Abstract. The promyelocytic leukemia (PML) nuclear body (also referred to as ND10, POD, and Kr body) is involved in oncogenesis and viral infection. This subnuclear domain has been reported to be rich in RNA and a site of nascent RNA synthesis, implicating its direct involvement in the regulation of gene expression. We used an analytical transmission electron microscopic method to determine the structure and composition of PML nuclear bodies and the surrounding nucleoplasm. Electron spectroscopic imaging (ESI) demonstrates that the core of the PML nuclear body is a dense, protein-based structure, 250 nm in diameter, which does not contain detectable nucleic acid. Although PML nuclear bodies contain neither chromatin nor nascent RNA, newly synthesized RNA is associated with the periphery of the PML nuclear body, and is found within the chromatin-depleted region of the nucleoplasm immediately surrounding the core of the PML nuclear body. We further show that the RNA does not accumulate in the protein core of the structure. Our results dismiss the hypothesis that the PML nuclear body is a site of transcription, but support the model in which the PML nuclear body may contribute to the formation of a favorable nuclear environment for the expression of specific genes.

Key words: transcription • nuclear structure • acetylated chromatin • correlative microscopy • electron spectroscopic imaging

Introduction

The promyelocytic leukemia (PML) nuclear body is a nuclear matrix-associated structure 250–500 nm in diameter that is present in the nucleus of most cell lines (A scoll and M aul, 1991; D e Graaf et al., 1992; Stuurman et al., 1992; Chang et al., 1995; Grande et al., 1996). There are approximately ten of these structures per nucleus, though this can vary considerably depending on the cell type, cell cycle, and other factors (A scoll and M aul, 1991). The first biochemical component to be identified was the Sp100 nuclear matrix associated protein, which is an autoantigen in some patients with primary biliary cirrhosis (Szostecki et al., 1990). This protein has been found to trans-activate a variety of promoters (Guldner et al., 1992; Xie et al., 1993) and has an MHC class I-like domain at its NH₂ terminus (Sternsdorf et al., 1997a). The promyelocytic leukemia gene product, PML, is also found in PML nuclear bodies. In some acute promyelocytic leukemias, a t(15;17) chromosomal translocation creates a fusion protein of PML and the retinoic acid receptor alpha (Borrow et al., 1990; de The et al., 1990, 1991; Kakizuka et al., 1991; Dyck et al., 1994; Koken et al., 1994). PML protein may have transcription regulation activity (Xie et al., 1993; V allian et al., 1997, 1998a,b), may suppress growth and transformation (L e et al., 1996; Fagioli et al., 1998), and is implicated in apoptosis (Wang et al., 1998). Int-6 is another protein that may localize to the PML nuclear body, reported in the context of studies of the HTLV-1 virus (D esbois et al., 1996). A ubiquitin homologue, PIC1/SUMO-1 can be covalently bound to PML and Sp100 protein, and partially colocalizes with the PML nuclear body (Sternsdorf et al., 1997b, 1999). This posttranslational modification has been proposed to be responsible for the targeting of proteins to the PML nuclear domain (M uller et al., 1998; D uprez et al., 1999).

PML nuclear bodies are also a target of a variety of viruses, and are thought to be involved in coordinating the expression and replication of viral genomes from phylogenetically distinct families (for a review, see Sternsdorf et al., 1997a). A number of DNA viruses, such as SV40, A d5, and H SV-1, begin replication in the vicinity of a few, but
not all, of the PML bodies in a nucleus. Increased amounts of infectious virus in a cell do not result in an increase in the number of nuclear sites of viral replication. This implies that PML nuclear bodies may not be identical (De Bruyn-Kops and Knappe, 1994; Mau1, 1998). Viral protein expression does not appear to be necessary for targeting viral genomes to these replication sites (Ishov and Mau1, 1996). Transcription of the viral genome also takes place at the periphery of the PML nuclear bodies, perhaps because they create a suitable environment (Ishov et al., 1997; Maui, 1998). Finally, many viruses disassemble or degrade PML nuclear bodies before proceeding to the late phase in infection cycles (Doucas et al., 1996; Mau1 et al., 1996; Ishov et al., 1997; Mau1, 1998). This may have a disruptive effect on host nuclear function.

