

Nuclear Envelope Composition Determines the Ability of Neutrophil-type Cells to Passage through Micron-scale Constrictions^{*[5]}

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Background: The unusual nuclear shape of neutrophils has been speculated to facilitate their passage through confined spaces.

Results: Levels of nuclear protein lamin A modulate cell passage through micron-scale pores.

Conclusion: The unique protein composition of neutrophil nuclei facilitates their deformation; lobulated nuclear shape is not essential.

Significance: Altered nuclear envelope composition, as reported in cancer cells, could impact cell passage through physiological gaps.

Neutrophils are characterized by their distinct nuclear shape, which is thought to facilitate the transit of these cells through pore spaces less than one-fifth of their diameter. We used human promyelocytic leukemia (HL-60) cells as a model system to investigate the effect of nuclear shape in whole cell deformability. We probed neutrophil-differentiated HL-60 cells lacking expression of lamin B receptor, which fail to develop lobulated nuclei during granulopoiesis and present an *in vitro* model for Pelger-Huët anomaly; despite the circular morphology of their nuclei, the cells passed through micron-scale constrictions on similar timescales as scrambled controls. We then investigated the unique nuclear envelope composition of neutrophil-differentiated HL-60 cells, which may also impact their deformability; although lamin A is typically down-regulated during granulopoiesis, we genetically modified HL-60 cells to generate a subpopulation of cells with well defined levels of ectopic lamin A. The lamin A-overexpressing neutrophil-type cells showed similar functional characteristics as the mock controls, but they

had an impaired ability to pass through micron-scale constrictions. Our results suggest that levels of lamin A have a marked effect on the ability of neutrophils to passage through micron-scale constrictions, whereas the unusual multilobed shape of the neutrophil nucleus is less essential.

The passage of cells through narrow spaces is critical in physiological and disease processes from immune response to metastasis. For example, neutrophils are required to rapidly traverse constrictions that are much smaller than their own diameter of 7–8 μm : during perfusion through capillaries with diameters as small as 2 μm or during migration through transendothelial and interstitial spaces ranging from 0.1 to 10 μm (1). The ability of neutrophils to transit through narrow constrictions is essential; increased cell stiffness results in retention of neutrophils in arteries and capillaries (2), as well as accumulation in postcapillary venules leading to inflammation in the vascular bed (3).

Although the mechanical properties of neutrophils can be regulated by cytoskeletal filaments such as actin (4–6) and microtubules (7), the hallmark multilobed nuclear morphology has long been thought to facilitate the deformation of neutrophils through narrow spaces (8, 9); a round-shaped nucleus could sterically hinder the deformation of a cell through a narrow pore, whereas the multilobed neutrophil nucleus could aid cell passage as individual lobes could be sequentially “threaded” through constrictions. Indeed, cells with lobulated nuclear shape show less retention in 8- μm porous membranes as compared with their progenitors with round nuclei (10). However, it remains unclear to what extent this hyperlobulated nuclear shape is required for neutrophils to deform through narrow gaps; tightly regulated modifications in nuclear envelope protein composition also occur during granulopoiesis. Specifically,

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during the process of granulopoiesis, as recapitulated *in vitro* using human promyelocytic leukemia (HL-60)⁴ cells, major alterations occur in the expression levels of two key nuclear envelope proteins; the integral nuclear membrane protein, lamin B receptor (LBR), is strongly up-regulated, whereas there is a concurrent decrease in levels of lamin A, a key structural protein that forms a network underlying the inner nuclear membrane and imparts the nucleus with mechanical stability (11–13). Thus, although the unique shape of the neutrophil nucleus could facilitate the passage of these cells through narrow constrictions, we hypothesized that reduced levels of lamin A could enhance nuclear deformability and thereby facilitate the passage of cells through micron-scale constrictions.

To dissect the role of nuclear shape and nuclear envelope composition in the passage of cells through constrictions that mimic physiological gaps, we used all-*trans*-retinoic acid (ATRA)-stimulated HL-60 cells to recapitulate granulopoiesis; this *in vitro* system is widely used for structural and functional assays of white blood cells (14–16). We probed the ability of cells to transit through micron-scale constrictions and investigated the effects of both altered nuclear shape and altered lamin A expression levels. Our results show that levels of lamin A have a predominant effect on the ability of cells to passage through narrow constrictions, whereas the altered shape of the neutrophil nucleus is not essential for rapid passage through micron-scale pores.

EXPERIMENTAL PROCEDURES

Cell Culture—HL-60/S4 cells were maintained in RPMI 1640 medium with L-glutamine (Invitrogen), 10% fetal bovine serum (FBS), and 1% penicillin:streptomycin (Gemini Bio-Products, West Sacramento, CA). We generated scrambled control cells to compare with HL-60/S4 cells with stable shRNA-mediated knockdown of LBR (LBR KD cells) (17). To induce differentiation into neutrophil-type cells, we added ATRA at a final concentration of 5 μM to 1×10^5 cells/ml; ethanol was used as vehicle control. We probed nuclear shape and nuclear envelope composition at days 0, 3, and 5 after ATRA treatment; we performed functional assays of neutrophil-type cells at 4 days after ATRA treatment, when cells display characteristics of neutrophils (11, 18).

