Lamin A/C-promoter interactions specify chromatin state–dependent transcription outcomes

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The nuclear lamina is implicated in the organization of the eukaryotic nucleus. Association of nuclear lamins with the genome occurs through large chromatin domains including mostly, but not exclusively, repressed genes. How lamin interactions with regulatory elements modulate gene expression in different cellular contexts is unknown. We show here that in human adipose tissue stem cells, lamin A/C interacts with distinct spatially restricted subpromoter regions, both within and outside peripheral and intra-nuclear lamin-rich domains. These localized interactions are associated with distinct transcriptional outcomes in a manner dependent on local chromatin modifications. Down-regulation of lamin A/C leads to dissociation of lamin A/C from promoters and remodels repressive and permissive histone modifications by enhancing transcriptional permissiveness, but is not sufficient to elicit gene activation. Adipogenic differentiation resets a large number of lamin-genome associations globally and at subpromoter levels and redefines associated transcription outputs. We propose that lamin A/C acts as a modulator of local gene expression outcome through interaction with adjustable sites on promoters, and that these position-dependent transcriptional readouts may be reset upon differentiation.

[Supplemental material is available for this article.]
We examine here the impact of LMNA association with promoter regions on the expression outcome of all genes, that is, genes not necessarily localized in peripheral LADs. We show in human adipose stem cells (ASCs) interaction of LMNA with thousands of promoters, on spatially restricted domains, and in a manner that specifies chromatin-dependent local transcription outcomes. LMNA influences chromatin modifications at sites it interacts with and beyond. Adipogenic differentiation remodels LMNA-promoter interactions and resets associated gene expression. Our results suggest a model of a variable modulatory function of promoter-associated LMNA on local chromatin environments and transcription outcomes.

Results

Promoters associate with LMNA in primary adipose stem cells

We mapped in human ASCs the occupancy of A-type lamins (LMNA) on promoters using chromatin immunoprecipitation (ChIP) and array hybridization. Lamins A and C are localized at the nuclear periphery and in the nuclear interior in ASCs (Fig. 1A) and cofractionate with chromatin prepared by formaldehyde cross-linking and sonication in a low-salt/EDTA/Triton X-100-based buffer (Fig. 1B; Supplemental Fig. 1A). Lamins A and C were immunoprecipitated (Fig. 1B) and promoter occupancy queried by hybridization of ChIP DNA to microarrays tiling 4 kb around the TSS of 17,790 RefSeq genes (Pearson correlation of log2 IP/input ratios $r = 0.975$ between ChIP replicates) (Fig. 1C).

Using a “model-based analysis of two-color arrays” (MA2C) algorithm with $P < 0.01$ for LMNA peak calling, we identify 4827 genes with a promoter enriched in LMNA, which represents 27% of RefSeq genes (Fig. 1D; see Supplemental Table 1 for a list of LMNA-enriched genes). Knockdown of lamins A and C (LMNA KD) with an shRNA to the $LMNA$ gene (Supplemental Fig. 1B) abolishes or markedly reduces LMNA occupancy (Fig. 1C,D), indicating specificity of the immunoprecipitation. Validation of ChIP-chip data by ChIP and quantitative (q)PCR for LMNA-enriched and depleted regions shows high concordance between data sets (Supplemental Fig. 1C,D). Gene ontology (GO) analysis of LMNA-associated genes identifies genes pertaining to development and differentiation, signaling, and immune response functions (Supplemental Table 2), suggesting that LMNA may be involved in the regulation of developmental functions.

Although nuclear lamins can bind DNA in vitro, LMNA-promoter interactions determined by ChIP provide no information on whether LMNA binds directly to promoter sequences or whether interactions are mediated by DNA-binding factors. Nonetheless, a search for DNA motifs enriched among LMNA-associated sequences identifies a recurrent GAGA motif underlying 21% of LMNA peaks ($E$-value $= 1.17 \times 10^{-15}$, Bonferroni-corrected $P$-value) (Fig. 1E), which is similar to the extended GAGA motif found in mouse lamin associating sequences (LASs) (Zullo et al. 2012). This extended mouse GAGA motif is bound by the transcription repressor CKROX, which mediates targeting of loci to the lamina (Zullo et al. 2012). The $ZBTB7B$ gene encoding CKROX is expressed in human ASCs (Boquest et al. 2005), suggesting a similar mechanism of association of LMNA with GAGA motifs in mouse and human cells. We also find an A/T-rich motif underlying 62% of LMNA peaks ($E$-value $= 1.15 \times 10^{-26}$) (Fig. 1E), in line with the A/T-richness of LADs (Meuleman et al. 2012). A/T-rich motifs match those of transcription factors such as ARID3A. ARID3A, also expressed in ASCs (Boquest et al. 2005), has been shown to tether loci to the nuclear envelope (Webb et al. 2011) and may confer lineage restriction in somatic stem cells (An et al. 2010), potentially linking LMNA-promoter interactions to developmental regulation.