The cellular role of the PML nuclear body remains uncertain, despite its obvious links to oncogenesis and viral infection (for reviews, see Sterndsford et al., 1997a; Slack and Gallagher, 1999). PML nuclear bodies may be sites of storage of transcription factors, sites of transcription, or sites of RNA accumulation. The presence of RNA, as well as fluorescein-UTP microinjection and EDTA-regressive staining, has been observed within the structure of the PML nuclear body (LaMore et al., 1999), indicating a potential role in the transcription of some genes. The high enrichment of the transcriptional coactivator/histone acetyltransferase, CBP, is further suggestive of transcriptional activity within this domain (LaMore et al., 1998; Doucas et al., 1999). Consequently, it is important to understand the composition of these structures and the composition and organization of the surrounding nucleoplasm. In this study, we used both indirect immunofluorescence microscopy and a novel analytical transmission electron microscopy method, electron spectroscopic imaging (ESI), to address the composition of PML nuclear bodies. PML nuclear bodies did not stain with antibodies recognizing highly acetylated histones nor those recognizing fluorine-substituted uridine (Fu). Furthermore, phosphorus mapping by ESI rules out the presence of chromatin or RNA within these structures. However, nascent RNA can be detected at the periphery, indicating that the surroundings of the PML nuclear bodies are sites of transcriptional activity.

**Materials and Methods**

**Immunodetection of Nascent RNA and PML Nuclear Bodies**

SK-N-SH cells were cultured directly on glass coverslips under conditions recommended by the American Type Culture Collection. Pulses of FU were performed by addition of FU to a final concentration of 2 mM in the culture medium. Cells were fixed with 1.0% paraformaldehyde in PBS, pH 7.5, at room temperature for 5 min. Subsequently, cells were permeabilized in PBS containing 0.5% Triton X-100 for 5 min. Antibodies against halogenated UTP (anti-BrdU; Sigma Chemical Co., catalog no. B-2531) were used to label nascent RNA. A cetylated chromatin was visualized using an antibody recognizing diacetylated histones H3 (Upstate Biotechnology, Inc., catalog no. 06-599). PML nuclear bodies were visualized using an anti-CBP NH2-terminal antibody (Santa Cruz, catalog no. sc-369; or Upstate Biotechnology, Inc., catalog no. 06-297) or an anti-PML antibody (5E10; Stuurman et al., 1992). Cells were then incubated with secondary antibody, goat anti-rabbit Cy3 (Chemicon International, Inc.) and goat anti-mouse (A lexa 488, Cedarlane Labs; or Cy2, Nycomed Amersham, Inc.). A riter rinsing, the samples were mounted in 1 mg/ml para-phenylenediamine in PBS/90% glycerol, containing DNA-specific staining DAPI (4',6-diamidino-2-phenylindole) at 1 µg/ml. Digital deconvolution to remove haze in optical sections was performed using a cooled CCD camera (Princeton Instruments) mounted on a Leica DMIRE immunofluorescence microscope. VayTek Micromote digital deconvolution software was used to remove out of focus contributions, and image stacks were projected into one image plane using Scion Image software. False coloring and superimposition was done with Adobe Photoshop 5.0.

**Correlative Microscopy and Electron Microscopy**

Detailed descriptions of the EM procedure are presented elsewhere (Bažetjones and Hendl, 1999). In brief, cells were grown on polypropylene caps, fixed with 1.0% paraformaldehyde in PBS, and labeled for immunofluorescence as described above. Cells were then reixed in 2% glutaraldehyde in PBS for 5 min, dehydrated in ethanol, starting at 30%, before embedding in Quetol 651 resin. We have found that this fixation procedure preserves fine structural detail in chromatin. Moreover, chromatin-based and protein-based fibers are routinely observed (Bažetjjones and Hendl, 1999; Bažethjones et al., 1997; Hendzel et al., 1999). To reveal the detail seen with this protocol, a section from the same block as that used for data in this paper was stained with uranyl acetate and imaged in bright-field (Fig. 1A and B). E last images recorded at 0-eV energy loss reveal well preserved heterochromatin masses along the nuclear envelope. Fine structural detail is particularly evident in the high magnification, uranium-enhanced image (recorded at 120 eV) of the region indicated in Fig. 1C.