Microfluidic Deformation—Soft lithography was used to fabricate microfluidic channels in polydimethylsiloxane (Sylgard 184 silicone elastomer, Dow Corning) (19). Devices were bonded to #1.5-thickness coverglasses. We drove the flow of cells by applying 28 kilopascals (4 p.s.i.) of pressure to a tube of 2.5×10^6 cells/ml with 0.1% F127 (Pluoronic F-127, Invitrogen) to minimize surface adhesion (20). Images were acquired at 300 frames/s with a high speed camera (Miro ex4, Vision Research, Wayne, NJ) mounted on an inverted light microscope (Zeiss Observer) with 10 \times /0.25 Ph1 objective (A-Plan, Zeiss). The resulting image sequences were analyzed using a custom-written program (MATLAB) to extract the time for cell passage through the first constriction.

⁴ The abbreviations used are: HL-60, human promyelocytic leukemia; LamA OE, lamin A-overexpressing; LBR, lamin B receptor; LBR KD, LBR knockdown; ATRA, all-*trans*-retinoic acid; Bis-Tris, 2-(bis(2-hydroxyethyl)amino)-2-(hydroxymethyl)propane-1,3-diol.

Retroviral Transduction—We generated the stably modified lamin A-overexpressing (LamA OE) cells from the parent HL-60/S4 cell line by retroviral transduction (21–23) with the bicistronic vector (pRetroX-IRES-ZsGreen1, Clontech) for lamin A and the fluorophore reporter *Zoanthus* green fluorescent protein (ZsGreen1) with the 5' Moloney murine leukemia virus LTR as the promoter. Cloning of the wild-type prelamins A into the bicistronic retroviral vector was performed as follows: the insert was generated by cutting pSVK3-prelamins A (24) (kind gift from Howard J. Worman) with SmaI and Sall; this was ligated to the vector obtained from cutting pEGFP-C1 (Clontech) with Ecl136II and Sall resulting in a shuttle vector, which was subsequently digested with XmaI, blunted with Klenow, and then cut with BglII. The insert from the latter digestion was then ligated to the vector generated from cutting the pRetroX-IRES-ZsGreen1 with BamHI and blunted with Klenow followed by BglII digestion. Transfection of the resultant pRetro-prelamins A-IRES-ZsGreen 1 expression vector into the 293GPG retroviral packaging cell line (kind gift from Richard C. Mulligan) was performed using Lipofectamine Plus reagent (Invitrogen) based on the manufacturer's specifications and previous protocols with minor modifications (21–23). A ZsGreen1 retrovector without lamin A insert was used to generate the mock control cells. Viral supernatant was collected daily for 6 consecutive days, filtered through 0.45- μm pores, and stored at -20°C . Later, the viral supernatants collected per batch were thawed and pooled, and viral titer was determined by viral infection of mouse embryo fibroblasts. Two rounds of viral transduction of HL-60/S4 cells were then performed using unconcentrated viral supernatant supplemented with 6–8 $\mu\text{g/ml}$ Polybrene (Sigma-Aldrich) at a multiplicity of infection of 25–50. Gene transfer efficiency was assayed 5 days after retroviral infection by flow cytometry probing ZsGreen1 levels; because ZsGreen1 and lamin A are derived from the same bicistronic mRNA transcript, we sorted individual cells based on ZsGreen1 levels into a subpopulation of cells with well defined, elevated expression levels by fluorescence-activated cell sorting (Aria II, BD Biosciences or MoFlo, Beckman Coulter) into calcium-free PBS buffer. The resulting subpopulation does not derive from a single clone, but is rather generated by the highest expressing cells that may contain multiple insertions; the 5' LTR promoter is relatively weak, and we observed an ~ 20 – 30 -fold increase in lamin A levels in comparison with the mock controls.

Analysis of Protein Expression—Cell lysates were prepared from 5×10^6 cells using urea lysis buffer with final concentrations of 9 M urea, 10 mM Tris-HCl (pH 8), 10 μM EDTA, 500 μM phenylmethylsulfonyl fluoride, 20 μl of β -mercaptoethanol, and 1 $\mu\text{l/ml}$ protease inhibitor mixture (Sigma). All steps were performed at 4°C . Proteins were separated on a 4–12% Bis-Tris gel with 1 \times MOPS running buffer and then transferred onto activated transfer membranes, blocked, and labeled using horseradish peroxidase-conjugated antibodies (Bio-Rad). We used protein standard (Invitrogen SeeBlue Plus2) for size calibration and used β -tubulin as a loading control because its levels remain constant throughout differentiation (11). Primary antibodies used for probing are described in the supplemental Methods.

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Expression levels were quantified by optical density analysis using ImageJ (National Institutes of Health).

Cell Surface Marker Analysis—To assay expression levels of CD11b, we used Fc receptor polyclonal human IgG (Sigma) as a blocking agent and labeled 10^6 cells with Alexa Fluor 700 mouse anti-human-CD11b (BD Pharmingen). We analyzed fluorescence levels by flow cytometry (LSR II, BD Biosciences).

Respiratory Burst Assay—We determined superoxide radical production of day 4/ATRA-treated cells using luminol-enhanced chemiluminescence (Diogenes reagent, National Diagnostics, Atlanta, GA) following stimulation by phorbol 12-myristate 13-acetate (25, 26), as per the manufacturer's instructions. Cells were activated by the addition of phorbol 12-myristate 13-acetate (stock solution 1 mg/ml dimethyl sulfoxide (DMSO), Sigma) to a final concentration of $11 \mu\text{M}$; we recorded luminescence values after 30 min using a plate reader (SpectraMax M5).

