

Tumorigenesis and Neoplastic Progression

Chromatin Organization Measured by *AluI* Restriction Enzyme Changes with Malignancy and Is Regulated by the Extracellular Matrix and the Cytoskeleton

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Given that expression of many genes changes when cells become malignant or are placed in different microenvironments, we asked whether these changes were accompanied by global reorganization of chromatin. We reasoned that sequestration or exposure of chromatin-sensitive sites to restriction enzymes could be used to detect this reorganization. We found that *AluI*-sensitive sites of nonmalignant cells were relatively more exposed compared to their malignant counterparts in cultured cells and human tumor samples. Changes in exposure and sequestration of *AluI*-sensitive sites in normal fibroblasts versus fibrosarcoma or those transfected with oncogenes, nonmalignant breast cells versus carcinomas and poorly metastatic versus highly invasive melanoma were shown to be independent of the cell cycle and may be influenced by proteins rich in disulfide bonds. Remarkably, regardless of degree of malignancy, *AluI*-sensitive sites became profoundly sequestered when cells were incubated with laminin, Matrigel, or a circular RGD peptide (RGD-C), but became exposed when cells were placed on collagen I or in serum-containing medium. Disruption of the actin cytoskeleton led to exposure, whereas disruption of microtubules or intermediate filaments exerted a sequestering effect. Thus, *AluI*-sensitive sites are more sequestered with increasing malignant behavior, but the sequestration and exposure of these sites is exquisitely sensitive to information conferred to the cell by molecules and biomechanical forces that regulate cellular and tissue architecture. (*Am J Pathol* 2005, 166:1187–1203)

It is well known from pathological specimens that normal and malignant cells differ in terms of nuclear structure. We have postulated that tissue organization depends on interactions between the extracellular matrix (ECM) and the nuclear structure and chromatin.^{1,2} Observations from *in vivo* and culture studies have provided evidence that the ECM may indeed influence higher order chromatin structure and gene expression.^{3–12} Furthermore, experiments on endothelial cells and fibroblasts have demonstrated that a mechanical tug to an integrin receptor at the cell surface could change the organization of both the cytoskeleton and the alignment of intranuclear molecules within 1 second, whereas force applied to non-integrin receptors such as LDL did not change nuclear organization.²

Using *AluI* restriction enzyme that cuts the widely distributed AG-CT sites, we asked five interrelated questions: 1) Do sensitivities to chromatin digestion differ in nonmalignant and malignant cells of different origins? 2) If so, can one use this information to distinguish between cells of low and high metastatic potential? 3) Does the microenvironment in general, and ECM in particular, affect chromatin structure regardless of degree of malignancy? 4) If so, is there some specificity with regards to the nature of the ECM molecules used? And finally, 5) Is the cytoskeleton involved? We find that answers to the above questions are affirmative.

Materials and Methods

Cell Culture

We used primary uveal melanoma cell lines of low (OCM1a) and high (M619) invasive potential, and a highly invasive

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metastatic uveal melanoma cell line (MUM2B); the characteristics of these cell lines have been described in detail previously.¹³ Determination of the modal chromosome number revealed the OCM1a and M619 lines to be in the tetraploid range while the MUM2B line was pseudo-diploid. These cell lines were also characterized by their differential ability to form laminin-rich patterns that appear to control melanoma morphogenesis.¹⁴ OCM1a uveal melanoma cells were a generous gift from Dr. June Kan-Mitchell, Karmanos Cancer Institute, Wayne State University, Detroit, MI, and UM54 normal uveal melanocytes were a generous gift from Dr. J. William Harbour, Washington University, St. Louis, MO. Melanoma cells and uveal melanocytes were plated in EMEM (BioWhittaker, Inc., Walkersville, MD), and supplemented with heat inactivated 15% fetal bovine serum (Fisher, Ontario, Canada) without the addition of exogenous extracellular matrix molecules or growth factors.

Normal WI-38 human fibroblasts were obtained from the ATTC (Rockville, MD), and were maintained in 10% serum in DMEM. Telomerase, large-T, and *ras* transfected fibroblasts were developed by Hahn et al¹⁵ and maintained according to their protocols, and were a generous gift from Dr. Igor Roninson (University of Illinois at Chicago). HT1089 fibrosarcoma cells were obtained from the ATTC. No antibacterial or antifungal drugs were used in the maintenance of cell lines or in experiments, as their chronic use has been shown to interfere with the differentiation potential of other primary cell types.⁷ MCF10A breast epithelial cells and MDA-MB231 breast carcinoma cells were obtained from the ATTC, and were maintained on DMEM plus heat inactivated calf serum. The isolation and phenotypes of HMT-3522 human mammary epithelial cells^{16,17} both nonmalignant (S1) and their tumorigenic counterparts designated T4-2¹⁸ were described previously. All cell cultures were determined to be free of mycoplasma contamination using the GenProbe rapid detection system (Fisher, Itasca, IL).

Restriction Enzyme Assays of Interphase Cells

We used three assays to compare sequestration and exposure of *AluI* binding sites in interphase cells of different invasive behaviors. In the permeabilized cell assay, we exposed cells in monolayer cultures to *AluI* restriction enzyme after permeabilization with Triton X-100. In the cell smear assay, we mechanically dislodged cells from culture, applied them to glass slides, allowed the cells to air dry, and exposed the cells to *AluI* restriction enzyme, thus simulating a diagnostic cytological examination performed in a clinical setting (eg, the examination of cells extracted by fine needle aspiration biopsy, brush biopsy, or scraping). In the flow cytometry assay, we quantified differential digestion of chromatin between cell lines by *AluI* restriction enzyme over time, again using methodology that could be easily adapted to a clinical diagnostic setting.

Permeabilized Cell Assay

Cells grown as monolayer cultures in the absence of exogenous matrix molecules were exposed to 0.1% Tri-

ton X-100 (Sigma, St. Louis, MO) for 90 seconds followed by a rinse of DMEM. The preparation was exposed to *AluI* restriction enzyme (0.5 μ l in 40 μ l DMEM; Promega, Madison, WI) for 60 minutes to 24 hours. The permeabilized cultures were then exposed to ethidium bromide (25 μ l, 1 μ g/ml; Sigma) and photographed with a Leica inverted microscope (Leica, Bannockburn, IL).

Cell Smear Assay

After mechanically dislodging cells grown under monolayer conditions, the slurry was suspended in 1X PBS or serum-free DMEM, and a drop containing 15 μ l of the suspension was placed onto a glass slide. The drops were allowed to evaporate over 30 minutes to 1 hour. *AluI* restriction enzyme (0.5 μ l in 40 μ l) or *MspI* restriction enzyme (0.5 μ l in 40 μ l DMEM; Promega) was applied to the dried cells, and the preparation was placed in a humidified 37°C chamber to optimize enzyme activity and minimize enzyme evaporation. Endonuclease digestions were terminated at pre-designated time points (30 minutes and hourly increments thereafter up to 24 hours) to determine the optimum digestion time that would allow for discrimination of differential chromatin digestion between cell lines. Ethidium bromide was added to terminate the digestion, and the preparation was photographed immediately. Touch preparations of normal human tissue and human tumor tissue were made and air dried. The preparation was incubated with *AluI* restriction enzyme and the reaction was terminated with ethidium bromide at 5 and 24 hours.

To test the influence of different soluble ECM molecules on *AluI* sensitivity, we mechanically dislodged cells that had been growing under monolayer conditions from flasks and suspended the slurry of dislodged cells in 1X PBS or serum-free DMEM. Two rows of 15 μ l drops of the suspension were applied to a glass slide, 3 drops per row. In some experiments as described below, we added soluble laminin, Matrigel, FBS, RGD-C, collagen Type I, bFGF, EGF (all reagents from Clontech, Palo Alto, CA), circularized RGD (RGD-C, a kind gift from Renata Pasquallini) to one of the two rows of drops. RGD-C is known to bind to integrin receptors.² The test and control drops were then permitted to evaporate at room temperature for at least 1 hour, leaving "smears" of dried cells that had or had not been incubated with a test molecule. For each assay, cellular permeability was checked using the trypan blue exclusion method.

The *AluI* restriction enzyme (0.5 μ l in 40 μ l DMEM; Promega) was applied to each drop in a humidified 37°C incubator for up to 24 hours. The buffers used in these assays did not contain either DTT or mercaptoethanol to avoid removing any proteins within the cytoplasm or nucleus that might sequester *AluI* binding sites from enzyme digestion. Incubation with *AluI* was terminated by adding ethidium bromide (Sigma; 250 μ l, 100 ng/ml) to each drop per slide at 30 minutes and at hourly intervals thereafter up to 24 hours of incubation.

Scoring Results

In the cell smear assay and the permeabilized cell assay, samples were photographed with a Leica inverted microscope and were scored qualitatively from the micrographs as follows: 1) Nuclei in which no fluorescence was detected except for nucleoli were scored as complete digestion; 2) Nuclei in which fluorescence intensity was equal to that detected at time before application of restriction enzyme was scored as no digestion; and 3) Detection of non-nucleolar nuclear fluorescence that was less intense than detected before application of restriction enzyme was scored as partial digestion.

Flow Cytometry

To provide for a quantitative measure of DNA digestion, we performed flow cytometry. Cells from monolayer cultures of UM54 normal uveal melanocytes, poorly invasive OCM1a primary uveal melanoma, highly invasive M619 primary uveal melanoma, and highly invasive MUM2B metastatic uveal melanoma were suspended in DMEM and spun down at 1400 rpm for 5 minutes. The pellet was re-suspended in 0.1% Triton X-100, incubated for 1 minute at room temperature, and spun down again at 1400 rpm for 5 minutes. After re-suspension in DMEM, 0.5 μ l of *Alu*I restriction enzyme in 40 μ l of DMEM was added and the preparation was incubated at following time points: 0 (baseline), 1, 3, and 5 hours at 37°C. Propidium iodide (10 μ l/ml; Molecular Probes, Eugene, OR) was added at the conclusion of each digestion period. The cells were analyzed in a FACS Calibur (BD Bioscience, San Jose, CA) equipped with a 488 laser, detectors for forward and side scatter, and 520, 575, and 675 nm detectors. 10,000 cells were counted and the results were analyzed with FACS dot-plots and histograms. Three *Alu*I digestions monitored by flow cytometry were performed for each of the three melanoma cell lines. The percentage of cells in M2–M4 (roughly corresponding to 10² to 10⁴ fluorescence intensity) was calculated and compared between cell lines at 5 hours of digestion with the *t*-test. The PI signal, representing stoichiometrically the amount of labeled undigested DNA,¹⁹ was recorded immediately after cell permeabilization and labeling with PI and used as a baseline (0 time point), and was then measured for each cell line after 1, 3, and 5 hours of exposure to *Alu*I restriction enzyme. CellQuest software (BD Bioscience) was used to generate overlays of the histograms for each cell line at each time point in the digestion experiment.

