H3.3K4M destabilizes enhancer H3K4 methyltransferases MLL3/MLL4 and impairs adipose tissue development

Younghoon Jang¹, Aaron Broun¹, Chaochen Wang¹, Young-Kwon Park¹, Lenan Zhuang¹, Ji-Eun Lee¹, Eugene Froimchuk¹, Chengyu Liu² and Kai Ge¹,*

¹Adipocyte Biology and Gene Regulation Section, Laboratory of Endocrinology and Receptor Biology, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, MD 20892, USA and ²Transgenic Core, National Heart, Lung, and Blood Institute, NIH, Bethesda, MD 20892, USA

Received April 19, 2018; Revised October 04, 2018; Editorial Decision October 07, 2018; Accepted October 10, 2018

ABSTRACT

Histone 3 lysine 4 (H3K4) methyltransferases MLL3 and MLL4 (MLL3/4) are required for enhancer activation during cell differentiation, though the mechanism is incompletely understood. We have attempted to address this issue by generating two mouse lines: one expressing H3.3K4M, a lysine-4-to-methionine (K4M) mutation of histone H3.3 that inhibits H3K4 methylation, and the other carrying conditional double knockout of MLL3/4 enzymatic SET domain. Expression of H3.3K4M in lineage-specific precursor cells depletes H3K4 methylation and impairs adipose tissue and muscle development. Mechanistically, H3.3K4M prevents enhancer activation in adipogenesis by destabilizing MLL3/4 proteins but not other Set1-like H3K4 methyltransferases MLL1, MLL2, SET1A and SET1B. Notably, deletion of the enzymatic SET domain in lineage-specific precursor cells mimics H3.3K4M expression, destabilizes MLL3/4 proteins, and prevents adipose tissue and muscle development. Interestingly, destabilization of MLL3/4 by H3.3K4M in adipocytes does not affect adipose tissue maintenance and thermogenic function. Together, our findings indicate that expression of H3.3K4M, or deletion of the enzymatic SET domain, destabilizes enhancer H3K4 methyltransferases MLL3/4 and impairs adipose tissue and muscle development.

INTRODUCTION

During cell differentiation, transcriptional enhancers are bound by lineage-determining transcription factors (LDTFs) and play a key role in regulating cell type-specific gene expression. Cell type-specific enhancers are marked by specific epigenomic features (1). Histone 3 lysine 4 (H3K4) mono-methylation (H3K4me1) is the predominant mark of a primed enhancer state. Histone 3 lysine 27 acetylation (H3K27ac) by H3K27 acetyltransferases CBP/p300 further follows H3K4me1 to mark an active enhancer state (2). There are six mammalian Set1-like H3K4 methyltransferases, each containing a catalytic SET domain that enables the deposition of methyl marks on H3K4: MLL1 (or KMT2A), MLL2 (or KMT2B), MLL3 (or KMT2C), MLL4 (or KMT2D), SET1A (or KMT2F) and SET1B (or KMT2G) (3). Among them, MLL4 is a major mammalian H3K4 mono-methyltransferase with partial functional redundancy with MLL3. MLL3 and MLL4 (MLL3/4) are required for CBP/p300-mediated enhancer activation in cell differentiation and cell fate transition (4–6). Deletion of Mll3/4 genes depletes H3K4me1 in cells and prevents the enrichment of CBP/p300-mediated H3K27ac, epigenome reader BRD4, Mediator coactivator complex, and RNA Polymerase II on enhancers. Consequently, Mll3/4 deletion prevents enhancer RNA production, cell type-specific gene induction, and cell differentiation (7). However, the role of H3K4me1 in enhancer regulation and cell differentiation, as well as H3K4me1 function in vivo, is poorly understood.

Adipogenesis and adipose tissue are useful model systems for studying cell differentiation as well as tissue development and function. Adipogenesis is mainly controlled by a cascade of sequentially expressed transcription factors (TFs) (8). Although many TFs have been implicated in adipogenesis, PPARγ and C/EBPβ are primary drivers of the induction of thousands of adipocyte genes (9,10). Adipose tissues, including white adipose tissue (WAT) and brown adipose tissue (BAT), are dynamic endocrine organs that regulate thermogenesis, energy metabolism and homeostasis (11). The study of adipogenesis and adipose tissue in vivo requires the isolation of tissues from the adipose lineage at particular developmental and functional stages. Myogenic factor 5 (Myf5) promoter-driven Cre (Myf5-
Cre) allows factors to be expressed or deleted in mouse preadipocytes, enabling a focus on adipogenesis. Conversely, Adiponectin promoter-driven Cre (Adiponectin-Cre, hereafter referred to as Adipoq-Cre) is generally expressed in differentiated adipocytes but not precursor cells, permitting study of adipocyte function (12). By crossing Mll4 conditional knockout (KO) mice with Myf5-Cre or Adipoq-Cre mice, we have shown that MLL4 is required for adipose tissue development but largely dispensable for adipose tissue maintenance (4,5). However, the roles of MLL3/4 enzymatic activities and MLL3/4-mediated H3K4me1 in adipose tissue development and function are unclear.