Nuclear Shape Analysis—We incubated cells with Hoechst 33342 ($1 \mu\text{g/ml}$, Invitrogen) for 30 min at 37°C . We then placed the cells on a glass slide pretreated with poly-L-lysine (0.01% w/v in water) by centrifuging a $20\text{-}\mu\text{l}$ drop of cell suspension at 1,000 rpm for 5 s. Images were acquired using a $20\times/0.5$ Ph2 objective (EC Plan Neofluar, Zeiss), DAPI filter set, and charge-coupled device camera (AxioCam MRm, Zeiss). Analysis of nuclear geometry was performed using ImageJ. Circularity for a nuclear cross-section is $4\pi A/P^2$, where A is the cross-sectional area and P is the perimeter.

Transwell Migration Assay—We used membranes with 3- and $8\text{-}\mu\text{m}$ pore sizes (Millipore) and FBS as chemoattractant (27, 28). Day 4/ATRA-treated cells were resuspended to 5×10^6 cells/ml in RPMI without FBS. We placed media with and without FBS in the bottom well and cells in the top well and then incubated the plate at 37°C , 5% CO_2 for 2 h. We then removed the membrane insert, labeled cells in the bottom well with Hoechst, and imaged each well by microscopy using a $10\times/0.25$ Ph1 objective (A-Plan, Zeiss), charge-coupled device camera (AxioCam MRm, Zeiss), and DAPI filter set. We determined the number of cells per well using image analysis (ImageJ).

Two-dimensional Migration Assay—Glass-bottomed dishes (World Precision Instruments, Sarasota, FL) were coated with human fibronectin ($10 \mu\text{g/ml}$ in Hanks' balanced salt solution without calcium and magnesium, Gemini Bio-products). Cells were seeded onto the dishes, and images were acquired at 1-min intervals over 3 h (5% CO_2 , 37°C) using a Zeiss microscope outfitted with an automated stage (Applied Scientific Instruments, Eugene, OR), $10\times/0.3$ Ph objective (EC-Plan Neofluar, Zeiss), and charge-coupled device camera (AxioCam MRm, Zeiss); x - y positions of cells were extracted from the resultant movies (ImageJ), and trajectory analysis was performed using MATLAB.

RESULTS

Lobulated Nuclear Shape Is Not Essential for Cell Transit—To probe the ability of neutrophil-type cells with round or lobulated nuclei to deform through narrow gaps, we designed a microfluidic device with precisely defined constrictions of $5\text{-}\mu\text{m}$ width (Fig. 1A); this width is less than the typical $7\text{--}10\text{-}\mu\text{m}$ diameter of HL-60 nuclei, such that nuclear deforma-

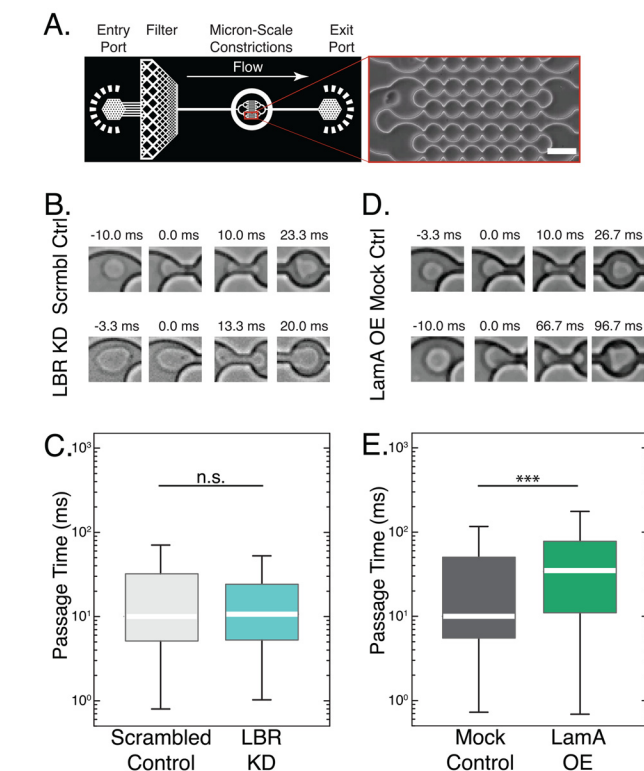


FIGURE 1. Ectopic expression of lamin A increases passage time through microfluidic constriction channels. A, schematic overview of the microfluidic device and close-up of the $5\text{-}\mu\text{m}$ constrictions. Pressurizing the reservoir drives the cell suspension through the inlet, and cells passage through the channels with $5\text{-}\mu\text{m}$ constrictions, as shown in the inset. Scale bar, $20 \mu\text{m}$. B, time-sequence images of day 4/ATRA-treated cells passing through $5\text{-}\mu\text{m}$ constrictions. *Scrambl Ctrl*, scrambled control; *Mock Ctrl*, mock control. C, LBR KD cells have similar passage times as the scrambled controls despite the round shape of their nucleus, which has been speculated to sterically hinder the passage of cells through narrow pores. D and E, LamA OE cells take longer to passage the constrictions than mock control cells. In all box plots, the white bar denotes the population median, boxes are the 25th and 75th percentiles, and lines show the 10th and 90th percentiles. n.s., $p > 0.05$ for LBR KD versus scrambled control; ***, $p < 0.001$ for LamA OE versus mock control. $n > 300$ cells for each cell type. Error bars represent S.E. over three independent experiments.

