fixed permeabilized cells (Fig. 11, k–m); fixation causes little apparent change to their morphology at the light microscope level.

We next observed the dynamics of the channels by incu-
bating cells with DiOC₆ and collecting repeated complete 3-D images over 35 min (Fig. 11, n–p); the channels clearly persist over this timescale. Some channels do not change position within the nucleus but do alter their morphology, while others alter their position.

Discussion

The results reported here suggest that the nucleoplasm of mammalian cells contains long, dynamic, branching membrane channels that are derived from the ER as deep, narrow invaginations of both membranes of the nuclear envelope. Nuclear invaginations with a range of morphologies have been reported in EM studies on mammalian nuclei (Bourgeois et al., 1979; Dupuy-Coin et al., 1986; Stevens and Trogadis, 1986), in insect nuclei (Hochstrasser and Sedat, 1987; Parke and de Boni, 1992), and in plant nuclei (Dickinson and Bell, 1972; Li and Dickinson, 1987).

Single image planes, whether optical sections in the confocal microscope or thin sections in the TEM, do not allow unambiguous identification of tubular channels. Consequently, all data presented here have been analyzed in three dimensions, after reconstruction from serial sections obtained using the confocal microscope, or TEM. This approach permits the unambiguous identification of tubular incursions, distinct from flat plates or folds, that would be difficult to identify without such a 3-D analysis. Furthermore, the nuclear channels are seen in live cells and so cannot be artifacts of the fixation or immunocytochemistry.

3-D analysis demonstrates that channels are entirely distinct from the folds that render the nuclei of some cell types lobulated (e.g., polymorphonuclear leukocytes). In particular, channels are unrelated to the constricting furrows in the surface of the nuclear envelope that may be associated with bands of intermediate filaments (Kamei, 1994).
These latter structures are characterized by loops of bundled intermediate filaments constricting the nuclear envelope; they do not form independent intranuclear membrane profiles separated from the nuclear envelope, even in confocal reconstructions (Kamei, 1994). The absence of detectable intermediate filaments may also result in irregularities of the nuclear envelope, characterized as prominent, multiple infoldings (Sarria et al., 1994). The infoldings found in these vimentin-negative cells are clearly distinct from the deep, narrow channels described in this study.

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The channels described here all consist of a cytoplasmic invagination bounded by a double membrane. This makes them quite distinct from the structures described in insect nuclei, where the invaginations are of the inner nuclear membrane only, producing a structure with a core in continuity with the ER (Hochstrasser and Sedat, 1987; Parke and de Boni, 1992). Furthermore, detailed statistical analysis demonstrates that the single-membrane invaginations found in Drosophila nuclei are always associated with an adjacent double-membraned evagination of the nuclear envelope into the cytoplasm (Parke and de Boni, 1992).

These differences, together with the observation that the core of the Drosophila invaginations are frequently filled with large electron-dense granules (Hochstrasser and Sedat, 1987) not seen in mammalian cells in this study, suggest that the structures we describe are unlikely to be closely related to the invaginations described in Drosophila.

Double-membraned invaginations of the nuclear envelope in plant cells have been described (Dickinson and Bell, 1972; Li and Dickinson, 1987). These structures are developmentally restricted to a short interval in postmeiotic gymnosperm microspores and have several features that distinguish them from the channels described here. Firstly, they are very short invaginations (maximum length, 0.9 μm); secondly, they are very numerous (200–400 per nucleus) and, thirdly, they are composed of membranes devoid of nuclear pores (Li and Dickinson, 1987). These features lead us to conclude that these plant cell structures are unlikely to be related to the large double-membraned channels found in mammalian nuclei.

Invaginations of the mammalian nuclear envelope may be single-membraned or double-membraned (Bourgeois et al., 1979). Double-membraned nuclear invaginations have previously been described in serial section EM studies as “nucleolar channels” because of an invariant association with nucleoli (Bourgeois et al., 1979; Dupuy-Coin et al., 1986). The results presented here confirm the presence of invaginating channels terminating in, or adjacent to, nucleoli in nuclei from a variety of cell types. However, it is clear that a combined CLSM and serial TEM analysis demonstrates additional categories of intranuclear invagination, in some cases unrelated to nucleoli and even in some cases having no intranuclear termination. Some channels (tubes) terminate within the nucleoplasm, others (tun-
nels) pass entirely through; some nuclei contain none, others many; some are simple, others branched. These overall features may be seen by confocal microscopy. More detailed structure is revealed by serial TEM sections, where the double membrane, the embedded nuclear pore complexes, the fine membrane processes, and the closed end of the channel are all clearly visible. One of the most unexpected findings of the present study is channels that traverse the nucleoplasm and have connections to two separate points on the nuclear envelope. This novel morphology means that the nucleoplasm forms an annulus around the channel. This is topologically quite distinct from any sheet- or plate-like infolding of the nuclear envelope. In particular, these traversing channels cannot be removed from the interphase nucleus without a membrane fusion or fission event.

