Latent nuclear antigen of Kaposi’s sarcoma herpesvirus/human herpesvirus-8 induces and relocates RING3 to nuclear heterochromatin regions

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LANA, the major latency-associated nuclear antigen of Kaposi’s sarcoma herpesvirus/human herpesvirus-8 (KSHV/HHV-8), binds RING3 protein, one of five human homologues of the fsh (female sterile homeotic) gene product of Drosophila. In KSHV/HHV-8-infected cells LANA and the viral episomes accumulate in heterochromatin-associated nuclear bodies. Here we show that in several KSHV/HHV-8-negative cell lines derived from carcinomas, sarcomas and lymphomas, RING3 was expressed at low levels, primarily localized to the euchromatin, and dissociated from the chromosomes during mitosis. In contrast, in KSHV/HHV-8-infected body cavity lymphoma cells the bulk of RING3 localizes to the LANA nuclear bodies and remains associated with the chromosomes during cell division. KSHV/HHV-8-infected body cavity lymphoma cells expressed RING3 at much higher levels than cells without the virus. Transfection of full-length LANA, but not the C terminus alone, greatly induced RING3 gene expression, and LANA and RING3 co-localized even in the transfected cells, in the absence of KSHV/HHV-8 viral DNA. High levels of LANA expression led to the disappearance of heterochromatin in both human and mouse cells. We suggest that LANA and RING3 may create a local euchromatic microenvironment around the viral episomes that are anchored to the heterochromatin.

Introduction

LANA is the major latency-associated nuclear antigen encoded by human herpesvirus-8 (Kedes et al., 1997; Kellam et al., 1997; Rainbow et al., 1997). It is expressed at high levels in Kaposi’s sarcoma herpesvirus/human herpesvirus-8 (KSHV/HHV-8)-infected Kaposi’s sarcoma, body cavity lymphoma (BC; also called primary effusion lymphoma) cells, and cells of lymphadenopathy associated with multicentric Castleman’s disease (Boshoff & Weiss, 1998; Rainbow et al., 1997; Dupin et al., 1999; Katano et al., 1999). It accumulates in distinct nuclear bodies in KSHV/HHV-8-infected cells. Previously we have shown that the nuclear bodies are associated with the borders of heterochromatin in KSHV/HHV-8-positive BC cell lines, human–mouse hybrid cells and derivatives of the hybrids that had lost most of the human genetic material (Szekely et al., 1999). LANA-containing nuclear bodies remain associated with the chromosomes during mitosis (Szekely et al., 1998). LANA binds and anchors the circular viral DNA to the host cell chromatin during both the interphase and mitosis. LANA binds to the latent origin of replication of HHV-8 (Ballestas et al., 1999). It is also able to bind histone H1, a possible target for the tethering of viral episomes to the mitotic chromosomes (Cotter & Robertson, 1999). In this respect it is reminiscent of the Epstein–Barr virus (EBV)-encoded latent nuclear antigen 1 (EBNA-1). However, LANA and EBNA-1 accumulate into distinct, non-overlapping, nuclear bodies in EBV/HHV-8 dually infected BC cells (Szekely et al., 1998).

LANA is a nuclear phosphoprotein encoded by orf73 as part of a polycistronic message (Rainbow et al., 1997). It is expressed in all latently infected cells and is considered to be one of the major viral proteins that contributes to cell transformation. It binds and inhibits the transcriptional trans-
activating function of p53 and through this may protect cells from p53-induced apoptosis (Friborg et al., 1999). LANA also binds the retinoblastoma (Rb) protein and activates E2F-dependent transcription. LANA alleviates the growth inhibitory effect of Rb on Saos cells and, together with HRas, transforms primary rodent fibroblasts (Radkov et al., 2000). LANA associates with co-repressor protein SAP30, part of the mSin3 co-repressor complex that directs histone deacetylases to specific DNA sites (Krithivas et al., 2000).