tion is required for a cell to passage through a pore (supplemental Fig. S1). We forced the neutrophil-type cells (day 4/ATRA-treated HL-60 cells) to transit through these micron-scale pores using pressure to drive a flow of cell suspension through the channels; we monitored the passage of cells as a function of time. When a cell arrives at a constriction, it is subjected to physical forces resulting from external stresses due to the pressure drop across the cell trapped in the constriction; these stresses cause the cell to deform and passage through the pore. Given the dimensions of a single pore, a pressure of 28 kilopascals corresponds to approximately micronewton-scale forces. The rate at which the cell deforms largely depends on the applied stress (driving pressure) as well as the global mechanical properties of the cell and nucleus (5, 29–32). As individual cells deformed through the $5\text{-}\mu\text{m}$ constrictions of the microfluidic device, we imaged their passage using a high speed camera (Fig. 1, B and D). By automated image analysis, we determined the time required for the cell to passage through the first $5\text{-}\mu\text{m}$ constriction, which we define as its passage time. Given these millisecond timescales of cell passage at a driving pressure of 28

kilopascals, this microfluidic assay primarily probes the passive mechanical behavior of the cell, as actin remodeling and protein expression changes occur on timescales of several minutes and more (33). Although actin can contribute to the cortical stiffness of neutrophils (4–6), we confirmed that the actin makes little contribution to these measurements by treating a subset of neutrophil-type cells with cytochalasin D to disrupt actin polymerization; this treatment had no effect on passage times (data not shown), indicating that the deformability of the nucleus has a pivotal role in the passage of cells through micron-scale pores.

To assess the effect of hypobulbated or round-shaped nuclei on the passage of neutrophil-type cells through micron-scale constrictions, we used LBR KD cells as an *in vitro* system. In contrast to the control cells that exhibit strong up-regulation of LBR during differentiation and develop lobulated nuclei, LBR KD cells show only trace levels of LBR expression and maintain round nuclei (17). Nevertheless, despite their round nuclei, LBR KD cells exhibited similar passage times as compared with the scrambled control cells (Fig. 1B). These observations suggest that the multilobed shapes of nuclei in mature neutrophils provide no significant advantage in the time required for cells to deform through 5- μ m constrictions.

Generating Neutrophil-type Cells with Increased Lamin A Expression—Because the above experiments indicate that lobulated nuclear shape is not essential for neutrophil-type cell passage through narrow constrictions, we hypothesized that the unique molecular composition of the nuclear envelope in neutrophils could determine the ability of cells to deform. One possible origin may be the low levels of the key structural protein of the nucleus, lamin A; this protein is normally down-regulated by over 90% in ATRA-stimulated HL-60 cells after 4–5 days of stimulation (Fig. 2B) (11, 14). Given the essential role of lamin A in nuclear mechanical stability (12, 13, 30), we postulated that preventing lamin A down-regulation could reduce nuclear deformability and impair cell passage through pores. Because the LBR KD neutrophil-type cells that have round nuclei have similar reduced lamin A expression levels as unmodified and mock-modified cells, this may also explain their unaltered passage times (17).

To test the effect of increased lamin A levels on cell passage through narrow constrictions, we generated a LamA OE HL-60 cell line by retroviral transduction. The resulting subpopulation of high expressing cells exhibits lamin A levels that are about 20–30-fold higher than the mock-modified cells (supplemental Fig. S2A). Although lamin A expression levels in the LamA OE cells are greater than those in unmodified HL-60 cells, they are comparable with physiological levels in other somatic cells such as mouse embryo fibroblast cells (supplemental Fig. S2B). To confirm that the ectopic lamin A is properly localized to the nuclear envelope, we conducted immunofluorescence and confocal imaging (supplemental Fig. S3).

Protein Composition of LamA OE Cells—To characterize how protein levels of the LamA OE cells change during granulopoiesis, we monitored expression levels of major structural proteins over the differentiation time course; we induced the HL-60 cells to differentiate into neutrophil-type cells by ATRA treatment, collected cell lysates at days 0, 3, and 5 following

