Mapping the micro-proteome of the nuclear lamina and lamina-associated domains

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The nuclear lamina is a proteinaceous network of filaments that provide both structural and gene regulatory functions by tethering proteins and large domains of DNA, the so-called lamina-associated domains (LADs), to the periphery of the nucleus. LADs are a large fraction of the mammalian genome that are repressed, in part, by their association to the nuclear periphery. The genesis and maintenance of LADs is poorly understood as are the proteins that participate in these functions. In an effort to identify proteins that reside at the nuclear periphery and potentially interact with LADs, we have taken a two-pronged approach. First, we have undertaken an interactome analysis of the inner nuclear membrane bound LAP2β to further characterize the nuclear lamina proteome. To accomplish this, we have leveraged the BioID system, which previously has been successfully used to characterize the nuclear lamina proteome. Second, we have established a system to identify proteins that bind to LADs by developing a chromatin-directed BioID system. We combined the BioID system with the m6A-tracer system which binds to LADs in live cells to identify both LAD proximal and nuclear lamina proteins. In combining these datasets, we have further characterized the protein network at the nuclear lamina, identified putative LAD proximal proteins and found several proteins that appear to interface with both micro-proteomes. Importantly, several proteins essential for LAD function, including heterochromatin regulating proteins related to H3K9 methylation, were identified in this study.

DOI 10.26508/lsa.202000774 | Received 12 May 2020 | Revised 8 February 2021 | Accepted 9 February 2021 | Published online 23 March 2021

Introduction

DNA and proteins are highly organized within the eukaryotic cell nucleus. Sequestration of proteins into nuclear sub-domains and the higher order organization of chromatin itself have been implicated in the regulation of the genome (1, 2, 3, 4, 5, 6). Association of chromatin with the nuclear periphery, in particular, has been implicated in gene regulation, specifically correlating with repression of developmentally regulated loci (4, 7, 8). DamID (DNA Adenine Methyltransferase Identification), a genome-wide technique to identify nuclear lamina-proximal chromatin, has allowed the identification of lamina-associated domains (LADs) (9). These 100 kb to megabase (Mb) sized domains are enriched for genes that are transcriptionally silent and enriched in histone modifications indicative of facultative heterochromatin, such as histone H3 lysine 9 di/tri-methylation (H3K9me2/3) and histone H3 lysine 27 trimethylation (H3K27me3) (10, 11, 12, 13). Moreover, recent studies have highlighted that both H3K9me2/3 and H3K27me3 are involved in LAD organization (12, 13, 14, 15, 16). LADs represent a large fraction of the genome (30–50%, depending upon the cell type) and are highly correlated with the so-called heterochromatic “B-compartment,” as identified by chromatin conformation capture assays (HiC) (13, 17 Preprint, 18 Preprint). Given their importance to cellular function and identity, it is important to understand how these large domains of heterochromatin are regulated, maintained, and formed to understand global genome regulation and organization. An important element of understanding LAD organization and function is identifying which proteins are present at LADs and the nuclear lamina.

In addition to H3K27me3 and H3K9me2/3, A-type lamins and the inner nuclear membrane (INM) proteins lamin-associated polypeptide β (LAP2β) and Lamin B receptor (LBR) have been implicated in organizing LADs (19, 20, 21, 22). The nuclear lamina is a proteinaceous network of type V intermediate filaments comprising A and B type lamins. These coiled-coil domain proteins provide a...
structural scaffold at the INM, with the A-type lamins being shown
to mediate LAD organization (12, 17 Preprint, 18 Preprint, 19).
Longstanding efforts have been undertaken to map and charac-
terize the local proteome of the nuclear lamina of the INM using
INM preparations, co-immunoprecipitation and, more recently,
BioID (Biotinylation Identification), a method for detecting proximal
protein interactions in living cells (23, 24, 25, 26, 27). However, these
efforts have exclusively focused on protein members of the INM/
lamina as baits and have not measured the protein landscape of
LADs themselves or the intersection of these proteome environ-
ments. To better understand these proximal protein compart-
ments, we have leveraged the BioID system to study the “micro-proteome”
of both LADs and the nuclear lamina using a multi-pronged ap-
proach. First, we have generated a chimeric protein comprising a
modified promiscuous biotin ligase, BirA*, fused to the nucleo-
plasmic N-terminus of LAP2β to pro-
file the INM/lamina proteome.
We posit that this, coupled with other published proteomic profiling
of the INM/lamina compartment, using other bait proteins, would
provide a more comprehensive overview of the INM/lamina (Fig 1A)
(24, 25). Second, we have taken a novel approach combining the
specificity of a DamID-based system to label LADs in live cells, the
m6A-tracer system, and coupled this with BioID to characterize
proteins proximal to LADs (Fig 1B and C) (16). Finally, we inte-
negrate these proteomic data with previous findings on lamin protein
interactions to generate a comprehensive mapping of the LAD/
lamina proteome interface.

Whereas a previous study used CRISPR-directed proximity-
labeling approach to identify proteomes associated with repeti-
tive sequences, this study represents, for the first time to our
knowledge, a chromatin-directed-BioID strategy that is indepen-
dent of identifying specific repetitive DNA sequences and targets a
specific nuclear compartment (28). Herein, we identify the pro-
teome of a chromatin compartment (LADs/B-compartment) by
leveraging two proximity labeling techniques: BioID and DamID. In
addition, by integrating published datasets, our new LAP2β inter-
actome data, and our LAD directed proteome data, we have

Figure 1. Strategy for investigating the nuclear periphery microproteomes.
(A) Representative images showing the biotinylated interactome of Lap2β (green) at the nuclear periphery in mouse embryonic fibroblasts (MEFs). (B, C) Pictorial representation of BirA* localization within the nucleus. Lamina-associated domains become methylated at GATCs upon Shield1 stabilization of Dam LmnB1, thereby recruiting BirA* mCherry m6A. Green shading in (C) depicts a putative biotinylation radius where available lysines on proximal proteins will be modified by BirA*. (D) Representative images showing the expected localization of the BirA*-m6a-tracer (red) and its biotinylated proteins (green) in the absence of DD-Dam LmnB1 (top row), in the absence (middle row) and presence (bottom row) of the shield ligand in MEFs. (E) Experimental workflow for mass spectrometry based BioID lamina associated domain microproteome analysis. MEF cells expressing BirA*-LAP2β, BirA*-m6A-tracer alone, BirA*-m6A-tracer/DD-Dam-LmnB1 plus/minus shield ligand were cultured with exogenous biotin then nuclei were extracted and Biotinylation Site Identification Technology (BioSiTe) coupled to liquid chromatography/tandem mass spectrometry analysis was used to identify biotinylated proteins.
identified different interaction zones at the nuclear periphery, thus mapping the differential and overlapping microproteomes of the peripheral nuclear compartment. In zone 1, we identified proteins that appear to be restricted to the INM/lamina that do not interact with LADs. In zone 2 we identified proteins that interact with both LADs and the INM/lamina and these may comprise the “middle-men” required to organize the LADs to the nuclear lamina. Finally, in zone 3 we identify proteins that are restricted to LADs, many of which are involved in regulating histone H3 lysine 9 methylation (H3K9me2/3) and cell cycle regulation.