Sections of ~30- and 90-nm thickness were obtained by ultramicrotomy with a diamond knife (Drukker), and were picked up onto finder grids. These lettered grids allow particular cells in the section, first imaged in the fluorescent microscope, to be found later in the electron microscope. Electron micrographs were obtained with a Gatan 650 slow scan cooled CCD detector on a Zeiss EM 902 transmission electron microscope equipped with an imaging spectrometer (Bažetzjones and Hendzel, 1999). All images were corrected by dark current subtraction before being used for analysis. Energy filtered images recorded before an inner shell ionization edge do not contain element-specific information, but serve as a reference for the element-enhanced image recorded at an energy loss on, or just beyond, the inner shell ionization edge. In the case of phosphorus imaging, for example, an image recorded at an energy loss of 120 eV serves as a reference for the phosphorus-enhanced image recorded at 155 eV. A phosphorus map can be formed by subtracting the 120-eV image from the 155-eV image, after alignment to within a pixel, and normalizing of the two images on a region that is known not to contain phosphorus or, by dividing the 155-eV image by the 120-eV image after alignment of the two images. Similarly, a reference image for nitrogen is recorded at 385 eV and the nitrogen-enriched image is recorded at an energy loss of 415 eV (Batetzjones and Hendzel, 1999). The sections are preirradiated before the images are recorded. Preirradiation stabilizes the sections and produces a state of terminal mass loss. Therefore, no mass loss occurs during the recording of the complete set of reference and element-enhanced images. All images are recorded with nearly equal exposures measured over the background resin to insure equal signal counting statistics at the detector level for all images (Bažetzjones and Hendl, 1999; Bažetzjones et al., 1999).

Examples of reference, element-enhanced, and element maps for phosphorus and nitrogen are shown in Fig. 1D-I. Blocks of chromatin on the nuclear envelope (top of image in Fig. 1, D-I), throughout the nucleoplasm, and at the surface of the nucleolus (Fig. 2, Nu), are clearly seen in the phosphorus map. The granular component of the nucleolus has less contrast than the blocks of chromatin, but more contrast than the nucleoplasm or the cytoplasm. Blocks of chromatin can also be seen in nitrogen maps, but distinguishing chromatin from the granular component of the nucleolus, for example, is more difficult because nitrogen is present in both protein and nucleic acid. The granular component of the nucleolus has a higher protein/nucleic acid ratio than that of chromatin.

In this paper we used the two window method (one reference image and one post-edge, element-enhanced image) to obtain the elemental maps. We have shown that unstained, 30-40-nm thick sections do not contribute to detectable multiple scattering events or produce mass-density artifacts in the net images (Bažetzjones and Hendl, 1999; Bažetzjones et al., 1999). It is always possible to find mass-dense structures, some of which contain no detectable element, and others with high levels of that element. Similarly, low mass-dense structures can be found that have either no detectable element or high amounts. Mass-density artifacts in the...
sections used in this study are only noticed where contaminating crystallites of salts are imaged.

A 600-μm condenser aperture and a 90-μm objective aperture were used. The energy-selecting slit aperture was set to correspond to 20 eV. To measure the relative amount of phosphorus or nitrogen in various morphologically defined structures in and around PML nuclear bodies, we used the technique called image EELS (electron energy-loss spectroscopy). Images of a selected field are collected at small steps (e.g., 10 eV) in the energy-loss spectrum, spanning, for example, the phosphorus L_{2,3} and nitrogen K edges. Masks are used to delineate objects of interest, and the mean signals under these masks are measured in every image. If a structure contains phosphorus, a steep increase in the energy-loss spectrum above 132 eV will emerge, along with a second steep increase beyond 150 eV, due to the delayed characteristic of this energy-loss event. The presence of nitrogen is more obvious than phosphorus, due to a sharper, less delayed event beginning at 385 eV (see Bazett-Jones et al., 1999). M ea-