Isolated Chromosome Set Assay

We have previously described a method by which the entire chromatin contents of nuclei can be extracted, producing a complete chromosome set for each cell thus manipulated.^{20,21} We removed chromatin sets from cells that had not been pretreated with any agent. All chromosome extractions and digestions were carried out under isotonic culture conditions in DMEM at pH 7.4; similar

results also were obtained using complete cell culture medium containing FBS.

Chromosome sets removed from metaphase cells of varying invasive potentials were studied under buffered media conditions that do not disturb the native structure of chromosomes.^{20–29} Cells from which chromosomes were to be removed were grown on small coverslips and placed in the center of a 35-mm plastic dish lid containing 2 ml of gassed DMEM. This preparation was allowed to equilibrate at 37°C in a 5% CO₂ buffered incubator before micromanipulation. For optimal micromanipulation conditions, cells were grown to near confluence to produce strong attachments to each other and ECM needed to resist the pulling forces associated with rapid (within 1 second) micropuncture and removal of chromosomes without causing cell detachment or death of the manipulated cell. Chromosome sets were obtained either in the presence or absence of colchicine and cytochalasin-B as reported previously.²⁰ All chromosome set extractions and digestions were carried out under isotonic culture conditions in DMEM at pH 7.4; similar results also were obtained using complete cell culture medium containing serum.

Chromosome sets were extracted from living metaphase cells by rapidly piercing the cell directly on the side of a mitotic plate with a glass microneedle and then laterally drawing out the chromosomes through the hole on the cell surface created by the micropuncture using a Leitz micromanipulator (Leica Microsystems). In certain studies which used treatments that cause cell detachment (mercaptoethanol, DTT, and proteinase K), chromosome sets were removed from the cell with a pipette and placed on culture substrata. The chromosome sets were then deposited to pre-designated areas defined by scratching the Petri dish lid with a fine needle. Narishige micropipettes (Narishige Scientific Instruments, Tokyo, Japan) were pulled with a Sutter micropipette puller (Sutter Instrument Company, Novato, CA) adjusted to produce long barrels approximately 1 to 5 μ m wide along a length of 40 to 100 μ m (tip widths were consistently less than 0.5 μ m).

In this isolated chromosome set assay, it is important to use restriction enzymes and proteases in the range of specific ion concentrations if they are to function efficiently. We used buffered media conditions that preserve physiological ion concentrations to maintain the extent of chromatin compaction observed in living cells while providing the appropriate concentrations of NaCl and MgCl₂ to permit enzymes to work.²⁰ The restriction enzymes were used in the absence of DTT or β -mercaptoethanol (which are typically included in commercially available kits used to digest DNA) to avoid interfering with DNA-associated proteins.

Restriction enzymes were dissolved in 2 ml of DMEM applied directly to 35 mm tissue culture dish lids on which the chromosome preparations were attached in the concentrations listed below. Changes in chromosome morphology were photographed by time-lapse video microscopy in real time. All microsurgical procedures were observed at 63 \times by phase and fluorescence microscopy with a Leica inverted microscope, and captured using a

time-lapse video recorder (Sony) and Pinnacle Image software (Pinnacle Systems, Modesto, CA).

We tested the following enzymes: *AluI* (5–50 units), Msp-I, (75 units, Promega), DNase I (1–20 units/droplet; Promega), micrococcal nuclease (1–10 units; Promega), *EcoRI* (5–50 units; Promega), *HindIII* (5–10 units; Promega), *BamHI* (5–10 units; New England Biolabs, Beverly, MA), RNase A (1–100 units; Sigma), RNase I (1–100 units; Promega), EMBO (3 units; Promega), Sau-III (20 units; Promega), *PstI* (30 units; Promega), XHO-1 (40 units; Promega). Ethidium bromide (25 μl , 1 $\mu\text{g/ml}$; Sigma) was added to the medium after 1 to 3 hours of digestion with restriction enzyme and the preparations were photographed with a Leica inverted microscope.

Permeabilized Cell Assay with Polymerized ECM Molecules

We designed a method to place different polymerized ECM molecules on glass slides to screen for changes in the sensitivity of *AluI*-sensitive sites in normal cells and tumor cells or tumor cells with varying degrees of invasive behavior. First, a coating of FBS or Type I collagen was applied to the glass slide and allowed to dry completely. Then a stencil was positioned on top of the dried FBS or collagen and clamped so that it would not move. The stencil was fashioned in such a way that it allowed application of a second ECM molecule over the dried FBS or collagen in square patches. Laminin or Matrigel were then applied to areas of the slide exposed by the stencil and the entire assemblage was baked in an oven for 1 hour at 90°C. The stencil was gently removed under sterile conditions, and cells were then seeded at subconfluent density onto the entire surface. The cultures were then exposed to 0.1% Triton X-100 (Sigma) for 90 seconds followed by a rinse of DMEM. *AluI* restriction enzyme (0.5 μl in 40 μl DMEM) was applied for 30 to 60 minutes after which time ethidium bromide (250 μl , 100 ng/ml) was placed on the preparation. Nuclei were photographed immediately and scored for digestion as described above.

Three-Dimensional Cultures

As reported previously, highly invasive primary and metastatic melanoma cells grow in monolayers on tissue culture plastic but form looping patterns when placed in three-dimensional (3D) culture conditions identical to those seen in tissue sections of primary aggressive and metastatic lesions resected from patients.¹⁴ Three-dimensional cultures were established by growing cells on Matrigel³⁰ (Collaborative Biomedical, Bedford, MA) or Type I collagen³¹ (Collaborative Biomedical) placed onto plastic tissue culture dishes to a depth of about 0.2 mm followed by polymerization for 1 hour at 37°C. The plates were then returned to the incubator and allowed to completely polymerize for several hours. Cells were then seeded at saturating densities (400 million cells/dish) on the polymerized 3D gel coatings, and the cultures were checked daily for the presence of looping patterns that

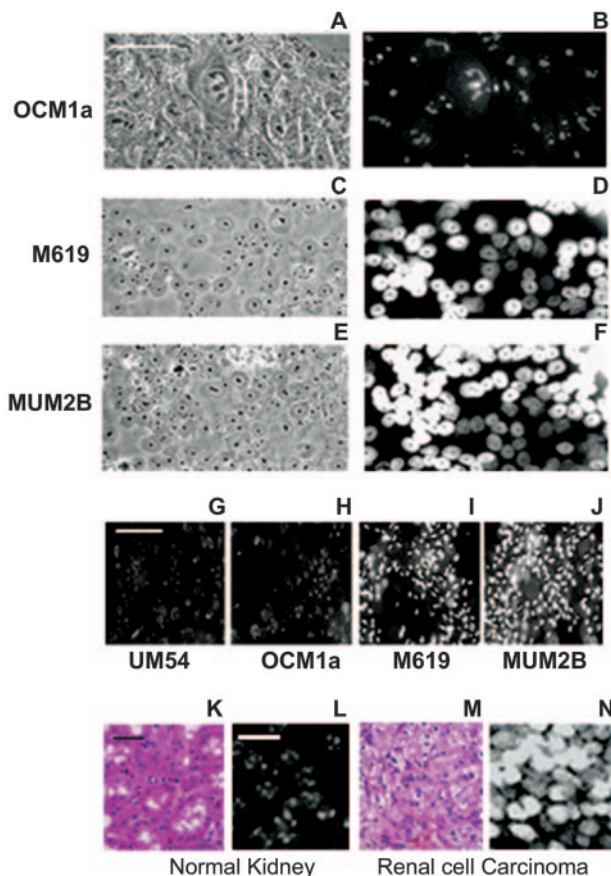


Figure 1. Differential sensitivity of interphase cells to *AluI* restriction enzyme. **A–F:** Permeabilized cell assays. **A:** Phase micrograph of intact poorly invasive OCM1a uveal melanoma cells grown on monolayers after 2 minutes of exposure to Triton X-100 for permeabilization followed by 30 minutes of exposure to *AluI* restriction enzyme. Cells were labeled with ethidium bromide. **B:** Fluorescent micrograph of the same preparation illustrated in **A**. DNA has been digested leaving only the *AluI* enzyme-resistant nucleoli fluorescent. **C:** Phase micrograph of highly invasive M619 uveal melanoma cells prepared as in **A**. **D:** Fluorescent micrograph of the same preparation illustrated in **C**. DNA is resistant to digestion by *AluI* restriction enzyme. **E:** Phase micrograph of highly invasive MUM2B metastatic melanoma cells after preparation identical to cells illustrated in **A**. **F:** Fluorescent micrograph of same preparation illustrated in **E** labeled and with ethidium bromide. **G–J:** Cell smear assays comparing UM54 normal uveal melanocytes, OCM1a, M619, and MUM2B melanoma cells. After 5 hours of exposure to *AluI* restriction enzyme, only the nucleoli are detected in the UM54 and OCM1a cells, but the chromatin of M619 and MUM2B cells is undigested and therefore brightly fluorescent. **K:** Normal renal tubular epithelium. **L:** After 5 hours of digestion of normal renal tubular epithelial cells from the region depicted in **K** with *AluI* restriction enzyme: only the nucleoli fluoresce. **M:** Renal cell carcinoma. **N:** After 5 hours of digestion with *AluI*, the chromatin of cells from the adjacent renal cell carcinoma depicted in **M** has resisted digestion and is brightly fluorescent. Scale bars: **A–F** = 50 μm ; **H–J** = 100 μm ; **K, L** = 50 μm ; **M, N** = 100 μm .

are characteristic for highly invasive melanoma cells.¹³ Techniques for growing mammary cells on 3D gels have been described^{30,31} (see review by Schmiechle and Bissell³²).

Transmission Electron Microscopy

Cells were fixed by adding in 4% buffered glutaraldehyde to the culture medium, followed by post fixation in osmium tetroxide. After dehydrations in ethanol, cultures were embedded in Epon. Ultra-thin sections were cut

Table 1. Summary of Response to DNA Digestions by *AluI* Restriction Enzyme Using Three Different Assays

Cell Line	Interphase cell assays		Metaphase cell assay isolated chromosome sets
	Permeabilized	Smear	
Melanoma			
OCM1a poorly invasive primary melanoma ^{13,71}	Sensitive	Sensitive	Sensitive
M619 highly invasive primary melanoma ¹³	Resistant	Resistant	Resistant
MUM2B highly invasive metastatic melanoma ¹³	Resistant	Resistant	Resistant
Fibroblast/fibrosarcoma			
WI-38 fibroblasts			Sensitive
HT1080 fibrosarcoma			Resistant
Fibroblasts + transfected oncogenes ¹⁵			
+ Telomerase			Sensitive
+ Telomerase + Large T			Sensitive
+ Telomerase + Large T + <i>ras</i>			Resistant
Breast epithelium/breast carcinoma			
MCF10A non-malignant breast epithelial cells	Sensitive	Sensitive	
MDA-MB231 metastatic breast carcinoma cells	Resistant	Resistant	
S1 non-malignant breast epithelial cells ³³	Sensitive		Sensitive
T4-2 tumorigenic breast cells derived from S1 ³³	Resistant		Resistant

perpendicular and parallel to the culture, stained with uranyl acetate and lead citrate, and were examined with a JEOL 1011 transmission electron microscope (Peabody, MA).