By tissue-specific ectopic expression of a histone H3.3 lysine-to-methionine mutant (H3.3K4M) in mice, we show that depletion of H3K4 methylation by H3.3K4M impairs adipose tissue and muscle development. By tissue-specific deletion of the enzymatic SET domain of MLL3/4 in mice, we further show that the SET domain is required for adipose tissue and muscle development. Mechanistically, expression of H3.3K4M or deletion of the SET domain prevents MLL3/4-mediated enhancer activation in adipogenesis by destabilizing MLL3/4 proteins. Interestingly, H3.3K4M does not affect adipose tissue maintenance nor the thermogenic function.

MATERIALS AND METHODS

Plasmids, antibodies and chemicals

The retroviral pQCXIP plasmids expressing FLAG-tagged wild type (WT) or mutants of histone H3.3 including K4M, K9M, K27M and K36 were described previously (13). The following homemade antibodies have been described: anti-MLL4#3 (14), anti-MLL3#3 (15) and anti-UTX (16). Anti-RbBP5 (A300-109A), anti-BRD4 (A301-985A100) and anti-MED1 (A300-793A) were from Bethyl Laboratories. Anti-SET1A/B antibodies were described previously (17). Anti-H3 (ab1791), anti-H3K4me1 (ab8895), anti-H3K4me2 (ab7766), anti-H3K27ac (ab4729) and H3K36me3 (ab9050) were from Abcam. Anti-SET1A/B antibodies were described (4).

Generation of mouse strains

To generate LSL-K4M transgenic mice, H3.3K4M and a 3' FLAG tag (H3.3-K4M-FLAG) were fused downstream of CAG promoter with a loxP-STOP-loxP cassette in the middle of the pBT346.6 plasmid (AST-3029, Applied StemCell) (Figure 2A). H3.3K4M-FLAG of pQCXIP-H3.3K4M was subcloned into the pBT346.6; after confirmation by DNA sequencing, the plasmid was linearized by Spel and Scal, gel purified and injected into zygotes harvested from C57BL/6 mice. Founder mice were identified by genotyping. For genotyping the LSL-K4M alleles, PCR was done using the following primers: 5'-CTAGCTGAGCTGCG ATGAAACATGTCG-3' and 5'-TTCGGCGCCGCGA ATCCCTAGGCGTAGTCG-3'. PCR amplified 524 bp from the LSL-K4M alleles. Total genomic DNA was extracted from mouse tail samples and PCR was done using Quick-Load Taq 2X Master Mix (NEB), following the manufacturers’ instructions.

MLL3f/f mice were obtained from Lee (18) (Figure 5A, left panel). To generate Mll4SET conditional KO mice, the loxP/FRT-flanked neomycin cassette was inserted at the 3' end of exon 51 and the single loxP site was inserted at the 5' end of exon 50 (Figure 5A, right panel). We electroporated the linearized targeted construct, which includes exons 50–51, into WT ES cells. After selection with G418, surviving clones were expanded for PCR genotyping to identify Mll4SETflox/flox ES cells, which were further micro-injected into mouse blastocysts following standard procedures. Mice bearing germ line transmission (Mll4SETfloxflox) were crossed with FLP1 mice (Jackson no. 003946) to generate Mll4SETf/f mice. For genotyping the Mll4 and Mll4SET alleles, PCRs were done using the following primers: Mll3 (5'-GTCATCGGTGTGCTGAATGA-3' and 5'-AAC CGGAAGAGAAGCTTATTG-3') and Mll4SET (5'-C AGTTGAGCTAGTCAAGTGATT-3' and 5'-TCTCAATG TGGAGGGTGTCA-3'). PCR amplified 174 bp from the wild-type Mll3 and 208 bp from the Mll3 floxed allele, or 277 bp from the wild-type Mll4 and 346 bp from the Mll4SET floxed allele.

LSL-K4M mice and Mll3f/f;Mll4SETf/f mice were crossed with Myf5-Cre (Jackson no. 007893), Cre-ER (Jackson no. 008463), or Adipoq-Cre (Jackson no. 028020) to generate LSL-K4M;Myf5-Cre, LSL-K4M;Cre-ER, LSL-K4M;Adipoq-Cre, Mll4SETf/f;Myf5-Cre, Mll3f/f;Mll4SETf/f;Myf5-Cre, or Mll3f/f;Mll4SETf/f;Cre-ER mice.

Histology and immunohistochemistry

E18.5 embryos were isolated and fixed in 4% paraformaldehyde, dehydrated in ethanol and embedded in paraffin for sectioning. Paraffin sections were stained with routine H&E or subjected to immunohistochemistry using anti-Ucp1 (ab10983; Abcam) and anti-Myosin (MF20; Developmental Studies Hybridoma Bank) antibodies as described (4).

Immortalization of primary brown preadipocytes and adipogenesis

Primary brown preadipocytes were isolated from interscapular BAT of newborn (P0) LSL-K4M;Cre-ER or Mll3f/f;Mll4SETf/f;Cre-ER pups, and immortalized by SV40T. Isolation, immortalization and adipogenesis assays of brown preadipocytes were done as described (19,20).