Figure 1. Energy-filtered electron micrographs of a 90-nm section stained with uranyl acetate (A–C). Elastic images, collected at 0-eV energy loss (A and B), show chromatin and other structures as black on a gray background. An energy-loss image recorded at a uranium edge (120 eV; C) shows chromatin and other structures as white on a black background. The inset (4×) shows a 10-nm-diam chromatin fiber with periodicity (vertical fiber at center of inset). Fibers of <50% of this thickness are also prevalent in this section. An unstained section from the same block used in A–C was used to map phosphorus and nitrogen. A phosphorus reference image (120 eV; D), a phosphorus-enhanced image (155 eV; E), and a phosphorus map (F) are shown. Nitrogen reference (385 eV), enhanced (415 eV), and map are shown (G–I, respectively).

Bosivet et al. PML Nuclear Bodies and Transcription
surements of signal intensities in the masked regions were made with E-rgoVista 4.4 image analysis software.

Results

Structure of the PML Nuclear Body by Correlative Microscopy

It has been reported that PML nuclear bodies are sites of transcription and RNA accumulation (LaMorte et al., 1998). We chose to test this model and to extend the model to higher resolution by direct visualization without using heavy atom contrast agents. We used ESI, a sensitive TEM method, to map protein-based and nucleic acid-based regions in and around PML nuclear bodies. One advantage of this technique is that comparisons of nitrogen and phosphorus maps provide the ability to directly distinguish protein from nucleic acids in situ. Second, it is possible to quantify mass contributions from both the protein and the nucleic acid components of a structure and to map the spatial relationships of protein- and nucleic acid-based structures within the complex nuclear environment (Bazett-Jones and Hendzel, 1999). Third, the heavy atom contrast agents, which limit resolution and stain different biochemical components in a nonuniform or predictable manner, can be avoided.

Positive identification of PML nuclear bodies amongst other nuclear structures with ESI alone is difficult. To identify these structures definitively, we first labeled cells with antibodies against the NH$_2$ terminus of CBP, which we have shown to colocalize with PML in PML nuclear bodies in SK-N cells, by immunofluorescence microscopy. Indeed, in this cell line, CBP is as reliable a marker of PML nuclear bodies as for the PML protein itself. After embedding and thin sectioning, the same section can first be examined in the fluorescence microscope to identify the location of PML nuclear bodies in individual cells. The same section is then imaged at high resolution in the energy-filtering electron microscope (Bazett-Jones et al., 1999). A thin section showing a nucleus imaged by immunofluorescence is shown in Fig. 2 A. Five PML nuclear bodies can easily be identified. Two are indicated (Fig. 2 A, D, and E, arrows) near the nucleolus (Fig. 2 A, D, and E, Nu). The same section imaged at 155-eV energy loss is shown in Fig. 2 B at low magnification and superimposed with the immunofluorescence image in Fig. 2 C, to correlate the fluorescently labeled structures with the underlying ultrastructure. Two PML nuclear bodies are then de-
tectable in the high magnification phosphorus (Fig. 2 D) and nitrogen (Fig. 2 E) maps. The cores of each of these nuclear bodies are well contrasted in the nitrogen map, whereas the phosphorus content is low. Notably, there is no evidence of small fibers or granules rich in phosphorus. Such signals are always observed when RNA or DNA is present, even as structures in the 2–10 nm range. These results indicate that the cores are mainly composed of protein.

