Nuclear lamins are not required for lamina-associated domain organization in mouse embryonic stem cells

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Abstract

In mammals, the nuclear lamina interacts with hundreds of large genomic regions, termed lamina-associated domains (LADs) that are generally in a transcriptionally repressed state. Lamins form the major structural component of the lamina and have been reported to bind DNA and chromatin. Here, we systematically evaluate whether lamins are necessary for the LAD organization in murine embryonic stem cells. Surprisingly, removal of essentially all lamins does not have any detectable effect on the genome-wide interaction pattern of chromatin with emerin, a marker of the inner nuclear membrane. This suggests that other components of the lamina mediate these interactions.

Keywords interphase chromosome organization; genome-wide mapping; nuclear architecture; nuclear lamina

Introduction

The nuclear lamina (NL) is thought to play a role in the spatial organization of the genome inside the interphase nucleus, by providing an interaction platform for hundreds of large genomic regions named lamina-associated domains (LADs). Genes in LADs are generally inactive, indicating that LADs form a transcriptionally repressive chromatin compartment [1]. It is likely that specific proteins of the NL recognize DNA sequences or chromatin features in LADs and thereby anchor LADs to the periphery [2]. However, the identity of these NL proteins is still largely unknown.

The primary components of the NL are lamins, which are intermediate filament-type proteins that form a fibrous network [3]. Most mammalian somatic cells express four lamin proteins derived from three genes: lamins A and C (LmA/C), encoded by the LMNA gene, and B1 and B2, encoded by LMNB1 and LMNB2, respectively. In addition to lamins, the NL harbors a range of other proteins, several of which are anchored in the inner nuclear membrane [4].

Several reports suggest that lamins may be involved in chromosome organization. Both A- and B-type lamins can directly interact with genomic DNA and chromatin in vitro [5]. In D. melanogaster and C. elegans, depletion of lamins can have effects on gene positioning and silencing [6]. Human fibroblasts with a dominant-negative mutation in the LmA gene causing Hutchinson-Gilford progeria syndrome exhibit altered chromatin–LmA/C interactions, loss of peripheral heterochromatin, and a perturbed overall chromosome structure [7,8]. Depletion of lamin B1 (LmB1) in somatic mouse and human cells was found to lead to changes in interphase chromosome morphology and positioning, concomitant with genome-wide changes in gene expression [9–12]. In contrast, no change in cell behavior and gene expression was observed in keratinocytes and murine embryonic stem (mES) cells lacking lamins [13–15].

While most of these data point to a role for lamins in certain aspects of interphase chromosome architecture, they do not directly address whether lamins are necessary for overall LAD organization. Here, we address this issue by generating a series of high-resolution maps of NL–genome interactions in mES cells upon systematic removal of all lamins. We correlate these results with microscopy observations and gene expression profiling. Surprisingly, these data indicate that lamins are dispensable for LAD organization of the genome, at least in mES cells.

Results

To address the role of lamins in the formation of LADs, we used the DNA adenine methyltransferase identification (DamID) technique [1] to construct genome-wide maps of NL interactions in wild-type (wt) and lamin B1/B2 double knockout (dKO) mES cells [13].

 Naturally, we could not perform DamID with a Dam-lamin fusion protein, as this would reintroduce a lamin into the null background. We instead selected the lamin-interacting inner nuclear membrane protein emerin (Emd) as Dam-fusion partner, for several reasons. First, Emd is endogenously present in mES cells and it is expressed at similar levels in wt and dKO mES cells, as indicated by mRNA sequencing (Supplementary Fig S1A). Second, by immunofluorescence microscopy, we found that the localization of Emd at the nuclear envelope is not perturbed in dKO mES cells (Supplementary
Fig S1B), which is consistent with recent reports [13,16]. Third, in human cells, the DamID profiles of Emd and LmB1 are virtually identical [1], indicating that Dam-Emd can be used instead of Dam-LmB1 to identify LADs. To verify that this is also the case in wt mES cells, we generated a genome-wide DamID profile for Emd and compared it to the one previously generated with LmB1 [17]. Indeed, the two profiles were similar, with an overall Pearson correlation coefficient of 0.87 and highly similar domain patterns along the chromosomes (Supplementary Fig S1C and D).