Evidence for peripheral and intra-nuclear lamin-rich domains

We next analyzed the genomic distribution of LMNA-interacting promoters. First, we determined whether genes enriched in LMNA
cluster or are uniformly distributed in the genome. To this end, we computed the number of LMNA-enriched genes in a 31-gene sliding window across the genome and attributed a gene to a “lamin-rich domain” (LRD) if at least 10 of the closest 30 genes were enriched in LMNA (Fig. 2A). This cut-off was significant in a Monte Carlo simulation with a null hypothesis of uniform distribution of LMNA (p < 10^{-6}, two-sided t-test), indicating that LMNA-enriched genes are nonrandomly distributed. LRDs are identified on nearly all chromosomes (Fig. 2A), consistent with a localization of clustered LMNA-interacting genes that extends beyond the nuclear periphery.

An overlap analysis of LRDs with LADs previously identified in human fibroblasts (Guelen et al. 2008) shows that 50% (n = 2113) of 4202 RefSeq genes found in LADs are enriched in LMNA in our study. Conversely, nearly 50% of LMNA-bound genes are located in LADs. Thus, the localization within LADs (Guelen et al. 2008) of LMNA-associated genes identified by ChiP is highly significant (p = 2.2 × 10^{-16}, Pearson’s χ² test). Our analysis provides, therefore, independent evidence for a clustering of LMNA-associated genes into wider lamin-enriched domains (Kind and van Steensel 2010).

Secondly, we examined the intra-nuclear localization of LMNA-enriched genes by 3D immunofluorescence and fluorescence in situ hybridization (immuno-FISH). We analyzed nine loci, seven of which are LMNA targets identified by ChiP and located within or outside LRDs (DEFA3, SCN10A, TCN1, ACTL7A, DNAL4, NANOG, ATPSE2), and two are not (HOXB9, LDHB) (Supplemental Fig. 2A,B). Several configurations were identified among these loci: (1) Genes strongly enriched in LMNA and found in an LRD display localization both at the nuclear periphery (defined as ≤0.5 μm from peripheral lamin A/C immune-labeling) and in the nuclear interior (DEFA3, TCN1), or intra-nuclear localization only (ACTL7A) (Fig. 2B,C; Supplemental Fig. 2C,D). This suggests that LRDs exist not only at the nuclear periphery but also in the nuclear interior. (2) A gene enriched in LMNA and close to (<500 kb) an LRD (SCN10A) also shows more peripheral positioning (Fig. 2B,C); thus, localization in an LRD is not required for peri-nuclear localization. (3) Genes enriched in LMNA away from an LRD (>500 kb) only or predominantly display intra-nuclear localization (DNAL4, NANOG), or both peripheral and intra-nuclear localization (ATPSE2) (Supplemental Fig. 2C,D). So, LMNA enrichment does not always entail peripheral localization. (4) A gene not bound by LMNA but localized in a “weak” LRD (LDHB) can also be peripheral and intra-nuclear (Supplemental Fig. 2C,D). Thus, absence of LMNA enrichment is compatible with localization at the nuclear periphery. (5) Lastly, a gene not enriched in LMNA and outside an LRD (HOXB9) is found only in the nuclear interior (Fig. 2B,C). We conclude that LMNA enrichment does not systematically imply peri-nuclear localization and that loci enriched in LMNA within or outside an LRD can be found at the nuclear periphery as well as in the nuclear interior. Moreover, our expression data indicate that these genes, except LDHB, are not expressed in ASCs; thus, for the LMNA-bound genes studied here (DEFA3, TCN1, DNAL4, NANOG, ATPSE2), expression status is not necessarily related to peri-nuclear positioning.