Results

Differential Chromatin Digestion by AluI Restriction Enzyme in Different Interphase Cells with Varying Malignant Behavior

The sequestration and exposure of *AluI* binding sites was tested qualitatively by exposing permeabilized cultured cells on glass slides to restriction enzyme (permeabilized cell assay, Figure 1, A–F) to maintain the structural integrity of the nucleus and cytoplasm. We also smeared cultured cells on slides and allowed them to dry (cell smear assay, Figure 1, G–J) to rapidly assess a large number of samples. Each assay was repeated 18 times with identical results.

In both the permeabilized cell assay and the cell smear assay, we detected striking difference in the susceptibility of chromatin to digestion by *AluI* in cells of different invasive potential (Table 1). For example, the chromatin of UM54 normal uveal melanocytes and poorly invasive OCM1a melanoma cells digested completely within 60 minutes of exposure to *AluI* restriction enzyme. Bright fluorescence was detected in nucleoli, but no fluorescence was demonstrated elsewhere in the nucleus. By contrast, the nuclei of highly invasive M619 and MUM2B melanoma cells resisted digestion by *AluI* restriction enzyme and were brightly fluorescent (Figure 1). In a parallel experiment, chromatin of MCF10A breast epithelial cells digested completely within 60 minutes whereas chromatin from MDA-MB231 breast carcinoma cells resisted digestion (data not shown).

In preliminary experiments, we tested the feasibility of detecting differences in the susceptibility of chromatin to *AluI* between normal and neoplastic human tissue in touch preparations. For example, the chromatin of normal

renal tubular epithelial cells digested completely within 5 hours, but chromatin from an adjacent focus of renal cell carcinoma resisted digestion by *AluI* restriction enzyme in the same time frame (Figure 1, K–N); the same differential digestion was detected between normal gastric epithelium and gastric carcinoma (data not shown).

To achieve quantitative assessment of chromatin digestion in the uveal melanocytic cell lines studied by cell smear assay (Figure 1, G–J), we measured DNA content in cell suspensions by flow cytometry after staining with propidium iodide (PI). Histograms comparing OCM1a, M619, and MUM2B melanoma cells with each other and the normal UM54 melanoma cell line are presented in Figure 2. There are striking differences in the percentage of cells in channels M2–M4 (corresponding to 10^2 - 10^4 fluorescence intensity on the x axis) over time. At 5 hours of digestion, only 1% of UM54 normal melanocytes are recorded in M2–4, compared with 52% of OCM1a cells, 73% of M619 cells, and 94% of MUM2B cells. The experiment illustrated in Figure 2 was repeated twice, and the percentage of cells in M2–M4 at 5 hours after digestion was compared between the three melanoma cell lines in three replicate assays using a *t*-test. There are significantly more MUM2B cells in M2–M4 than OCM1a cells ($P = 0.006$) and significantly more M619 cells than OCM1a cells in M2–M4 ($P = 0.02$). There were also more MUM2B cells than M619 cells in M2–M4 after 5 hours of digestion by *AluI* ($P = 0.054$). Thus, the quantitative flow cytometric assays showed that chromatin in highly invasive melanoma cells was more resistant to digestion with *AluI* restriction enzyme than poorly invasive melanoma cells and was consistent with the qualitative permeabilized cell and cell smear assays (Figure 1).

Differential Digestion of Chromatin by AluI Restriction Enzyme Is Independent of the Cell Cycle

The previous assays tested the digestion of chromatin by *AluI* in asynchronous populations of interphase cells. To

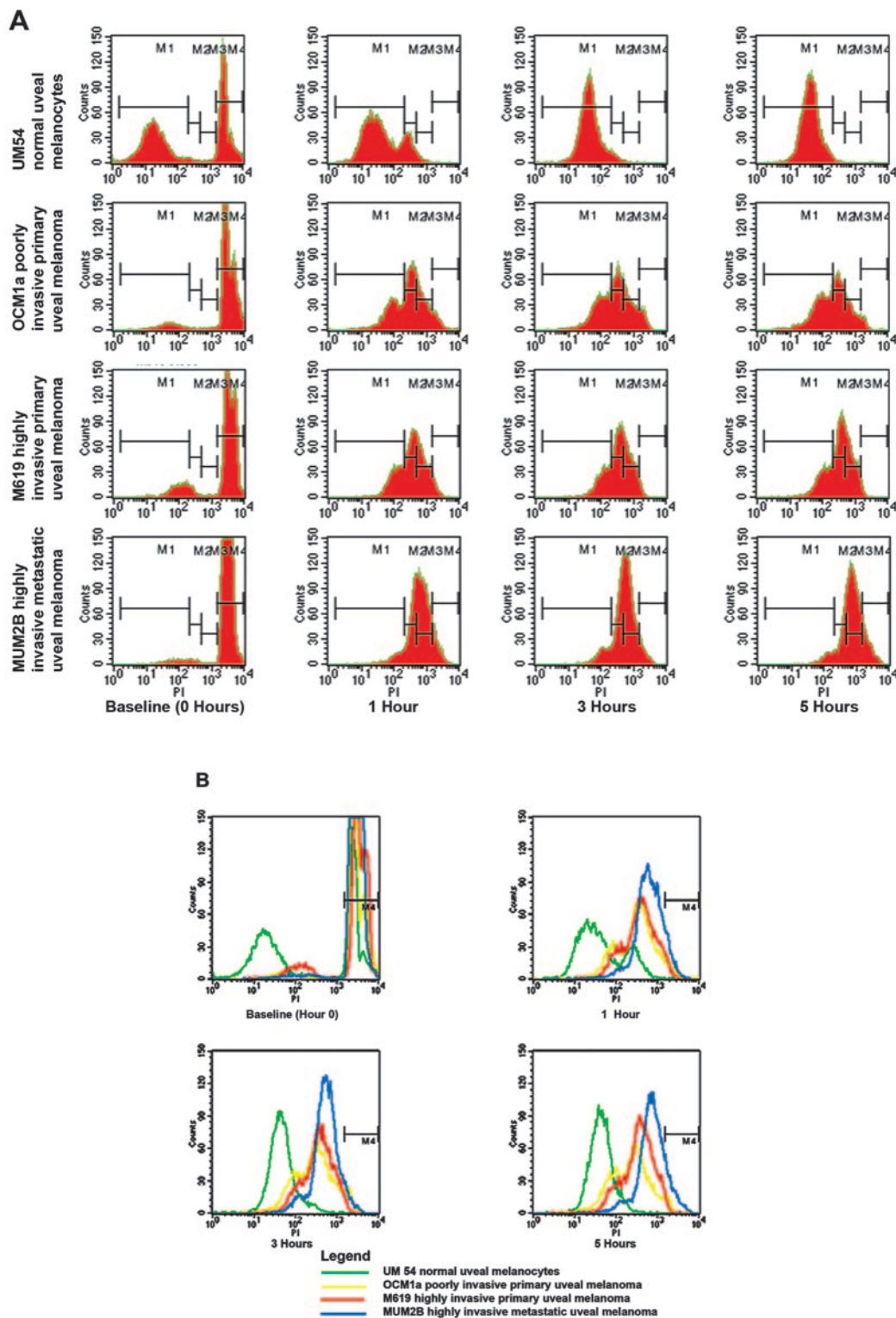


Figure 2. Flow cytometric study of UM54 normal uveal melanocytes, OCM1a poorly invasive primary melanoma cells, M619 highly invasive primary melanoma cells, and MUM2B highly invasive metastatic melanoma cells displaying the propidium iodide (PI) fluorescence intensity signal in the x axis, and cell count in the y axis for each cell line at each of the four time points. **A:** Gating for M1-M4 is indicated in the histogram for UM54 at time 0. The percentage of cells in each gate is reported for all four cell lines at time 0 and 5 hours of digestion. Note the differences between the percentages of cells in M2-4 between the four cell lines at 5 hours. The histograms at 5 hours of digestion parallel the cell smear assay performed in these cell lines and illustrated in Figure 1, G-J. **B:** For each time-point (baseline, 1, 3, and 5 hours), the histogram outline of each cell line is superimposed, color coded according to cell line as indicated in the legend. Note that the PI signal for UM54 normal melanocytes degrades rapidly at 1 hour and continues to degrade throughout the time-course. The PI signal for OCM1a cells degrades rapidly at 1 hour and gradually throughout the 5-hour time course of digestion by *AtaI*. The PI signal for M619 cells degrades at 1 hour but shows less degradation at 5 hours than OCM1a cells. The PI signal for MUM2B cells degrades the least at 1 hour of all cell lines and there is little signal degradation thereafter.