Results

Establishment of BioID system to map the local proteomes at the nuclear periphery

The BioID system was initially developed using the Lamin A protein as the bait allowing a robust interactome of this insoluble protein (25). The BioID system relies on an engineered biotin ligase, BirA* which, when expressed in cells, has a small practical in vivo biotinylation radius of ~10 nm (29), thus labeling lysines on both proximal and directly interacting proteins. BioID, among other methods, has been used to analyze the local proteome of the INM using Lamin A/C proteoforms, Lamin B1 (LMNB1), SUN domain-containing protein 2 (SUN2), and nuclear pore complex (NPC) members as baits (23, 25, 29, 30 Preprint). To expand on these efforts and to focus on an INM protein implicated in LAD organization, we chose to identify vicinal proteins of the LAP2β in a BioID study. The LAP2β protein results from alternative splicing of the gene thymopoietin (TMPO) and was initially observed in nuclear envelope isolations and shown to bind lamin proteins (31). LAP2β is an integral INM protein that is thought to link the nuclear membrane to chromatin and also to regulate transcription factor functions (32). LAP2β is a member of the LEM (Lap2-Emerin-Man1) domain family of proteins and contains a LEM domain as well as a LEM-like domain, both of which have been thought to mediate interactions between protein/DNA complexes with the lamin binding region. To leverage the BioID system for analysis of the LAP2β proximal interactome we have tagged the nucleoplasmic N terminus with BirA* (Fig 1A).

BioID is closely related to the DamID technique in that both techniques rely on in-cell labeling of proximal molecules (9). In DamID, instead of modifying proteins, a Dam-X fusion protein modifies interacting DNA segments by methylating GATC stretches (GmATC). This modification can be used to isolate interacting DNA regions by cutting with the methyl-specific restriction enzyme, DpnI. In a typical DamID experiment, these fragments are amplified and subjected to DNA sequencing to generate genomic maps of interacting regions. The m6A-tracer system is an adaptation of the DamID technology for visualizing LADs in live-cell imaging (16). Our variation of the m6A-tracer system relies on the demarcation of LADs by an inducible and interphase restricted methylation of LADs underlying the nuclear lamina by Dam-LMNB1 (DD-Dam-HCdt-LMNB, Fig 1B). The inducibility and restriction to interphase labeling is conferred by the destabilization domain (DD) (16, 17 Preprint, 33 Preprint) and the Cdc10 dependent transcript 1 (hCdt) domains (17 Preprint, 33 Preprint, 34), respectively, with the DD domain causing degradation of Dam-LMNB1 in the absence of the shield ligand and the hCdt domain causing its degradation in all phases of the cell cycle with the exception of G1. Detection of the Gm6AATC-marked DNA (the LADs) is performed by the m6A-tracer, a catalytically inactive Gm6AATC binding domain of the DpnI enzyme, fused to a fluorescent protein (such as mCherry) (16, 17 Preprint, 33 Preprint).

To map the local proteome of the INM and LADs we generated three independent MEF cell lines expressing either BirA*-Lap2β (to map the INM/lamina, Fig 1A), BirA*-m6A-tracer alone (control), or BirA*-m6A-tracer with DD-Dam-LMNB1 (to map the LAD proteome) (Fig 1D). These cells were grown in the presence of exogenous biotin to enable efficient labeling of proximal proteins by BirA*. Cells harboring both BirA*-m6A-tracer and DD-Dam-LMNB1 constructs were split into two sets of cultured cells, one in the presence of the shield ligand and one without, as an additional control for the LAD proteome experiments (Fig 1E). To remove cytoplasmic contamination and limit our interrogation to the nucleus, a crude nuclear extraction preparation was performed on all cells. To detect biotinylation on candidate proteins, we used Biotinylation Site Identification Technology (BioSiTe) and liquid chromatography-tandem mass spectrometry (LC-MS/MS) (Fig 1E) (35).

Analysis of the LAP2β interactome using BioID

To identify proximal proteins of LAP2β, we used BirA* tagged LAP2β (BirA*-Lap2β) containing cells (Fig 1D). As a background control we used the BirA*-m6A-tracer construct expressed alone, without the presence of the DD-Dam-LMNB1 construct, which resulted in diffuse nuclear localization (red signal, Fig 1B) and subsequently similar biotinylation pattern (green signal, Fig 1D). A previous study using LAP2β tagged with BirA* found that the expressed protein was mislocalized to the ER and cytoplasm. This is not very surprising because it has been previously shown that over-expression of LEM domain proteins leads to their accumulation in the ER and cytoplasm, and such mis-localization occurs with disease variants of lamin proteins as well (31, 36, 37). We therefore sought to express our protein at low levels to preserve normal localization (Fig S1). In addition, this low expression ensures minimal disruption to normal function of Lap2β and its partners and minimizes cellular stress due to overexpression (and mislocalization). To detect where our low-expressing BirA*-Lap2β construct localized, we measured biotinylation signals using immunofluorescence on fixed MEFs expressing BirA*-Lap2β. As evidenced by the strong nuclear rim staining, we did not observe any gross mislocalization of LAP2β (Fig 1B). Some evidence of ER and mitochondrial expression is evident by the streptavidin signals, and this is to be expected because LAP2β transits the ER and there are endogenously biotinylated proteins in mitochondria. Therefore, as an additional step to maximize bona fide nuclear interactions, we performed a nuclear extract before protein isolation. Although this extra step may result in the loss of some true nucleoskeletal–cytoskeletal interactions, it increases the rigor in detecting nuclear interactions. The correct localization of our LAP2β-BirA* (Fig 1B), its low expression (1–2% of endogenous levels of LAP2B, Fig S1) that obviates perturbations to the INM due to overexpression of a key INM protein, and a nuclear
isolation step allowed for enriched detection of Lap2β-proximal proteins.

Using BioSiTe to detect biotinylated proximal proteins in our BioID screen we identified 334 total biotinylation sites in the BirA*-LAP2β containing cells and 684 sites in the BirA*-m6A-tracer alone containing cells (control). MS1 level quantitation was applied to obtain relative abundance differences between duplicate LC-MS/MS analyses of BirA*-LAP2β and BirA*-m6A-tracer cells. Replicate agreement was plotted (Fig 2A) and proteins enriched >2-fold were considered for LAP2β proximity/interaction (Table S1). Among the proteins enriched over control were expected and known interactors of LAP2β: LEM-domain containing protein 3 (MAN1), Emerin (EMD), and Lamins A and B1/2. Other known nuclear lamina proteins that were biotinylated were SUN domain contain proteins 1 and 2 (SUN1 and SUN2), LBR, and many NPC members (4, 38).

In an effort to rank the biotinylated proteins enriched over the control, in terms of potential closer proximity, we calculated the degree of biotinylation (number of biotinylated lysines/total lysines) and ordered proteins from greatest to least (Fig 2B) (35). We detected that the bait LAP2β has the greatest degree of biotinylation followed by Emerin, MAN1, LBR, LMNB1/2, LMNA, and SUN1/2. We hypothesize that these proteins represent the most proximal and/or abundant interactors of LAP2β on the nucleoplasmic side of the INM.

To verify that our list of proteins was in fact enriching for nuclear envelope proteins we also submitted detected biotinylated proteins to cell component analysis using the Enrichr Web-based analysis tool (39). Some of the top cellular component terms included nuclear periphery (GO:0034399), nuclear lamina (GO:0005652), and nuclear matrix (GO:00016363) (Fig 2C). To further examine if our data were consistent with previous studies, we benchmarked our data on four published studies of nuclear envelope interactomes (23, 24, 25, 29, 30 Preprint, 40, 41) (Table S3): (1) Bar et al combined a novel antibody based proximity labeling strategy and a meta-analysis of many Lamin A and B interactome studies that included methods such as BioID, yeast two hybrid, and co-immunoprecipitation to obtain a list of higher confidence “true” proteins at the NE/lamina. (2) Kim et al used BioID on many NPC members to build out the proximal proteins within the NPC and (3) Xie et al used BioID on both A-type lamins. (4) Birendra et al also used BioID to examine the interactomes of Lamin A and Sun2. We found 36 proteins to be overlapping between the union of these four studies (protein must have been observed in three of the four studies) and our LAP2β interactome analysis (Fig 2D), including...
most of our proteins showing the highest degrees of biotinylation. We believe that these proteins represent very high confidence nuclear envelope proteins.