Having established that Emd can substitute for LmB1 to identify LADs, we then generated DamID profiles in wt and dKO mES cells. Dam-Emd was expressed at very low levels using the low leaky activity of a non-induced *Drosophila* heat-shock promoter [18] in order to avoid interference with endogenous NL components. Amplification of the DNA methylated by Dam-Emd did not show differences in yield between the cell lines (Supplementary Fig S1E), further indicating that in dKO cells, there is no major relocation of Dam-Emd from the inner nuclear membrane to a cytoplasmic compartment. This is consistent with previous observations that Emd is largely retained in the nuclear envelope of mES cells lacking lamins, which contrasts with Emd behavior in differentiated cell types lacking lamins [16,19]. For both wt and dKO mES cells, we obtained Emd interaction maps by combining the data from two independent DamID experiments.

Strikingly, the Emd interaction patterns of wt and dKO mES cells were highly similar in genome-wide correlation, amplitude of the signals, and overall appearance (Fig 1A and B). We used a domain detection algorithm [1] to determine for each cell line the number, size, and genome coverage of the LADs. While the total number of LADs was slightly reduced in dKO cells, their total coverage along the genome was nearly identical (38.4% vs 38.8%), and there was a strong concordance between their positions in wt and dKO cells (Fig 1C and D). Taken together with a general lack of off-diagonal data points in the scatterplot analysis (Fig 1A), these data indicate that overall LAD organization is largely retained in dKO cells.

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**Figure 1.** No detectable changes in LADs organization in dKO mES cells.

A Scatterplot of \( \log_2 \) ratio of Dam-Emd over Dam for wt compared to dKO mES cells. Pearson correlation coefficient \( r \) is indicated.

B Emd interaction profile along chromosome 18 in wt and dKO mES cells.

C Distribution of LAD sizes in wt (black) and dKO (red) mES cells. Dashed lines mark the median LAD sizes. Percentage of total genomic DNA covered by LADs (% of gDNA), the median size of LADs, and the total number of LADs are listed for wt (black) and dKO (red) cells.

D Concordance between wt and dKO cells for LADs and iLADs as defined in (C).

E Distribution of Emd DamID probe values in wt and dKO mES cells, divided into facultative (f) and constitutive (c) LADs and iLADs. Horizontal lines of boxes depict percentiles 25, 50, and 75, vertical lines extend from the box edge to the highest or lowest value that is within 1.5× inter-quartile range of the edge.

F Concordance between wt and dKO cells for facultative (fLAD, fiLAD) and constitutive (cLAD, ciLAD) regions, as identified in (E).

Data information: Data in (A) and (B) were smoothed (median) with a running window of 11 probes. DamID data are averages of two independent experiments.
We then investigated whether specific subsets of LADs were affected, which may not be noticeable in the bulk analyses above. Specifically, we tested whether the previously identified facultative (cell-type specific) or constitutive (cell-type invariant) LADs and inter-LADs were affected [20]. Given their different dynamics during cell differentiation, it was possible that they would respond differently to the loss of B-type lamins. However, these regions showed a high overall concordance, with almost identical interactions of the constitutive regions (Fig 1E and F). A somewhat lower concordance was observed in facultative LADs, but this should be interpreted with caution because these regions have somewhat weaker DamID signals overall in wt cells, and therefore, the signal/noise ratio may be lower in these regions. Finally, we applied a specially designed statistical test to identify genes with significantly altered DamID signals (Fig 2C) or with their constitutive or expression changes between wt and dKO mES cells did not correlate with NL interaction changes (Fig 2C) or with their constitutive or facultative status (Supplementary Fig 2B and C), again indicating that LADs remain largely unaffected in dKO mES cells.