LANA can modulate the expression levels of several different synthetic or natural promoters of both viral and cellular origin. LANA up-regulates its own expression along with v-cyclin and v-FLIP and a set of interferon-responsive cellular genes. On the other hand it down-regulates expression of v-cyclin and v-FLIP and a set of interferon-responsive cellular genes. LANA up-regulates its own expression along with v-cyclin and v-FLIP and a set of interferon-responsive cellular genes. LANA up-regulates its own expression along with v-cyclin and v-FLIP and a set of interferon-responsive cellular genes.

### Methods

#### Cell culture and transient transfection

The cells were grown at 37 °C with 5% CO₂ in Iscove's modified Dulbecco's cell culture medium containing 10% foetal bovine serum, 2 mM l-glutamine, 100 U/ml penicillin and 100 U/ml streptomycin. The absence of mycoplasma contamination was examined routinely by Hoechst 33258 staining. The cell lines used in this study were the following: HHV-8-carrying human BC lines BC1, BCBL1; MCF7 human breast carcinoma; HeLa human cervical carcinoma; SW480 human colon carcinoma; ECV human endothelial cells; EBV-immortalized lymphoblastoid lines LCL 94C1 and LCL Nadia; Jurkat human T cell lymphoma; NKL92 human NK cell lymphoma; Burkitt lymphoma lines with different states of EBV latency, such as DG75 (EBV⁺), Rael (EBV⁺, type I), Raji (EBV⁺, type III), Mutu I (EBV⁺, type I), Mutu III (EBV⁺, type III), Akata(−) (EBV⁻) and Akata(+) (EBV⁺, type I), as well as L mouse fibrosarcoma cells. Transient transfections of full-length LANA cDNA, inserted into a pCDNA1 vector were done according to the manufacturer's instructions using lipofectamin Plus reagent (GIBCO). Empty vector or a pBabe-EBNA-5 construct was used for control transfections. Construct containing GFP–LANA C terminus was described by Szekely et al. (1999).

#### Immunofluorescence microscopy

Cells grown on coverslips or centrifuged (cytospin) onto glass slides were fixed in methanol:acetone (1:1) at −20 °C for 20 min and then re-hydrated in PBS for 20 min at room temperature. The primary antibodies used in this study were human anti-LANA (antisera collected from patients suffering from the classical form of Kaposi’s sarcoma at the Dermatology Unit of Debrecen Medical School, Hungary; a gift from Attila Juhasz) and rabbit polyclonal anti-RING3 antibody (Platt et al., 1999). Rhodamine-conjugated rabbit anti-human (DAKO) or FITC-conjugated swine anti-rabbit (DAKO) antibodies were used as secondary antibodies. Control transfection of pBabe-EBNA-5 was stained with a mouse monoclonal, JF186 (Finke et al., 1987). Rhodamine-conjugated horse anti-mouse (Vector) was used as secondary antibody. The sera were diluted in blocking buffer (2% BSA, 0.2% Tween-20, 10% glycerol, 0.05% NaN₃ in PBS). Following a 1 h incubation at room temperature with primary antibody, the cells were washed three times in PBS and then incubated for 30 min with the secondary antibody. Double staining between LANA and RING3 was performed in the following order: rabbit anti-RING3, FITC-conjugated

### Table 1. Cellular distribution and staining intensity of RING3 in different cell lines

<table>
<thead>
<tr>
<th>Cell line</th>
<th>RING3 staining pattern</th>
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<tr>
<td>BCBL1 (HHV-8, EBV)</td>
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</tr>
<tr>
<td>BC1 (HHV-8)</td>
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<tr>
<td>MCF7</td>
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<tr>
<td>HeLa</td>
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<td>ECV</td>
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<td>SW480</td>
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<td>LCL 94C1</td>
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<tr>
<td>LCL Nadia</td>
<td></td>
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<tr>
<td>Akata (EBV⁻)</td>
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<td>Akata (EBV⁺)</td>
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<tr>
<td>Mutu I</td>
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<td>Mutu III</td>
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<td>Rael</td>
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<td>Raji</td>
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<tr>
<td>DG75</td>
<td></td>
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<tr>
<td>Jurkat</td>
<td></td>
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<td>NKL92</td>
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<td>L cells (mouse)</td>
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#### Weak (homogeneous or fine speckled)