Western blot, qRT-PCR

Western blot of histone modifications using acid extracts or of nuclear proteins using nuclear extracts were done as described (21). Total RNA was extracted using TRIzol (Invitrogen) and reverse transcribed using ProtoScript II first-strand cDNA synthesis kit (NEB), following the manufacturers’ instructions. qRT-PCR was done using the following
Figure 1. Histone H3.3K4M and H3.3K36M mutations impair adipogenesis. (A) A schematic of the process of preadipocyte derivation from interscapular BAT of newborn pup, followed by adipogenesis assay in culture. (B–D) Immortalized brown preadipocytes were infected with retroviral vector (Vec) expressing FLAG-tagged wild type (WT) or K-to-M mutant histone H3.3, followed by adipogenesis assay. (B) Histone extracts from preadipocytes were subjected to western blot analyses using antibodies indicated on the left. Long exposure of histone H3 Western blot reveals the relative levels of ectopic H3.3 and endogenous H3. (C) Six days after induction of differentiation, cells were stained with Oil Red O. Upper panels, stained dishes; lower panels, representative fields under microscope. (D) qRT-PCR of Pparg, Cebpa, and Fabp4 expression at day 0 (D0) and day 6 (D6) of adipogenesis. Quantitative PCR data in all figures are presented as means ± SEM. Statistical comparison between groups was performed using Student’s t-test in all figures. WT at D7 was compared to K4M and K36M (∗∗P < 0.01; ∗P < 0.05) from three biological replicates (n = 3).

SYBR green primers: H3.3K4M (forward, 5′-AACCTGGGTGCCATCCACG-3′, and reverse, 5′-CGACTCTTCACGTGCCTCCT-3′), Mll3 (forward, 5′-GATTGACGCCACACTACAG-3′, and reverse, 5′-TTTCTGTATCTTGTTGG-3′), and Mll4SET (forward, 5′-GGGTGGACAGCTGTCAGAATTATT-3′, and reverse, 5′-CATAGCAGAGCTGTCAGAATTATT-3′). SYBR green primers for other genes were described previously (22, 23), and PCR was done using Luna Universal qPCR Master Mix according to the manufacturers’ instructions (NEB).

RNA-seq and ChIP-seq

RNA-seq and ChIP-seq were performed as described in detail previously with the use of Illumina HiSeq 2500 (4,6,7,24). Global differences in H3K4me1 and H3K27ac levels between control and K4M-expressing cells are not captured in ChIP-seq analysis if one uses naïve normalization, which simply normalizes the read counts by library size with the assumption that control and K4M cells have similar global histone modification levels. Therefore, H3K4me1 and H3K27ac ChIP-seq was done in the presence of Drosophila spike-in chromatin and antibody following the manufacturers’ instructions (Active Motif). For RNA-seq, we purified mRNAs using Dynabeads mRNA purification kit (Invitrogen) then synthesized double-stranded cDNAs using SuperScript Double-stranded cDNA Synthesis kit (Invitrogen), following the manufacturers’ instructions. For ChIP-seq, we collected ChIP-DNA using Dynabeads Protein A (Invitrogen) then purified DNA using QIAquick PCR Purification Kit (QIAGEN), following the manufacturers’ instructions. Library
construction for RNA-seq and ChIP-seq was completed using NEBNext Ultra II DNA Library Prep Kit (NEB), following the manufacturers’ instructions.

Computational analysis
For differentially expressed genes from RNA-seq data, gene ontology (GO) analysis was carried out using DAVID (https://david.ncifcrf.gov). For ChIP-seq data analysis with spike-in normalization, sequences were aligned to either mouse genome (mm9) or Drosophila genome (dm6). Normalization factors were determined by counting Drosophila tags and applying to mouse tags to generate normalized heat maps and profiles. To identify H3K4me1 or H3K27ac enriched regions at day 4 (D4) of adipogenesis in LSL-K4M;CreER brown preadipocytes, we used ‘SICER’ method (25) with window size of 200 bp and with an estimated false discovery rate (FDR) threshold of 10−10. Previously published ChIP-seq data sets for MLL4, PPARγ, C/EBPα, CBP, BRD4 and MED1 at day 2 (D2) of adipogenesis were used (GSE50466, GSE74189, and GSE99101) (4,6,7) and the window size was chosen to be 50 bp. To define MLL4 active enhancers during adipogenesis, we compared 13 871 MLL4 binding sites at D2 with 38 618 active enhancers (H3K4me1*H3K27ac* at D4). 8777 MLL4+ (22.7%, 8777MLL4+ active enhancers) at the center. Average profiles were plotted according to the intensity of 8777MLL4+ active enhancers at the center. Results indicate that histone H3.3K4M and H3.3K36M mutations impair adipogenesis.