Next, we investigated whether B-type lamins are involved in repressing genes at the NL. We generated mRNA expression profiles of wt and dKO mES cells and averaged two biological replicates for each cell line. In wt mES cells, the genes that interact with the NL (high DamID log2-ratios) generally exhibit low mRNA expression (Fig 2A), as it was reported previously for various cell types [1,17]. This correlation was also observed in dKO mES cells, indicating that the NL remains a repressive environment regardless of the presence of B-type lamins (Fig 2B). The wt and dKO mES cell mRNA profiles showed an overall Pearson correlation coefficient of 0.99, with only 94 genes changing expression (P ≤ 0.05 and fold change ≥ 2) out of 37,991 genes that were analyzed (Supplementary Fig S2A). Of the differentially expressed genes, 18 are in LADs and 76 in inter-LADs, a distribution that is not statistically different from what may be expected by chance (P = 0.7, Fisher’s exact test). Indeed, gene expression changes between wt and dKO mES cells did not correlate with NL interaction changes (Fig 2C) or with their constitutive or facultative status (Supplementary Fig 2B and C), again indicating that the rather modest changes in gene expression are generally not linked to altered NL interactions.

One possible caveat to our analyses is that B-type lamins could be functionally compensated by LmA/C. While LmA/C was initially thought not to be present in mES cells [13,21], it was later established with multiple antibodies that it is expressed in mES cells, including the cell lines studied here [15,16,22]. In our cultures, we could detect moderate levels of LmA/C transcripts by RT–PCR and RNA sequencing and of LmA/C proteins by Western blot using two different antibodies, one detecting both isoforms and one specific for LmA (Supplementary Fig S3A–C). Co-staining with an Oct4 antibody confirmed that the cells are in an undifferentiated state (Supplementary Fig S3D).

Immunofluorescence microscopy of LmA/C showed the classic nuclear rim localization in wt mES cells. In dKO mES cells, LmA/C also localized to the nuclear rim, but it showed a more punctuated distribution. This we observed with the two different antibodies described above (Supplementary Figs S1B and S3E), and is in agreement with recently reported observations [16]. In addition, we observed the patchy distribution of LmA/C in LmB1–/– mES cells, but not in LmB2–/– cells, pointing to LmB1 as being responsible for the even distribution of LmA/C along the nuclear envelope (Supplementary Fig S3F).

We could recapitulate the LmA patchy pattern by lentivirus-mediated expression of GFP-tagged LmA in dKO mES cells, while GFP-LmA exhibited a normal homogeneous rim staining in wt mES cells (Supplementary Fig S3G). Surprisingly, GFP-tagged LmC also accumulated in bright spots in the nuclear interior of dKO mES cells, while it was properly located at the nuclear periphery in wt mES cells (Supplementary Fig S3H), indicating that LmC requires one or both B-type lamins for its proper integration in the NL of mES cells.

Previous studies have shown that LmA and B-type lamins interact with the same LADs [20,23,24]. To test whether LmA requires B-type lamins for its LAD interactions, we generated DamID maps using Dam-LmA in wt and dKO mES cells. Because the DamID protocol requires the Dam-LmA protein to be expressed at very low levels [18], it is unlikely that this small amount of fusion protein perturbs the distribution of endogenous LmA. In the rare cells in which the Dam-LmA is weakly detectable by immunofluorescence microscopy, we found it to show a similar patchy appearance at the nuclear rim as endogenous LmA in dKO cells (Supplementary Fig S3I).