The Core of the PML Nuclear Body Is a Protein-based Structure

Qualitative analysis of such images (Fig. 2, D and E) indicates that the core of the PML nuclear body is depleted in nucleic acids. Fig. 3 shows high magnification images of a PML nuclear body (Fig. 3, left, phosphorous map, right, nitrogen map). The presence of RNA or DNA in this structure would likely resemble phosphorus-rich granules, as seen in the core of the interchromatin granule clusters and throughout the nucleoplasm (Hendzel et al., 1998, 1999), or more extended fibrils, reflecting noncompacted nucleic acids (Hendzel et al., 1999). Such structures are not apparent, even at high magnification (Fig. 3). To extend the morphological analysis, we quantified the phosphorus and nitrogen content in different regions around the PML nuclear body (Fig. 3, B and D). The core of the PML nuclear body, a region corresponding to condensed chromatin, a background region in the nucleoplasm, and a background outside of the cell comprised of the embedding resin. The signal over the resin was used to normalize all images and was adjusted to the same value for each image. Therefore, these represent background-stripped spectra on the basis of this normalization. The phosphorus map is shown in A and the nitrogen map in C. Energy-loss spectra of these regions spanning the phosphorus L_{2,3} edge (B) and the nitrogen K edge (D) are presented at 10-eV intervals. Structures are represented as white objects on a black background.

Figure 3. Quantification of phosphorus and nitrogen content of a PML nuclear body at high magnification. The three indicated areas were used for quantification of phosphorus and nitrogen. The left region corresponds to chromatin, the middle region to nucleoplasm, and the right region to the core of a PML nuclear body. A fourth region (not shown) corresponds to a region outside the cell, containing only the embedding resin. The signal over the resin was used to normalize all images and was adjusted to the same value for each image. Therefore, these represent background-stripped spectra on the basis of this normalization. The phosphorus map is shown in A and the nitrogen map in C. Energy-loss spectra of these regions spanning the phosphorus L_{2,3} edge (B) and the nitrogen K edge (D) are presented at 10-eV intervals. Structures are represented as white objects on a black background.
resin alone (not shown) were delineated with masks. The mean integrated signals from the energy-loss spectrum spanning the phosphorus L_{2,3} and nitrogen K edges were measured over the regions delineated by the masks (Fig. 3, B and D). No phosphorus or nitrogen can be detected in the embedding resin, indicated by the flat line in both spectra. The standard deviation in the phosphorus and nitrogen signal intensities was always <2% of the mean for the PML nuclear body, the chromatin, and the nucleoplasm. Error bars indicating these values would not show outside of the symbols on the graphs. The masks for the embedding resin, the chromatin, and the core of the PML nuclear body are approximately equal in area, whereas the mask for the nucleoplasm had to be smaller, ~0.15% of the area of the others. However, the standard deviation of the means of numerous small regions of the nucleoplasm was also <0.5%, indicating that differences in sampling between nucleoplasm and nuclear structures do not account for the differences in element concentrations measured. The energy-loss spectra of regions in the nucleus are consistent with the presence of phosphorus, characterized by the strong delayed edge above 150 eV. The phosphorus content (155 eV) of the core of the PML nuclear body was 1.4% above the nucleoplasmic background, whereas the chromatin was 6% above the nucleoplasmic background, corresponding to a signal 11 times higher for the chromatin compared with the PML nuclear body. In contrast, the nitrogen signal (415 eV) for chromatin is 9% over the nucleoplasm and the PML body core is 16% over the nucleoplasmic background. The phosphorus signal in the nucleoplasmic background is likely derived predominantly from phosphorylated proteins. Because of a comparable phosphorus signal in the PML nuclear body core and the nucleoplasm, and the absence of morphologically recognizable phosphorus-rich complexes, we conclude that the core of this subnuclear structure does not contain nucleic acid (DNA and RNA) and is composed only of protein.

**Phosphorus-rich Fibers Are Present at the Periphery of the Core of the PML Nuclear Body**

Although the use of thin sections (30 nm) is necessary for quantitative elemental analysis by ESI (electron accelerating voltage of 80 kV), we have found that thicker sections stained with uranyl acetate still contain qualitatively useful elemental information (Boisvert, F.-M., preliminary observations). Thicker sections allow one to follow fibers that would otherwise rapidly leave the plane of the section if they are not parallel to the section. Fig. 4 shows consecutive 90-nm sections through a PML nuclear body. The center of the PML body core is characterized by a hole in the nitrogen map (Fig. 4, 1, arrow). Blocks of condensed chromatin surrounding the core are evident in the phosphorus-enhanced maps (Fig. 4, 1, 2, arrows; Fig. 3 A). Extended thin fibers measuring as little as 2 nm in diameter are consistently visualized at the immediate periphery of the core of the PML nuclear body when imaged with high resolution (Fig. 5). Nascent RNA is the most abundant subnuclear structure that has been identified in an extended conformation (Malatesta et al., 1994), and it is likely, therefore, that these phosphorus-rich fibers are RNA-based.