H3.3K4M impairs adipose tissue and muscle development
Next, we investigated whether H3.3K4M affects adipose tissue development in vivo. We generated a conditional H3.3K4M transgenic mouse line, lox-STOP-lox-H3.3K4M (LSL-K4M) for tissue-specific expression of H3.3K4M. The insertion of four copies of SV40 stop signals (STOP) flanked by two loxP sites prevents the CAG promoter-driven expression of FLAG-tagged H3.3K4M in the absence of Cre (Figure 2A and B). We crossed LSL-K4M mice with Myf5-Cre mice to induce H3.3K4M expression specifically in somitic precursor cells of brown adipose tissue (BAT) and skeletal muscle (4). No LSL-K4M;Myf5-Cre mice were observed at the weaning age. Newborn (P0) LSL-K4M;Myf5-Cre pups were obtained but died immediately after birth from breathing malfunctions due to deficient muscle groups in the rib cage (Figure 2C–D). Immunohistochemical analysis of cervical regions of LSL-K4M;Myf5-Cre E18.5 embryos and P0 pups revealed moderate reduction in BAT and back muscle mass (Figure 2E, F and Supplementary Figure S1), indicating that expressing H3.3K4M in progenitor cells impairs adipose tissue and muscle development in vivo.

H3.3K4M destabilizes MLL3/4 proteins in adipogenesis
To confirm that H3.3K4M impairs adipogenesis in a cell autonomous manner, we crossed LSL-K4M with Cre-ER mice to obtain primary LSL-K4M;Cre-ER brown preadipocytes. After immortalization, cells were treated with 4-hydroxytamoxifen (4OHT) to delete the STOP cassette and induce FLAG-tagged H3.3K4M expression. As expected, induction of H3.3K4M expression decreased global levels of endogenous H3K4me1/2/3. Interestingly, H3.3K4M expression also decreased H3K27ac levels (Figure 3A). Consistent with our previous findings observed in embryonic stem (ES) cells (21), expressing H3.3K4M destabilized endogenous MLL3/4 as well as the MLL3/4-associated protein UTX in cells. However, H3.3K4M did not affect protein levels of other mammalian Set1-like H3K4 methyltransferases SET1A, SET1B, MLL1 and MLL2 (15) (Figure 3B).

Ectopic expression of H3.3K4M had little effect on cell proliferation (Figure 3C), but prevented adipogenesis and the induction of adipocyte marker genes such as Pparg, Cebpa, and Fabp4 (Figure 3D and E). We confirmed these H3.3K4M-driven adipogenesis defects using independent brown preadipocyte lines stably expressing WT H3.3 or H3.3K4M (Supplementary Figure S2A–C). To investigate how H3.3K4M prevents adipogenesis, we further performed RNA-seq analysis of LSL-K4M;Cre-ER preadipocytes treated with or without 4OHT. Using a 2.5-fold cut-off for differential gene expression from RNA-seq analysis, we defined genes up-regulated (5.0%, 605 genes) or down-regulated (5.8%, 709 genes) by H3.3K4M at D7 of differentiation (Figure 3F and G). Gene ontology (GO)
Figure 2. H3.3K4M impairs adipose tissue and muscle development. (A) Schematic of lox-STOP-lox-H3.3K4M (LSL-K4M) transgene and breeding strategy. The LSL-K4M transgene consists of the following elements from 5′ to 3′: a CAG promoter, quadruple copies of SV40 stop signals flanked by two loxP sites, H3.3K4M with C-terminal FLAG and HA tags, and polyadenylation signal. LSL-K4M transgenic mice were crossed with Myf5-Cre to generate mice expressing ectopic H3.3K4M in progenitors of brown adipose tissue (BAT) and muscle. The locations of PCR genotyping primers P1 and P2 are indicated by arrows. (B) PCR genotyping of LSL-K4M transgenic mice. (C) Genotype of progeny from crossing between LSL-K4M mice and Myf5-Cre mice at 3 weeks of age, post-natal day 0 (P0) and embryonic day 18.5 (E18.5). LSL-K4M; Myf5-Cre mice died soon after birth from breathing malfunction due to defects in muscles of the rib cage. (D) Representative morphology of P0 pups. (E) Representative morphology of E18.5 embryos. (F) Histological analysis of E18.5 embryos, including LSL-K4M; Myf5-Cre embryos and matched LSL-K4M controls from two different litters (#1 and #2). Sagittal sections of cervical/thoracic area were stained with H&E (left panels) or with antibodies against the BAT (B) marker UCP1 (green) and the muscle (M) marker Myosin (red) (right panels). Scale bar = 80 μm.

Analysis showed that down-regulated genes were strongly associated functionally with fat cell differentiation and lipid metabolism (Figure 3H). Because MLL3/4 are essential for adipogenesis (4), these data suggest that H3.3K4M prevents adipogenesis at least in part by destabilizing MLL3/4 proteins.