**LMNA association with spatially restricted promoter subregions is linked to distinct transcription outcomes**

Given the wide genome distribution of LRDs, we next assessed the impact of LMNA enrichment on expression of genes associated with, or in proximity to, LMNA. We also mapped transcriptionally permissive (H3K4me3) and repressive (H3K9me3, H3K27me3) histone modifications to provide a chromatin context to LMNA enrichment. LMNA-bound genes are weakly or not expressed (p < 10^{-6}, Wilcoxon rank sum test compared to RefSeq genes) (Fig. 3A), and accordingly, they are enriched in H3K9me3 and/or H3K27me3 (p < 10^{-6}, χ² with Yates’ correction) (Fig. 3B). However, not all genes associated with LMNA are repressed, as illustrated by the wide range of expression levels of genes co-enriched in H3K4me3 in the absence of repressive marks (Fig. 3A). Thus, a fraction of LMNA-bound genes escapes repression, suggesting that LMNA association is by itself not conducive of a repressive state.

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**Figure 2.** 3D immuno-FISH unveils peripheral and intra-nuclear LMNA-associated loci. (A) LMNA enrichment in sliding 31-gene windows (x ± 31) across chromosomes. Scale shows the numbers of LMNA-enriched genes in the window. (B) Distribution of FISH signals from LMNA-enriched (DEFA3, SCN10A, DNAL4) and nonenriched (HOXB9) loci relative to peripheral LMNA (n = 30 loci per gene). (C) 3D immuno-FISH (green; arrows) of loci shown in B relative to peripheral LMNA labeling (red). DNA is stained with DAPI. Scale bars, 5 μm.
We next evaluated the impact of LMNA enrichment in a genomic “neighborhood” (defined as 1-Mb bins throughout the genome) on the expression of genes bound by LMNA or not bound by LMNA. We find that proportions of expressed LMNA-bound genes are negatively correlated with LMNA enrichment in 1-Mb bins (Supplemental Fig. 3). However, proportions of expressed genes not bound by LMNA do not correlate with LMNA enrichment of neighboring LMNA-bound genes (Supplemental Fig. 3). Thus, localization of a gene in an area rich in LMNA does not imply transcriptional repression, particularly if the gene is itself not tethered by LMNA.

Our findings that not all LMNA- and H3K4me3-marked genes are repressed suggest that an additional component may modulate expression of these genes, such as positioning of LMNA relative to the TSS. Mapping LMNA peak density across promoters (see Methods) shows that LMNA most frequently occupies “upstream-distal” (1.4 kb and upstream) and “TSS/downstream” regions, and, less frequently, “upstream-proximal” regions (Fig. 3C; Supplemental Table 1). Moreover, median LMNA peak size within promoter regions is relatively short (1.5 ± 0.47 kb) (data not shown), suggesting that LMNA tends to associate with discrete and spatially restricted subregions.

A gene expression heat map corresponding to LMNA peak positioning, generated from gene expression ratios across promoter regions, shows that LMNA peak position corresponds to a distinct gene expression output. LMNA association at the TSS or upstream-proximal of the TSS coincides with essentially no expression (Fig. 3D). In contrast, upstream-distal association is compatible with gene expression (Fig. 3D), implying that these loci are marked by H3K4me3. Indeed, 2D maps of LMNA and H4K3me3 enrichment show that the most frequently expressed LMNA- and H3K4me3-enriched genes harbor H3K4me3 at the TSS/downstream and LMNA upstream-distal (Fig. 3E). However, genes with both LMNA and H3K4me3 at the TSS/downstream are not expressed. Altogether, these data suggest that LMNA association with promoters is overall spatially restricted and linked to distinct transcription outcomes in a manner dependent on local chromatin organization.

LMNA influences chromatin modifications at and beyond loci with which it interacts

Down-regulation of LMNA results in dissociation of LMNA from most promoters (Fig. 1C). Strikingly however, genes that lose LMNA are not transcriptionally activated (Fig. 4A), despite the low proportion marked by H3K9me3 or H3K27me3 (Fig. 4B). In fact,
overall expression levels are not profoundly altered by loss of LMNA binding (cf. Figs. 3A, 4A, RefSeq), even though the extent of enrichment in repressive modifications globally decreases \( (P < 10^{-4}) \) (Fig. 4C). In contrast, LMNA KD results in an increase in the proportion of genes enriched in H3K4me3 (Fig. 4C); this implies that, in addition to LMNA-bound genes, LMNA KD affects H3K4me3 on genes initially not associated with LMNA. Thus, LMNA down-regulation impacts chromatin organization by globally reducing the incidence of repressive marks and increasing the incidence of permissive marks. This occurs at loci beyond those that LMNA interacts with, mostly without affecting overall gene expression patterns.