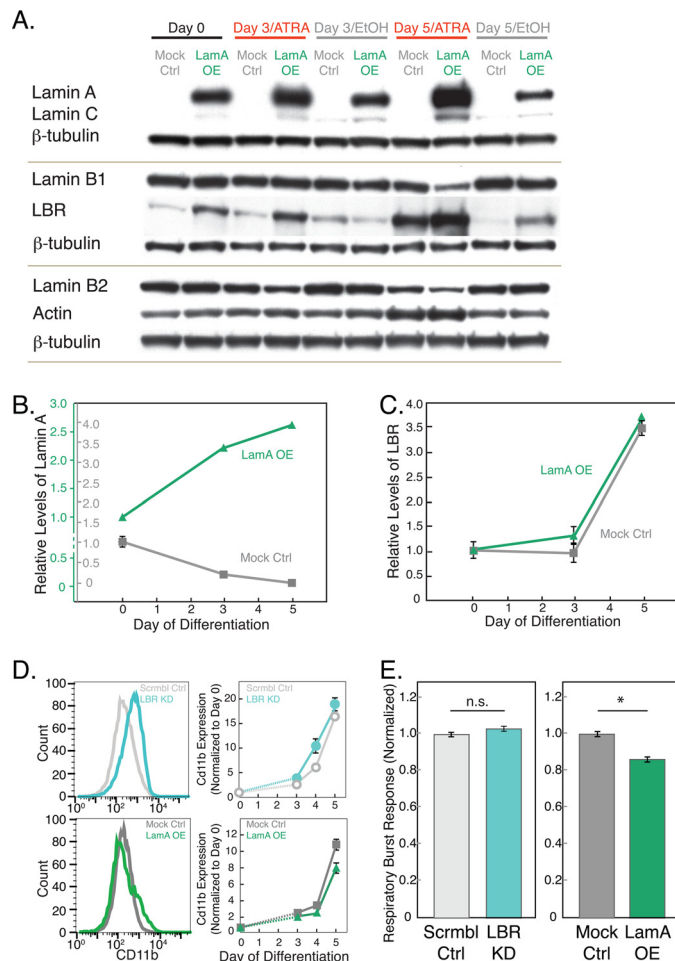


FIGURE 2. Genetically modified HL-60 cells show typical characteristics of neutrophils after ATRA stimulation. A, representative immunoblots for lamins A/C, B1, B2, and LBR with β -tubulin as loading control. Cell lysates are collected from LamA OE and mock control (*Mock Ctrl*) cells at days 0, 3, and 5 after ATRA stimulation. B and C, quantitative analysis of lamin A and LBR protein levels normalized first to β -tubulin and then to day 0 for each protein in each cell line. Error bars represent S.E. of 3–5 independent experiments; where not visible, they are smaller than the symbols. Based on immunoblot analysis, base-line levels of lamin A are estimated to be ~ 20 – $30\times$ greater in the LamA OE cells as compared with the mock control cells (supplemental Fig. S2); for this reason, two separate axes are plotted for each cell line. D, expression levels of the cell surface antigen, CD11b, a hallmark of neutrophils, increase during differentiation for all cell lines. Left, representative histograms of data from a single flow cytometry experiment showing the distribution of CD11b expression levels at day 4 after ATRA stimulation. Right, graphs showing median values of CD11b after ATRA treatment with the values for each cell line normalized to day 0 for each independent experiment. Error bars represent S.E. over three independent experiments. *Scrambl Ctrl*, scrambled control. E, respiratory burst assay that probes superoxide production using a luminescence assay 30 min after stimulation by phorbol myristate acetate, indicating that all cells show normal functional characteristics of neutrophils. Luminescence values are relative to the mock and scrambled control for the left and right panels, respectively. Data represent the average of three independent experiments; error bars represent the S.E. n.s., $p > 0.05$ for LBR KD versus scrambled control; *, $p < 0.05$ for LamA OE versus mock control.

ATRA treatment, and performed immunoblotting (Fig. 2A). As expected, unmodified and mock-modified cells displayed a strong up-regulation of LBR during granulopoiesis with a concurrent decrease in lamin A levels (Fig. 2, B and C), confirming previous observations (11). In contrast, LamA OE cells have increased levels of lamin A that further increased during granulopoiesis (Fig. 2, B and C), possibly due to an ATRA-sensitive element in the ectopic promoter region. LamA OE cells showed

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elevated basal levels of LBR as compared with the mock controls, with a similar ~4-fold increase in LBR levels during granulocytic differentiation. We also probed levels of other structural proteins that could contribute to cell deformability (4–6); during differentiation in both the mock-modified and LamA OE cells, actin levels showed minor variations, and other structural nuclear proteins, including lamin B1 and B2, showed a decrease in expression levels (Fig. 2) (34). Although we cannot exclude the possible contribution of lamin B1 and B2 down-regulation to altered cellular mechanical properties, we anticipate that the observed changes in lamin B1/B2 levels would have little effect on nuclear mechanical properties in comparison with the lamin A up-regulation; lamins A/C have a predominant role in nuclear shape stability and stiffness (12, 13, 30), whereas lamin B1 does not have any significant effect on nuclear mechanical stability (13).

Genetically Modified Cells Display Characteristics of Neutrophils—To test whether the genetically modified HL-60 cells still undergo normal granulopoiesis, we assayed essential functional, biochemical, and proteomic characteristics that define neutrophils. One metric to assess the differentiation of HL-60 cells into neutrophil-type cells is to measure cell density following ATRA stimulation; decreased proliferation rates are an indicator of successful differentiation as cells exit the cell cycle to commit to their differentiation into neutrophils (35). Both LBR KD and LamA OE cells showed a similar progressive decrease in proliferation rates over the days following ATRA treatment as compared with the scrambled and mock controls (supplemental Fig. S4). As a more direct assay of differentiation into neutrophil-type cells, we measured expression levels of the cell surface marker, CD11b, a subunit of a heterodimeric adhesion glycoprotein, which is widely used as a marker for neutrophils (36). After 4 days of ATRA treatment, CD11b levels were increased for all cell types as compared with the undifferentiated HL-60 cells and vehicle-treated controls (supplemental Fig. S5). These results confirmed that the HL-60 cells are differentiating into neutrophil-type cells. Importantly, we observed that all cell lines show significant increase in CD11b levels following ATRA stimulation, with levels varying slightly between cell lines (Fig. 2D); this demonstrates that the changes in nuclear envelope composition do not markedly affect differentiation efficiency.