**Figure 4.** Serial sections of 90-nm thickness of a PML nuclear body stained with uranyl acetate. Images in the left column are phosphorus-enhanced, recorded at 155 eV, and those in the right column are nitrogen-enhanced, recorded at 415 eV. The middle of the PML nuclear body is shown in section 1. The protein core is visible in the nitrogen-enhanced image, and a hole on the core is indicated with the arrow in the nitrogen-enhanced image. Arrows in the phosphorus-enhanced images indicate blocks of condensed chromatin. Structures are represented as white objects on a black background. The next section (2) still shows the protein core, whereas section 3 is now out of the core of the nuclear body. The last section (4) shows the chromatin closing back over the structure.
Localization of Nascent RNA at the Periphery of the PML Nuclear Bodies

Our ESI analysis indicates that the core of the PML nuclear body is composed mainly of protein, whereas the periphery contains DNA and/or RNA. In a previous report (LaMorte et al., 1998), the entire PML body was reported to contain RNA, based on fluorescently labeled uridine in permeabilized cells. To determine whether newly transcribed RNA is present in the vicinity of PML bodies, we chose to label unperturbed cells with FU. Incorporation of FU in nascent RNA is rapid and specific and does not require a cell permeabilizing step. Signal visualized with antibodies against halogenated nucleotides can be detected throughout the nucleoplasm after two-minute pulses, with increasing signal over time (Fig. 6, C, H, and M). Strong, but less punctate, nucleolar labeling can also be seen after ten minutes, concordant with the high level of transcription therein. The diffuse nature of this signal causes an apparent attenuation of the signal relative to the punctuate foci found outside of the nucleolus in deconvolved optical sections. Before deconvolution, the nucleolar signal appears dominant. This nucleolar signal indicates that the penetration of the antibody and the FU is not inhibited by dense, compartmentalized structures, such as the nucleolus. To determine the relationship between PML nuclear bodies and RNA synthesis and accumulation, we labeled cells with anti-CBP NH₂-terminal antibodies (Fig. 6, B, G, and L, red) and anti-FU antibodies (Fig. 6, C, H, and M, green) after 2-, 10-, and 60-min pulses of FU. After only two minutes, most PML nuclear bodies have FU-incorporated RNA on their peripheries. This FU incorporation appears as a small number of foci (between 1 and 3) on the periphery of a PML nuclear body. When pulses were extended to longer times, RNA is still seen on the periphery in a small number of foci, and does not accumulate inside the structure as visualized by CBP labeling (Fig. 6, E, J, and O). The FU labeling is consistent with the ESI data, leading us to conclude that the core of the PML nuclear body is a protein-dense structure where RNA synthesis and RNA accumulation do not occur. Transcription does appear to occur near the periphery of PML nuclear bodies, but not within.

Highly Acetylated Chromatin Surrounds PML Nuclear Bodies, but Is Not Found Within

Phosphorus mapping by ESI and FU incorporation detected by fluorescence microscopy indicated that there is no nucleic acid in PML nuclear bodies. Nevertheless, we wished to strengthen our conclusion that there is no chromatin within the PML nuclear body, but also to determine whether transcription occurs on the periphery. For this experiment, we imaged PML nuclear bodies by deconvolution of optical sections, using an antibody recognizing the transcription-associated, highest acetylated species of histone H3 (Fig. 6, Q and R; Boggs et al., 1996). As expected from the FU incorporation experiments and ESI analysis, the cores of PML nuclear bodies do not show any evidence for the presence of highly acetylated euchromatin. All of the PML nuclear bodies, however, are within one diameter (250 nm) of a block of highly acetylated chromatin (Fig. 6 S). Moreover, some PML nuclear bodies are associated with multiple blocks of acetylated chromatin (Fig. 6 T; also see blocks of chromatin surrounding the PML nuclear body in the phosphorus map in Fig. 3 A). Further studies are required to determine whether transcriptionally active chromatin is indeed functionally associated with PML nuclear bodies, or whether the association is a random event. For example, it would be necessary to determine whether specific gene sequences associate with the periphery of PML nuclear bodies. From the ESI data and from the absence of highly acetylated chromatin or DAPI-stained chromatin within the core of the PML nuclear body, we conclude that the core of the structure is not a site of transcriptional activity.