LMNA down-regulation also affects chromatin organization at the sub-promoter level. Promoters retaining LMNA show preferential association upstream-distal of the TSS and much less frequently at the TSS/downstream (Fig. 4D). H3K4me3 distribution is also affected, with a marked increase upstream of the TSS, not only on promoters that retain LMNA but also on all H3K4me3 promoters (Fig. 4E). In contrast, profiles of H3K9me3 and H3K27me3 enrichment are not affected by LMNA loss (Supplemental Fig. 4). We conclude that alteration in the nuclear lamina by down-regulation of LMNA results in a remodeling of chromatin on promoters, particularly H3K4me3 marking.

**Adipogenic differentiation remodels promoter interactions with LMNA and associated transcription outcomes**

Our findings suggest thus far a view of LMNA positioning on a promoter linked to a given transcription outcome. To further test this possibility, we mapped LMNA-promoter interactions in ASCs differentiated into adipocytes. Adipogenic differentiation leads to a marked nuclear compaction (Fig. 5A, arrows), which is likely to affect lamin-genome interactions, particularly since levels of lamins A and C are similar (Supplemental Fig. 5A). Accordingly, adipogenic differentiation remodels LMNA-promoter interactions at several levels. (1) Loss and gain of LMNA occurs on, respectively, ~4000 and 2000 genes (Fig. 5B; Supplemental Table 3). (2) LMNA loss affects entire LRDs; however, no de novo LRDs are detected in adipocytes (Supplemental Fig. 5B), suggesting that interactions with LMNA in adipocytes occur on stand-alone or weakly clustered loci. (3) Nearly 800 genes remain associated with LMNA (Fig. 5B). These genes are notably involved in bone morphogenic protein signaling and ectodermal and endodermal differentiation pathways (Supplemental Table S3), suggesting a selective retention of LMNA on developmentally important promoters whose function is no longer required in adipocytes. (4) LMNA-promoter interactions are also remodeled at the subpromoter level, with LMNA mainly occupying proximal promoter and TSS regions (cf.
Figs. 5C and 3C). It appears, therefore, that remodeling of LMNA-promoter interactions at the genome-wide and subpromoter levels is part of the adipogenic program. Strikingly, ~83% of genes losing or gaining LMNA after differentiation retain their expression status of undifferentiated cells (Fig. 5D); thus, association with, and disengagement from, LMNA are largely uncoupled from changes in gene expression. Yet, ~17% of genes escape this uncoupling and display expression changes as they lose or gain LMNA; among these, gain and loss of LMNA is, respectively, associated with transcriptional down- and up-regulation (Fig. 5D). These genes notably include genes essential for induction of adipogenesis (see below). Maintenance of expression status of genes that gain or lose LMNA after differentiation suggests that the relationship we identified between LMNA positioning and expression outcome may be altered. Indeed, in adipocytes, LMNA peak enrichment upstream-proximal or at the TSS/downstream is compatible with expression (Fig. 5E). Thus, in these cells, LMNA occupancy at the TSS or downstream appears to be no longer inhibitory for gene expression.

Collectively, these observations indicate that adipogenic differentiation results in a remodeling of subnuclear architecture. This is manifested by a readjustment of LMNA-promoter interactions at the genome-wide and subpromoter levels and by a resetting of the expression capacity of a subset of genes that interact with LMNA in adipocytes.

**LMNA-promoter interaction dynamics on adipogenic control genes**

A feature of LMNA-promoter interaction dynamics upon adipogenic differentiation is that it affects genes essential for induction of the adipogenic program. For instance, LMNA dissociates from the P2 promoter of the adipogenic master regulator PPARγ and from the FABP locus (Fig. 5F). This disengagement from, LMNA may prime their transcriptional induction, in line with the activation of these genes in adipocytes (Mikkelsen et al. 2010). Not all regulators of adipogenesis bind LMNA in ASCs, however (e.g., CEBPβ, SREBF1 [also known as SREBF2]), suggesting that these may already be “primed” for activation. Additionally, genes essential for differentiation into nonadipogenic lineages are bound by LMNA in ASCs, and strikingly, most retain LMNA in adipocytes (Fig. 5G; Supplemental Fig. 5C). This promoter drives expression of the RUNX2 transcription factor to promote osteogenesis over adipogenesis (Long 2011); thus, LMNA interaction may reflect a loss of RUNX2 activation potential in adipocytes. Similarly, genes linked to pluripotency, such as POU5F1, NANOG, KLF4 and SOX2, associate with LMNA in ASCs and adipocytes (Supplemental Fig. 5D), suggesting that long-term repression may be associated with tethering to LMNA. These observations suggest that LMNA association of promoters important for lineage commitment is under developmental control and regulated in a locus-specific manner.