Another hallmark of neutrophil cells is their respiratory burst response upon exposure to phagocytotic stimuli, such as yeast or bacteria. To probe this functional characteristic of the modified neutrophil-type cells, we stimulated cells with phorbol myristate acetate and measured subsequent superoxide production (Fig. 2E). LBR KD cells showed similar response as compared with the scrambled controls. LamA OE cells exhibited a small yet statistically significant 15% reduction in superoxide production as compared with the mock controls. Overall, these experiments suggest that the functional and biochemical characteristics of the genetically modified neutrophil-type cells are generally maintained despite their altered nuclear envelope composition.

Lamin A Expression Alters Nuclear Lobulations during Granulopoiesis—A key hallmark of granulopoiesis is the transition from round to multilobed nuclear shape, which is observed in

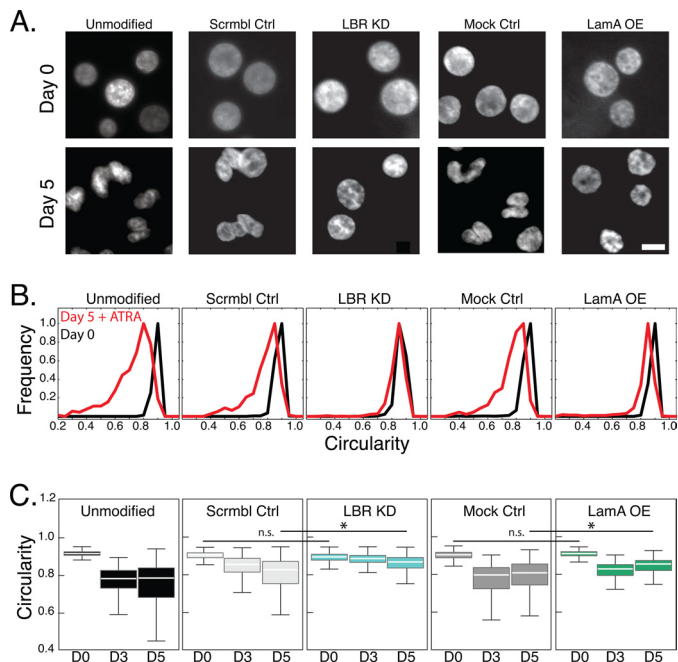


FIGURE 3. Nuclear shape transition during granulopoiesis requires lamin A down-regulation and LBR up-regulation. *A*, fluorescent images of Hoechst-stained nuclei acquired at day 0 and day 5 after ATRA treatment. All images were acquired at the same magnification. *Scale bar*, 5 μ m. *B*, to quantitatively describe nuclear shape, the circularity of the nucleus is defined as $4\pi A/P^2$. Histograms show the distribution for each cell type at days 0 and 5 after ATRA treatment. *C*, box plots show the circularity of nuclei at days 0, 3, and 5 after ATRA treatment. The *white bar* denotes the population median, *boxes* are the 25th and 75th percentiles, and *lines* show the 10th and 90th percentiles. To evaluate statistical significance, we compared the medians of at least 3 independent experiments for each cell type. Day 0, unmodified to scrambled control. *n.s.*, $p > 0.05$; *, $p < 0.05$. Nuclei from over 300 individual cells were analyzed for each cell type.

HL-60 cells after 3–5 days following ATRA treatment (11, 18). To investigate the effect of altered nuclear envelope composition on this shape transition, we imaged Hoechst-stained nuclei by fluorescence microscopy over the differentiation time course. To quantify changes in nuclear shape, we analyzed the circularity of nuclei, defined as $4\pi A/P^2$, where A is the cross-sectional area and P is the perimeter of an individual nuclear cross-section. For a perfect circle, the circularity value equals one; lower values reflect deviations from a circular shape. In the undifferentiated state, all cell lines have nuclei with predominantly circular shape and similarly high circularity values, representative of round nuclei (Fig. 3, A–C).

After 3 days of ATRA treatment, the unmodified, mock, and scrambled control cells exhibited nuclei with large invaginations; circularity values correspondingly showed a lower median and greater variability, reflecting these irregular nuclear shapes. By contrast, LBR KD cells retained their round shape, as reflected by the higher circularity values, even after 5 days of ATRA treatment (Fig. 3A) (17). The nuclei of LamA OE cells showed some morphological changes but failed to develop the characteristic lobulations seen in the unmodified and mock control cells (Fig. 3A, supplemental Fig. S3); the lack of severe lobulation that is typical for normal neutrophil cells illustrates that down-regulation of lamin A expression during neutrophil differentiation could also be required for the lobulated nuclear shape of mature neutrophils.

Increased Lamin A Expression Delays Cell Passage through Pores—Lamin A is a crucial modulator of nuclear deformability (13, 30, 37). To probe the effects of increased lamin A levels on the ability of cells to deform through physiological gaps, we measured the passage time of LamA OE neutrophil-type cells and mock controls when forced through the 5- μm constrictions of our microfluidic device (Fig. 1, A and D). The LamA OE neutrophil-type cells exhibited a 3-fold increase in median passage time as compared with the mock controls (Fig. 1E); these results indicate that increased density of lamin protein at the nuclear envelope may impair the ability of LamA OE neutrophil-type cells to passage through the 5- μm constrictions. Taken together, our results show that lamin A levels have an important effect on the ability of cells to passage through 5- μm constrictions; physiological down-regulation of lamin A following ATRA-induced differentiation of HL-60 cells results in faster passage through the micron-scale constrictions, whereas ectopically increased expression of lamin A results in slower passage of LamA OE-neutrophils through the 5- μm constrictions.

