Textural analysis of Nuclear Mitotic Apparatus antigen (NuMA) spatial distribution in interphase nuclei from human drug-resistant CEM lymphoblasts

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In tumour cell lines, the resistance of cancer cells to a variety of structurally unrelated chemotherapeutic drugs is termed multidrug-resistance or MDR. We reported previously \cite{6} that MDR leukemic cells displayed nuclear texture changes, as assessed by image cytometry. The nature of these changes remained uncertain but they could be associated with alterations of the nuclear matrix which could serve an important role in DNA organization and chromatin structure. Therefore, we have compared the textural features observed in G\textsubscript{0}/G\textsubscript{1} nuclei from human leukemic CEM cells and their MDR variant CEM-VLB, after staining of either DNA by Feulgen method or nuclear matrix by immunodetection of NuMA antigen on DNase treated samples. Chromatin or NuMA distributions within the nucleus were evaluated by image cytometry. Changes in textural parameters indicate that modifications of NuMA distribution observed in MDR cells are parallel to those observed at the whole chromatin level (i.e., a more decondensed and coarse texture with increase of Energy and Long-run sections and decrease of Contrast and Short-run sections). Moreover, Optical Densities measurements indicate that MDR cells seem to contain less NuMA, a datum confirmed by immunoblotting of nuclear proteins. In conclusion, chromatin changes observed by image cytometry in drug-resistant human leukemic CEM cells appear associated with modifications of the nuclear matrix structure.

Keywords: NuMA, nuclear matrix, drug resistance, texture, image analysis

1. Introduction

The development of resistance to chemotherapeutic agents is a limiting factor in the management of neoplastic diseases and results in treatment failure. One type of resistance, induced by structurally unrelated drugs, such as vinca alkaloids, anthracyclins, epipodophyllotoxins, or taxanes, is termed multidrug-resistance (MDR). The mechanisms underlying this phenomenon have been widely studied using \textit{in vitro} models, and MDR appears as a complex and multifactorial process \cite{20}. In many tumour cell lines with acquired MDR, drug resistance is associated with the expression of a membrane protein, P-glycoprotein (P-gp), the product of the \textit{mdr1} gene. This protein acts as an energy dependent efflux pump preventing accumulation of the drug in the cell \cite{10}. However, other mechanisms of MDR have been described and, in some cells, several mechanisms may operate simultaneously \cite{20}. This multifactorial process led to the suggestion that these complex actions might have multiple consequences on the MDR cell phenotype. Indeed, cellular concomitants of MDR are numerous but less information is currently available on the cytological changes associated with MDR. Among them, specific cytological alterations, as evaluated by image cytometry were reported in animal and human MDR cells \cite{5,16,17,22}. Most of these alterations concerned the nu-
clear texture which accounts for the condensation and distribution of the chromatin within the nucleus. They were not provoked by *mdr1* gene overexpression alone but appeared to be merely a consequence of other alterations occurring during selection of cells by cytotoxic drugs [6]. However, these alterations could have consequences on cell functions as they appear associated with a different sensitivity of chromatin to enzymatic digestion [31].

The nature of these additional changes remains unknown, but one possibility could be the presence of alterations of the nuclear matrix which could serve an important role in DNA organization and chromatin structure. Among matrix proteins, Nuclear Mitotic Apparatus Antigen (NuMA) [18] is a 236–240-kDa phosphoprotein coded by genes on chromosome 11q13 [27]. It consists of a central rod domain (207 nm -90%  α-helical) flanked by globular end domains and forms one of the longest coiled-coil currently known [12]. This antigen is distributed in the nucleus during interphase but translocates to the spindle poles in mitosis [4]. In interphase nuclei, NuMA is found on a subset of “core” filaments of the nuclear matrix [32] and appears to be present on the “diffuse skeleton” of the matrix [14]. NuMA plays a functional role in the organization of the microtubules of the mitotic spindle [7]. It appears to be a structural support to the nucleus during interphase and represents the structural component involved in the association of the pre-mRNA splicing apparatus with the nucleoskeleton [33].

In order to evaluate a role for the nuclear matrix in the chromatin textural alterations observed in MDR cells, we investigated by image cytometry the distribution of NuMA antigen in interphase nuclei from MDR cells.