#### Intense (distinct nuclear bodies)
swine anti-rabbit serum, normal rabbit serum, human anti-LANA and rhodamine-conjugated rabbit anti-human serum. DNA was stained by Hoechst 33258 or by propidium iodide. Each incubation was followed by three washes in PBS. The coverslips or glass slides were mounted with 70% glycerol containing 2.5% DABCO anti-fading agent (Sigma). Images were collected using a Leica DM RB microscope, equipped with Leica PL Fluotar 100× and 40× oil immersion objectives. Composite filter cubes were used for the FITC, Texas red/rhodamine and Hoechst 33258 fluorescence, respectively. The pictures were captured with a Hamamatsu dual mode cooled CCD camera (C4880), recorded and analysed on a Pentium PC computer equipped with an AFG VISIONplus-AT frame grabber board using Hipic5.1.0 (Hamamatsu), Image-Pro Plus (Media Cybernetics). Digital images were assembled using Adobe Photoshop software. Optical sectioning and threedimensional reconstitution was carried out using a Zeiss Axiophot microscope, equipped with 16× oil Plan-Neofluar NA 0.5, 63× oil Plan-Apochromat NA 1.4 and 100× oil Plan-Neofluor NA 0.7–1.3 objectives, illuminated with an Osram HBO 100 W mercury short arc lamp. The following excitation filters, mounted in the computer-controlled filter wheel, were used in this study: single band UV exciter for Hoechst (84360), single band blue exciter for FITC (84490), single band green exciter for TRITC (84555). The emission filter was a multiple band pass filter (84000) mounted on a static stage. All filters were purchased from Chroma Technology. The filter wheel, dual shutter and Z axis motor were controlled through a LEP MAC2000 Communication Interface 73000400 using an RS-232 serial connection (all devices from Ludl Electric Products). Images were captured with a PXL cooled CCD camera (Photometrix) operating at −25 °C, using 12 bit (4096 greyscale level) capture mode. The hardware control and image processing was provided by a Pentium PC computer equipped with a 600 MHz Pentium III processor and
Fig. 3. The bulk of LANA co-localizes with the bulk of RING3 in the nuclear bodies of BCBL1 cells as shown on the three-dimensional reconstitution of an optically sectioned nucleus (11 sections 0.2 μm apart). LANA (red), RING3 (green), DNA (blue), co-localization (yellow). To preserve the depth perspective, the maximum intensity projection algorithm was used on the individually de-blurred optical sections, creating stereo pictures. The effect of a stereo-image is achieved by crossed eyes, producing a third image between the left and right panels, or with the help of stereoscopic glasses.

512 Mb RAM. The operation system was LINUX Mandrake 7.0. The imaging programs were written and operated using the Isee 5.1 graphical programming system (Inovision). Our program ST-FITC-Rhodamine-Hoechst-bin1 was a novel hybrid version of trooper3 and stereotrooper (Holmvall & Szekely, 1999) that produces both single and stereo-projected three-colour images from a series of wide-field pictures where the out-of-focus blur was removed by nearest neighbour de-convolution and the resulting images were built up using a maximum intensity projection algorithm. This novel program also automatically compensated for any shining through from FITC, TRITC and Hoechst 33258 colour channels. The program chromat (Holmvall & Szekely, 1999) was used on individual optical sections created by the program ST-FITC-Rhodamine-Hoechst t-bin1.