We do, however, note the representation of ER proteins and nuclear pore proteins (NUPs) in our LAP2β data set which includes vesicle-associated membrane protein-associated protein B (VAPB) and nucleoporin NDC1 (NDC1). We hypothesize that this could be due to the NPC presenting itself as a kinetic bottleneck for the relatively large BirA*-LAP2β and therefore, longer residence times in the ER and NPCs. In addition, notably absent from our Lap2β interactome analysis is Barrier to Autointegration factor (BANF1) which is a known interactor of Lap2β, as well as HDAC3, another putative interactor (42, 43, 44). We hypothesize that the small size of BAF (≈10 kDa) makes detection of this protein by mass spectrometry based approaches particularly challenging. It is also important to emphasize that BiolD is a label transfer technique and hence, the biotinylation of potentially transient interactors such as HDAC3, coupled with limited material because of the sub-physiological expression level of BirA*-LAP2β, it is possible that some of these might have been missed by our analysis.

Given that LAP2β contains a transmembrane domain which is localized within the INM, we next examined our data for transmembrane domain-containing proteins. We observed that 34 of the 58 LAP2β proximal proteins contained a transmembrane domain (Table S1). It has been shown that membrane topology can be predicted with biotinylation site level data, which BioSITe provides (35). Many of these proteins exhibited biotinylation on areas of the protein consistent with lumen/nucleoplasmic annotated topology (35). Many of these proteins were involved in negative regulation of mitotic metaphase plate congression. Notable cell cycle–related genes include many members of the cell division cycle–associated protein family, including 2 (CDC2), 5 (CDC5), 7 (CDC7), and 8 (CDC8) and many kinesin family members such as 4A (KIF4A), 18B (KIF18B), 20A (KIF20A), and 20B (KIF20B). We also observed that many microtubule/spindle attachment proteins related to the kinetochore including the serine/threonine kinase that regulates segregation of chromosomes during mitosis Aurora kinase B (AurkB), nucleolar and spindle associated protein 1 (Nups1), regulator of chromosome condensation 2 (RCC2), microtubule nucleation factor (TPX2), sister chromatid cohesion protein (PDSS), and inner centromere protein (INCENP). In addition, the replication timing factor Rif1, was identified as LAP2β-interacting, in agreement with previous studies (51, 52).

As expected, another highly represented class of proteins enriched on LADs were proteins of the INM. These included many known INM proteins such as LAP2β, MAN1, Emerin, Lamins A and B1/2, LBR, and SUN1 (Fig 3C). Strikingly, NUP proteins were not enriched in the LAD proteome, suggesting that these chromatin domains are indeed not proximal to NPCs, in agreement with cytological data.

LADs are particularly enriched in chromatin modifying and binding proteins

As mentioned above, one of the major classes of proteins enriched on LADs were proteins involved in negative regulation of transcription. Specifically, many of the identified proteins have roles in the establishment/regulation of H3K9 methylation or bind to H3K9 methylated histones, such as heterochromatin protein 1 α and β.
Figure 3. Lamina associated domain (LAD) microproteome analysis. 
(A) Plot of replicate runs of MS1 level quantitation ratios of mass spectrometry identities of BirA*-m6A-tracer/DD-Dam-LMNB1 plus shield over minus shield ligand. 
(B) Gene set enrichment analysis of m6A-tracer BioID LAD proteome. 
(C) Integrative venn diagram of published nuclear lamina proteomic analyses, our current LAP2β BioID interactome analysis and the LAD-ome analysis. Proteins marked * have been validated by various groups, including ours. The experiments and results are summarized in Table 1. Proteins marked ** have been bioinformatically validated in-house. Note: not all proteins in these overlaps are shown for display purposes. Please see Table S3 for a full list of proteins. Data from C57BL/6 3T3-derived WT MEFs.

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(HP1α and HP1β), polycomb repressive complex 2 subunit Suz12, TRIM28 (Kap1), SMCHD1, PRR14, and DEK (Fig 3C). Of particular interest were the abundance of LAD-interacting proteins involved in establishment and maintenance of heterochromatin, specifically euchromatic lysine methyltransferase 2 (EMH2T2) and its binding partners chromodomain Y like (CDYL) and the transcription factor euchromatic lysine methyltransferase 2 (EMHT2) and its binding establishment and maintenance of heterochromatin, specifically euchromatic lysine methyltransferase 2 (EMH2T2) and its binding partners chromodomain Y like (CDYL) and the transcription factor euchromatic lysine methyltransferase 2 (EMHT2) and its binding establishment and maintenance of heterochromatin, specifically euchromatic lysine methyltransferase 2 (EMH2T2) and its binding partners chromodomain Y like (CDYL) and the transcription factor euchromatic lysine methyltransferase 2 (EMHT2) and its binding.

Finally, we also observed many additional chromatin modiﬁng and DNA binding proteins such as AT-hook containing transcription factor 1 (AHCTF1, also known as ELYS, which has also been implicated as a bona ﬁde NPC protein (53, 54)), ATRX chromatin remodeler (ATRX), MECP2, tyrosine-protein kinase BAZ1B (BAZ1B), high mobility group nucleosomal binding domain 2 (HMGN2), and PIN2/TERF1 interacting telomerase inhibitor 1 (PINX1) (53, 55, 56, 57, 58). We also identiﬁed a few transcription factors not known to be involved in heterochromatin such as MAFA (MAFG), zinc ﬁnger motifs (WIZ). These proteins have been shown to facilitate mono- and dimethylation of H3K9.

Integration of laminome and LADome data identify unique and overlapping micro-proteomes

We next sought to determine the overlapping proteomes between the nuclear lamina and LADs. We focused on using existing data and our new Lap2β proteome to maximize potential proteins interacting with chromatin. There were 54 proteins that overlapped between the published laminome data and our Lap2β proteome or the m6A-tracer LAD proteome (Fig 3C, Venn diagram overlaps). Importantly, the well-known nuclear lamina proteins such as LBR, LMNA/C, LMNB1, and SUN1 all showed enrichment on LADs as well (Fig 3C, yellow box). In contrast, whereas the published laminome and Lap2β showed interactions with NPCs (NUPs, Fig 3C, blue box), these were largely missing from the LAD interactome data. This ﬁnding supports cytological and DamID studies, suggesting that the chromatin underlying the NUPs is more euchromatic and distinct from the peripheral heterochromatin that comprises LADs.

In total, we identiﬁed 29 proteins, including MECP2, AHCTF1 (also known as ELYS), and PRR14, that were found to be biotinylated in the BirA*-m6A-tracer experiment that were also detected in the published nuclear lamina interactomes and/or our Lap2β interactome (Fig 3C, purple and yellow boxes). These are potential candidates for proteins that are at the interface of the INM and LADs, perhaps linking them together and regulating dynamic LAD organization (4).

The 47 proteins detected only in the Lap2β-BioID experiment, as well as most proteins in the published laminome data that are not LAD-enriched (Fig 3C, blue box and light and dark blue Venn regions), likely represent more INM proximal proteins that do not interact with and are farther from the underlying chromatin (LADs). These include the previously described NUP proteins. Other examples include, Progesterone Receptor Membrane Component 2 (PGRMC2), Torsin-1A-interacting protein 1 (Tor1aip1), Myoferlin (MYOF), transmembrane protein 214 (TMEM214), metadherin (MTDH also called protein LYRIC), cytoskeleton associated protein 4 (CKAP4), vesicle-associated membrane protein-associated protein B/C (VAPB); these are all transmembrane-containing proteins that may not have large enough nucleoplasmic domains to be in proximity to the LADs, and, for the Lap2β proximal transmembrane proteins, could also represent transient interactions from transit through the ER. Nonetheless, these proteins appear to be more distal to the underlying LAD chromatin. Interestingly, SYNE1 and SYNE3 which code for Nesprins 1 and 3 are classiﬁed in this group. Although there is evidence that nucleoplasmic nesprins exist (59, 60), it is also possible that these proteins were biotinylated en route to the INM.