**Discussion**

**Peripheral and intra-nuclear lamin-rich domains**

We identify LMNA-associated promoters for over 25% of RefSeq genes in human adipose stem cells. A feature of lamins A and C is the existence of a nucleoplasmic detergent-soluble pool (Kolb et al. 2011), in addition to a peripheral pool, because they lack C-terminal farnesyllation mediating attachment to the inner nuclear membrane (Burke and Stewart 2013). Under conditions used in our study, ChIP significantly enriches lamin C over lamin A, likely as a result of greater solubility (Kolb et al. 2011). This enrichment may yield an underrepresentation of loci associated with peripheral, less soluble A-type lamins. Nonetheless, ChIP and 3D immuno-FISH data demonstrate interactions of LMNA with promoters and are consistent with interactions taking place at the nuclear periphery and in the nuclear interior. In fact, some LMNA-enriched loci are found exclusively in the nuclear interior (e.g., ACTL7A, DNAL4), and LMNA-interacting loci with alleles at the nuclear periphery also display alleles in the interior (e.g., DEFA3, SCN10A, TCN1, ATSEP2).

Imaging of single cells based on tagging of sequences in proximity to the lamina reveals that only 38% of LADs are at the nuclear periphery (Kind et al. 2013). Similarly, in our study, only ~30% of alleles of loci associated with LMNA and located within LRDs are detected at the nuclear periphery. This implies that a significant proportion of LRDs (or LADs) localize in the nuclear interior and that associations with the periphery are transient (Kind et al. 2013). One may speculate that some of the loci detected by FISH close to the periphery (e.g., DEFA3, SCN10A, TCN1, ACTL7A) could, at some other time point, locate at the periphery. In contrast, loci localized in the nuclear interior (e.g., DNAL4, NANOG) and associated with LMNA may be anchored to intra-nuclear LRDs (or LADs) because they may be too far from the periphery. It will be important to identify DNA sequences associated with nucleoplasmic and peripheral lamins, address whether lamin association with the genome is direct or mediated by transcription factors, and determine whether peripheral and intra-nuclear LRDs are functionally different.

**Association of LMNA with distinct promoter subregions**

LMNA interactions often appear to be confined to promoter subregions rather than to entire promoter regions. This suggests that even when LMNA-interacting promoters are contiguous, as within LRDs, interactions with the lamina may occur through multiple spatially restricted regions. This view is consistent with the interaction of nuclear lamins with short LASs (in the kb range) able to target sequences to the nuclear periphery (Zullo et al. 2012) and with focal genomic domains associating with the nuclear membrane in Caenorhabditis elegans (Ikegami et al. 2010). LMNA may also be anchored directly or indirectly with loci outside LRDs through targeting DNA motifs (Ahmed et al. 2010; Brickner et al. 2012; Zullo et al. 2012). Of note, A/T-rich motifs may be contained within so-called scaffold/matrix attachment regions (S/MARs) that have previously been shown to directly bind lamins in vitro (Luden et al. 1994; Baricheva et al. 1996; Zhao et al. 1996).

LMNA can be differentially positioned on promoters, and our data suggest that this is linked to distinct transcription outputs of the corresponding genes. Upstream-proximal and TSS/downstream LMNA interaction correlates with gene inactivity in undifferentiated ASCs, regardless of histone methylation mark. Repression of genes with LMNA and H3K4me3 at the TSS may be due to inhibition of nucleosome turnover (Henikoff 2008), facilitated recruitment of repressive chromatin modifiers, or inhibition of transcription by blocking access of the TSS for RNA polymerase II. Expansion of H3K4me3 upstream of the TSS on promoters after LMNA loss supports a view of LMNA hindering the targeting of factors to the TSS. RNA polymerase II and H3K4me2 have been observed at LAD borders, and some genes within LADs are transcribed (Guelen et al. 2008). Our data show co-enrichment in LMNA and H3K4me3 on
genes that are expressed in undifferentiated ASCs, when these marks do not spatially coincide. We suggest, therefore, that lamin-promoter interactions, per se, do not have a causative role on gene repression but may be able to modulate transcription in a manner dependent on local chromatin marks.