■ Ratio RT–PCR assay. mRNA was isolated from pCDNA4-LANA-transfected and pCDNA4 MCF7 cell lines after 0 h, 24 h, 36 h and 72 h using Dynabeads (Dynal) according to the manufacturer’s recommendation. The mRNA samples were reverse-transcribed with Superscript II (Life Technologies) according to the manufacturer’s recommendation in a 20 μl volume. Ten μl of the mRNA samples was reverse-transcribed with oligo(dT)$_{18}$ primer. The RT–PCR reactions were performed on a Rapidcycler (Idahotech) capillary PCR machine in a 10 μl volume. The amplification reactions contained 0.5 μl of cDNA template, 50 mM Tris pH 8.0, 500 μg/ml BSA, 3 mM MgCl$_2$, 200 μM dNTP, 0.3 μM of GAPDH primers (GPDH5 5’ ACCACAGTCCATGCCA-TCAC 3’, GPDH3 5’ TCCACCCCTTGCTGTA 3’), 1 μM of RING3 primers (RING35 5’ AGGCCCTGCATCTGCTGGA 3’, RING33 5’ GTGCTGGCCTAGGCCCTCAAGA 3’) and 0.4 U Platinum Taq polymerase (Life Technologies). The reactions were cycled 27, 30 and 33 times, after an initial 15 s denaturation at 94 °C, under the following conditions: 94 °C, 0 s; 55 °C, 0 s; 72 °C, 30 s. The PCR products were separated on a 2.5% agarose gel in 1 × TAE buffer in the presence of 10 μg/ml ethidium bromide. The bands were visualized in an Intelligent Dark Box II (FujiFilm) and quantified using ImageGauge 3.122 software (FujiFilm).

Results
RING3 accumulates in nuclear bodies in HHV-8-positive BC cells but not in virus-negative cells

Immunofluorescence staining of different cell lines with rabbit antibodies raised against the RING3 protein showed either weak homogeneous or a fine speckled pattern in most of the lines studied (Table 1 and Fig. 1). In KSHV/HHV-8-negative cell lines RING3 predominantly localized to nuclear areas with low DNA condensation levels (Figs 1 and 2a).
RING3 dissociates from the chromosomes of the mitotic cell of uninfected SW480 and MCF7 cell lines but remains associated with the chromosomes of the KSHV/HHV-8-positive BCBL1 cells, where it co-localizes with LANA. In contrast HHV-8-positive BC lines BC1 and BCBL1 showed intense RING3 staining localizing to LANA like nuclear bodies that associated with the heterochromatin. This differential localization was further confirmed by analysing HHV-8-negative (SW480) and -positive (BC1) nuclei with the help of our computer program CHROMATIN. We used out-of-focus blur-free optical sections of nuclei as input for the analysis. The program created a 40 step scale of the blue fluorescence intensities of bisbenzidine-stained nuclei and calculated the total amount of green (RING3), red (LANA) and blue (DNA) signals corresponding to the pixels that belonged to the individual levels of the scale. In the SW480 cells the majority of the RING3 staining was associated with areas with low DNA condensation levels (on the left side of the DNA distribution curve) (Fig. 2a), whereas in BC1 nuclei the majority of the RING3 signal was primarily associated with the high DNA density areas (had the maximal peak at the right side of the DNA distribution curve). In BC1 cells RING3 showed an almost identical distribution to LANA (Fig. 2b). Double immunofluorescence staining of LANA and RING3 showed an almost complete co-localization between the two proteins. The co-localization was further confirmed by optical sectioning and three-dimensional reconstitution of double-stained cell nuclei (Fig. 3).

Controls included in the double fluorescence staining experiments involved replacing the LANA-specific primary antibody with normal human serum or human serum reacting with nucleolar antigens and alternatively replacing the antibody to RING3 with normal rabbit serum.

As previously reported (Szekely et al., 1999; Ballestas et al., 1999), LANA remains associated with the chromosomes during mitosis. In view of the co-localization of LANA with RING3 in interphase cells we investigated the distribution of RING3 in KSHV/HHV-8-infected and -uninfected metaphase cells. As shown in Fig. 4, RING3 relocates from the euchromatin-containing regions of the interphase nucleus of KSHV/HHV-8-uninfected cells to the cytoplasm during mitosis. In marked contrast, in KSHV/HHV-8-infected cells RING3 remains associated with mitotic chromosomes where it co-localizes almost perfectly with LANA. These findings
Fig. 5. (a) Transfection of LANA induces the increased expression of RING3 in an MCF7 human cell line as shown by double immunofluorescence staining. LANA (red), RING3 (green), DNA (blue). (b) Transfection of LANA induces increased expression of RING3 mRNA as shown by RING3/GAPDH ratio RT–PCR as a function of time (x axis) and RING3/GAPDH ratio (y axis).
indicate that the previously demonstrated (Platt et al., 1999) interaction of LANA with the ET domain of RING3 is strong enough to relocate RING3 almost completely to mitotic chromosomes during anaphase.