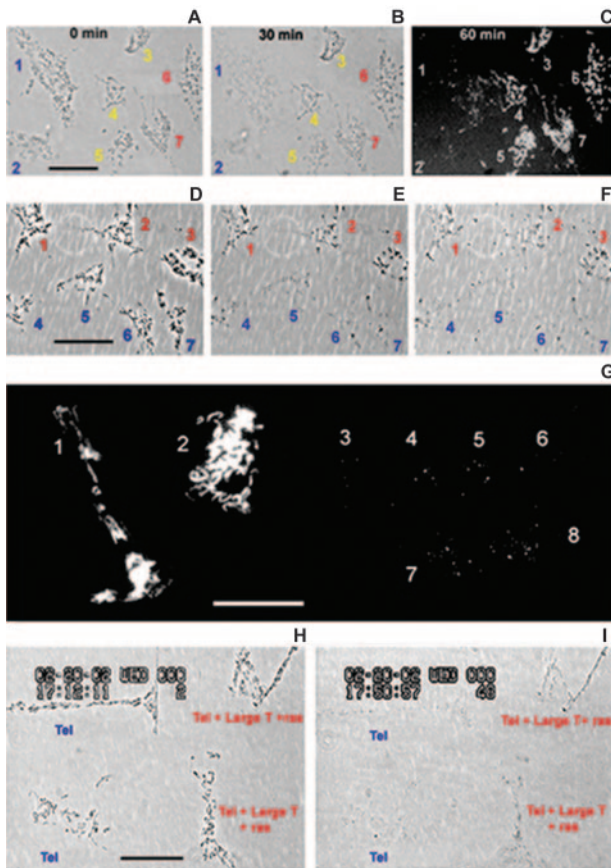


Figure 3. Differential sensitivity of isolated sets of chromosomes extracted from primary and metastatic uveal melanoma cells. **A:** Phase micrograph of chromosomes isolated from poorly invasive primary OCM1a uveal melanoma cells (labeled 1 and 2 in blue), highly invasive primary M619 uveal melanoma cells (labeled 3, 4, and 5 in yellow), and highly invasive MUM2B metastatic uveal melanoma cells (labeled 6 and 7 in red) are placed side-by-side on a glass slide. **B:** Same preparation as **A** after 30 minutes of digestion with *AluI* restriction enzyme. Chromosomes from the poorly invasive OCM1a primary uveal melanoma cell line are digesting whereas chromosomes from the more invasive cell lines remain intact. **C:** Fluorescence micrograph of same preparation as in **A** and **B** after 60 minutes of digestion with *AluI* restriction enzyme and labeling with ethidium bromide. Chromosomes extracted from the poorly invasive primary OCM1a uveal melanoma cells have almost completely digested leaving only trace amounts of ethidium-positive material whereas chromosome sets from the more invasive cell lines have largely resisted digestion. **D–F:** Differential effects of *AluI* restriction enzyme digestion on chromosomes extracted from metaphase fibroblasts, fibroblasts transfected with oncogenes, and fibrosarcoma cells. **D:** Phase micrograph of three sets of metaphase chromosomes extracted from fibrosarcoma cells (labeled 1–3 in red) and attached to charged plastic next to 4 sets of metaphase chromosomes from normal fibroblasts (labeled 4–7 in blue). **E:** Phase microscopy of the same preparation as in **A** after 30 minutes of digestion with *AluI* restriction enzyme. Normal fibroblast chromosomes have dissolved while chromosomes from the fibrosarcoma cells are relatively intact. **F:** Phase microscopy of the same preparation as in **A** and **B** after 60 minutes of digestion with *AluI* restriction enzyme. Normal fibroblast chromosomes have dissolved while chromosomes from the fibrosarcoma cells are relatively intact. **G:** Fluorescence microscopy of two sets of chromosomes extracted from fibrosarcoma cells (labeled 1, 2) and 6 sets of chromosomes extracted from normal WI-38 fibroblasts (labeled 3–8). The preparation was exposed to *AluI* restriction enzyme for 2 hours and labeled with ethidium bromide. The chromosomes from the fibrosarcoma cells (labeled 1, 2) resist digestion. Note that individual chromosomes remain intact. By contrast, the chromosomes from the normal WI-38 fibroblasts (3–8) have digested. **H:** Phase video microscopy of chromosomes extracted from fibroblasts transfected with telomerase (labeled Tel in blue) and from fibroblasts transfected with telomerase plus SV-40 Large T antigen plus *ras* (labeled Tel + Large T + ras in red). **I:** Same preparation as in **D** after 38 minutes of exposure to *AluI* restriction enzyme. Chromosomes extracted from fibroblasts transfected with only telomerase have dissolved whereas chromosomes extracted from fibroblasts transfected with telomerase plus SV-40 plus Large T have largely resisted digestion. Scale bars: **A–C** = 30 μm ; **D–F** = 40 μm ; **G** = 20 μm ; **H, I** = 20 μm .

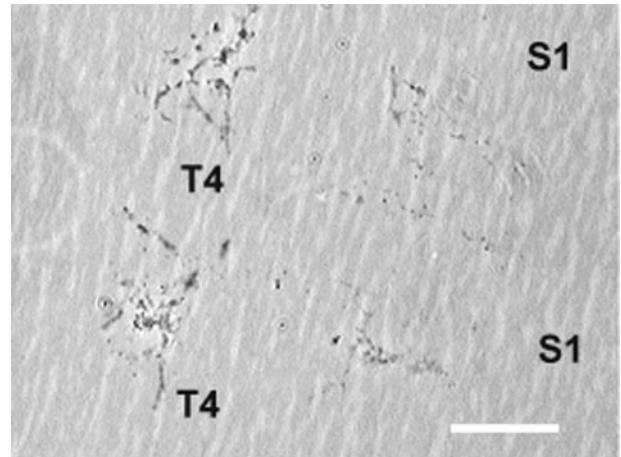


Figure 4. Phase micrograph of differential effects of *AluI* restriction enzyme digestion of chromosomes extracted from S1 breast epithelial cell and T4 cells derived from this line. After 60 minutes of exposure to *AluI* restriction enzyme, T4 chromosomes are resistant to digestion compared with S1 chromosome sets. Scale bar = 20 μm .

test the susceptibility of chromatin digestion by *AluI* in cells in mitosis, we used a method to extract complete sets of mitotic chromosomes from cells.^{7,20–29} We studied WI-38 fibroblasts, oncogene transfected cells, and cancer cell lines of varying degrees of invasive behavior (Table 1).

Metaphase chromosome sets from the poorly invasive OCM1a, highly invasive M619, and the highly invasive and metastatic MUM2B were extracted and placed next to one another so that chromosomes from OCM1a were next to chromosomes from M619 and MUM2B cells (Figure 3A). After 30 minutes of digestion by *AluI*, the OCM1a chromosomes had digested to a much greater extent than the chromosome sets extracted from highly M619 or MUM2B cells (Figure 3B). After 60 minutes of digestion by *AluI* restriction enzyme, chromosomes were stained with ethidium bromide to visualize undigested DNA (Figure 3C). Chromosomes from the most highly invasive and metastatic cells digested the least, while the poorly invasive cell chromosomes digested nearly completely. The relative sensitivities of the chromosomes extracted from metaphase cells to *AluI* restriction enzyme were therefore identical to the sensitivities observed within

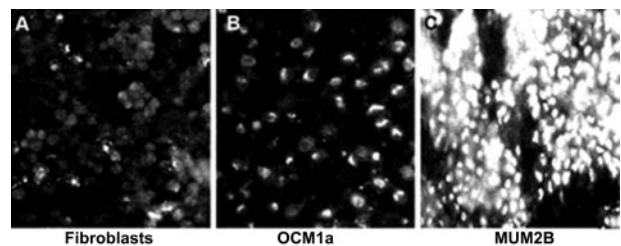


Figure 5. Fibroblasts, poorly invasive melanoma cells, and highly invasive and metastatic melanoma cell nuclei were tested for their differential sensitivity to *MspI* applied for 24 hours in a humidified 37°C chamber. Note that the highly invasive cell nuclei do not digest to the extent that the poorly invasive or normal cell nuclei digest under the same conditions and in the same time period. Scale bar = 40 μm .

the nuclei of corresponding asynchronous interphase cells (Figures 1 and 2).

Differential Sensitivity of Isolated Chromosome Sets from Normal Fibroblasts, Fibroblasts Transfected with Multiple Oncogenes, and Fibrosarcomas to AluI Restriction Enzyme

We compared the sensitivity to digestion by *AluI* for chromosomes extracted from WI-38 fibroblasts with chromosomes extracted from HT-1080 fibrosarcoma cells: the latter two resisted digestion by *AluI*, but chromosomes extracted from normal fibroblasts were digested rapidly by 30 minutes (Figure 3, D–G).

Normal fibroblasts and fibrosarcoma cells represent two extremes on a spectrum in tumor progression. To compare cells with similar genetic backgrounds, we therefore tested sensitivity to *AluI* restriction enzyme digestion on chromosomes of fibroblasts developed by Hahn et al¹⁵ that differ by single or multiple defined oncogene insertions: fibroblasts transfected only with telomerase, fibroblasts transfected with telomerase plus SV-40 Large T antigen, and fibroblasts transfected with telomerase plus Large T antigen plus *ras*. The chromosomes extracted from fibroblasts transfected with only the telomerase gene were digested by *AluI* within 38 minutes. Fibroblasts transfected with telomerase plus SV-40 Large-T digested within the same time frame. However, chromosomes from fibroblasts transfected with telomerase plus SV-40 plus *ras* (Figure 3, H and I) resisted digestion after the same time exposure to *AluI* restriction enzyme. Thus, in this fibroblast progression model, the insertion of three oncogenes induced a shift in the sensitivity of the entire genome to *AluI* digestion.

Differential Sensitivity of Isolated Chromosomes from a Breast Cancer Progression Series

To compare normal and spontaneously-induced malignant epithelial cells from the same genetic background, we used an extensively characterized series of breast and breast cancer cell lines (HMT3522). The nonmalignant cells referred to as S1 breast epithelial cells were isolated from reduction mammoplasty and a series of cell lines, including those malignant in nude mice, were derived in defined media after removal of EGFR as described previously.^{16–18} S1 cells form acini in three-dimensional cultures that reproduce the polarized structure of the breast acinus with high fidelity. The T4–2 line forms disorganized acini as do cells from primary human breast tumors³³ Chromosomes were extracted from S1 and T4–2 cells grown in monolayer cultures and exposed to *AluI* restriction enzyme. After 60 minutes, there were clear differences in digestion rates between chromosomes extracted from S1 cells and those removed from T4–2 cells (Figure 4).

Differential Digestion of Chromatin with Other Restriction Enzymes

To determine whether DNA digestion by enzymes other than *AluI* discriminated between cells of low and high invasive behavior, we tested *EcoRI*, *SauI*, *PstI*, *XhoI*, *HindIII*, and *MspI* in isolated chromosome digest assays using WI-38 fibroblasts, and melanoma cells of melanoma cells of increasing invasive potential (OCM1a, M619, and MUM2B). *MspI* (Figure 5) and DNase-I (not shown) also discriminated between cells of different invasive potential.

The Importance of Proteins in the Structural Sequestration of AluI Sites

We hypothesized that nuclear proteins are responsible for the sequestration and exposure of *AluI* sites. Therefore, we tested the sensitivity of chromosomes from non-invasive and highly invasive cells to proteases. We have previously shown that heparin and trypsin-sensitive proteins play key roles in chromosome length and compaction in normal cells.^{20,21} Exposure of chromosomes to heparin or trypsin did not discriminate between non-invasive and highly invasive cells (data not shown). However, these cells could be discriminated when chromatin from these cells was exposed to proteinase K. Under these conditions, chromosomes from highly invasive HT1080 cells dissolved within 6 minutes while chromosomes from non-invasive normal fibroblasts, normal human umbilical endothelial cells (not shown), and poorly invasive melanoma cells de-condensed but did not digest in the same time period (Figure 6, A–C). These data suggest a fundamental difference in the sensitivity of chromatin-associated proteinase K-sensitive proteins between normal and highly invasive cells.

It is not possible to study the role of proteins in the sequestration and exposure of *AluI* sensitive tissues using the nonspecific enzyme, proteinase K because this nonspecific protease would also digest the *AluI* restriction enzyme. Because there is evidence that disulfide-rich nuclear matrix-associated proteins play a key role in the sequestration and exposure of genes,³⁴ we tested the digestion of chromatin by *AluI* in the presence of β -mercaptoethanol or DTT, agents that disrupt proteins rich in disulfide bonds but do not interfere with *AluI* enzyme activity. Additionally, commercially available restriction enzyme digest kits first remove proteins from DNA with reagents such as β -mercaptoethanol or DTT.

We therefore repeated the extraction of chromosomes from fibroblasts transfected with telomerase only and from fibroblasts transfected with telomerase plus Large T and *ras*. After exposure of these chromosomes to *AluI* and either β -mercaptoethanol or DTT, the chromosomes extracted from telomerase plus Large T and *ras*-transfected fibroblasts digested more rapidly than the chromosomes from fibroblasts transfected only with telomerase (Figure 6, D–F2). Thus, digestion by disulfide disrupting agents renders highly invasive cell chromatin more sensitive to *AluI* restriction enzyme.