Discussion

There has been an increasing interest in the PML nuclear bodies in the past few years because of their involvement in viral infection (Sternsdorf et al., 1997b; Maul, 1998) and their disruption and disappearance in acute promyelocytic leukemia (Borrow et al., 1990; de The et al., 1990, 1991; Kakizuka et al., 1991; Dyck et al., 1994; Koken et al., 1994). Whether this structure serves only as a storage site of regulatory factors, functions as a site for transcription by creating a suitable environment, or plays some other role will only be determined as its structure and composition are characterized. In this study, we have addressed several questions about the structure, organization, and function of the PML nuclear body.

First, we have shown that the core of the PML nuclear body is composed entirely, or almost entirely, of protein. There is no detectable nucleic acid in the 250-nm-diam core of the structure. This core is not a uniform sphere of...
Figure 6. Digital deconvolution microscopy of SK-N cells pulsed with FU for 2 (A–E), 10 (F–J), or 60 (K–O) min. FU was labeled with an anti-BrdU antibody (green, C, H, and M) and anti–CBP-NT antibody (red, B, G, and L). Merged images are shown in D, I, and N, and a high magnification of one PML nuclear body for each time course is displayed in E, J, and O. DNA stained with DAPI shows the nucleus of each cell (A, F, K, and P). Some cells were labeled with an antibody against the highest acetylated form of histone H3 (green, R) to reveal transcriptionally active/competent chromatin. PML nuclear bodies were detected with an antibody against PML protein (5E10) shown in Q. Images Q and R are merged to form S, and the indicated PML nuclear body is magnified in T.
Because the sensitivity of these procedures has been shown to be very high (Bazett-Jones and Hendzel, 1999; Bazett-Jones et al., 1999; Hendzel et al., 1999), our results rule out the possibility that the intense staining of these structures using the EDTA-repressive staining method is the result of an abundance of RNA within the structure. Similarly, the nuclear run-on experiment, which shows labeling of the PML nuclear bodies, has a greater potential for experimental artifacts than does the direct incorporation into RNA through addition of FU into the tissue culture medium.

We present a number of independent methods that demonstrate an absence of: bulk chromatin; acetylated chromatin; newly synthesized RNA; or phosphorus-rich nucleic acids; in the core of the PML nuclear body. Therefore, we conclude that it is highly unlikely that the PML nuclear body represents a higher-order nuclear structure that nucleates or aggregates around specific clusters of transcriptionally active genes or even single, highly active genes. Instead, the presence of transcription regulatory factors within the PML nuclear body and the blocks of chromatin and nascent RNA in the immediate vicinity of the PML nuclear body, raise the importance of further experimentation to test whether these structures are responsible for creating or establishing a transcriptional domain. A nuclear-neighbor hypothesis for regulation of nuclear events has been presented to explain a number of similar relationships involving subnuclear domains (Schul et al., 1998).

Subnuclear compartments that function to concentrate regulatory factors and complexes may be important in modulating gene expression (Schul et al., 1998). The interchromatin granule cluster (IGC), for example, is able to concentrate transcriptional coactivators in the protein-rich core, and is surrounded by transcriptionally active chromatin (Hendzel et al., 1998). Indeed, the peripheries of IGCs are sites of transcription of particular genes (Smith et al., 1999). PML nuclear bodies, which, incidentally, are generally associated with IGCs (Ishov et al., 1997), may also serve to concentrate regulatory factors that service transcriptional events on their surfaces. Genes that are implicated in requiring PML nuclear bodies for expression are early viral genes (Guldner et al., 1992; Chelbi-Alix et al., 1999). 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