**LANA induces increased expression of RING3**

In addition to its relocation to LANA-containing nuclear bodies RING3 appeared to stain much more strongly in KSHV/HHV-8-infected than in -uninfected cells. This could suggest either a stabilization and accumulation of RING3 in these structures, or that the presence of the virus or even LANA itself might induce RING3 expression. As shown in Fig. 5(a), transfection into MCF7 cells of a plasmid directing the expression of full-length LANA from a CMV promoter resulted in increased immunofluorescence staining for RING3. The same result was obtained when LANA was introduced into SW480 human colon carcinoma or L mouse fibrosarcoma cells. No increased RING3 expression was observed when the C-terminal 182 amino acids were expressed as a part of a GFP fusion protein (data not shown).

LANA has recently been shown to induce several cellular genes (Renne et al., 2001) and RING3 was shown to be up-regulated in response to parasitic infections (Lau et al., 2001). To examine the possibility that LANA might induce expression of the RING3 gene, we carried out ratio PCR by quantifying RING3 and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) RT–PCR products from the same reaction mixture after different numbers (30, 33, 36) of amplification cycles. These experiments were carried out at different time-points (24, 48, 72 h) after transfection and were compared with RING3/GAPDH ratios from empty vector-transfected cells. These experiments clearly showed an increase of RING3 mRNA levels in LANA-transfected cultures (Fig. 5b).

**LANA and RING3 form nuclear bodies in transfected cells even in the absence of the viral genome and lead to the dissolution of heterochromatin**

LANA anchors HHV-8 DNA to the host chromatin. Introduction of a plasmid with the KSHV/HHV-8 latent origin of replication into LANA-expressing BJAB cells leads to the formation of distinct nuclear bodies (Ballestas et al., 1999). After transfecting LANA into MCF7 and SW480 cells we observed that a fraction of cells produced a stippled nuclear staining pattern. Double immunofluorescence staining and three-dimensional reconstitution from a series of optical sections showed that in these KSHV/HHV-8-negative cells transfected LANA also co-localized with RING3 in focal aggregates reminiscent of the nuclear bodies that are present in latent virus-harbouring cells (Fig. 6a). These focal aggregates regularly associated with the perinucleolar heterochromatin in human cells. High magnification images of the perinucleolar regions of the transfected cells showed decreased local DNA staining at the sites of focal LANA aggregates (Fig. 6b). Interestingly, in cells that showed high levels of LANA expression with homogeneous nuclear distribution, the peri-
nucleolar heterochromatin completely disappeared. Because it is difficult to clearly demarcate the euchromatin/heterochromatin borders in human cells we examined mouse L cells, in which the heterochromatin forms very distinct blob-like structures. Transfection of LANA produced mainly intense homogeneous nuclear staining in the L cells with the almost complete disappearance of heterochromatin (Fig. 7). Interestingly both mouse and human cells that expressed high levels of exogenous LANA and induced RING3 showed decreased overall staining with the DNA-binding dye bisbenzidine (Hoechst 33258), which binds to the AT-rich minor groove (Fig. 7a, b) but not with the intercalating dye propidium iodide (Fig. 7c, propidium iodide staining in red), indicating that the decrease in staining intensity was not due to DNA degradation. As a control we overexpressed EBNA-5 in MCF7 cells (Pokrovskaja et al., 2001). EBNA-5 did not induce any visible change in the chromatin distribution or bisbenzidine staining intensity (Fig. 7d). Control experiments were performed in which Hoechst 33258 was added in various incubation steps to exclude the possibility that the staining order affected the binding of Hoechst to DNA. Independently of staining order, the overall DNA staining was decreased in the LANA-expressing cells.

Discussion

Yeast two-hybrid screening identified RING3, a human homologue of the Drosophila fsh gene, as a binding partner of LANA. The binding was further confirmed by GST pull-down and co-immunoprecipitation assays. Association of RING3 with LANA leads to the phosphorylation of LANA through the recruitment of an as yet unidentified kinase (Platt et al., 1999).