The significance of LMNA-promoter interaction may also be cell type–dependent. LMNA association with the TSS region in ASCs is frequently coupled to transcriptional repression, whereas in adipocytes, it is more frequently coupled to expression. How may we reconcile this LMNA association with gene expression in adipocytes? Interestingly, although LMNA association with the TSS is still detected in adipocytes, the density of LMNA peaks in this subregion is significantly reduced, highlighting a dose-dependent effect of lamin binding. In addition, the majority of expressed genes bound to LMNA in adipocytes is already expressed in ASCs and thus may already be in a permissive state compatible with LMNA-TSS interaction. Upon differentiation, promoter-associated LMNA may also interact with a new set of regulatory partners enabling transcription or with chromatin remodelers with dual activator or repressor functions depending on subunit composition (Euskirchen et al. 2011).

LMNA neighborhoods are, per se, not transcriptionally repressive
LMNA enrichment level in a genomic “neighborhood” (which may contain one or several LRDs) correlates with transcriptional repression in that neighborhood. However, within a LMNA neighborhood, expression of genes not bound by LMNA is not affected by LMNA level in that neighborhood. Thus, a LMNA neighborhood is, in itself, not conducive of transcriptional repression. Accordingly, depletion of the H3K9 histone methyltransferase EHMT2 (also known as G9A) relieves repression of genes in the vicinity of the lamina without affecting their perinuclear location (Yokochi et al. 2009). So, LMNA may participate in genomic recruitment to, or stabilization within, a repressive compartment and not confer repression per se. This view is consistent with SET-domain methyltransferases mediating recruitment and silencing of loci at the lamina in C. elegans (Towbin et al. 2012) and in mammalian cells (Kind et al. 2013) and with peripheral targeting of the hunchback locus in Drosophila neuroblasts after its inactivation (Kohwi et al. 2013). In addition, through chromatin looping in the nuclear space (McCord et al. 2012), intra-nuclear lamins may cluster linearly distant loci, and chromosomal interactions (Kind and van Steensel 2010) may bring a lamin-deprived locus to the nuclear periphery.

Cell lineage–specific interaction of LMNA with promoters
Interactions of the genome with the nuclear lamina are dynamic (Kind et al. 2013) and developmentally modulated (Pickersgill et al. 2006; Meister et al. 2010; Peric-Hupkes et al. 2010). Adipogenic differentiation relocates promoters from and to LMNA. Adipogenic differentiation requires a proliferation step during which mitotic disassembly and reassembly of the lamina likely provokes a redistribution of lamin-interacting domains (Kind et al. 2013). Nonetheless, the conservation of some LRDS after differentiation suggests the retention of an LRD “memory” on mitotic chromosomes, which would be consistent with a model of “constitutive LADs” maintained between cell types (Meuleman et al. 2012). We detect a loss of LMNA from adipogenic promoters that are activated after induction of differentiation. This contrasts with promoters important for pluripotency or differentiation into nonadipogenic lineages which retain LMNA. We propose a model of cell type– and lineage-specific association of loci with LMNA and disengagement of these loci from LMNA in a lineage-specific manner (Fig. 6). This model is consistent with a locus-specific developmental regulation of lamin interactions with promoters important for differentiation and lineage commitment.

Methods
Adipose stem cells
ASCs were cultured as a pool from three donors and used at passages 8–9 as described (Boquest et al. 2005). Adipogenic differentiation was induced for 21 d with 1-methyl-3-isobutylxanthine, dexamethasone, insulin, and indomethacin, and lipid-stained with Oil red-O (Boquest et al. 2005).

LMNA silencing by shRNA
An shRNA targeting the 5′ UTR of LMNA transcripts and a control shRNA were designed with the following sequences (LMNA-specific and control sequences are underscored): shRNA-LMNA – 5′-tgctggtaagagaagctgctcggagtgcctactgcctcgga-3′; control shRNA – 5′-tgctggtaagagaagctgctcggagtgcctactgcctggga-3′. Sequences were cloned into BamHI/Mlu sites to a Lentiviral pGIPZ vector (ThermoFisher Scientific). Lentiviruses were produced in 293T cells. Cells were transduced and cultured with 1.5 μg/mL puromycin for 8 d to select shRNA-expressing cells, and maintained in culture with 0.1 μg/mL puromycin. shRNA expression was induced with 1 μg/mL doxycyclin in the presence of 0.1 μg/mL puromycin for 6 d.