Eighty-three proteins were identiﬁed as associating uniquely with LADs (Fig 3C, red Venn region and pink box). Because the LADs and the lamina are in such close proximity, we were surprised to ﬁnd such a clear distinction between the LADome and the lamina. The LAD-speciﬁc proteins include the previously described chromatin modiﬁer complexes such as EHMT1/2, cell cycle regulators and chromatin interactors. This is particularly interesting, given the recent evidence that chromatin state directs LAD organization (12, 14, 15, 16, 17 Preprint, 18 Preprint). These data suggest, and support previous ﬁndings, that these regions are enriched for chromatin complexes that initiate and maintain a heterochromatic state (10, 12). These data also support a model in which chromatin state is established independently from association of these chromatin domains with the lamina because these proteins do not interact with the INM/lamin directly (15, 17 Preprint).

Validation and other supporting data

To our knowledge, this presents the ﬁrst comprehensive characteriza- tion of the local proteome associated with LADs. Many of the proteins identiﬁed have already been extensively validated, with some of these validations corroborating the placement of the proteins in speciﬁc interaction groups: INM/lamina, interface, or chromatin restricted. Published and new validation experiments and data that directly show association with the INM/lamina and/or involvement in LAD establishment/maintenance have been summarized and compiled in Table 1.

Proteins at the INM/lamina that do not interact with LADs

Proteins at the INM/lamina that showed no enrichment likely represent nuclear envelope proteins that are spatially removed from LADs and would therefore be restricted from interacting with LAD chromatin. To test this supposition, we interrogated the interaction of one such protein, Nup153, with LADs using publicly available data (Fig 4) (10, 61). Nup153 has previously been shown to interact with chromatin, but our data suggest that this protein is excluded from LAD chromatin domains. We ﬁnd that most Nup153 interactions with chromatin occur outside of LADs and, intriguingly, Nup153 peaks that appear to be within LADs (at a gross scale) coincide with regions that have low Lamin B1 signal, which we have previously termed DiPs (Depleted in Peripheral signal (18 Preprint, Fig 4). Interestingly, these DiPs correlate with active transcription start sites and enhancers, with a distinct and discrete switch from the inactive B to the active A chromatin compartment. These combined data support our integrated BioID data which suggest that the Nup153 interactions with chromatin at the lamina are more distal to LADs (Fig 4).
Table 1. Table detailing experiments and results from published work directly validating protein hits from our BioID study.

| Protein | Experiment | Results |
|---------|------------|---------|
| Lap2b (20) (Fig 5) | DamID | Lap2b DamID show traces almost identical to LMNA DamID (molecular association with LADs) |
| Emerin (10) | DamID | EMD DamID profiles are virtually identical to Lmnb1 DamID (molecular association with LADs) |
| LBR (19) | Developmental characterization | LBR is generally expressed more predominantly in progenitor cells switching to Lamin A/C later in development |
| | Functional characterization of chromatin in Lamin A/C and LBR double KO mice | All post mitotic cells in Lamin A/C, LBR double knockout mice exhibit loss of peripheral chromatin and inverted chromatin configuration (histological) |
| | Ectopic LBR expression in photoreceptor rod cells in mice | Inverted chromatin configuration found in rod cell nuclei can be counteracted with ectopic LBR expression |
| Lamin A/C (12, 17 Preprint, 18 Preprint, 19, 20, 21, 22, 65; Fig 5) | DamID | LMNA DamID profiles are virtually identical to Lmnb1 DamID (molecular association with LADs) |
| | Knockdown and Immunofluorescence | Fragments that by default localized at the periphery were found to have loss peripheral localization. |
| | Knockdown studies and 3D-immunoFISH | Disrupted association of LADs with the nuclear periphery AND overall chromosome territorial architecture |
| | Developmental characterization of expression | LMNA expressed more predominantly in more differentiated cells (compared with LBR) |
| | Histological characterization of chromatin in KO mice | Removing Lamin A/C from cells lacking LBR expression resulted in loss of LADs and an inverted chromatin organization. |
| | All post mitotic cells in Lamin A/C, LBR double knockout mice exhibit the inverted chromatin organization. |
| Rif1 (51, 52) | Electron microscopy | Gold labelled beads (as readout Rif1 localization) on heterochromatin at the periphery |
| | Nuclear subfractionation. | Found in DNAse-resistant as well as salt-resistant nuclear fraction |
| | Knockout affected replication timing | Affected replication timing and nuclear architecture |
| | Rif1 ChIP | Correlates with LADs |
| | Immunofluorescence | Localizes at the nuclear periphery |
| | Co-immunoprecipitation | Rif1 immunoprecipitates Lmnb1 |
| | ChIP | Highly correlated with LAD traces |
| | Immunofluorescence and colocalization studies | MeCP2 and LBR colocalizes at the nuclear periphery |
| | Biochemical fractionation of nuclei | MeCP2 exists in the MNase and salt-resistant nuclear pellet |
| | Coimmunoprecipitation assays | MeCP2 coimmunoprecipitates LBR and vice versa |
| | Bimolecular Fluorescence Complementation | MeCP2 and LBR interacts at the nuclear periphery |

(Continued on following page)
Proteins that bind the INM/Lamina network and LADs—the “middlemen”

Proteins that interface the INM/Lamina and LADs are candidates for linking them together and regulating dynamic LAD organization (4). This group is composed of both INM/Lamina proteins and chromatin binding proteins. It is hardly a surprise that lamin proteins fall into this category as they have been shown to interact with LADs and to be important for their localization to the nuclear periphery (Table 1 and Fig 5). In addition to the lamin proteins, some INM proteins were also identified as spanning the INM/lamina and interacting with chromatin. For instance, LBR has previously been shown to interact with chromatin and is required for normal chromatin domain organization in early development (Table 1, 19) as well as Emerin, a LEM domain protein, displays interactions with chromatin, as measured by DamID, which are highly correlated with LADs (10). In addition, both LBR and emerin have been identified in multiple BioID and biochemical experiments to interact with the same lamin network (10). These combined data support our findings here that these proteins span the lamina network and chromatin interface. We find that the LEM domain protein Lap2β also falls into this category. We note, however, that we failed to detect another LEM domain protein, Man1, in our LAD proteome. We speculate that this is due to a lower abundance of this protein, but this could also be because it does not have lysine residues proximal to LADs. Nonetheless, Lap2β has been shown, through multiple experiments, to interact with the lamina and INM

References and figures describing the role of these proteins at the INM/lamina or on LADs are in parentheses.

### Table 1. Continued

| Protein          | Experiment                                      | Results                                                                 |
|------------------|-------------------------------------------------|-------------------------------------------------------------------------|
| TRIM28 (KAP1)    | • Coimmunoprecipitation                         | • Interacts with Lamin A                                                |
|                  | • Ectopic recruitment to genomically integrated | • Acts as obligate corepressor of hormone-responsive KRAB zinc finger proteins; represses genes by associating with HP1 and SETDB1 |
|                  | transgene via hormone-responsive KRAB zinc      |                                                                         |
|                  | finger proteins                                 |                                                                         |
| RRP1B (66, 70)   | • Bimolecular Fluorescence Complementation      | • RRP1B interacts with Sun2 at the nuclear periphery                   |
|                  | ChiP-reChIP                                     |                                                                         |
| PRR14 (63, 64, 73, 98) | • Yeast 2 hybrid                                  | • Interacts with Trim28 and HP1                                         |
|                  | • Functional and deletion mapping coupled to     | • PRR14 is a binding partner for HP1α                                    |
|                  | immunofluorescence                              | • N-terminal 135 amino acids (containing NLS- and HP1-binding site) sufficient for nuclear peripheral localization |
|                  | • Knockdown studies                             | • Mutation of HP1-binding motif disrupts nuclear peripheral targeting |
|                  | • Cell cycle studies by immunofluorescence      | • Central region required for lamina localization                       |
| EHMT2/ G9a (12, 13, 14, 15, 16, 17 Preprint, 18 Preprint) | • Drug (BIX-01924) inhibition of G9a and/or siRNA against G9a followed by DamID or immunofluorescence | • Assemble on chromatin at anaphase                                     |
|                  | • G9a overexpression                            | • Fragments that by default localized at the periphery were found to have loss peripheral localization (by immunofluorescence) and association (by DamID) |
|                  | • Drug inhibition of G9a followed by 3D-         | • Endogenous LADs were also shown to have a lower relative nuclear association when compared with non-treated or control-treated samples. |
|                  | or immunofluorescence                           | • Increased relative nuclear lamina association of LADs compared with non-treated or control-treated samples. |
|                  | • Disrupted association of LADs with the nuclear periphery AND overall chromosome territorial architecture | • Disrupted association of LADs with the nuclear periphery AND overall chromosome territorial architecture |
| CBX1 (HP1β) (72; Fig 6) | • DamID                                      | • Highly correlated with LAD traces                                     |
|                  |                                                 | • Important for X-chromosome structure                                 |
| SMCHD1 (72; Fig 6) | • DamID                                      | • Highly correlated with LAD traces                                     |
|                  |                                                 | • Important for X-chromosome structure                                 |
network. In addition, several studies have suggested that this protein is important for LAD organization and regulation (12, 20). We therefore decided to interrogate the interaction of Lap2 with LADs to verify that this protein indeed interacts across the INM/chromatin interface, as our BioID data suggest, and to test the breadth of such interactions. Using a DamID approach, we find that Lap2 interaction with chromatin occurs on LADs; the Lap2 interaction signatures are virtually identical to LADs (Figs 5A and B and S2).