H3.3K4M prevents MLL3/4-mediated enhancer activation in adipogenesis

Next, we investigated whether H3.3K4M affects MLL3/4-mediated enhancer activation in adipogenesis. We performed ChIP-seq analyses of enhancer mark H3K4me1 and active enhancer mark H3K27ac in LSL-K4M; Cre-ER cells treated with or without 4OHT at D4 of adipogenesis. Since ChIP-seq analysis did not consider the global differences between samples, Drosophila spike-in control was used to normalize H3K4me1 and H3K27ac ChIP-seq data. Cell type-specific enhancers can be computationally defined by the binding of lineage-determining transcription factors and the enrichment of epigenomic features including H3K4me1, H3K27ac, MLL4, CBP, BRD4 and MED1 (1,2,4,6,7,29). By comparing with the published MLL4 ChIP-seq data during adipogenesis (4,6), we identified 38,618 MLL4+ active enhancers during adipogenesis. 4OHT-induced H3.3K4M expression prevented H3K4me1 and H3K27ac accumulation on MLL4+ active enhancers during adipogenesis (Figure 4A and B). H3.3K4M also prevented H3K4me1 and H3K27ac accumulation on representative MLL4+ active enhancers (e1–e5) located on gene loci of master adipogenic regulators PPARγ and C/EBPα (7). These active enhancers were defined by the binding of
Figure 3. H3.3K4M destabilizes MLL3/4 proteins in adipogenesis. Immortalized LSL-K4M;Cre-ER brown preadipocytes were treated with 4-hydroxytamoxifen (4OHT) to induce ectopic H3.3K4M expression, followed by adipogenesis assay. (A, B) H3.3K4M destabilizes MLL3/4 proteins. Histone extracts (A) or nuclear extracts (B) were analyzed by Western blot using antibodies indicated on the left. (C–E) H3.3K4M prevents adipogenesis and induction of adipocyte genes. (C) Cell growth rates. 10^5 preadipocytes were plated at D0 and the cumulative cell numbers were determined every day for 5 days. (D) Oil red O staining at day 7 (D7) of adipogenesis. (E) qRT-PCR of H3.3K4M-FLAG, Pparg, Cebpα and Fapb4 expression at D0 and D7 of adipogenesis. –4OHT at D7 was compared to +4OHT (\*P < 0.01, *P < 0.05; n = 3). (F–H) RNA-seq analyses were performed at D7 of adipogenesis. (F, G) Identification (F) and heat maps (G) of down- or up-regulated genes in H3.3K4M-expressing cells. The cut-off for differential expression is 2.5-fold. (H) Gene ontology (GO) analysis of gene groups defined in (F).
Figure 4. H3.3K4M prevents MLL3/4-mediated enhancer activation in adipogenesis. 4OHT-treated LSL-K4M;Cre-ER brown preadipocytes were collected at D4 of adipogenesis for ChIP-seq of H3K4me1 and H3K27ac. *Drosophila* spike-in control was used to normalize ChIP-seq data. (A–C), ChIP-qPCR of MLL4, H3K4me1, H3K27ac, BRD4, MED1 and Pol II, and qRT-PCR of eRNAs (D, E). (A, B) Heat maps (A) and average profiles (B) around MLL4+ active enhancers during adipogenesis. (C) Genome browser view of PPARγ/H9253, C/EBPα/H9251, MLL4, CBP, BRD4, MED1, H3K4me1 and H3K27ac (data obtained from published data sets GSE50466, GSE74189, and GSE99101 (4,6,7), as well as current data set GSE110972) on Pparg and Cebpa gene loci during adipogenesis with schematic of genomic locations of representative MLL4+ active enhancers (e1–e5). (D) ChIP-qPCR analyses of indicated factors are shown on enhancers e1–e5 at D0 and D4 of adipogenesis. An enhancer of constitutively expressed gene Jak1 was chosen as a negative control (n). (E) qRT-PCR of eRNA transcription on enhancers e1–e5 at D0 and D4 of adipogenesis. −4OHT at D4 was compared to +4OHT (∗∗∗P < 0.01, ∗∗P < 0.05; n = 3). (F) Proposed model showing that H3.3K4M prevents enhancer activation in adipogenesis by destabilizing MLL3/4.
PPARγ and C/EBPα, and by the enrichment of H3K4me1, H3K27ac, ML4, CBP, BRD4 and MED1 during adipogenesis (4,6,7) (Figure 4C). Further, ChIP-qPCR analyses revealed that, on representative ML4 active enhancers (e1–e5), H3.3K4M markedly reduced the occupancy of ML4, BRD4, MED1 and Pol II in addition to ML3/ML4-mediated H3K4me1 and CBP/p300-mediated H3K27ac (Figure 4D). H3.3K4M also decreased eRNA production from ML4 active enhancers (Figure 4E). Together, these results suggest that H3.3K4M prevents ML3/4-mediated enhancer activation in adipogenesis.

Deletion of the enzymatic SET domain of ML3/4 prevents adipose tissue and muscle development

ML3 and ML4 are partially redundant as major H3K4 mono-methyltransferases on enhancers in cells (4). In a separate attempt to investigate the functional role of ML3/4 enzymatic activity in differentiation and development in vivo, we used two conditional mouse KO lines targeting the enzymatic SET domain of ML3 and ML4 (ML3f and ML4f, Figure 5A–B). In the ML3f mice, exons 57 and 58, which encode critical amino acids of the SET domain, were flanked by two loxP sites (18). In the ML4f mice, exons 50 and 51, which encode the entire SET domain, were flanked by two loxP sites. Cre-mediated deletion of these exons would result in the production of enzyme-dead ML3/4.