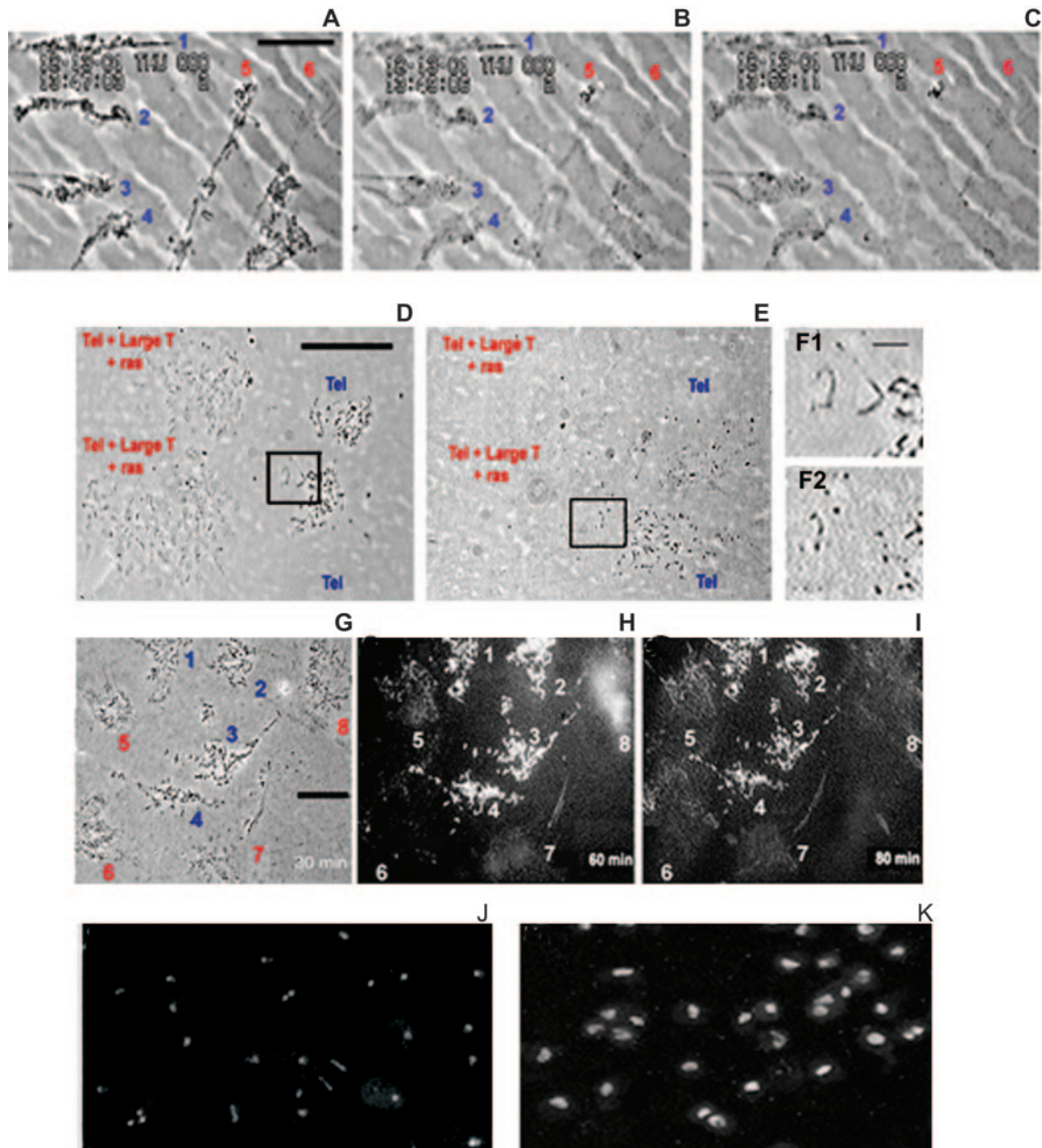


Figure 6. **A:** Phase micrograph of four sets of chromosomes extracted from WI-38 fibroblasts (labeled 1–4 in blue) and from two fibrosarcoma cells (labeled 5, 6 in red). **B:** Same preparation as in **A** after 2 minutes of exposure to proteinase K. **C:** Same preparation as in **A** and **B** after 6 minutes of exposure to proteinase K. Chromosomes from the fibrosarcoma cells have digested but chromosomes from the normal endothelial cells are relatively intact. **D–K:** Effects of disrupting protein disulfide bonds on sensitivity of chromatin to *AluI* restriction enzyme. **D:** Phase micrograph of two sets of chromosomes extracted from fibroblasts transfected with telomerase (labeled Tel in blue) are placed on the same slide as two sets of chromosomes extracted from fibroblasts transfected with telomerase plus Large T plus *ras* (labeled Tel+Large T+*ras* in red). **Box** identifies a portion of the *Tel* only transfected fibroblast chromosomes. **E:** Phase micrograph of same preparation illustrated in **A** after 80 minutes of exposure to β -mercaptoethanol and *AluI* restriction enzyme. Chromosomes from the fibroblasts transfected with telomerase plus Large T plus *ras* have digested completely but chromosomes transfected only with telomerase are only minimally digested, a sensitivity opposite that seen when *AluI* restriction enzyme is used alone (Figure 3, E and F). As the chromosomes from the *Tel* only transfected fibroblasts digest, they also shift position in the preparation. Note the relative positions of the **boxed areas** in **D** and **E**. **F1:** Phase microscopy high magnification of **boxed region** in **D** showing details of chromosomes. **F2:** phase microscopy of same chromosomes from the **boxed area** in **E** (after 80 minutes of digestion). Note the partial dissolution of the chromosomes. **G:** Phase micrograph of chromosomes extracted from poorly invasive OCM 1a primary melanoma cells (labeled 1–4 in blue) and highly invasive M619 primary melanoma cells (labeled 5–8 in red). **H:** fluorescence micrograph of same preparation as in **G** after 60 minutes of exposure to β -mercaptoethanol and *AluI* restriction enzyme, labeled with ethidium bromide. Chromosomes from the poorly invasive OCM 1a cell line (1–4) resist digestion, but chromosomes from the invasive M619 cell line are beginning to digest. Note specifically the fluorescent “cloud” of labeled DNA in chromosome set 8. **I:** Fluorescence micrograph of same preparation as in **G** and **H** after 80 minutes of exposure to β -mercaptoethanol and *AluI* restriction enzyme: Chromosomes from the poorly invasive OCM 1a cell line (1–4) still resist digestion while chromosomes from the aggressive M619 (5–8) cell line are completely digested. **J:** Fluorescence micrograph of smear preparation of invasive M619 melanoma cells after 30 minutes of digestion with *AluI* restriction enzyme in the presence of DTT. **K:** Fluorescence micrograph of smear preparation of poorly invasive OCM 1a melanoma cells after the same conditions and time point as in **J**. In the presence of DTT, the nuclei in highly invasive cells are more sensitive to digestion with *AluI* restriction enzyme, an effect opposite that seen when *AluI* is used alone (compare with Figure 1). Scale bars: **A–C** = 20 μ m; **D, E** = 40 μ m; **F1, F2** = 10 μ m; **G, H** = 20 μ m; **J, K** = 20 μ m.

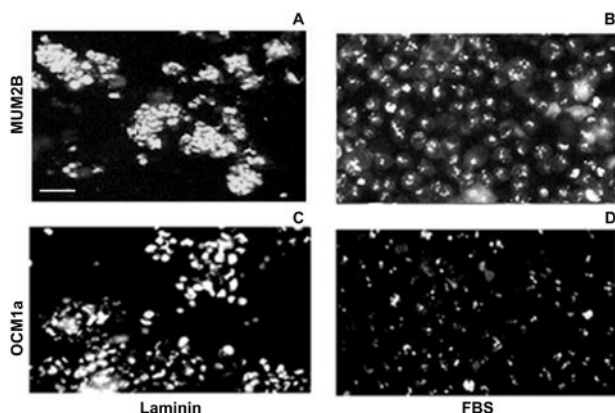


Figure 7. Laminin increases sequestration of *AluI* sensitive sites in cell smear assays. **A:** Highly invasive MUM2B cells pre-incubated with soluble laminin for 60 minutes, digested with *AluI* for 1 hour and labeled with ethidium bromide: chromatin resists digestion. **B:** MUM2B cells pre-incubated with FBS for 60 minutes, digested with *AluI* for 1 hour and labeled with ethidium bromide. Except for nucleoli, the chromatin is digested. **C:** Poorly invasive OCM1a cells pre-incubated with laminin for 60 minutes, digested with *AluI* for 1 hour and labeled with ethidium bromide. **D:** OCM1a cells pre-incubated with serum for 60 minutes, digested for 1 hour with *AluI* and labeled with ethidium bromide. Except for nucleoli, the chromatin is digested. Scale bars: **A–D** = 50 μ m.

Although chromosomes from M619 are relatively resistant to *AluI* restriction enzyme compared with chromosomes from OCM1a cells in the absence of β -mercaptoethanol or DTT (Figure 3), chromosomes from the former digested more rapidly than those from the OCM1a cells when these disulfide bond-disrupting agents were present (Figure 6, G–I). Interphase nuclei of highly invasive M619 uveal melanoma cells were also more sensitive to *AluI* restriction enzyme in the presence of DTT after 30 and 60 minutes of digestion than were nuclei of poorly invasive OCM1a cells (Figure 6, J and K). Thus, disulfide bond disrupting agents— β -mercaptoethanol or DTT—expose *AluI* binding sites in highly invasive cells.

Regulation of Susceptibility to *AluI* Restriction Enzyme by Matrix Composition in 2D Conditions by the Cell Smear Assay

There is now compelling evidence that the microenvironment including the ECM regulates gene expression.^{1,7,13,35–39} We therefore reasoned that different ECM molecules would induce changes in the sequestration and exposure of DNA.

We tested for differential digestion of chromatin in intact cells by *AluI* after cells were exposed to different soluble ECM molecules with a cell smear preparation that allowed high through-put analysis of cells dried onto glass slides. Given the results described above, the buffers used in these assays did not contain either DTT or mercaptoethanol to avoid removing any proteins within the cytoplasm or nucleus that might sequester *AluI* binding sites from enzyme digestion.