In addition to well-known resident proteins of the INM/Lamina, other proteins less often associated with INM/Lamina studies were also classified in this group as potential INM/LAD links. An example would be methyl-CpG binding protein (MECP2), which we have found to interact with both LADs and in published laminome data. MECP2 is a known repressor of DNA that binds to methylated CpG regions in the DNA. Recently, a study demonstrated that CpG methylation demarcates the heterochromatic "B-compartment" of the genome, which is highly correlated with LADs (17 Preprint, 18 Preprint, 74). Consistently, bioinformatic analysis of publicly available MeCP2 ChIP-seq data shows a high coincidence of MeCP2 binding domains with LADs—two thirds of MeCP2 occupancy coinciding with LADs and almost half of the LADs potentially regulated by MeCP2 binding events (Fig SC–E (10, 71)). Furthermore, an interaction between LBR and MeCP2, which has been observed, further implicating MeCP2 physically interacting with the INM proteome and the LAD proteome (62, 75, 76, 77, 78, 79). Other examples in this category include TRIM28; RRP1B, which have both been documented to interact with nuclear lamina components and shown to be associated with H3K9 methylated locations in the genome; PRR14, which has been shown to provide a link between LADs and the INM through its association with HP1; and RIF1, a known chromatin modifier involved in replication timing and previously shown to be enriched on LADs (Table 1).

**Proteins associating uniquely with LADs**

We found a large number of LAD-restricted proteins that did not overlap with the laminome or lap2b datasets. These proteins potentially regulate/modulate LADs independent of their association with the nuclear periphery. Proteins in this group fall under three sub-categories consisting of cell cycle regulators, proteins that bind and/or regulate DNA and proteins that bind to or modulate H3K9 methylation. We were particularly interested in the proteins that affect H3K9 methylation because the methylation of H3K9 is an important feature of heterochromatin and has been shown to be enriched at and required for LADs (10, 11, 12, 14, 15, 16, 17 Preprint, 18 Preprint, 80, 81, 82, 83). This includes the HP1 proteins, chromobox 5 (CBX5) also known as heterochromatin protein 1α (HP1α) which binds to H3K9me3 (84) and heterochromatin protein 1β (HP1β or CBX1). CBX1 serves a similar function as HP1α and as shown, in our bioinformatic analysis, to be highly coincident with LADs (Table 1 and Fig 6A–C (72)). We find that 80% of CBX1 binding peaks reside within LADs (Fig 6B). In addition, we detected other chromobox family proteins: chromobox 3 (CBX3 or HP1γ) and 8 (CBX8) in our BioSiTe data. HP1 proteins, particularly HP1α, bind to methylated H3K9 and mediate gene silencing by maintaining and spreading heterochromatin. Alongside these HP1 proteins, we also identified...
the histone chaperone and proto-oncogene DEK, which has been shown to prevent access to transcription machinery and to interact with HP1, enhancing its binding to H3K9me3 (85, 86). Furthermore, we found the SUZ12 polycomb repressive complex 2 subunit (Suz12) which has been found to both stabilize HP1 and increase H3K27me3, which in turn increases the HP1α/β/γ binding of H3K9me3 (87).

Another intriguing protein that we identified in this group is the euchromatic lysine methyltransferase 2 (EHMT2, or often called G9a). EHMT2 is an important methyltransferase known to methylate H3K9 and H3K27, facilitating H3K9me1 and H3K9me2 in particular, and subsequent gene silencing (88). Disruption of this protein by either epigenetic drugs or shRNA-mediated silencing disrupts peripheral heterochromatin and LAD formation (Table 1) (12, 16, 17 Preprint, 18 Preprint). Intriguingly, this protein was also recently implicated in methylating Lamin B1, an event suggested to be important for LAD organization (although, we note that we did not detect interactions of Lamin B1 and EHMT1/2 across multiple datasets) (89). Moreover, EHMT2 was found to interact with and is recruited by chromodomain Y like (CDYL), another protein we
identified to be enriched at LADs, to facilitate H3K9 dimethylation and repression of the neurogenesis master regulator, REST, target genes (90, 91). EHMT2 and EHMT1 have been found to form a complex with widely interspaced zinc finger motifs (WIZ), another protein we found in our data, which acts to stabilize this complex on chromatin and prevent degradation of EHMT1 (92, 93).

Interestingly, we also identified structural maintenance of chromosomes flexible hinge domain containing 1 (SMCHD1) protein as LAD enriched—a protein that had not been previously identified on LADs. On inactive X chromosomes, SMCHD1 and HP1 are found to colocalize at areas of H3K9 methylation (72, 94 Preprint, 95). SMCHD1 has also been shown, by co-immunoprecipitation, to interact with ligand dependent nuclear receptor interacting factor 1 (LRIF1), another protein we found in our data, which acts to stabilize this complex on chromatin and prevent degradation of EHMT1 (92, 93).

Interestingly, we also identified structural maintenance of chromosomes flexible hinge domain containing 1 (SMCHD1) protein as LAD enriched—a protein that had not been previously identified on LADs. On inactive X chromosomes, SMCHD1 and HP1 are found to colocalize at areas of H3K9 methylation (72, 94 Preprint, 95). SMCHD1 has also been shown, by co-immunoprecipitation, to interact with ligand dependent nuclear receptor interacting factor 1 (LRIF1), another protein identified in this study, which helps target the protein to H3K9me3 regions (96). To test whether SMCHD1 interacts with LADs, as indicated by our proteomic data, we compared published SMCHD1 DamID data (from a study focused on the structural regulation of the inactive X chromosome and not LADs) with our Lap2β DamID data. This analysis showed a remarkably high coincidence rate of 86%, of SMCHD1 binding domains with LADs, suggesting that SMCHD1 potentially regulates LADs on top of its known regulatory roles in establishing and/or maintaining the inactive X chromosome (Fig 6A, D, and E (72)).