Composition of the Channels

EM of serial sections showed direct connections between the double-membraned intranuclear tubes and the double membrane of the nuclear envelope. The double membranes of the nuclear envelope and the channels both contain nuclear pores. This topology is consistent with a deep invagination of both the outer and inner nuclear membranes. Three lines of evidence support this view. Firstly, both the ER lumen and the luminal space between the two membranes forming the boundary of the channels contain protein disulphide isomerase (by CLSM, Fig. 5, and by EM immunolabeling, Fig. 2), Con A binding sites (Figs. 4 and 5), and glucose 6 phosphatase (Fig. 3). Secondly, the membrane proteins of the ER were found in the peripheral ER, the bulk nuclear envelope, and the nuclear channels (Fig. 5). Thirdly, membranes of both ER and the nuclear channels accumulated the lipophilic dye DiOC6 (Fig. 11). These results identify the channels as deep invaginations of both inner and outer nuclear membranes, but cannot exclude the possibility that the composition of the membranes forming the channel differs in detail from that of the bulk nuclear envelope.

Our data strongly suggest that the majority of tubular membranes within the nucleus contain a core of material that is continuous with the bulk of the cytoplasm. The space at the center of the double-membraned channels has the electron density of cytoplasm in EM images, and such an identity would be topologically plausible. This space could be separate from bulk cytoplasm or accessible to it, depending on the organization of the junctions between the intranuclear channels and the nuclear envelope. High molecular weight tracers that are too large to enter the nucleoplasm through nuclear pores have access to the lumen of the channels after scrape-loading into the cytoplasm, confirming that the lumen is continuous with the cytoplasm. The lumen of nuclear channels is not always empty; ribosomes, small vesicles, and cytoskeletal elements have all been seen. Thus, one effect of these structures is to bring cytoplasmic space closer to the interior of the nucleus.

The nucleoplasm around the nuclear channels has an increased electron density similar to that seen in peripheral heterochromatin adjacent to the nuclear envelope (Fig. 1). Light and EM immunocytochemistry confirm that in both cases these electron-dense areas include a lamina containing nuclear lamins (Figs. 2 and 9). Several recent reports suggest that the nuclear lamina proteins are not confined to a lamina at the nuclear periphery, but may be found at focal sites throughout the nucleoplasm (Goldman et al., 1992; Bridger et al., 1993). We confirm the presence of focal
lamin accumulations in the nucleoplasm and show that some of these sites correspond to channels, while others do not (Fig. 9), confirming that there are at least two classes of focal lamin sites within the nucleus, besides the peripheral nuclear lamina (Belmont et al., 1993; Bridger et al., 1993). A functional significance for intranuclear lamin B sites has been suggested by colocalization of these sites with sites of DNA replication in mid to late S phase (Moir et al., 1994). Interestingly, this association is confined to lamin B even though intranuclear lamin A–C foci are also found (Goldman et al., 1992), suggesting a still higher order of complexity for intranuclear distribution of lamins.

**Distribution and Dynamics of the Channels**

Nuclear channels were found in all cell types examined, including cultured primary cells (macrophages explanted from the peritoneal cavity). The number of channels and their morphological complexity vary widely but remain characteristic for a given cell type and do not change significantly over time (Fig. 6). The organization of the channels with respect to the growth substrate is characteristic for each cell type, and this also remains stable for a given cell type over time in culture. The abundance of channels orientated vertically with respect to the growth substrate could depend in part on the better visibility of such structures by light microscopy, but this cannot be the whole explanation since the same bias was seen in oriented serial EM section analyses.

Channels are not restricted to transformed cell lines in culture, as they are also found in cultured primary peritoneal exudate cells (Fig. 5 D), which are predominantly postmitotic, differentiated macrophages in these preparations, and in nuclei of normal hepatocytes in tissue sec-

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**Figure 11.** Nuclear channels are detectable in live cells and correspond to the Con A–labeled structures seen in fixed cells. a shows a single plane from the dataset for two DiOC₆ labeled live cells; e shows the dihydroethidium label in the same plane. b and f are from data collected 20 min later. c shows the loss of the DiOC₆ signal at the end of methanol fixation, while g shows that the cells remain intact and the dihydroethidium label persists, although reduced in intensity. d shows the same cells and plane after 20 min of Con A labeling. h–j show sequential rotations of reconstructions of DiOC₆ labeled cells, and underneath in k–m, respectively, the same cell is shown after Con A labeling. Cutaway views of a reconstructed DiOC₆ labeled HeLa nucleus from a separate experiment are shown at three time points in n–p. Note that the morphology of the larger left hand tube alters over the 35-min period, and the smaller channel shows a progressively increasing separation from the large channel. Bars: (a–m) 25 μm; (n–p) 2 μm.
tions (Fig. 7). Furthermore, the presence of channels in these postmitotic nuclei suggests that such structures are maintained in cells that are not cycling.