Remarkably, chromatin of both highly invasive (MUM2B) and low invasive (OCM1a) melanoma cells exposed to laminin for 30 to 60 minutes resisted digestion by *AluI* at 1 hour (Figure 7, A and C), whereas the chromatin from the same

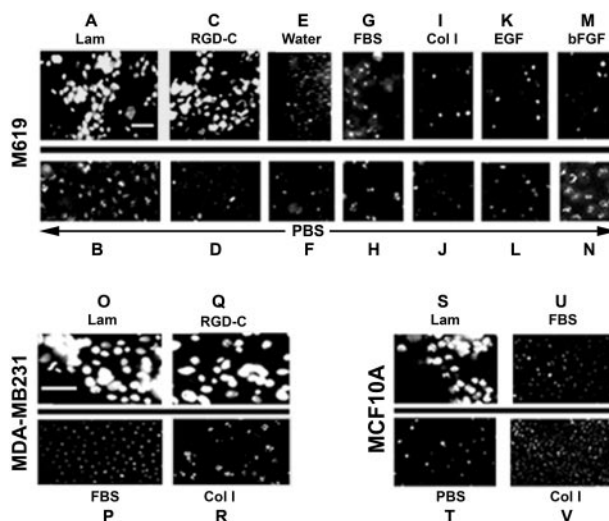


Figure 8. Exposure of cells (invasive melanoma, invasive breast cancer, and breast epithelial cells) to different matrix molecules and growth factors on the cell smear assay. **A–M:** Paired comparison of highly invasive M619 primary uveal melanoma cells treated with matrix molecules, serum, H₂O, or growth factors on upper row (**A, C, E, G, I, K, M**) and control incubations in PBS on lower row (**B, D, F, H, J, L, N**). Note resistance to digestion by *AluI* restriction enzyme on laminin (Lam) and RGD, but digestion on FBS, collagen Type I (coll), water, epithelial growth factor (EGF), and basic fibroblast growth factor (bFGF). Comparison of aggressive breast carcinoma cell line MDA-MB231 on different matrix molecules: **O**, Laminin; **P**, FBS; **Q**, RGD-C; and **R**, collagen Type I. Note the resistance to digestion on laminin and RGD, but susceptibility to digestion on FBS and collagen Type I (Col I). Comparison of MCF10A breast epithelial cells on matrix molecules: **S**, laminin; **T**, PBS control; **U**, serum; **V**, collagen Type I. Scale bars: **A–N** = 20 μ m; **O–U** = 40 μ m.

cells exposed to collagen 1 and FBS was completely digested (Figure 7, B and D) in the same time period (the same results were obtained with purified fibronectin, data not shown). Nucleoli, which are made of ribonucleoprotein and thus resistant to *AluI*, were not digested. The composition of ECM molecules affected the differential digestion of chromatin in the cell smear assay in multiple cell lines. Similar results were found when DNase was substituted for *AluI* restriction enzyme (data not shown).

The chromatin of highly invasive M619 melanoma cells when exposed for 30 minutes to soluble laminin and RGD-C peptide resisted digestion with *AluI* restriction enzyme. RGE had no effect on chromatin sequestration and exposure (data not shown). Chromatin of the same cells completely digested in the same time period after exposure to FBS, water (to ensure that exposure to ions did not affect chromatin structure), collagen I, EGF, and bFGF, a response identical to cells exposed only to PBS (Figure 8, A–N).

We then tested MCF10A nonmalignant breast cells and MDA-MB231, a metastatic breast cancer cell line with the cell smear assay. Similar to melanoma cells, 30 minutes of incubation with laminin or RGD-C before exposure to the *AluI* restriction enzyme rendered the chromatin of both MB321 and MCF10A cells resistant to digestion, whereas incubation of the same cells with collagen I and RGD-C rendered the chromatin susceptible to digestion (Figure 8, O–V).

Permeabilized Cell Assay with Polymerized ECM Molecules

In vivo, cells are exposed to both soluble and polymerized ECM molecules. Therefore, to test the exposure and sequestration of *AluI* sensitive sites on cells exposed to polymerized ECM molecules, we developed a permeabilized cell model system in which cells were situated on heat-denatured and localized thin ECM films of varying compositions. These heat-denatured ECM films resisted penetration and rapid digestion by highly invasive, matrix invading tumor cells, allowing the cells to rest on the ECM surface where they were accessible to various reagents after permeabilization with Triton X-100.

For this assay, we constructed a grid of different ECM molecules on glass slides: laminin or Matrigel deposited as squares, and dried FBS or Type I collagen in the rows between squares. Cells were then distributed evenly over the slide before exposure to the *AluI* restriction enzyme (see diagram in Figure 9, A and B).

Polymerized laminin or Matrigel, but not FBS or Type I collagen, greatly increased the resistance of chromatin digestion by *AluI* restriction enzyme in MDA-MB231 cells (Figure 9, C and D). To ensure that cells had complete access to the enzymes and dyes added after the cells were permeabilized with Triton X-100, we dislodged the cells using a microneedle, and verified that they were indeed on the heat-denatured polymerized ECM surfaces instead of being buried within them (Figure 9E).

Identical ECM composition-dependent sensitivities to *AluI* digestion were observed with MCF10A breast epithelial cells (Figure 9, F–I). Small groups of these cells that were situated along a laminin-dried FBS interface provided the most compelling evidence that ECM contact itself, and not access to restriction enzyme or dye, influenced the digestion of chromatin by *AluI*. When MCF10A cells made contact with even the edge of a laminin square, DNA resisted digestion by *AluI* restriction enzyme for up to 60 minutes of exposure to the enzyme (Figure 9F), but cells not in contact with laminin digested in this time frame (Figure 9G). When clusters of MCF10A cells straddled the interface between laminin and dried FBS, the nuclei of cells in contact with laminin resisted digestion by *AluI* restriction enzyme within 60 minutes, but those cells situated entirely on dried FBS digested completely by 60 minutes (Figure 9, H and I). Matrix-dependent alterations in chromatin sensitivity to *AluI* restriction enzyme were also observed in melanoma cell lines (Figure 9, J–L).

In all permeabilized cell assays, nucleoli, which are made of ribonucleoprotein and thus resistant to *AluI*, were used as an internal control and were not digested (Figure 9L). Brownian motion by the ethidium-positive nucleolar residues (Figure 9, M and N) further confirmed digestion of DNA, suggesting disruption of the constraints that DNA imposes on the nucleoli.

To test if matrix composition on *AluI* site influences sequestration and exposure in mitotic cells, we extracted chromosomes^{20,21} from aggressive M619 primary uveal

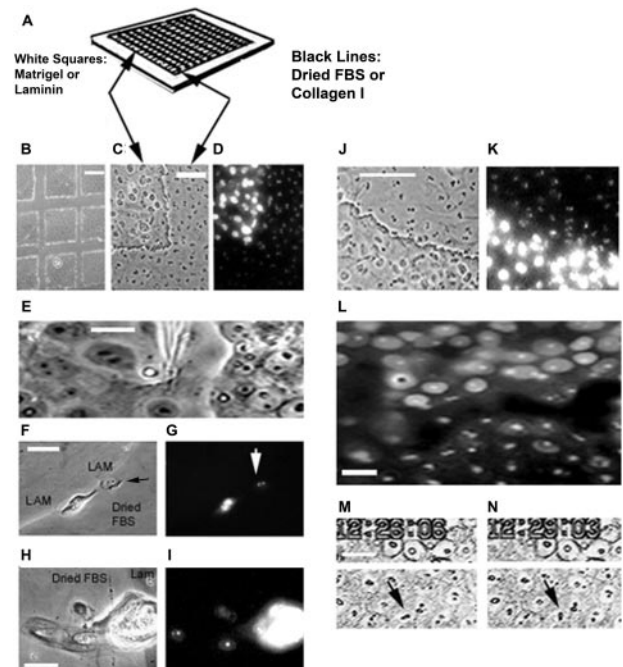


Figure 9. Effect of polymerized ECM molecules on sensitivity of *AluI* sites to *AluI* restriction enzyme with the permeabilized cell assay. **A:** Representation of a matrix chip on which squares of laminin or Matrigel are separated from each other by rows of pre-adsorbed FBS or Type I collagen (**thick black lines**). **B:** Phase-contrast micrograph of the matrix grid patterns. The squares in this grid are composed of laminin, and the intersquare rows are composed of FBS. **C:** Phase-contrast micrograph of highly invasive MDA-MB231 breast carcinoma cells seeded on laminin squares and FBS intersquare rows. **D:** Same field as **C** showing MDA-MB231 cells labeled with ethidium bromide after 3 hours of digestion with *AluI* restriction enzyme. The cells on the laminin square with brightly fluorescent nuclei have resisted digestion by *AluI* restriction enzyme. The small fluorescent dots in the intersquare rows are nucleoli. **E:** MDA-MB231 cells are easily dislodged from the grid surface with a microneedle, demonstrating that cells are on the surface of the laminin square and are therefore completely accessible to the *AluI* restriction enzyme. **F:** Three MCF10A breast epithelial cells at the interface between a laminin square and the FBS intersquare row. Two cells touch the edge of the laminin square, but one cell (**black arrow**) is situated within the FBS intersquare row. **G:** Corresponding fluorescence micrograph of the same field as in **F**. Only the two cells in direct contact with edge of the laminin square are not digested by *AluI* restriction enzyme. The nucleus of the one cell not in contact with the laminin square is digested (**white arrow**), and only its nucleolus (**small dots**) labels with ethidium bromide. **H:** Phase contrast of a cluster of MCF10A cells treated with *AluI* restriction enzyme for 60 minutes; the cluster of cells straddles the interface (**arrows**) between dried FBS (**left**) and laminin (**lam; right**). The cells on dried FBS (**arrowheads**) have flattened and spread, while the cells on laminin are round. **I:** Corresponding fluorescence micrograph of the same field as in **H**. The nuclei of three flattened cells on FBS (**arrowheads**) digest with *AluI* restriction enzyme, leaving only nucleoli undigested, while the cells on laminin resisted to *AluI* digestion. **J:** Phase contrast of OCM1a cells treated with *AluI* restriction enzyme for 60 minutes; the cells are distributed both on the laminin square and on the FBS intersquare row. **K:** Corresponding fluorescence micrograph of the same field as in **J**. DNA of OCM1a melanoma cells on laminin resisted digestion by the *AluI* restriction enzyme, but DNA in the same cells situated on FBS digested completely; only the nucleoli are fluorescent. **L:** Fluorescence micrograph of metastatic MUM2B melanoma cells on laminin (**top**) collagen (**bottom**) grid. DNA in the melanoma cells on laminin resisted *AluI* enzyme digestion, while DNA from the same cells on collagen digested completely. **M:** Phase micrograph of same preparation illustrated in **L** showing MUM2B melanoma cells on laminin (**top**) and collagen (**bottom**) during digestion with *AluI* restriction enzyme. Note the horizontal orientation of the nucleolus marked by the arrow. **N:** Same field as in **M** but 3 minutes later. Note the vertical orientation of the nucleolus marked by the arrow indicating Brownian motion. Note that the nuclear membrane of the melanoma cells on laminin is sharply demarcated but the nuclei of the same cells on collagen appear to have no border; the chromatin of cells on collagen has been completely digested.