Discussion

We sought to identify the microproteome zones or regions at the INM/lamina/chromatin interface to identify potential mediators of LAD organization by mapping the LAD-proteome and comparing that to new and existing lamina/INM proteome datasets (Fig 1). Previous studies have shown that both chromatin state of the LADs and Lamin A/C are critical for LAD organization, suggesting the potential for chromatin scaffolding protein “middlemen” in mediating interactions between LADs and the lamina (97). In support of this idea, PRR14, a protein that interacts with HP1α (heterochromatin protein 1α), has been shown to interact with both the lamina and chromatin (64, 98). Previous studies have identified proteins enriched at the nuclear lamina (the “laminome”) through multiple methods, including BioID (23, 24, 25, 26, 27). Here we use BioID with BioSITe (BioID coupled with biotinylation site enrichment and...
To find LAD-enriched proteins, we used our BioID with BioSITe pipeline coupled with Lamin B1–directed DamID. We leveraged a modified version of a live cell LAD visualization system to recruit biotin ligase directly to LAD chromatin (Fig 1B–D). As with the Lap2B-directed BioID experiments, we verified that the biotin ligase marked proteins at the periphery of the nucleus (Figs 1B and S1). LAD-enriched proteins included cell cycle related proteins, proteins of the INM/lamina and chromatin interactors and modifiers (Fig 3B and C). Strikingly, most of the proteins identified in the LADome were specifically LAD-enriched, that is to say, they were not identified in either previous lamin interactome data or in our Lap2B interacting protein set. Importantly, LADs did not generally interact with NUPs, which is in agreement with cytological data and DamID data showing that NUPs interact primarily with transcriptionally active regions of the genome (99). NUP98, out of all the NUPs identified in the laminome, is the only NUP that displayed LAD interactions. We speculate that NUP98 interacts with LAD border regions given that it has been shown to interact with highly transcribed genes including the highly active genes that flank LADs, demarcating their borders (10).

The LAD-specific interactions highlight that although LADs and the INM/Lamina are spatially proximal, they are indeed different, but overlapping, microenvironments (Figs 3 and 7). It is especially intriguing that a large fraction of the proteins that are identified as LAD-specific are related to modulating chromatin state, particularly H3K9me2/3. Recent data from our laboratory have shown that, through the cell cycle, LADs within a chromosome self-interact before their organization at the nuclear lamina, suggesting that the higher order interactions of these chromatin domains are independent of their lamina association (17 Preprint). However, the inactive chromatin state, particularly H3K9me2/3 and H3K27me3 are required for lamina association (12, 13, 14, 15). These data are compatible with a two-step model in which chromatin state mediates self-association of these domains and that lamina association is then mediates by proteins which interact with (and help maintain) these specific chromatin modifications or states.

Of particular interest, then, are the proteins that sit at the interface of the INM/lamina and LADs. LADs were initially identified as domains of chromatin in molecular proximity to Lamin B1, using the proximity labeling method DamID. These “middlemen” proteins include Lamin A/C, emerin, and LBR, which have been previously shown to be necessary for LAD organization. In this and other studies, Lamin A/C and emerin have also been used to identify LADs, thus their identification in our screen as overlapping LADs and the INM/lamina is not surprising (65, 100, 101) (Figs 3C and 7, 13 of 20.

Figure 7. Putative model of the local proteome at the inner nuclear membrane (INM)/lamina/lamina-associated domain (LAD) interface.

The model depicts the nuclear periphery resolved into three zones based on data from this work as well as previous studies. Zone 1 is the most distal from chromatin comprising proteins that do not interact with LADs, such as nuclear pore complexes, some nesprins, and regulators of LEM domain proteins. Zone 1 then transitions into Zone 2 consisting of lamin proteins (shaded gray) and membrane-spanning proteins (colored teal) that traverse the lamina/LAD interface. In this region, we also find a unique set of chromatin/DNA binding proteins that display interactions with both the INM/lamina network and LADs (purple and gold). BAF, although not found in our datasets, is included in this zone based upon numerous in vivo and in vitro studies demonstrating its position as a linker between chromatin and the nuclear envelope. Thus, Zone 2 is made up of a lamin scaffold, some INM proteins, and a specific subset of DNA interacting proteins that may facilitate LAD interactions with the nuclear lamina. Finally, Zone 3 shows LAD-specific proteins (red) identified in our screen. These are highly enriched in chromatin-binding and modifying proteins.
zone 2). These “middlemen” proteins are all potential mediators of LAD organization at the lamina (12, 17 Preprint, 18 Preprint, 19). Thus, our analysis has identified known mediators of LAD scaffolding as proximally interacting with LADs.

Importantly, our BioSITe analysis also uncovered additional proteins that sit at this interface and are, heretofore, untested potential mediators of LAD organization to the nuclear lamina (Figs 3C and 7 zone 2). These include the previously implicated PRRT4, as well as RIFT, AHCFT1, and MEC2P and H2AFX. RIFT binding, which demarcates late replicating domains, has previously been identified as overlapping with LADs and in regulating genome organization (52). AHCFT1 (also known as ELYS) is implicated in nucleopore complex assembly, proper exit from mitosis, and genome stability (53, 54). MEC2P, a methyl CpG binding protein, has clear roles in establishing and maintaining heterochromatin. MeCP2 partners with HDACs and histone methyltransferase (as well as RNA) and is involved in higher order chromatin remodeling and silencing (55, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111). We have also shown bioinformatically using published MeCP2 ChIP datasets that MeCP2 is enriched within LADs (Fig 5). Intriguingly, although our proteomic analysis was not suited to identifying chromatin modifications, we identified histone H2AFX, which is phosphorylated to form γH2AX. γH2AX is found on double strand breaks on chromatin, which become heterochromatinized and possibly shunted to the γ form (113).

H2Ax may also be key in establishing de novo LAD organization because one study found that during epithelial to mesenchymal transition, regions that demarcate newly formed LAD borders become enriched in γH2AX (113).

The synthesis of this data allows us to better refine a putative model of the micro-proteome of the INM/lamina and LAD interface into “zones” of interaction (Fig 7). Whereas our model in Fig 7 depicts an INM/lamina separated into three “zones” of interaction (from a LAD point-of-view), more refined and future studies will likely uncover further subdivision of the NE into additional spatio-temporal regions, perhaps enriched with specific INM proteins, with the potential to differentially regulate chromatin and other nuclear processes. Taken together, our findings suggest that LADs and the INM/Lamina have distinct, but overlapping proteomes. Several of the proteins that are shared between LADs and the INM/lamina have already been shown to be important for organization of LADs to the lamina. We propose that several newly identified proteins spanning these two proteomic domains are likely involved in establishment or maintenance of LAD chromatin. Moreover, our findings are consistent with LADs themselves, supporting a heterochromatic architecture that is independent of their association with the nuclear lamina, but important for mediating interactions at the lamina.

Mapping the micro-proteome of the nuclear lamina and LADs

It is important to note that our proteomic analysis was not suited to identifying chromatin datasets that MeCP2 is enriched within LADs (Fig 5). Intriguingly, although our proteomic analysis was not suited to identifying chromatin modifications, we identified histone H2AFX, which is phosphorylated to form γH2AX. γH2AX is found on double strand breaks on chromatin, which become heterochromatinized and possibly shunted to the γ form (113).

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Materials and Methods

Plasmid construction

To clone Fu CMV DD-Dam–LMNB1, Dam LmnB1 was first amplified from Plgw Dam LmnB, kindly provided by the Van Steensel laboratory, with overhanging restriction sites. The amplicon was then TA cloned into pGEM-t easy (Promega). The cdt1 fucci tag, kindly provided by the Van Steensel labo-

To generate Fu-BirA-mCherry-m6A, the m6A was synthesized using codon-optimized, overlapping oligonucleotides that span the last 109 amino acids of the restriction endonuclease DpnI and amplified to be flanked by BsrGI and BstBI restriction sites. The amplicon was then cloned downstream of mCherry in the Fu-mCherry-hLMNA-bsr vector (unpublished) using BsrGI and BstBI, generating Fu-mCherry-m6A. Generation one BirA* flanked by Nhel and XhoI was amplified from mycBioID-PBAE-puro, a gift from Kyle Roux, and cloned into the Nhel site upstream of mCherry in Fu-mCherry-m6A.

To generate Fu-BirA* Lap2β, Lap2β was amplified from cDNA material from MEFs and TA cloned into pGEM-t easy (Promega). Lap2β along with a linker sequence corresponding to the multiple cloning site of pCDNA3.1 was subcloned downstream of BirA* in Fu_BirA*_hLMNA using the XhoI and BstBI restriction sites.