RING3 is a member of the family of bromodomain-containing proteins that include the SWI/SNF-type chromatin remodelling proteins, histone acetylases and CBP/p300 CREB-binding co-activators. NMR studies on the structure of bromodomain suggested that four of them form alpha-helix bundles that may serve as a chromatin-targeting module that can interact specifically with acetylated lysines (Dhalluin et al., 1999).

RING3 belongs to the BET subgroup of bromodomain proteins, members of which contain two bromodomains and an ET motif, a protein–protein interactive surface. This group also includes, besides the Drosophila fsh, the yeast BDF proteins that regulate transcription through interaction with general transcription factors.

The human genome encodes at least four additional BET subgroup proteins: BRDT, ORFX, MCAP and HUNK1 (Jones et al., 1997; Dey et al., 2000). Interestingly the gene encoding RING3 is the only gene of the MHC class II region of human 6p21.3 that is not obviously associated with the immune system (Beck et al., 1996).

The function of RING3 is as yet poorly understood. In Drosophila, fsh is implicated in the establishment of the segments in the early embryo (Beck et al., 1992) by interacting with the trithorax pathway (Mozer & Dawid, 1989). The murine RING3 homologue (Fsrg1) may play a regulatory role in folliculogenesis and in the hormone-dependent remodelling of epithelia (Rhee et al., 1998). RING3 is expressed at high levels in testis and may also play a role in the regulation of spermatogenesis (Taniguchi et al., 1998). A recent report found that RING3 expression was increased in several organs as a result of parasite infection, suggesting that
it could represent an early response gene involved in the innate immune response (Lau et al., 2001). In the present paper we also show that LANA activates RING3 gene expression.

We also show that in KSHV/HHV-8-uninfected cells RING3 localizes to euchromatin regions in the interphase nucleus. During mitosis, RING3 appears to be released to the cytoplasm, similar to other transcription regulators, e.g. CBP, SP1, TFIIIB (Dey et al., 2000). In this respect RING3 differs from another recently described human member of the fsh family, MCAP, which has recently been reported to be located in heterochromatin regions and to remain associated with condensed chromosomes during mitosis (Dey et al., 2000). This difference in the sub-cellular localization of RING3 and MCAP may point to different functional roles. Interestingly, MCAP contains a long C-terminal extension, absent in RING3, distal from the ET domain, which characterizes all members of the fsh family.

We found that the bulk of RING3 co-localizes with the bulk of LANA in the nuclear bodies of KSHV/HHV-8-positive BC cells. These LANA-positive nuclear bodies are located on the euchromatin/heterochromatin interface in BC cells or in human–mouse hybrids. This localization is preserved when LANA is produced from an exogenously introduced expression vector. This markedly different localization of RING3 in KSHV/HHV-8-infected versus -uninfected cells strongly suggests that its interaction with LANA, previously demonstrated biochemically (Platt et al., 1999), is strong enough to quantitatively relocate RING3 to sub-nuclear structures containing KSHV/HHV-8 episomal DNA and LANA.

Importantly, however, high level expression leads to the dissolution of both human and mouse heterochromatin with accompanying bisbenzidine (Hoechst 33258) but not propidium iodide hypochromazia. Bisbenzidine binds to the minor grooves of AT-rich DNA. Culturing mouse cells in the presence of bisbenzidine inhibits the chromatin condensation of the pericentromeric heterochromatin. The staining difference between bisbenzidine and the intercalating dye propidium iodide indicates that DNA is preserved in the transected cells and bisbenzidine hypochromazia is not a result of selective DNA degradation. Another possible explanation for this phenomenon is that association of LANA or RING3–LANA complexes with DNA leads to decreased accessibility of minor grooves. However, the images presented in Figs 5(a), 6, 7 are more likely to indicate that LANA–RING3-containing heterochromatin regions show signs of heterochromatin dissolution.

We suggest that in virus-infected cells the LANA–RING3 nuclear bodies create a local microenvironment where the viral DNA is anchored to host heterochromatin but heterochromatization is inhibited in the immediate neighbourhood of the viral DNA LANA and/or RING3.

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References


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