Inspection of the DamID profiles revealed that in wt mES cells, the detected LmA interactions were similar to the Emd interactions...
Interestingly, LmA–genome interactions in dKO cells were highly similar to those in wt cells (Fig 3B–G). Thus, even though B-type lamins are absent and LmA is not homogeneously distributed along the nuclear periphery, LmA in dKO mES cells can still interact with all LADs that are contacted by Emd in dKO mES cells and by LmA and Emd in wt mES cells. The difference in DamID signal between LADs and inter-LAD regions (iLADs) is lower in dKO mES cells (Fig 3F), possibly since only a random subset of LADs in each single cell can interact with the patches of LmA.

To address whether LmA/C are required for LAD formation in the absence of B-type lamins, we further reduced LmA/C expression in dKO mES cells by RNA interference using a short hairpin RNA (shRNA). By RT–qPCR, Western blot, and immunofluorescence labeling, we found that the LmA/C transcripts and proteins were substantially reduced (Supplementary Fig S4A–C). For simplicity, we will refer to the resulting cells as tKO mES cells. We confirmed that in the absence of all lamins, mES cells maintain their undifferentiated phenotype (Supplementary Fig S4D). To perform DamID, we again used Dam-Emd, since Emd is clearly present at the NE of mES cells also in the absence of LmA/C (Supplementary Fig S4E) and amplification of the DNA methylated by Dam-Emd did not show differences in yield between the cell lines (Supplementary Fig S4F). Again, the DamID maps show that chromatin organization in wt and tKO mES cells is virtually indistinguishable (Fig 4A–F).

Statistical analysis [17] did not detect any gene with a significant change in DamID log-ratios. Thus, removal of nearly the entire lamin meshwork has no detectable influence on the interaction pattern of the genome with the residual NL in mES cells.

Complementary to this DamID mapping in tKO mES cells, we inspected the distribution of DNA inside the nucleus of dKO mES cells by confocal microscopy. We noticed that not only LmA has a patchy appearance, but also the density of DAPI staining is non-uniform along the nuclear rim (Fig 5A–C). We reasoned that if LmA participates in the tethering of heterochromatin, then the LmA-rich patches might preferentially bind regions with more dense DAPI staining, compared to the intervening LmA-poor areas of the NL. We designed a quantitative image analysis approach to test this (Fig 5B–D). Analysis of 54 nuclei revealed that there is no significant positive correlation between the presence of LmA and local DNA intensity (Fig 5D), again suggesting that the presence of LmA does not detectably help in tethering DNA to the NL.

Discussion

Although lamins are able to interact with chromatin in vitro and in vivo [5], we report here that lamins are to a very large extent dispensable for the LAD organization of the genome in mES cells.
Because Dam-Emd produces in wt cells essentially the same genome-wide DamID profile as Dam-LmB1, and because it has previously been demonstrated that Dam-LmB1 methylation signals correlate with NL proximity in the nucleus [1,17,23], it is reasonable to interpret the DamID profiles obtained here with Dam-Emd as maps of NL contact probabilities in the nucleus. We found that this genome-wide NL interaction pattern remains virtually unchanged in the absence of LmB1, LmB2, and LmA/C. Moreover, only a handful of genes exhibit altered expression in the absence of LmB1 and LmB2, but these genes are not enriched in LADs, indicating that B-type lamins are not involved in silencing genes at the NL, which is in agreement with a previous study [13].

Our results contrast with results obtained in flies and worms, where depletion of lamins was found to affect the expression and the peripheral positioning of specific genomic loci [6,25,26]. Loss of LmB1 has also been reported to result in changes in nuclear organization in differentiated mouse and human cells. For example, in mouse fibroblasts, the loss of LmB1 caused relocation of chromosome 18 from the periphery toward the nuclear interior [9]. Depletion of LmB1 in a human colon cancer cell line resulted in overall decondensation of chromosome territories [10], while in HeLa cells, it was reported to lead either to an overall reduction in transcription followed by relocation of chromosomes toward the nuclear periphery [11], or to the formation of nuclear blebs that have a euchromatic appearance yet are devoid of transcription activity [12]. These results were all based on microscopy observations. A study in human fibroblast expressing progerin, a dominant-negative form of lamin A, also indicates that the disruption of the nuclear lamina results in a loss of peripheral heterochromatin, change in H2K27me3 distribution, and a global loss of chromatin compartmentalization [7].