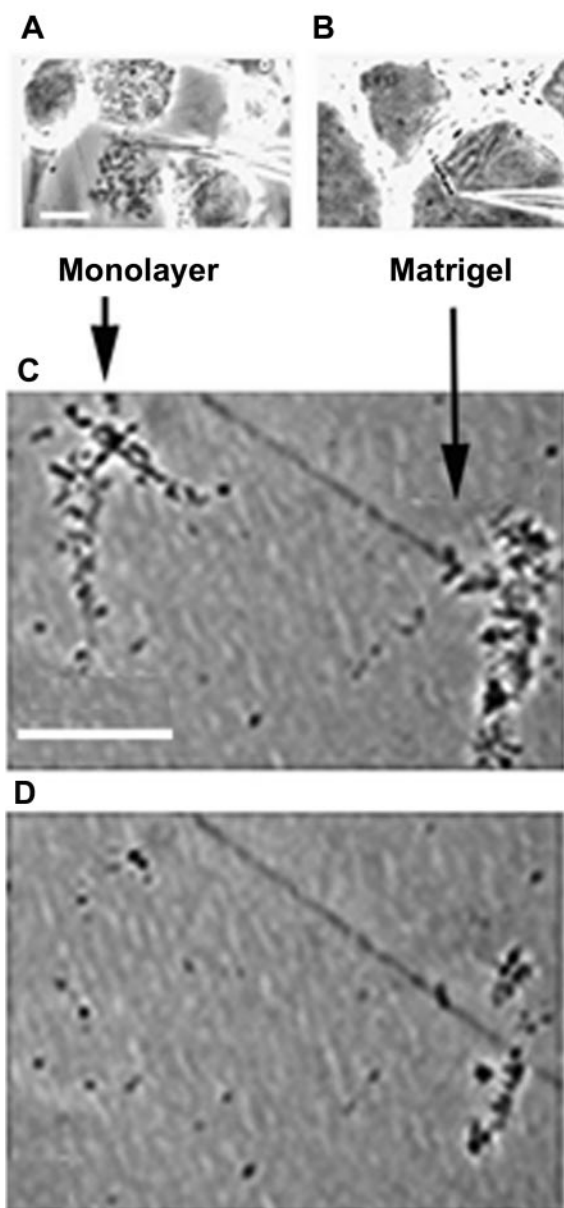


Figure 10. Comparing the sensitivity to *AluI* restriction enzyme of chromosomes from aggressive M619 uveal melanoma cells grown in monolayer conditions or on Matrigel. **A:** Chromosomes are extracted from an M619 cell grown in monolayer conditions. **B:** Chromosomes are extracted from an M619 uveal melanoma cell in a culture forming networks of cords on the surface of Matrigel. **C:** The chromosomes extracted in **A** and **B** are placed side by side on plastic. **D:** The chromosomes from M619 cells grown as monolayers after 60 minutes of exposure to *AluI* restriction enzyme, but the chromosomes from M619 cells grown on the surface of Matrigel resist digestion in this time period. Scale bars: **A, B** = 10 μm ; **C, D** = 5 μm .

melanoma cells grown as a monolayer and placed them side by side with chromosomes extracted from the same cells grown on the surface of polymerized Matrigel. These chromosome sets were then exposed to the *AluI* restriction enzyme for 60 minutes, labeled with ethidium bromide, and photographed. Chromosomes from the aggressive melanoma cells grown as monolayers digested more rapidly than those extracted from the same cells grown on polymerized Matrigel (Figure 10).

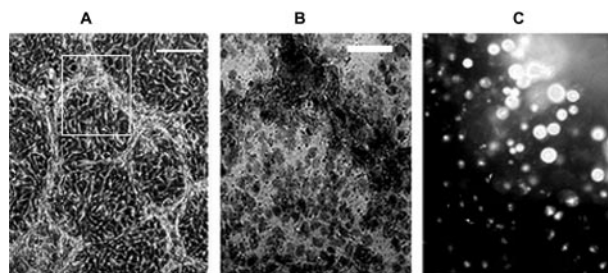


Figure 11. Resistance of *AluI* sites in lysed aggressive uveal melanoma cells to *AluI* digestion in three-dimensional cultures. **A:** Phase micrograph of highly invasive melanoma cells exhibiting looping vasculogenic mimicry patterns. **B:** Bright field photomicrograph taken of the same preparation as in **A** after 3 minutes of Triton X-100 treatment, and 10 minutes of Trypan blue exposure; all cells exhibit uptake of this dye. Note that the areas where the matrix patterns have formed vasculogenic mimicry patterns (**box** in **A**), Trypan blue dye appears to be most concentrated. **C:** Fluorescent micrograph of ethidium bromide uptake in the same area as shown in **B** after a 2 hour digestion with *AluI*: the tumor cell nuclei resist digestion with *AluI* restriction enzyme. Scale bars: **A** = 50 μm ; **B, C** = 20 μm .

Regulation of Susceptibility to *AluI* Restriction Enzyme Digestion by Thickness of ECM Gels

The cell smear and polymerized ECM assays provide a limited approximation of the behavior of tumor cells *in vivo* where these cells are often embedded within the complex ECM of different compositions. Additionally, it is well known that there are large shifts in gene expression between 2D and 3D culture conditions,⁴⁰⁻⁴² suggesting a possible rearrangement of chromatin structure that might be detectable to differential chromatin digestion to *AluI*. To determine whether the susceptibility and resistance of chromatin to *AluI* digestion is influenced by the thickness of ECM, we prepared cultures using 0.3 mm thick preparations of native Matrigel.

Highly invasive M619 and MUM2B uveal melanoma cells formed looping patterns in thick matrix culture conditions as described previously.^{13,14} The 3D cultures (Figure 11A) were treated with Triton X-100 and were exposed to Trypan for 10 minutes to ensure that cells buried within matrix were permeabilized (Figure 11B). These cultures were exposed to the *AluI* restriction enzyme for 1 hour, labeled with ethidium bromide, and immediately photographed. The chromatin of pattern-forming cells in thick ECM resisted digestion with *AluI* restriction enzyme as indicated by bright nuclear staining with ethidium bromide (Figure 11C). A comparison of gene expression by Affymetrix microarray between MUM619 and MUM2B cells in corresponding flat and 3D culture conditions produced a list of 990 transcripts expressed differentially (651 genes differentially expressed in cells grown in flat culture conditions, and 339 genes differentially expressed by cells in 3D cultures, not shown). A detailed listing and analysis of this differential gene expression will be reported separately.

The Role of the Cytoskeleton in Resistance and Susceptibility to *AluI* Digestion

Both biochemical and mechanical signaling pathways have been implicated in the remodeling of chroma-

tin.^{2,20,21,37,43–45} We have previously shown that specific cytoskeletal components alter nuclear shape, molecular alignment of structures in the nucleus, and nucleolar positioning in normal endothelial cells, fibroblasts, and breast cells.^{2,43} Therefore, we investigated the function of the cytoskeleton in modulating the sensitivity of chromatin digestion by *AluI* to matrix composition.

We studied normal fibroblasts instead of tumor cells because the cytoskeleton of tumor cells tends to be abnormal.^{46–48} Cell smear assays were used to assess the response of fibroblast nuclei to *AluI* digestion under different conditions because fibroblasts tend to spread when serum is present thus facilitating observation of specific cytoskeletal components. Most fibroblasts in the assay flattened and spread in 30 minutes, but rare cells remained round (Figure 12A). After 30 minutes of exposure to *AluI* restriction enzyme, the chromatin of the flat spreading fibroblasts was digested as detected by ethidium bromide labeling; the chromatin of the fibroblasts that did not spread resisted digestion (Figure 12B). However, after exposure to colchicine for 30 minutes, most of fibroblasts failed to flatten and spread (Figure 12, C and D), and the chromatin of these cells resisted 30 minutes of digestion by *AluI* restriction enzyme. The chromatin of colchicine-treated fibroblasts that did not flatten and spread remained resistant up to 3 hours of digestion with *AluI* (not shown). These findings were consistent with the role of the cytoskeleton in regulating cell and nuclear shape^{2,49} and suggested that microtubules participated in regulating the sequestration or exposure of *AluI* binding sites.

To determine whether other components of the cytoskeleton—actin filaments or intermediate filaments—were involved, each component was disrupted in turn. Monolayer cultures of normal fibroblasts were either left untreated or treated for 30 minutes with cytochalasin B to disrupt actin filaments or acrylamide to disrupt intermediate filaments.⁵⁰ Cells were prepared for a cell smear assay as described above and each preparation was exposed to *AluI* for 30 minutes before labeling with ethidium bromide. Compared with normal fibroblasts (Figure 12E), the chromatin of fibroblasts treated with cytochalasin B digested completely, whereas the chromatin of fibroblasts treated with either colchicine or acrylamide resisted digestion. Thus, disruption of the actin cytoskeleton exposes chromatin while disruption of intermediate filaments and microtubules sequesters the chromatin.

To investigate structural continuity between the ECM, the cytoskeleton, and the nuclear envelope, we attached 5- μm beads coated with RGD-C peptide to the surfaces of the fibroblasts in serum-free media to form a focal adhesion.² Within 5 to 30 minutes of bead binding we documented an ultrastructural contiguity between the RGD-bead-induced focal adhesion complexes, microtubules, and the nuclear surface and its pore complexes (Figure 13A). Identical ultrastructural connections between the cell cortex, microtubules and intermediate filaments, and the nuclear surface were identified in normal fibroblasts exposed to laminin (Figure 13B). Whereas microtubules appeared to be predominantly associated with nuclear pore complexes, intermediate filaments in

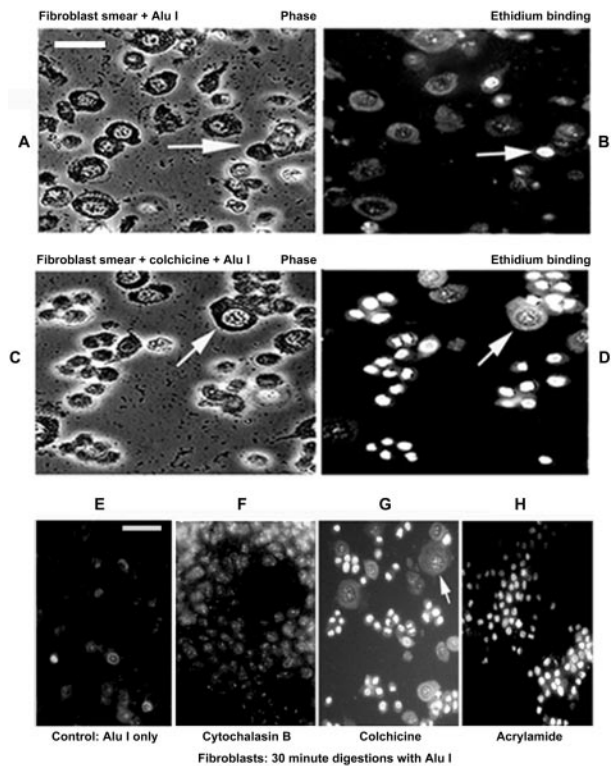


Figure 12. Pharmacological inhibition of *AluI* sensitivity. **A:** Phase contrast micrograph of WI-38 fibroblasts dried and digested for 30 minutes with *AluI* restriction enzyme. Most cells are well spread. Note the two non-spreading cells (arrow) **B:** Fluorescence micrograph of the same cells shown in **A** after labeling with ethidium bromide. Note that all of the spread cells exhibit nuclei that completely digest after 30 minutes of treatment with *AluI*, but the nuclei in the cells that do not spread (arrow) resist digestion. **C:** Phase micrograph of fibroblasts treated with colchicine for 30 minutes, incubated with *AluI* for 30 minutes, and labeled with ethidium bromide. Note that most of the colchicine-treated cells have not spread (arrow points to a cell that has spread) **D:** Fluorescence micrograph of the same cells shown in **C** after labeling with ethidium bromide. Note that all of the nuclei of non-spread cells resist digestion, while the nuclei of the cells that spread (arrow) are completely digested. **E:** Fluorescence micrograph of WI-38 fibroblasts after drying, digestion with *AluI* restriction enzyme for 30 minutes, and labeling with ethidium bromide. The nuclei of all cells except for one (bright spot at left) are digested. **F:** Fluorescence micrograph of WI-38 fibroblasts after 30 minutes of treatment with cytochalasin B followed by 30 minutes of incubation with *AluI* restriction enzyme. Note that all of the nuclei are completely digested; only nucleoli are labeled. **G:** Fluorescence micrograph of WI-38 fibroblasts after 30 minutes of treatment with colchicine, followed by a 30-minute incubation with *AluI* and labeling with ethidium bromide. Note that every nucleus in each spread cell is completely digested while the nuclei in cells not spread completely resist digestion with *AluI* restriction enzyme. **H:** Fluorescence micrograph of WI-38 fibroblasts after 30 minutes of treatment with acrylamide, followed by 30 minutes incubation with *AluI* restriction enzyme and labeling with ethidium bromide. All cells failed to spread during the 30-minute drying period due to the presence of acrylamide, and the nucleus of each cell resists digestion with *AluI*. Scale bars: **A–D** = 20 μm ; **E–H** = 20 μm .

colchicine-treated fibroblasts appeared to associate with the nuclear envelope (Figure 13C).