Active Migration through Pores Is Impaired in LamA OE Cells—The results of our microfluidic experiments illustrate that altered expression of lamin A can substantially alter the passive deformability of cells. However, a critical function of neutrophils is their ability to actively migrate through narrow constrictions. To test migration efficiency, we used a transwell migration assay to probe the ability of cells to migrate through 3- and 8- μm pores; we monitored the number of cells that migrate through the pores after 2 h and determined the migration efficiency relative to the respective control cells. As seen in the passive deformation results obtained by microfluidic assays, the LBR KD cells exhibited similar migration efficiency as the scrambled control cells (Fig. 4, A and B), further substantiating that neutrophil-type cells with round nuclei can exhibit equivalent passage efficiency through micron-scale pores. By contrast, the LamA OE cells showed a marked reduction in migration through 3- μm pores (Fig. 4, D and E). The impaired migration was less severe in the experiments with 8- μm pores (Fig. 4, D and E); because deformation through 8- μm pores requires smaller deformations of nuclei, these results are consistent with our observations that nuclear deformation rate-limits the passage of cells through micron-scale constrictions.

To address the possibility that a general migration defect underlies the impaired transwell migration efficiency of the LamA OE cells, we performed two-dimensional migration assays; cells exhibited velocities from 2 to 5 $\mu\text{m}/\text{min}$, consistent with previous observations of neutrophil migration (38). The LBR KD cells showed a slightly increased velocity as compared with the scrambled control cells (Fig. 4C). Importantly, LamA OE cells exhibited similar migration velocities as the mock control cells (Fig. 4F), indicating that the observed differences in the transwell assay cannot be attributed to general defects in their migration. Taken together, our experiments indicate that the density of lamin A at the nuclear envelope is crucial in facilitating the passage of cells through micron-scale constrictions.

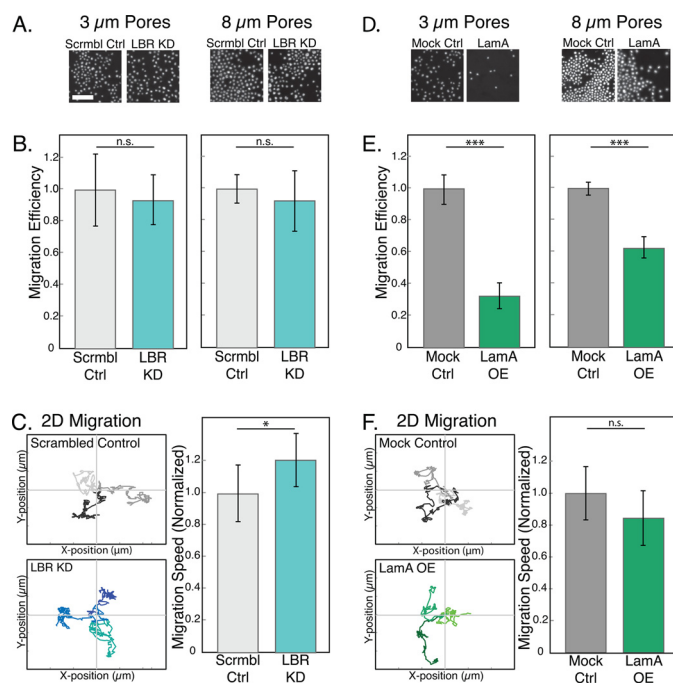


FIGURE 4. Impaired migration of LamA OE cells through narrow constrictions. To probe the active migration of cells through micron-scale pores, we use a Transwell migration assay. A and D, representative images from a single experiment showing Hoechst-stained nuclei of cells that have passed through 3- or 8- μm pores. Scale bar, 100 μm . Ctrl, control; Scrambl Ctrl, scrambled control; Mock Ctrl, mock control. B and E, migration efficiency is defined as the number of cells that passed through the porous membrane relative to the corresponding scrambled or mock control. Bars represent averages from at least three independent experiments; error bars represent S.E. ***, $p < 0.001$ for LamA OE versus mock control. C and F, two-dimensional migration experiments were performed by tracking the positions of individual cells at 1-min intervals over 3 h. Traces of three representative cells for each cell type show the total distance traveled and the directionality of movement over the three-hour time-lapse experiment. Axes are 150 μm with 50- μm increments. Migration speed over the entire trajectory is computed from the individual traces of over 50 cells for each cell type. Mean values for each LBR KD and LamA OE cells are normalized to their respective controls. n.s., $p > 0.05$; *, $p < 0.05$; ***, $p < 0.001$. Absolute velocities of cells are: LBR KD, 4.4–5.1 $\mu\text{m}/\text{min}$; scrambled control, 3.5–4.2 $\mu\text{m}/\text{min}$; LamA OE, 2.2–4.7 $\mu\text{m}/\text{min}$; mock control, 2.8–4.8 $\mu\text{m}/\text{min}$.

DISCUSSION

It has long been speculated that the lobulated shape of the neutrophil nucleus is "a special adaptation for passing through vessel walls" (8). However, here we show that nuclear shape alone does not always determine the timescale for neutrophil deformation through micron-scale pores; neutrophil-type cells with round nuclei resulting from LBR knockdown (17) show unaltered passage efficiency through pores down to 3 μm , as probed using both passive deformation through 5- μm microfluidic constrictions, as well as active migration through 3- and 8- μm porous membranes.