We first crossed ML4f mice with Myf5-Cre mice. The resulting ML4SETf;Myf5-Cre mice survived until birth. E17.5–18.5 ML4SETf;Myf5-Cre embryos were unable to breathe and died immediately after isolation, displaying an abnormal hunched posture and impaired back muscles (Supplementary Figure S3A and B). These embryos showed similar BAT mass but altered BAT morphology and impaired muscle mass compared to WT (Supplementary Figure S3C), possibly due to a compensatory effect of ML3 as ML4 KO alone does not affect ML3 protein level (Supplementary Figure S4A and B). To eliminate the compensatory effect, we crossed ML4SETf with ML3f and Myf5-Cre mice to delete the SET domain-coding portion of both ML3 and ML4 genes in progenitor cells of BAT and muscle lineages. The resulting F18.5. ML3f,ML4SETf;Myf5-Cre (conditional double KO, DKO) embryos were unable to breathe and died immediately after isolation. These embryos showed profound reduction of BAT as well as muscle mass (Figure 5C–E). These data indicate that deletion of the enzymatic SET domain of ML3/4 prevents adipose tissue and muscle development.

Deletion of the enzymatic SET domain inhibits adipogenesis by destabilizing ML3/4

To confirm that deletion of the enzymatic SET domain of ML3/4 prevents adipogenesis in a cell autonomous manner, we crossed ML3f,ML4SETf with Cre-ER mice to obtain primary ML3f,ML4SETf;Cre-ER brown preadipocytes. After immortalization, cells were treated with 4OHT to delete exons encoding the SET domain of ML3/4. Consistent with our previous finding that ML3 and ML4 are major H3K4 mono- and di-methyltransferases in cells (4,5), deletion of the enzymatic SET domain in preadipocytes decreased global levels of endogenous H3K4me1/2 but not H3K4me3 (Figure 6A).

In addition, deleting the enzymatic SET domain destabilized endogenous ML3/4 and reduced protein levels of ML3/4-associated UTX in cells (Figure 6B), which is consistent with a recent report that deleting the SET domain destabilizes ML3/4 in ES cells (30).

Deletion of the enzymatic SET domain of ML3/4 had little effect on cell proliferation (Figure 6C), but prevented adipogenesis and the induction of Pparγ, Cebpα, and Fabp4 (Figure 6D and E). RNA-seq analysis at D7 of differentiation confirmed the deletion of target exons of both ML3 and ML4 loci (Figure 6F). We further performed RNA-seq analysis of ML3f,ML4SETf;Cre-ER preadipocytes treated with or without 4OHT. Using a 2.5-fold cut-off for differential expression from RNA-seq analysis, we defined genes up-regulated (51%, 623/12 184) or down-regulated (7.3%, 890/12 184) by deletion of the enzymatic SET domain at D7 of differentiation (Figure 6G and H). GO analysis showed that down-regulated genes were strongly functionally associated with fat cell differentiation and lipid metabolism (Figure 6I). These data suggest that deletion of the enzymatic SET domain inhibits adipogenesis by destabilizing ML3/4.

H3.3K4M expression mimics ML3/4 SET domain deletion in preventing adipogenesis

Next, we generated scatter plots of gene expression changes using RNA-seq data from cells with deletion of ML3/4 SET domain (DKO) and cells with expression of H3.3K4M (K4M) (Figure 7A). A combined scatter plot revealed that down- or up-regulated genes were highly correlated between DKO and K4M cells (Figure 7B). We found that the majority of down or up-regulated genes at D7 of differentiation were shared by DKO and K4M (Figure 7C and D). The shared down-regulated genes were highly associated functionally with fat cell differentiation and lipid metabolism, whereas shared up-regulated genes were highly associated with cell proliferation and development (Figure 7E and F). These data suggest that ectopic H3.3K4M mimics the effect of ML3/4 SET domain deletion in preventing adipogenesis.

H3.3K4M does not affect adipose tissue maintenance and thermogenic function

We also investigated the role of H3K4 methylation in adipose tissue maintenance and function in mice. For this purpose, we crossed LSL-K4M mice with Adipoq-Cre mice to achieve adipocyte-selective expression of H3.3K4M in vivo. Histone western blot of BAT from LSL-K4M;Adipoq-Cre mice showed that ectopic expression of H3.3K4M depleted endogenous H3K4me1/2/3 and ML3/4 in BAT (Figure 8A and B). LSL-K4M;Adipoq-Cre mice did not show any discernable differences in body weight, fat/lean mass, or adipose tissue mass relative to control (LSL-K4M) mice at 8–10 weeks of age (Figure 8C–E). In the BAT, induction of H3.3K4M expression was successful in LSL-K4M;Adipoq-Cre mice, but the expression levels of adipocyte identity genes Pparγ, Cebpα, and Fabp4...
Figure 5. Deletion of the enzymatic SET domain of MLL3/4 prevents adipose tissue and muscle development. (A) Conditional KO mouse lines targeting the SET domain of MLL3 and MLL4. Schematics of WT allele, targeted allele, conditional KO (flox) allele and KO allele are shown in upper panels. Deletion of neomycin selection cassette by FLP recombinase generates the flox allele. The locations of PCR genotyping primers P1-P4 are indicated by arrows. Expected protein domains and molecular weights are shown in bottom panels. (B) PCR genotyping of flox and WT alleles using P1-P4 primers. (C) Genotypes of progeny at E18.5 from crossing between Mll3<sup>f</sup;/<sup<f</sup>;Mll4SET<sup>f</sup>/<sup<f</sup> and Mll3<sup>f</sup>/+;Mll4SET<sup>f</sup>/+;Myf5-Cre. Mll3<sup>f</sup>/+;Mll4SET<sup>f</sup>/+;Myf5-Cre (conditional double KO, DKO) mice died immediately after cesarean section from breathing malfunction due to defects in muscles of the rib cage. (D) Representative morphology of E18.5 embryos. (E) Histological analysis of E18.5 embryos. Sagittal sections of cervical/thoracic area were stained with H&E (upper panels) or with antibodies against the BAT (B) marker UCP1 (green) and the muscle (M) marker Myosin (red) (lower panels). Scale bar = 80 μm.