Cell line generation, reagents, and antibodies

MEFs were purchased from ATCC (CRL-2752) and cultured according to their establish protocols. These MEFs were transduced with lentiviral particles from the described plasmids. Specifically, viruses were generated in HEK 293T/17 cells (CRL-11268; ATCC) by cotransfecting VSV-G, delta 8.9, and the indicated construct. 10 mM sodium butyrate was then added to the transfected cells 3 h post transfection for an overnight (~16 h) incubation at 37°C, 5% CO2. The transfection media containing sodium butyrate was removed the following day and the cells were washed with 1× PBS. Opti-MEM was then added back to the cells which were then incubated at 37°C, 5% CO2. Viral supernatant was collected every 12 h (up to four collections), and the supernatant of all collections were pooled. MEFs were incubated overnight with fresh viral supernatants supplemented with 4 μg/ml polybrene and 10% FBS. Fresh MEF medium was then added to the cells after the virus was removed and selected with 10 μg/ml blasticidin or 1 μg/ml puromycin (or both).

Antibodies used in this study include an anti-LMNB antibody (sc-6217, goat IgG; Santa Cruz Biotechnology, Inc.) and anti-biotin antibody (A150-109A; Bethyl Laboratories, Inc.).

Imaging and immunofluorescence

Cells were prepared for immunofluorescence by plating on sterilized 25-mm round coverslips (German borosilicate glass #15; Harvard Apparatus) in six-well tissue culture dishes. Immunofluorescence was carried out as previously described (12, 114). The nuclear lamina was visualized using an anti-LMNB antibody (sc-6217, goat IgG; Santa Cruz Biotechnology, Inc.) and Alexa Fluor 647 AfiniPure Donkey Anti-Goat (#705-606-147; Jackson Immunoresearch), and anti-biotin antibody (A150-109A; Bethyl Laboratories, Inc.).
inverted fluorescence microscope (AxioVision; Carl Zeiss) fitted with an ApoTome and camera (AxioCam MRm; Carl Zeiss). The objective lens used was a 63× apochromat oil immersion (Carl Zeiss) with an NA of 1.5 (Immersol 518; Carl Zeiss). All immunofluorescence was performed at room temperature on #1.5 coverslips. AxioVision software (Carl Zeiss) was used for image acquisition. Images were exported as TIFFs to (FIJI ImageJ, National Institutes of Health) for further analyses (115).

**BioID with BioSiTe**

NIH3T3 cells expressing LAP2β-BioID, BirA*-m6A-tracer alone, BirA*-m6A-tracer + DD-Dam-LMN1B1 constructs were cultured overnight (~16 h) with 50 μM exogenous biotin. Cells expressing BirA*-m6A-tracer + DD-Dam-LMN1B1 were cultured with 2 μM shield-1 ligand (AOBIOS, #AOBI1848) for 24 h before the addition of exogenous biotin for a total of 48 h. Cells were trypsinized, washed in large volume PBS washes, then resuspended in a hypotonic lysis buffer (5 mM PIPES, 85 mM KCl, 1% NP-40, and protease inhibitors) for 10 min to separate cytoplasmic fraction from nuclear fraction. The resulting nuclei were pelleted and protein extraction was carried out by sonication (three rounds, duty cycle 30%, 20 s pulses) in 50 mM TEABC and 8 M urea. The protein concentration of samples was measured by BCA assay. A total of 10 mg of lysate per replicate was then reduced and alkylated by incubation with 10 mM DTT for 30 min followed by 20 mM IAA for 30 min in the dark. The lysate was diluted to 2 M urea by adding three cell lysate volumes of 50 mM TEABC. The proteins were digested with trypsin (1:20 of trypsin to protein) at 37°C overnight (~16 h). The resulting tryptic peptides were desalted using a Sep-Pak C18 column and subsequently lyophilized. Protein G agarose beads (#16-266; Millipore Sigma) were exported as TIFFs to (FIJI ImageJ, National Institutes of Health) for transparency. The peptide concentration of samples was measured by BCA assay. A total of 10 mg of lysate per replicate was then reduced and alkylated by incubation with 10 mM DTT for 30 min followed by 20 mM IAA for 30 min in the dark. The lysate was diluted to 2 M urea by adding three cell lysate volumes of 50 mM TEABC. The proteins were digested with trypsin (1:20 of trypsin to protein) at 37°C overnight (~16 h). The resulting tryptic peptides were desalted using a Sep-Pak C18 column and subsequently lyophilized. Protein G agarose beads (Wang et al. 115) were coupled to 120 μl of protein G bead slurry, overnight (~16 h) at 4°C. Antibody-coupled beads were further washed with PBS once and BioSiTe capture buffer (50 mM Tris, 150 mM NaCl, and 0.5% Triton X-100) twice. Lyophilized phosphatidyls were dissolved in 1 ml of BioSiTe capture buffer and pH solution was adjusted to neutral (7.0–7.5). Peptides were subsequently incubated with anti-biotin antibody-bound protein G beads for 1 h at 4°C. The bead slurry was washed three times with BioSiTe capture buffer, three times with 50 ml of Tris, and two times with ultrapure water. Biotinylated peptides were eluted with four rounds of 200 μl elution buffer (80% acetonitrile and 0.2% trifluoroacetic acid in water). The eluents were dried, desalted, and concentrated using homemade C18 reversed-phase column.

**Mass spectrometry**

The fractionated peptides were analyzed on an Orbitrap Fusion Lumos Trubrid Mass spectrometer coupled to the EASY-nLC 1200 nanoflow liquid-chromatography system (Thermo Fisher Scientific). The peptides from each fraction were reconstituted in 20 μl of 0.1% formic acid and loaded onto an Acclaim PepMap 100 Nano-Trap Column (100 μm × 2 cm; Thermo Fisher Scientific) packed with 5-μm C18 particles at a flow rate of 4 μl per minute. Peptides were separated at 300 nL/min flow rate using a linear gradient of 7–30% solvent B (0.1% formic acid in 95% acetonitrile) over 95 min on an EASY-Spray column (50 cm × 75 μm ID; Thermo Fisher Scientific) packed with 2 μm C18 particles, which was fitted with an EASY-Spray ion source that was operated at a voltage of 2.3 kV. Mass-spectrometry analysis was carried out in a data-dependent manner with a full scan in the mass-to-charge ratio (m/z) range of 300–1,800 in the “Top Speed” setting, 3 s per cycle. MS and MS/MS were acquired for the precursor ion detection and peptide fragmentation ion detection, respectively. MS scans were measured at a resolution of 120,000 (at m/z of 200). MS/MS scans were acquired by fragmenting precursor ions using the higher energy collisional dissociation (HCD) method and detected at a mass resolution of 30,000 (at m/z of 200). Automatic gain control for MS was set to 1 million ions and for MS/MS was set to 0.05 million ions. A maximum ion injection time was set to 50 ms for MS and 100 ms for MS/MS. MS data were acquired in profile mode and MS/MS data in centroid mode. Higher energy collisional dissociation was set to 32 for MS/MS. Dynamic exclusion was set to 35 s, and singly charged ions were rejected. Internal calibration was carried out using the lock mass option (m/z 445.120025) from ambient air.