Expression microarrays
Three replicate array experiments were done for ASCs, LMNA-KD ASCs, and adipocytes using Illumina HumanHT-12 v4 expression BeadChip arrays. The Gene Expression module from Illumina GenomeStudio was used to compute a detection P-value for each probe. Signal intensities between replicates were normalized using the lumi package (Du et al. 2008) and replicate signals combined by using median signal intensity. A gene was considered expressed when probe intensities were significantly stronger (P < 0.01) than background signals. Adipocyte gene expression data have been
published earlier (Sørensen et al. 2010) and are available in NCBI GEO with accession number GSE19773.

**Chromatin immunoprecipitation**

ChIP of modified histones (H3K9me3, Diagenode, pAb-056-050; H3K27me3, Millipore, 07-449; H3K4me3, Abcam, Ab8580) and processing for array hybridization were as described (Sørensen et al. 2010). For LMNA ChIP, cells were cross-linked for 8 min with 1% formaldehyde and chromatin prepared by lysis (5 min on ice) and sonication into 200- to 400-bp fragments for 4 × 10 min on a Bioruptor (Diagenode) in lysis buffer (10 mM Tris-HCl, pH 7.5, 10 mM KCl, 2 mM EDTA, 1% Triton X-100, and protease inhibitors). Chromatin (100 μl at 5 A260 units) was incubated on a rotator overnight at 4°C with 2.4 μg anti-lamin A/C antibody (Santa Cruz, sc-7292) coupled to Dynabeads Protein A (Invitrogen). ChIP material was washed three times in lysis buffer, the cross-link was reversed, and DNA was purified by phenol-chloroform isoamyl alcohol extraction, RNase-treated, and processed for microarray hybridization (Sørensen et al. 2010). For ChIP-qPCR, purified LMNA ChIP DNA was eluted in 30 μl H2O and 2.5 μl used as a template for qPCR. Duplicate qPCR were run on a MyiQ Real-time machine with SYBR Green (BioRad) using primers listed in Supplemental Table 4. PCR conditions were 95°C for 3 min and 40 cycles of 95°C for 30 sec, 60°C for 30 sec, and 72°C for 30 sec.

**Array hybridization and data preprocessing**

ChIP and input DNA fragments were, respectively, labeled with Cy5 and Cy3 and hybridized to human HG18 RefSeq Promoter arrays (Roche) tiling 3.2 kb upstream of and 0.8 kb downstream from the TSS of 17,790 RefSeq genes. A promoter region was defined as the interval spanning –2200 bp to +500 bp of a TSS. Each probe was assigned an MA2C score (Song et al. 2007) to reflect normalized and window-averaged log2 ChIP/Input ratios, with peak calling at P < 0.01 using a 1-kb sliding window. Peaks within 500 bp of each other were merged.

**LMNA-enriched domains and bins**

Each chromosome was viewed as a sequence of genes, ordered by the position of their TSS. A sliding window of 31 genes (“size 31”) with step size 1 was used to count genes enriched in LMNA in the vicinity (+/- 15 genes) of the middle gene at each iteration. Repeating occurrences of the same gene were collapsed. We performed 10,000 trials in a Monte Carlo simulation to test whether LMNA enrichment clusters at the gene level. In each trial, the number of LMNA-enriched genes per chromosome was kept constant and identical to our experimental results, but the order of genes was shuffled. The sample set of the number of LMNA-enriched genes in the vicinity of more than 10 LMNA-enriched genes was fitted to a normal distribution.

**Motif discovery**

Motif discovery within sequences underlying LMNA peaks was done using the RSAT peak-motifs pipeline (Thomas-Chollier et al. 2011) with the oligo-analysis module.

**Promoter density maps**

The number of transcripts with a LMNA or modified histone peak at a given position as a function of distance from the TSS determined whether a mark had a nonuniform distribution of peaks in promoter regions, and if so, where. This was computed by the following equation:

\[
f(x) = \frac{\sum_{i=1}^{n} T_{ix}}{\sum_{i=1}^{n} x_{i}}
\]

where \( T_{ix} \) equals 1 if transcript \( i \) has a peak of a given type at position \( x \) relative to the TSS and equals 0 otherwise.

The relationship between relative peak position and transcription can similarly be expressed as the ratio of expressed transcripts as a function of peak position relative to the TSS:

\[
f(x, y) = \frac{\sum_{i=1}^{n} T_{i,xy} E_{i}}{\sum_{i=1}^{n} T_{i,xy}}
\]

where \( E_{i} \) equals 1 if transcript \( i \) is transcribed and equals 0 otherwise. \( T_{i,xy} \) is as defined above.