were similar between LSL-K4M;Adipoq-Cre and control mice (Figure 8F). This phenotype is consistent with what we have reported for the Mll4SET<sup>f</sup>/Adipoq-Cre mice, in which MLL4 is dispensable for the maintenance of differentiated adipocytes in vivo (5).

We further asked about the functional consequence of ectopic H3.3K4M expression in adipocytes in vivo. For this purpose, we acutely exposed the mice to environmental cold (6°C) up to 6 h. LSL-K4M;Adipoq-Cre mice maintained normal body temperatures, were cold tolerant and behaved similarly to control mice in the cold tolerance test (Figure 8G). After cold exposure, expression levels of thermogenesis genes Ucp1, Dio2, and Elovl3 were similarly induced in BAT of LSL-K4M;Adipoq-Cre and control mice (Figure 8H). Together, our data indicate that while H3.3K4M impairs adipose tissue development, it does not affect the maintenance nor the thermogenic function of adipose tissues.
Figure 6. Deletion of the enzymatic SET domain inhibits adipogenesis by destabilizing MLL3/4. Immortalized Mll3f/f, Mll4SETf/f, Cre-ER brown preadipocytes were treated with 4OHT to induce deletion of the exons encoding the SET domain of MLL3 and MLL4 proteins, followed by adipogenesis assay. (A, B) Deletion of the enzymatic SET domain destabilizes MLL3/4 proteins. Histone extracts (A) or nuclear extracts (B) from preadipocytes were analyzed by western blot using antibodies indicated on the left. (C–E) Deletion of the SET domain of MLL3/4 prevents adipogenesis. (C) Cell growth rates. (D) Oil red O staining at D7 of adipogenesis. (E) qRT-PCR of Mll3, Mll4SET, Pparg, Cebpa and Fabp4 expression at D0 and D7 of adipogenesis. (F–I) RNA-seq analyses were performed at D7 of adipogenesis. (F) Genome browser views of RNA-seq analysis on Mll3 and Mll4 loci. The targeted exons are highlighted in red boxes. (G, H) Schematic of identification (F) and heat maps (H) of down- or up-regulated genes in double KO (DKO) cells. The cut-off for differential expression is 2.5-fold. (I) GO analysis of gene groups defined in (G).
Figure 7. H3.3K4M expression mimics MLL3/4 SET domain deletion in preventing adipogenesis. (A) Scatter plots of down- or up-regulated genes in H3.3K4M-expressing or MLL3/4 SET domain DKO cells compared to the respective control (Ctrl) cells. RNA-seq data were from Figures 3 and 6. The cut-off for differential expression is 2.5-fold. Down- or up-regulated genes are depicted in green and red, respectively. (B) Combined scatter plot showing correlation of fold changes of gene expression by H3.3K4M expression and by MLL3/4 SET domain DKO. (C, D) Venn diagram showing the overlap of down- or up-regulated genes in H3.3K4M-expressing and MLL3/4 SET domain DKO cells. (E, F) GO analysis of gene groups defined in (C, D).

**DISCUSSION**

Site-specific histone methylations are generally correlated with gene activation or gene repression. To investigate the role of site-specific histone methylations in cell differentiation and development, we screened several K-to-M mutants of H3.3 and found that H3.3K4M and H3.3K36M mutations impair adipogenesis in cell culture. Mechanistically, H3.3K4M destabilizes MLL3/4 proteins but not other mammalian Set1-like H3K4 methyltransferases SET1A, SET1B, MLL1 and MLL2. Consequently, H3.3K4M prevents MLL3/4-mediated enhancer activation in adipogenesis. Using tissue-specific expression of H3.3K4M in mice, we next showed that H3.3K4M inhibits adipose tissue and muscle development. However, H3.3K4M does not affect adipose tissue maintenance nor the thermogenic function of BAT. Using tissue-specific deletion of the enzymatic SET domain of MLL3/4 in mice, we demonstrated that the SET domain is required for adipose tissue and muscle development. Mechanistically, deletion of the SET domain destabilizes MLL3/4 proteins. Notably, H3.3K4M expression mimics MLL3/4 SET domain deletion in preventing adipogenesis. Together, our findings suggest that H3.3K4M destabilizes enhancer H3K4 methyltransferases MLL3/4 and impairs adipose tissue development (Figure 4F).