Our data suggest that mES cells do not require lamins for overall LAD organization. It is possible that mES cells employ a unique mechanism for LAD organization, because mES cells appear to have a more dynamic chromatin architecture [27,28], exhibit an extremely rapid cell cycle [29], have a more plastic NL architecture in which LmB1 is less stably incorporated [30], and have a higher AT content in its LAD sequences compared to other differentiated cells [20].

It thus seems likely that one or more non-lamin components of the NL help to organize LADs. The lamin B receptor (Lbr) was reported to interact with chromatin in vitro [31–33] and recently to
Lamins are dispensable for LAD organization

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Materials and Methods

Cell culture, lentivirus transductions, and DamID

mES cells were cultured as described in Supplementary Methods. The wt and dKO mES cells showed no major differences in proliferation rate, cell cycle distribution, and undifferentiated status (Supplementary Fig S6–D). Comparison of our mRNA-seq data to published microarray-based RNA expression profiles from the same mES cells [13] showed good correspondence (within the limits of such a cross-platform comparison), with similar expression levels of LmA/C and various mES cell markers (Supplementary Fig S6E and F). The mRNA-seq data also confirmed a > 100 and > 30-fold reduction in LmAB1 and LmB2 expression, respectively, in dKO cells compared to wt cells (Supplementary Fig S6E and F). Cloning, production, and transduction of lentiviral vectors (LVs) are described in the Supplementary Methods. DamID was performed as previously described [17]. Briefly, mES cells were plated and transduced the following day with LV expressing the Dam-fusion protein of interest. gDNA was collected 2 days after transduction, and it was processed and hybridized to NimbleGen genomic arrays as previously described [17]. The DamID data used in Figs 1–4 are the average of two independent biological replicates for each condition, which in a previous study [17] was sufficient to detect changes in NL interactions at hundreds of sites throughout the genome.

Immunofluorescence

mES cells were plated on matrigel (BD Bioscience, 356231)-coated glass coverslips. The following day, cells were fixed, stained, and analyzed as described in Supplementary Methods.
Data analysis

All genome-wide analyses were performed using the R statistical software. DamID data normalization was done as previously described [17]. LADs and iLADs were defined as described previously [1]. Separately, we defined cLAD, fLAD, iLAD, and cLAD status for each array probe using a hidden Markov model approach as described [20], but using the Emd DamID profile obtained in wt mES cells in addition to the four cell types used previously. Facultative probes were subdivided in facultative iLAD (iLAD) and facultative LAD (fLAD) if identified as iLAD or LAD, respectively, in Emd DamID performed in wt mES cells.

The concordance score is defined as: the amount of sequence overlap between LAD or iLAD domains of two cell types (domain defined as in [1]) or the percentage of all calls in agreement (e.g., LAD vs. LAD or inter-LAD vs. inter-LAD) between two cell types (calls defined according to the HMM as in [20]).

DamID values per gene were calculated by averaging the DamID scores of all probes overlapping with the gene. As a consequence, the comparison of gene expression levels to DamID scores (Fig 2) focuses on 14,157 genes. A previously reported statistical test to identify genes with significant changes in DamID signals [17] was applied using default settings (minimum gene size 5 probes; FDR = 5%). For other analyses of gene expression where the position of genes relative to LADs is considered (but not the DamID scores of the genes), all 37,991 genes are included.

Data availability

All data are available from the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE62685) under accession number GSE62685.

Primary data


Supplementary information for this article is available online: http://embor.embopress.org

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Author contributions

MA performed the experiments and analyzed the data; BvS and MA conceived the study, designed the experiments, and wrote the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

References