### 2. Material and methods

#### 2.1. Cells

Lymphoblastic leukemia CCRF-CEM and CEM/VLB100 (MDR) cells were obtained from W.T. Beck, Memphis, TN and Z. Steplewski, Philadelphia, PA, respectively. These were maintained in RPMI-1640 (Gibco, Grand Island, NY) containing 10% FCS (Gibco) in a 5% CO₂ atmosphere at 37°C. CEM-VLB were grown in the presence of 0.4 µg · ml⁻¹ vinblastin. All experiments with resistant cells were performed on cells cultured in drug-free medium for more than 48 h. Resistance indices of these cells have been previously reported [6].

#### 2.2. DNA staining

Cells in the exponential growth phase were smeared by cytocentrifugation (Shandon cytopsin, Life Science Intern., Eragny, France) and air dried. Cell smears were fixed in an alcohol/formalin mixture (3 : 1) for 10 min. Hydrolysis was performed in 5 N HCl for 30 min at 22°C. The cells were then stained in freshly prepared Schiff’s reagent (pararosanilin chloride, C.I. 42500, Gurr, BDH, Poole, UK, 0.5% (w/v) in sulphite water) for 2 h in the dark at 22°C. After rinsing in sulphite water and distilled water, the samples were mounted in Acrypot (refraction index: 1.5, Surgipath, Richmond, Canada).

#### 2.3. NuMA staining

Cells in exponential growth phase were synchronized by incubation with thymidine (2 mM, 12 h), washed, incubated in culture medium for 8–10 h and smeared by cytocentrifugation. About 85% of cells were in G0/G1 phase. Before NuMA staining, DNA was removed from preparations by treatment with DNase I (Boehringer, Mannheim, Germany) 200 µg · ml⁻¹ in buffer (100 mM NaCl, 10 mM Pipes, pH 6.8, 300 mM sucrose, 3 mM MgCl₂, 4 mM vanadyl riboside complex, 10 mM leupeptin, 1.2 mM PMSF, 0.5% Triton ×100 (50 µl per slide)) for 5 min [13]. Quality of DNA removal was checked on randomly selected slides by Feulgen staining and image cytometry. Cells were then fixed in cold acetone for 5 min. Endogenous peroxidase was blocked by preincubating slides in 3% H₂O₂ for 5 min, followed by a 3% BSA wash for 20 min. Slides were then incubated overnight in the presence of anti-NuMA-1 primary antibody (Oncogene Science, Cambridge, MA) at 10 µg · ml⁻¹ in 0.5% BSA at 4°C. Antibody fixation was revealed either with FITC labelled goat anti-mouse Ig (1 : 40, 1 h, RT) or peroxidase labelled antibody (Jackson Immunoresearch, West Grove, PA – same conditions) followed by DAB staining.

#### 2.4. Image cytometry

After DNA or NuMA staining, image cytometry was performed with an image analysis system (SAMBA 2005, Unilog, Grenoble, France) coupled to a colour 3CCD camera (XC-007P, Sony, Japan) and a microscope (Axioskop, Karl Zeiss, Oberkochen, Germany). Measurements were made with a plan-achromat ×40 objective. At least 200 cells were analyzed on each
microscope slide and three separate experiments were performed. Twelve parameters were computed from each nuclear image: two geometric features (nuclear area and nuclear perimeter), one densitometric feature (integrated optical density IOD), and nine textural parameters. As the camera interface provides an optical density image at the acquisition step, the IOD is the summation of all pixels in the image in the nucleus and quantitatively measures the total intensity of staining by the DNA- or NuMA binding dye. Because DNA staining with Feulgen reaction was stoichiometric, DNA-IOD was directly related to the total nuclear DNA content. The distribution of nuclei according to DNA-IOD was plotted to provide cell-cycle distribution. The margins defining G0/G1 and G2 peaks were defined as modal IOD ± 15% [6]. At least 100 reference cells (mouse hepatocytes or human lymphocytes) were measured under the same conditions for the calibration of the normal diploid (2c) value. Nine textural features were calculated on the nuclear image after reduction to 16 grey levels by linear rescaling. Four features were calculated on the grey-level co-occurrence matrix: local mean of grey levels (LM), energy (E), entropy (ENT), and inertia (I). Five parameters were calculated on the run-length matrix: short run-length emphasis (SRE), long run-length emphasis (LRE), grey level distribution (GLD), run-length distribution (RLD), and run-length percentage (RPC) [8, 9, 11].

Univariate analysis was used to obtain the distribution, mean and SD of the nuclear parameters for each cell population. For a given parameter, histograms were computed and the significance of the differences between histograms from various cell populations was calculated using the non parametric Kolmogorov–Smirnov test on cumulative frequencies. For comparison between different cell lines and staining techniques, variations between sensitive and resistant cells were expressed as the deviation index, according to the formula: DI = (Value in resistant cells/Value in sensitive cells) – 1. This index value is 0 if no variation occurs between cells [6].