Consistent with our observation in preadipocytes that the SET domain is required for MLL3/4 protein stability, Dorighi et al. showed recently that MLL3/4 proteins with
Figure 8. H3.3K4M does not affect adipose tissue maintenance and thermogenic function. All data were from 8 to 10-week-old male mice fed with regular diet. (A, B) Histone extracts (A) or nuclear extracts (B) from BAT were analyzed by Western blot using antibodies indicated on the left. (C) Representative pictures of BAT, inguinal WAT (iWAT), epididymal WAT (eWAT), and liver. (D) Fat mass, lean mass, and total body weight were measured by MRI (n = 9 per group). (E) The average tissue weights are presented as % of body weight (n = 6 per group). (F) qRT-PCR of H3.3K4M, Pparg, Cebpa and Fabp4 expression in BAT (n = 6 per group). (G, H) Cold tolerance test. Mice were housed at room temperature (RT, 22°C) and then in a cold room (6°C) for 6 h (n = 6 per group). (G) Body temperatures. (H) qRT-PCR of gene expression in BAT after 6 h. LSL-K4M was compared to LSL-K4M; Adipoq-Cre (**P < 0.01).

SET domain deletion are unstable in ES cells (30). However, Rickels et al. reported that MLL3/4 proteins with SET domain deletion are stable in ES cells (31). In our study, deleting exons 57 and 58 results in a destabilized MLL3 protein in cells (Figure 6B). In contrast, Rickels et al. deleted exons 56, 57 and 58 without affecting MLL3 protein stability in cells (31). The full-length mouse MLL4 protein has 5588 aa. Dorighi et al. showed that a truncated MLL4 protein containing amino acids 1–5482 is unstable in ES cells (30). In contrast, Rickels et al. showed that a more severely truncated MLL4 protein containing amino acids 1–5402 is stable in ES cells (31). Future studies are needed to confirm that the SET domain is required for MLL3/4 protein stability in cells.

A previous report showed that H3K27M inhibits the H3K27 methyltransferase activity of Polycomb repressive complex 2 through interaction with the enzymatic subunit EZH2 (28). Our data (Figure 1B and Figure 3A) and previous reports (13, 28) show that H3.3K4M reduces global levels of H3K4me1/2/3, suggesting that H3.3K4M inhibits the enzymatic activities of multiple SET1-like H3K4 methyltransferases including MLL3/4. The detailed biochemical mechanisms remain to be determined.
Consistent with our previous report that ectopic expression of H3.3K4M reduces endogenous levels of MLL3/4 and MLL3/4-associated protein UTX in ES cells (21), we now show that H3.3K4M destabilizes MLL3/4 and UTX proteins in preadipocytes and BAT. How does H3.3K4M specifically destabilize MLL3/4 but not other Set1-like H3K4 methyltransferases in ES cells? (i) Particular sequence similarities between the H3.3K4M-interacting SET domains of MLL3/4 that are not conserved in other Set1-like H3K4 methyltransferases could make MLL3/4 more susceptible for proteasome degradation. (ii) Alternatively, we described previously the possibility that MLL3/4 protein stability could be regulated by cellular H3K4 methylation levels (21). (iii) Another possibility is that MLL3/4 protein stability is regulated by auto-methylation. H3.3K4M may inhibit MLL3/4 enzymatic activity, thereby reducing auto-methylation and causing their destabilization. (iv) Also, MLL3/4 may methylate associated proteins or other non-histone proteins, which regulate their stability. (v) Finally, it is possible that the reduced MLL3/4 protein levels by H3.3K4M are due to the expression changes of genes regulated by MLL3/4. Further study will be needed to determine mechanistically how H3.3K4M regulates MLL3/4 protein stability.

Although H3K4me1 is the predominant mark of a primed enhancer state, it is unclear whether H3K4me1 affects or simply correlates with enhancer activation in cell differentiation and development. Interestingly, the latest two studies from one group reported that MLL3/4-dependent H3K4me1 has an active role at enhancers to facilitate binding of the chromatin remodeler SWI/SNF complex and recruitment of the chromatin organization regulator cohesin complex to orchestrate long-range chromatin interactions in ES cells (32,33). These findings imply that H3K4me1 is not simply a correlate outcome in enhancer regulation. Conversely, using CRISPR to generate catalytically inactive MLL3/4, another recent study surprisingly reported that MLL3/4 proteins, rather than MLL3/4-mediated H3K4me1, are required for enhancer activation and gene transcription in undifferentiated ES cells. However, the role of MLL3/4-mediated H3K4me1 in ES cell differentiation was not elucidated (30). Thus, despite various efforts to uncover the role of H3K4me1 in enhancer function, it is still unclear whether H3K4me1 controls enhancer activation in cell differentiation. Therefore, future work will be needed to clarify the role of MLL3/4-mediated H3K4me1 in enhancer function during cellular differentiation and animal development.

DATA AVAILABILITY

RNA-seq (D7) and H3K4me1/H3K27ac ChIP-seq (D4) datasets generated in the paper have been deposited in NCBI Gene Expression Omnibus under accession number GSE110972.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

We thank Zhiguo Zhang for retroviral plasmids expressing WT or mutant histone H3.3, NIDDK Genomics Core for sequencing, Victoria Noe-Kim and Weiqun Peng for helping with computational analyses, and Jae W. Lee for Mll3+/− mice.

FUNDING


REFERENCES