**Database searching, quantification, and post-processing of MS data**

Proteome Discoverer (v 2.2; Thermo Fisher Scientific) suite was used for quantitation and identification of peptides from LC–MS/MS runs. Spectrum selector was used to import spectrum from raw file. During MS/MS preprocessing, the top 10 peaks in each window of 100 m/z were selected for database search. The tandem mass spectrometry data were then searched using SEQUEST algorithm against protein databases (mouse NCBI RefSeq 73: 580397 entries) with the addition of fasta file entries for BirA*-m6A-tracer and LAP2β-BioID constructs with common contaminant proteins. The search parameters for identification of biotinylated peptides were as follows: (i) trypsin as a proteolytic enzyme (with up to three missed cleavages); (ii) peptide mass error tolerance of 10 ppm; (iii) fragment mass error tolerance of 0.02 Da; and (iv) carbamidomethylation of cysteine (+57.02146 D) as a fixed modification and oxidation of methionine (+15.99492 D) and biotinylation of lysine (+226.07759 D) as variable modifications. Peptides and proteins were filtered at a 1% false-discovery rate at the peptide spectral match (PSM) level using percolator node and at the protein level using protein false-discovery rate validator node, respectively. For the MS1 level quantification of the peptides, the Minora Feature Detector, using the program’s standard parameters, was used and all of the raw files from the two replicates were quantified together. Unique and razor peptides both were used for peptide quantification, whereas protein groups were considered for peptide uniqueness. Identified protein and PSM level data were exported as tabular files from Proteome Discoverer 2.2. We used an in-house Python script to compile the peptide level site information mapped to RefSeq databases. We eliminated all non-biotinylated peptides from our analysis. The summary count on the number of supported peptides, PSMs, number of biotinylation sites and quantification was then calculated at the protein level. Quantitation of replicate agreement was plotted and average values between replicates were calculated to total biotinylated protein abundance. Transmembrane domain analysis was carried out as previously described (PMID:
We obtained annotated transmembrane domains, topological domains, and subcellular localization from Uniprot. Sites of biotinylation were then compared with annotated topologies to identify the location of biotinylation with respect to lumen, nucleoplasmic, and cytoplasmic portions of proteins. For gene set enrichment analysis, gene lists were uploaded to the Web portal Enrichr (http://amp.pharm.mssm.edu/Enrichr/) (PMID: 27141961).

**DamID**

DamID was performed as described previously (9, 10, 12, 20, 116). Cells were transduced with murine retroviruses harboring the Dam constructs. Self-inactivating retroviral constructs pSMGV Dam-V5 (Dam-Only), pSMGV Dam-V5-Lamin A (Dam-Lamin A), and pSMGV Dam-V5-Lap2β (Dam-Lap2β) were transfected using Fugene 6 transfection reagent (E2691; Promega) into the Platinum-E packaging line (RV-101; Cell Biolabs) to generate infectious viral supernatants in DMEM complete media were added to transduce murine embryonic fibroblast (MEF) cells. These viral supernatants were added to 16 h preincubated MEFs overnight (~16 h) with either Dam-only, Dam-Lamin A, or Dam-Lap2β viral supernatant and 4 μg/ml polybrene. Cells were allowed to expand for 2–4 days before being passaged.

MEFs were collected by trypsinization and DNA isolation was performed using the QiAamp DNA Mini Kit (51304; QIAGEN), followed by ethanol pre-harvest. Cells were allowed to expand for 2 days then harvested. Retroviral infections were carried out by incubating MEFs overnight (~16 h) with Dam-only, Dam-Lamin A, or Dam-Lap2β viral supernatant and 4 μg/ml polybrene. Cells were allowed to expand for 2–4 days then pelleted for harvest.

MEFs were collected by trypsinization and DNA isolation was performed using the QiAamp DNA Mini Kit (51304; QIAGEN), followed by ethanol precipitation and resuspension using 1.0 μl/m M Tris, pH 8.0. The DNA was dissolved in 10 mM Tris, pH 8.0. Digestion was performed overnight (~16 h) using 0.5–2.5 μg of this genomic DNA and restriction enzyme DpnI (R0176; NEB) and then heat-killed for 10 min at 65°C for 10 min to generate positive DamID fragments. These large fragments were sonicated in a volume of 200 μl, 10 mM Tris, pH 8.0) to generate fragments suitable for sequencing using a Bioruptor UCD-200 at high power, 30 s ON, 30 s OFF for 1 h in a 15 ml LoBind microfuge tube (02243100; Eppendorf). The DNA was then transferred to 15 ml TPX tubes (C30010010-1000; Diagenode) and sonicated for four rounds of 10 min (high power, 30 s ON and 30 s OFF). The DNA was transferred to new TPX tubes after each round to prevent etching of the TPX plastic. The sonication procedure yielded DNA sizes ranging from 100 to 200 bp. After sonication, the DNA was precipitated by adding 2 μl of 3 M sodium acetate, pH 5.5 and 500 μl ethanol, supplemented with 3 μl of glycerol (molecular biology grade, 20 mg/ml), and kept at −80°C for at least 2 h. The DNA mix was centrifuged at full speed for 10 min to pellet the sheared DNA with the carrier glycerol. The pellet was washed with 70% ethanol and then centrifuged again at full speed. The DNA pellet was then left to air dry. 20 μl of 10 mM Tris–HCl was used to resuspend the DNA pellet. 1 μl was quantified using the Quant-it PicoGreen dsDNA kit (P7589; Invitrogen). Sequencing library preparation was performed using the NEBNext Ultra DNA library prep kit for Illumina (E7770S; NEB), following the manufacturer’s instructions. Library quality and size was determined using a Bioanalyzer 2100 with DNA High Sensitivity reagents (5067–4626; Agilent). Libraries were then quantified using the Kapa quantification Complete kit for Illumina (KK4824; Kapa Biosystems) on an Applied Biosystems 7500 Real Time qPCR system. Samples were normalized and pooled for multiplex sequencing.

**DamID-seq data processing**

DamID-seq analysis—DamID-seq reads were processed using LADetector (available at https://github.com/thereddylab/LADetector), an updated implementation of the previously described LADetector (12) with incorporated sequence mapping. Specifically, 5‘ ends of reads were quality trimmed using a sliding window quality score average over three bases and a minimum score cutoff of 30. This was followed by trimming any matching overlap between read-ends and sequencing or DamID adaptor-primer sequence. Reads containing a DamID adaptor-primer sequence were split and adaptor-primer sequence removed. Total aligned reads were then filtered for DamID-seq reads using LADetector—DamID-seq reads were processed using LADetector, an updated implementation of the previously described LADetector (12) with incorporated sequence mapping. Specifically, 5‘ ends of reads were quality trimmed using a sliding window quality score average over three bases and a minimum score cutoff of 30. This was followed by trimming any matching overlap between read-ends and sequencing or DamID adaptor-primer sequence. Reads containing a DamID adaptor-primer sequence were split and adaptor-primer sequence removed. All resulting reads greater than 20 bp were aligned to mm9 using Bowtie (117) with parameters "--very-sensitive-m 1" Unaligned reads had 13 bases trimmed from the 5‘ end and remapped, and the resultant unmapped reads were trimmed 13 bases from the 3‘ end and remapped. Total aligned reads were assigned to DpnII bins, with reads straddling bin boundaries counting toward both. Before scoring, a value of 0.5 was added to bins with no reads. Bins falling in unaligned regions were removed before analysis. DamID scores were calculated for all non-zero bins as the log2 ratio of Dam-Lamin B1 over unfused Dam. Scores were partitioned using circular binary segmentation using the DNAcopy package in R (118, 119, 120). LADs were classified as regions >100 kb in size of positive signals, allowing for smaller regions of negative signal <10 kb in size.

**Data Availability**

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org).
Supplementary Information

Supplementary Information is available at https://doi.org/10.26508/lsa.2020000774.

Acknowledgements

We thank Rakel Tryggvadottir and Sinan Ramazanoglu for invaluable help with sequencing. This work was funded in part by (NIH) National Institutes of Health grant R21AG050132. X Wong and VE Hoskins were supported in part by NIH grant R21AG050132. VE Hoskins and JA Cutler were supported in part by (NIH) National Institute of General Medical Sciences Training Grant 5T32GM07814. M Gordon was supported by NIH Training Grant T32GM007445. Kl Reddy and VE Hoskins were supported in part by NIH grant R01GM132427.

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Conflict of Interest Statement

The authors declare that they have no conflict of interest.

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