It is thus possible to examine the subset of promoters enriched by at least two marks to determine whether interplay between the two marks alters the typical distribution of peaks for at least one of the marks. To gain information on how peaks of two marks are positioned on the same promoter, analysis was extended into two dimensions as follows:

\[
f(x, y) = \frac{\sum_{i=1}^{n} T_{i,xy} E_{i}}{\sum_{i=1}^{n} T_{i,xy}}
\]

where \( x \) is the position of \( X \) peaks relative to the TSS, \( y \) is the position of \( Y \) peaks relative to the TSS, and \( T_{i,xy} \) equals 1 if transcript \( i \) has a peak of type \( X \) at position \( x \) and a peak of type \( Y \) at position \( y \) relative to the TSS.

The transcription ratio as a function of peak position relative to the TSS can similarly be extended to two dimensions:

\[
f(x, y) = \frac{\sum_{i=1}^{n} T_{i,xy} E_{i}}{\sum_{i=1}^{n} T_{i,xy}}
\]

where \( T_{i,xy} \) and \( E_{i} \) are as above.

**Immunolabeling and fluorescence in situ hybridization**

ASCs cultured on glass slides were washed in PBS before fixation in 3% paraformaldehyde/PBS for 15 min. After three washes in PBS, cells were permeabilized for 30 min in 0.01% Tween/0.1% Triton X-100/2% BSA/PBS. Cells were incubated with anti-lamin A/C antibodies (1:400; Santa Cruz sc-7292) for 1 h in 0.01% Tween/2% BSA/PBS, washed 3× in PBS, and incubated with anti-mouse Cy3-conjugated antibodies (1:200; Jackson ImmunoResearch) for 1 h in 0.01% Tween/2% BSA/PBS. Cells were washed and refixed in 4% paraformaldehyde/PBS for 10 min for FISH. Fosmid clones (Supplemental Table S) from BACPAC (http://bapac.chori.org/) were prepared and nick-translated with digoxigenin-11-dUTP. The HOXB9 clone was described previously (di Pietro et al. 2012). One hundred to 150 ng digoxigenin-labeled probes were used per slide with 8–12 μg human Cot-1 DNA and 12 μg herring sperm DNA. Slides were denatured in 70% (vol/vol) formamide/2×SSC at 80°C for 20 min. Probes were denatured at 80°C for 5 min, reannealed with Cot1 DNA for 15 min at 37°C, and hybridized to the slides overnight. Washes and detection were as described (Eskeland et al. 2010). DNA was stained with 0.5 μg/ml 4,6-diamidino-2-phenylindole (DAPI), mounted in Vectashield (Vector), and examined with a 100× oil objective (NA 1.4) on a PersonalDV (Delta Vision) wide-field imaging station (Applied Precision, auxiliary magnification of 1.6 and optical step size of 0.2 μm through the whole nucleus). Images were deconvoluted with the integrated software (softWoRx v04.1.2).
Image analysis
For measurements of FISH spots and of their distance to peripheral LMNA, we used ImageJ (http://imagej.nih.gov/ij/) (Schneider et al. 2012) with the Sync_Measure_3D plug-in (http://rsb.info.nih.gov/ij/), default threshold and line tool. Measurement of the shortest distance of each of the two FISH signals to the nuclear boundary in three dimensions was done by defining the outline of the LMNA stain in each frame of the z-stack and measuring the distance in the stack with the highest level of intensity from the center of each fluorescent spot to the inner boundary of LMNA (on the x-y-axis). Nuclei with FISH signals near the top or bottom in the z-axis were not evaluated due to fuzziness of FISH signals and LMNA immunolabeling, making measurements inaccurate.

Immunofluorescence and immunoblotting
Immunofluorescence labeling of lamin A/C was done as described (Stein and Collas 2001) using anti-lamin A/C antibodies (Santa Cruz, sc-7292). Immunoblotting was done as described (Duband-Goulet et al. 2011) using antibodies to lamin A/C (1:2000 dilution; Santa Cruz, sc-7292), lamin B1 (1:1000; Santa Cruz, sc-6216), and γ-tubulin (1:10,000; Sigma, T5326), and horseradish peroxidase-conjugated secondary antibodies.

Data access
LMNA ChIP, histone ChIP, and ASC expression data have been deposited in the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE42560.

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