Discussion

There are significant precedents for the use of nucleases to study chromatin organization in normal cells and cancer cells. Weintraub and Groudine⁵¹ demonstrated that active genes in normal cells are preferentially digested by DNase-I. Puck et al⁵² demonstrated that the reversion of malignant cells to a normal phenotype by exposure to

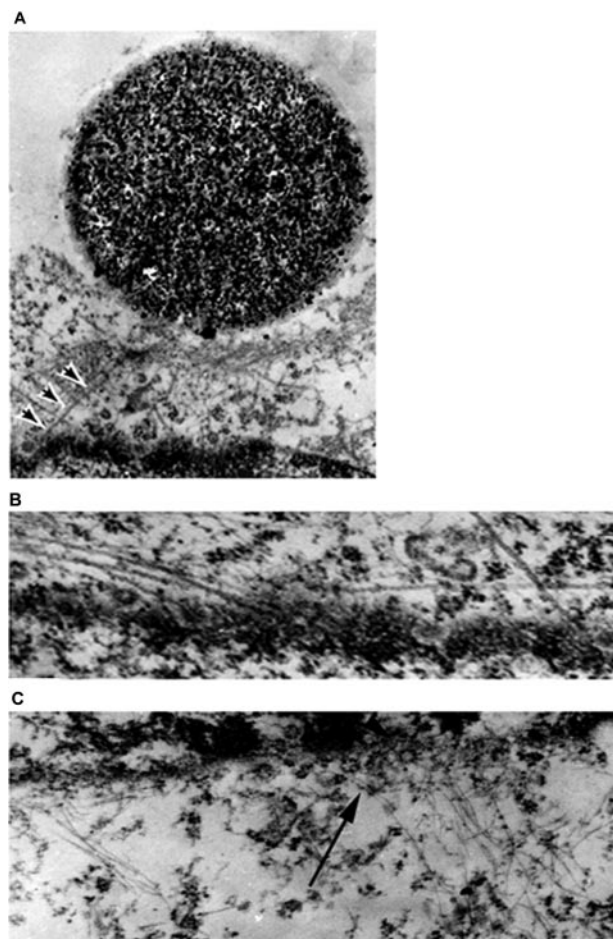


Figure 13. Ultrastructural characterization of structural contiguity between an RGD receptor site, the cytoskeleton, and the nucleus. **A:** RGD-C-coated 5 μm bead attached to a fibroblast for 5 minutes. Note microtubules (**arrows**) communicating between the focal adhesion forming around the bead and the nuclear pores. **B:** Normal fibroblast at higher magnification in the vicinity of nuclear pores reveals bowed microtubules apparently flanking the sides of nuclear pores (**arrow** identified a pore). The microtubular bowing may be the result of compression on these structures. **C:** Fibroblast photographed in the presence of colchicine to eliminate microtubules, showing intermediate filaments (**arrow**) associated with the nuclear envelope and possibly nuclear pores. Scale bars: **A** = 1 μm ; **B**, **C** = 100 nm.

cyclic AMP derivatives was accompanied by differences in the sequestration and exposure of DNase I-sensitive sites. We tested the hypothesis that chromatin organization as measured by *AluI* restriction enzyme differs in benign and malignant cells and is regulated by the ECM and the cytoarchitecture.

We demonstrated that there are differences in chromatin digestion by *AluI* in between normal melanocytes and melanoma cells, normal breast epithelium and breast carcinoma cells, and fibroblasts and fibrosarcomas. These differences are independent of the cell cycle as demonstrated by the identical differential digestion of chromosomes extracted from cells and nuclei from intact cells of varying malignant behavior. The insertion of oncogenes into fibroblast DNA¹⁵ resulted in a decrease in the sensitivity of the entire genome to *AluI* restriction enzyme. Differential sensitivity of chromatin to *AluI* restriction enzyme is also independent of ploidy.

In performing *AluI* restriction enzyme chromatin assays, it is important to ensure that the reagents used are free of DTT or β -mercaptoethanol. Both of these reagents disrupt proteins rich in disulfide bonds that play key roles in maintaining the sequestration of *AluI* sensitive sites. Indeed, alterations in higher order chromatin structure may be mediated by the proteinaceous disulfide-rich nuclear matrix proteins.^{3,53-57} We further discovered that chromatin isolated from poorly invasive and highly invasive cells are digested at different rates by the non-specific protease, proteinase K, thus suggesting that proteins, in addition to DNA,^{20,29} play a critical role in maintaining the structural organization of the genome, especially in highly invasive cancer cells.

The differential response of chromatin to *AluI* restriction enzyme among normal cells and cancer cells of varying levels of invasive behavior suggests that it may be possible to extend this observation to a clinical setting. The cell smear assay used to discriminate between cell lines of different malignant behavior is similar to a diagnostic cytological smear or touch preparation. Preliminary studies indicate that differential chromatin digestion can be used to discriminate between cancer cells and the normal adjacent tissue in human diagnostic specimens (Figure 1), and additional studies are planned to test the sensitivity and specificity of this technique. Additionally, the quantitative discrimination between melanoma cells of different invasive potential in suspension by flow cytometry (Figure 2) suggests another possible and easily standardized translational approach to classification of patient diagnostic samples.

We investigated the possible effects of ECM on the sequestration and exposure of *AluI* sensitive sites in non-malignant, benign, and malignant cells because there is considerable evidence that the ECM profoundly influences gene expression.^{37,45,58} The first evidence for the existence of an ECM-responsive element in the promoter of tissue-specific genes was produced more than a decade ago in one of our laboratories.⁵⁹⁻⁶² In particular, we showed that activation of the ECM-response element could occur only in the context of chromatin and not by transient transfection, and that ECM (laminin, in particular) induced changes in histone acetylation of the ECM-response element (as reported previously⁶² and in unpublished data).

In this study, we demonstrate for the first time a complete and global reorganization of chromatin in response to ECM composition and thickness. This reorganization was accompanied by differential changes in the expression of 990 genes (data to be reported separately) and is consistent with reports of significant ECM-induced shifts in gene expression.^{37,45,58-62} In all degrees of malignancy, *AluI*-sensitive sites became profoundly sequestered when cells were incubated with laminin, Matrigel, or a circular RGD peptide (RGD-C), but became exposed when cells were placed on serum or collagen I.

The mechanism by which the ECM induces changes in *AluI* binding site sequestration and exposure appears to depend on the cytoskeleton. Disruption of selective elements of the cytoskeleton—microtubules, actin, and intermediate filaments—resulted in profound changes in

cell shape as expected, but also in sequestration or exposure of *Alu* binding sites. We previously demonstrated that mechanical pulling on integrin receptors affected nuclear architecture, and that this effect was mediated differentially by microtubules, intermediate filaments, and actin: birefringence microscopy of living cells demonstrated that molecular alignment of structures within the nucleus changed in 1 second when integrin receptors, but not acetylated LDL receptors had a force applied to them.² Over longer time frames than 1 second, it has been widely established that different types and quantities of ECM molecules can induce cells and nuclei to rapidly change their shape and differentiation programs simultaneously, and that these changes persist for months in culture.^{4–6,9–11,63–65}

Pharmacological disruption of actin by cytochalasin B enhanced digestion of chromatin by the *AluI* restriction enzyme. Enhanced sensitivity of *AluI*-sensitive sites in the presence of cytochalasin may result from disruption of cytoplasmic organization, or because monomeric and polymeric actins are present in the nucleus, they may play a role in regulating chromatin organization and hence transcription.⁶⁶

The function of intermediate filaments in mediating the susceptibility and resistance of chromatin to *AluI* restriction enzyme is less certain. In this study, we used acrylamide to solubilize the intermediate filament lattice completely in the cytoplasm while leaving the actin and microtubule lattices unaffected;² it has been established in living cells that vimentin filaments help stabilize nuclear structure by acting as guy-wires.^{50,66–70} By solubilizing intermediate filaments, we may have also affected intermediate filaments within the nucleus.^{67,70}

The disruption of microtubules resulted in a profound change in cell shape and the sequestration of *Alu* binding sites. The chromatin of colchicine-treated cells was resistant to digestion by the *AluI* restriction enzyme (Figure 12, G and H). The physical coupling of microtubules with nuclear pores observed at the ultrastructural level in this study (Figure 13) is consistent with biophysical findings that associate this component of the cytoskeleton with stabilizing nuclear structure. Microtubules together with cell-ECM adhesions are postulated to hold the cell in place against the inward pull of actin.⁹ Because of the strong effect of microtubule and intermediate filament inhibitors on *Alu* sequestration or exposure, and because of the association of microtubules with nuclear pores, the results here suggest that the mechanism for *Alu* exposure may involve the positioning of microtubules at the nuclear pores. Microtubules may act as compression-resistant structures,⁹ distorting the surface of the nucleus thereby allowing the nuclear pores to widen and permit the influx of chromatin remodeling enzymes and ions.^{2,21} After pharmacological disruption of microtubules with colchicine, the nucleus does not distort to the extent observed when microtubules are present.² It is therefore possible that nuclear pores would assume a more closed status, thereby affecting the exposure of *Alu* binding sites.

ECM control of chromatin organization as detected by sequestration and exposure of *AluI* binding sites appears to be a mechanical signaling pathway mediated by the

cell cortex, the cytoskeleton, and higher order chromatin structure consistent with a tensegrity model of the cell. This mechano-genomic response to ECM may operate alongside ECM-mediated chemical signaling pathways.^{1,45} Therefore, changes in the sequestration and exposure of *AluI* binding sites that are mediated by the cytoskeleton and induced by the ECM suggest novel mechanisms of drug resistance for those agents that bind to or disrupt DNA.

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