These LBR KD cells also provide an *in vitro* model for Pelger-Huët anomaly; the nuclei from neutrophils of these individuals are round or bilobulated (36, 37) due to a complete or partial lack of functional LBR. The extent to which the altered nuclear shape of Pelger-Huët anomaly neutrophils affects their ability to passage through micron-scale constrictions has been inconclusive (26, 39–41). Some previous studies of these neutrophils discovered altered migration (26, 39, 40); however, these primary neutrophils also exhibited bilobular nuclei, and differ-

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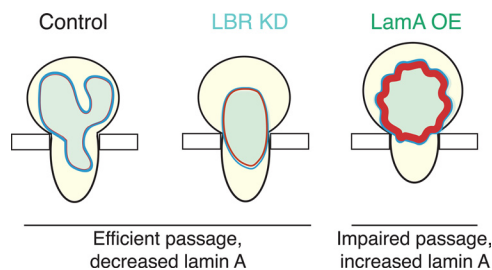


FIGURE 5. Lamin A levels, rather than nuclear shape, are a primary determinant of the efficiency of cell passage through narrow constrictions. A schematic illustration summarizing the effects of nuclear shape and lamin A expression levels on the ability of cells to deform through narrow constrictions is shown. The ratio of LamA to LBR expression levels is estimated from immunoblots. Undifferentiated (unmodified or mock-modified) HL-60 cells, as well as the LBR KD neutrophil-type cells, exhibit efficient passage, despite their round nuclear shape. By contrast, lamin A overexpression results in impaired passage, both through the constricted 5- μm channels of a microfluidic device, as well as the 3- and 8- μm pores of the transwell migration assay. Undifferentiated LamA OE cells with more circular nuclei and lower levels of LBR require even longer time to passage through narrow constrictions.

ences in migratory ability could result from other phenotypic differences. Here we used LBR KD cells as an *in vitro* system to specifically investigate the effect of the hypobolulated nucleus on cell passage through micron-scale constrictions. Despite the round shape of LBR KD nuclei, which could sterically hinder the passage of nuclei through constrictions, these cells exhibited similar passage efficiencies through micron-scale constrictions as compared with the scrambled control cells with multi-lobed nuclei (Fig. 1).

Although LBR KD neutrophil-type cells have an atypical round nuclear shape, they have similarly low levels of lamin A as the unmodified controls (17). Here, we show that lamin A expression levels, rather than the shape of the cell nucleus, can be a major determinant of the timescale of cell passage through micron-scale gaps (Fig. 5). By contrast, nonmechanical functions of these cells are not substantially affected by changes in nuclear envelope composition. Although other types of white blood cells with ovoid-shaped nuclei, such as macrophages, also undergo transendothelial migration, their deformations occur on a slower timescale as compared with neutrophils (31). Indeed, monocyte/macrophage-differentiated HL-60 cells also show increased levels of lamin A/C expression relative to neutrophil-type cells (34).

If irregular nuclear shape is not essential for the deformability of neutrophil cells, then why do their nuclei exhibit this distinct shape? One possibility is that the multilobed nucleus could simply result from the marked changes in nuclear envelope protein composition. Indeed, lamin A levels impact the mechanical stability of the nuclear envelope, whereas ectopic overexpression of LBR can increase nuclear membrane surface area (42, 43) (Fig. 5). Alternatively, the unusual multilobed nuclear shape may facilitate other neutrophil functions, such as phagocytosis, the formation of neutrophil extracellular traps, or migration through even smaller $<1\text{-}\mu\text{m}$ constrictions of the endothelium, either between or through cells (44).

Here we have used HL-60 cells, which are a well established *in vitro* model system to study white blood cell lineages; for example, the resulting neutrophil-type cells show similar structural and functional characteristics as primary neutrophils

(14–16). HL-60 cells also exhibit similar mechanical properties; recent measurements of cell compliance using an optical stretcher confirmed that *in vitro* differentiation of HL-60 cells into neutrophil-type cells recapitulates the 3–6-fold increase in cell deformability observed in primary neutrophils and their CD34+ precursor cells (31). A direct comparison of the absolute passage times through micron-sized constriction between primary neutrophils and HL-60-derived neutrophil-type cells is complicated by the fact that HL-60 cells are typically larger than primary neutrophils ($\sim 12\text{-}\mu\text{m}$ versus $7\text{--}8\text{-}\mu\text{m}$ median diameter, respectively) and exhibit substantially larger transit times through microfluidic constriction channels (29). Consequently, we have focused our study on HL-60 cells and vary protein levels within the same cell type; this has enabled us to clearly illustrate the importance of nuclear envelope composition, particularly the levels of lamins A/C, on the ability of cells to pass through narrow constrictions during perfusion and migration.

It is intriguing to speculate that changes in levels of lamin A expression may have implications for cellular deformability in a variety of physiological processes and diseases (45). For example, certain types of cancer cells have reduced levels of lamin A expression as compared with their nonmalignant progenitors (46, 47). Akin to neutrophils, large deformations of cancer cells and their nuclei are required during deformation through micron-scale constrictions (48) in extravasation and metastasis. Ultimately, a deeper knowledge of the molecular basis of cellular and nuclear deformability will provide unique insights into the mechanical aspects of cell biology and possibly new therapeutic approaches.

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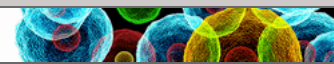
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Cell Biology:

**Nuclear Envelope Composition Determines
the Ability of Neutrophil-type Cells to
Passage through Micron-scale
Constrictions**

CELL BIOLOGY



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