2.5. Immunoblotting

Proteins from lysed cells were separated by 12.5% SDS-PAGE and transferred to a nitrocellulose membrane (Biorad, Hercules, CA). Nitrocellulose sheets were blocked with Tris buffer containing 0.5% non fat dry milk and 0.1% Tween 20 for 2 h, incubated overnight at 4°C with 10 μg·ml⁻¹ anti-NuMA-1, washed 5 times, and incubated with HRP-conjugated goat antimouse antibody (1 : 2000, Calbiochem, La Jolla, CA). After 5 final washes, proteins were visualized by an ECL chemoluminescence system (Amer sham, Les Ulis, France) [25].

3. Results

3.1. DNA content

The DNA content of the sensitive and resistant CEM cell sublines was estimated by measurement of the Integrated Optical Density (IOD) of cell nuclei stained by the Feulgen method and appears in Fig. 1. The DNA-IOD modal value (i.e., the location of the major peak of the DNA histogram) decreased significantly in the resistant cell line as compared to sensitive counterparts. Moreover, in these CEM cells, drug-resistance is associated with a decrease in cells with very high DNA content, more than 5c. However, this decrease in DNA-IOD values in resistant cells obtained by drug-selection was not accompanied by variations in the distribution of cells among the cell cycle phases (data not shown).

![Fig. 1. DNA histograms in drug-sensitive CEM and drug-resistant CEM-VLB cells. DNA distributions in the two cell populations are significantly different (p < 0.001, Kolmogorov–Smirnov test on cumulative frequencies).](image-url)
3.2. Nuclear texture analysis after DNA staining

Textural features estimated by image cytometry are closely related to the location of a given cell in the cell cycle. Comparisons of texture have therefore been performed on cells selected within a given cycle phase. For this purpose, G0/G1 cells have been isolated on the basis of their DNA-IOD value and analyzed for chromatin texture. The results appear in Fig. 2. The total numbers of analyzed cells were 296 and 268 for CEM and CEM-VLB, respectively. CEM-VLB drug-resistant cells display significant variations of nuclear texture as compared with sensitive cells: a more decondensed chromatin (decrease of LM, ENT and I), with an increase in the distance between isodensity points (decrease of SRE and increase of LRE) confirming the appearance of a coarse chromatin texture with more homogeneous distribution of chromatin within the nucleus (increase of GLD measuring the uniformity of distribution of optical densities).

3.3. Nuclear texture analysis after NuMA staining

NuMA distribution was highly correlated with cell position within the cell cycle. For example, distribution is very different between interphase and mitotic cells (Fig. 3). For these reasons, NuMA textural analysis was performed on cells synchronized in G1 phase by thymidine block. Results of staining and of nuclear texture analysis after immunocytochemical NuMA staining of these cells appear in Figs 4 and 5. When compared to the data obtained on Feulgen stained nuclei, NuMA texture measured in drug-resistant CEM-VLB cells, and expressed as deviation index from sensitive cells’ values, display modifications similar to those observed in the whole chromatin after DNA staining (Fig. 5A). Moreover, these alterations could be also observed in cell nuclei pretreated with DNAse I (Fig. 5B), indicating that DNA does not seem to play a significant role in these changes. These spatial changes are not related to alterations in nuclear geometry, as surface and perimeter remain unchanged (28.8 ± 0.7 and 3.0 ± 0.1 in CEM, 27.8 ± 0.6 and 3.0 ± 0.1 in CEM-VLB, respectively – NS).

Finally, when NuMA expression was analysed by immunoblotting, drug-resistant CEM-VLB displayed a lower level of expression of this nuclear matrix protein (Fig. 6). Although the immunocytochemical staining
Fig. 4. Detection of NuMA antigen by immunocytochemistry in nuclei from drug-sensitive and drug-resistant CEM cells. NuMA was detected on synchronized cells, without prior DNase treatment, using anti-NuMA-1 monoclonal antibody and peroxidase-labelled goat anti-mouse secondary antibody. Texture analysis of NuMA distribution was performed by absorption cytometry on these stained nuclei. Magnification ×400. A: sensitive cells. B: resistant cells.

was not strictly quantitative, this data was confirmed by the values of the Integrated Optical Densities of NuMA staining which were consistently lower in drug-resistant cells (Table 1) when compared to the values observed in the drug-sensitive counterparts.