Within individual tissue culture cells, although the channels may persist for hours in interphase nuclei, they may also rapidly change their morphology and position within the nucleoplasm on a timescale of minutes. We cannot exclude the possibility that some of these changes are the indirect result of gross changes in nuclear shape. However, de novo production of channels in interphase nuclei was not observed.

Possible Functional Implications of Nuclear Channels

Collectively, these data leave little doubt that there are widespread incursions of double ER membranes into the depths of the nucleoplasm in many cell types. Since the nuclear envelope is disassembled into vesicles during mitosis, it is most likely that the channels are formed during nuclear envelope reassembly. Chromatin becomes coated with bound nuclear envelope vesicles while decondensation is progressing; such vesicles penetrate between chromosomes and produce a deeply convoluted layer. These convolutions may be resolved by chromatin decondensation to produce a uniform convex vesicle layer; fusion of such a layer would produce a nuclear envelope devoid of deep channels. However, if the vesicle fusion event precedes complete decondensation, a complex, channel-rich nuclear envelope may be produced. Thus, the channels may arise as a result of incomplete resolution of invaginations during nuclear envelope reassembly; differences in channel morphology between cell types could then result from differential rates of chromatin decondensation and nuclear envelope vesicle fusion. It is possible that the slow resolution of channels could continue after vesicle fusion and, indeed, perhaps continue into interphase. In this case, the channels described might simply represent nonfunctional remnants of the process of nuclear envelope reassembly; their presence might even be deleterious for the cell.

There are three pieces of evidence that suggest that, although channels may originate in this way, they are not nonfunctional remnants. First, channels persist for long periods in interphase cells. More than 90% of CHO cells in a nonsynchronized culture have one or more channel, and this proportion suggests that most interphase CHO cells still have not resolved their channels, whether they are in G₁, S or G₂ phases of the cell cycle. Second, channels may be found in postmitotic populations (such as the primary peritoneal exudate cells and normal tissue hepatocytes examined in this study), again implying that the structures described here are not resolved in such populations. Third, if the channels represented mitotic remnants, it is not easy to explain the association with nucleoli that has been described (Bourgeois et al., 1979; Dupuy-Coin et al., 1986). Although an answer to this question must await further functional studies, we suggest possible functions that the channels might serve.

One plausible function of such invaginations is to bring a larger proportion of the nucleoplasm close to a nuclear pore. For a spherical nucleus 5 μm diam, 60% of the nucleoplasm lies within 0.5 μm of the nuclear envelope, and the center of the nucleoplasm lies at 2.5 μm away from it. However, a single tube traversing the nucleus halves the maximum distance from any point in the nucleoplasm to the envelope and brings a further 12% of the nucleoplasm within 0.5 μm of the envelope. Three or four appropriately placed tubes would bring almost all of the nucleoplasm within a distance of 0.5 μm of the envelope.

This means that even nucleoli which are buried within the nucleus may lie close to the envelope. A function of nuclear channels in transport processes has been suggested in measles virus–infected hamster cells, where an association between tubular invaginations and nucleoli was reported (Dupuy-Coin et al., 1986). Indeed, a substantial number of channels contact the nucleolus or terminate within it, both in tissue culture cells and normal hepatocytes. However, we also find a clear category of nuclear channels of varying morphology that have no apparent relationship with nucleoli, even when the entire nuclear volume is reconstructed in three dimensions and examined in animated rotation. Such channels may be genuinely dissociated from nucleoli, with attendant functional implications. Alternatively, the channels detected at the LM level might represent only the larger “trunks” of branched structures, leaving undetected fine branches to communicate with apparently distant nucleoli. The serial section EM data confirms that there are very fine lateral branches associated with some nuclear channels, as may be clearly seen in Fig. 1.

Taken together, these results and the data presented here suggest that the traditional view of the nuclear envelope as an approximately spherical coat defining the nucleoplasm is at best an incomplete one. Deep tubular invasions into the nucleoplasm occur sufficiently often in the population to warrant inclusion in any discussion of structure–function relationships in the envelope of mammalian interphase nucleus.

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