4. Discussion

Previous studies in human and animal MDR sublines have revealed numerous phenotypic alterations (for a review, see [2]). Among these alterations, the consequences of the resistance on the cellular morphology are still unclear. At this cytological level, we previously studied the phenotype of several hu-

Fig. 5. Variations of textural parameters in G0/G1 drug resistant CEM-VLB cells. Variations are expressed as deviation indices, as described in Material and methods. Panel A: □ DNA staining; ● NuMA staining. Panel B: NuMA staining without (●) or after (△) DNase I digestion of nuclei. Please note that the connecting lines between single unrelated parameters values were added for better legibility only in order to define a visually accessible nuclear “profile”.

Fig. 6. Immunoblot analysis of NuMA. Nuclear proteins from CEM cells (lane a), and CEM-VLB cells grown in presence (lane b) or absence (lane c) of vinblastin were resolved by SDS-PAGE, transferred to nitrocellulose membranes, and probed with anti-NuMA-1 monoclonal antibody.

Table 1

| Cell population | Number of cells analyzed | IOD after NuMA staininga |
|-----------------|--------------------------|--------------------------|
| CEM             | 196                      | 3.93 ± 0.12              |
| CEM-VLB         | 177                      | 3.37 ± 0.09b             |

aIODs are expressed in arbitrary units as mean ± SD.
b$p < 0.001$ (Kolmogorov-Smirnov test on cumulated frequencies).
man MDR cell lines. We showed that MDR cell lines obtained by drug selection displayed decreased DNA Feulgen stainability and a decondensation of chromatin with small but compact chromatin clumps [6]. The variations we observed in this study in CEM-VLB cells are similar to those previously described in these and other MDR cell lines (K562-Dox and HL60-Vinc).

As these changes did not appear to be associated with microphotometric negative errors related to progressive heterochromatization or to alterations of DNA lability [21], we tried to evaluate a role for nuclear matrix in these textural modifications. Such an implication could be suspected because, in rat resistant cells, resistance to chlorambucil was accompanied by a reduction of phosphoproteins in the nuclear matrix [28].

This nuclear matrix has been supposed to provide structural support for the nucleus and to play a role in the spatial organization of the genome and in the control of DNA replication and transcription [1]. Numerous studies have been devoted to the characterization of nuclear matrix in normal and neoplastic cells, either by biochemical or ultrastructural methods [15,23]. However, according to Brasch [3], it could be difficult to relate structure directly with biochemical characterization of the nuclear matrix fraction. Therefore, one analytical approach whereby isolation artifacts should be minimized is to monitor individual nuclear proteins in situ by immunomicroscopy, using monoclonal antibodies. Moreover, this approach could be optimized by using computer image analysis [1].

Taking these points into account we have analyzed directly by image cytometry the spatial distribution of NuMA in the nuclei from drug-sensitive and drug-resistant leukemic cells. This antigen has been chosen because numerous data support the possibility that NuMA serves a structural function either in the nucleus during interphase or at the polar ends of the mitotic spindle [7,14,26,33]. In this study, we observed that changes in NuMA texture in interphase MDR cells were similar to those observed at the whole chromatin level. These data suggest that previously observed nuclear changes could be explained by alterations of the higher order DNA organization mediated by the nuclear matrix. Moreover, a decrease in NuMA expression was observed in these MDR cells. Interestingly, in radioresistant rat embryonal cells, a progressive loss of nuclear matrix-associated proteins was observed, which correlated with increasing radioresistance and tumourigenicity in these cell lines. These authors therefore suggested that increased resistance associated with increasing tumourigenicity could be due to changes in nuclear matrix-mediated DNA organization [19]. Similarly, a significant decrease in nuclear matrix-associated topoisomerase II has been described in drug-resistant CHO cells [29].

However, these data on the spatial distribution of NuMA did not preclude changes in the functional characteristics of nuclear matrix, particularly its stability. For example, using DNase I digestion, it was suggested that more differentiated cells have a less stable nuclear matrix while the less differentiated ones have a more stable nuclear matrix. This cell differentiation stage-dependent stability of the nuclear matrix could probably be related to the nuclear activity and gene expression [30]. Since an unstable nuclear matrix may contribute to genetic instability [24] and selective pressure of drug exposure resulted in enhanced genetic stability in resistant cells [28], it could be supposed that nuclear matrix from MDR cells could display increased stability. Work is in progress in our laboratory to assess DNase I sensitivity of nuclear matrices from cell lines with different mechanisms of drug resistance.

In conclusion, chromatin changes observed by image cytometry in drug-resistant human leukemic CEM cells appear associated with modifications of the nuclear matrix structure.

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