Mechanism for DPY30 and ASH2L intrinsically disordered regions to modulate the MLL/SET1 activity on chromatin

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Recent cryo-EM structures show the highly dynamic nature of the MLL1-NCP (nucleosome core particle) interaction. Functional implication and regulation of such dynamics remain unclear. Here we show that DPY30 and the intrinsically disordered regions (IDRs) of ASH2L work together in restricting the rotational dynamics of the MLL1 complex on the NCP. We show that DPY30 binding to ASH2L leads to stabilization and integration of ASH2L IDRs into the MLL1 complex and establishes new ASH2L-NCP contacts. The significance of ASH2L-DPY30 interactions is demonstrated by requirement of both ASH2L IDRs and DPY30 for dramatic increase of processivity and activity of the MLL1 complex. This DPY30 and ASH2L-IDR dependent regulation is NCP-specific and applies to all members of the MLL/SET1 family of enzymes. We further show that DPY30 is causal for de novo establishment of H3K4me3 in ESCs. Our study provides a paradigm of how H3K4me3 is regulated on chromatin and how H3K4me3 heterogeneity can be modulated by ASH2L IDR interacting proteins.
Cells are complex, information-processing centers that handle an immense flow of signals often leading to fine tuning the expression of genes. To achieve exquisite regulation, chromatin post-translational modifications (PTMs) have evolved to demarcate, among a mosaic of functions, actively transcribed genes from the inactive ones. The mixed-lineage leukemia (MLL) family of histone methyltransferases (HMTs) catalyzes the deposition of histone H3 lysine 4 methylation (H3K4me) associated with active transcription. H3K4 methylation is highly enriched at gene promoters and distal regulatory enhancers, and plays a pivotal role in the recruitment of basal transcription machinery and chromatin remodeling complexes. It also promotes long-range chromatin interactions and higher-order chromatin organization. The dynamic interplay between H3K4me and co-transcriptional processes has been reported. Human genetic studies have corroborated ordered regions in the chromatin-modifying complexes may exert regulatory functions amongst themselves. Recent cryo-EM studies of the MLL1/3-NCP complexes show that DPY30 does not make direct contact with the NCP, but rather engages with a physiological substrate (i.e., NCP). These studies, together with solution structures of MLL1, show a surprising dynamic nature of the MLL1 core complex, especially the MLL1 domain and the RbBP5-NCP interface. Despite these studies, regulation of structural dynamics of the MLL1 complex on the NCP and its functional implication remain largely unknown.

Compared to the well-studied WDR5, RbBP5, and ASH2L SPRY proteins, the function of DPY30 and the extended intrinsically disordered regions (IDRs) of ASH2L in the MLL1 complex remains a mystery. The biochemically defined minimal core complex showed negligible DPY30 contribution to the activity of the MLL1/SET1 family of enzymes using recombinant histone H3 or peptic acid as substrates. On the other hand, DPY30 is capable of regulating global H3K4 methylation in cells and knockdown or knockout leads to global reduction of H3K4me3 in embryonic stem cells (ESCs) and hematopoietic stem cells. It is proposed as a potential therapeutic target for MLL1-rearranged leukemia. The conflicting reports of the minimal in vitro DPY30 activity versus its importance in regulating H3K4me3 in cells remain unresolved.

Here we show that DPY30 greatly stimulates MLL1 activity on the NCP. By combined NMR, SAXS, cryo-EM and biochemical approaches, we find that DPY30 functions through the extended IDRs of ASH2L to restrict the rotational dynamics of the MLL1 complex on the NCP, thereby promoting H3K4 methylation at higher methylation states. The NCP-specific regulation by DPY30 and ASH2L IDRs generally applies to all MLL/SET1 family enzymes. Cellular studies further confirm the importance of DPY30 in de novo establishment of H3K4me3 on chromatin. Taken together, we have established a paradigm of how the disordered regions in the chromatin-modifying complexes may exert loci-specific histone methylation and confer heterogeneity in the cellular epigenetic landscape.

Results
Activity of the MLL/SET1 family enzymes on the NCP requires DPY30. To examine the regulation of the MLL1 methyltransferase activity on the NCP in vitro, we performed the HMT assays using either recombinant histone H3 or NCP as substrates. The overall activity of the MLL1 core complex was much higher on the NCP as compared to recombinant histone H3 (Fig. 1a and Supplementary Fig. 1a). DPY30 was essential for the drastic increase of H3K4 methylation on the NCP (Fig. 1a), especially for higher H3K4 methylation states (i.e., H3K4me2 and H3K4me3) at the expense of H3K4me1 (Fig. 1a). In contrast, DPY30 had no effect on MLL1 activity or processivity when recombinant H3 was used as the substrate (Fig. 1a and Supplementary Fig. 1b), consistent with the previous studies. To test whether DPY30-dependent regulation on the NCP is a general mechanism for all MLL/SET1 family enzymes, we examined H3K4 methylation by NCP-2 and SET1A/1B in the presence or absence of DPY30. As shown in Fig. 1b and Supplementary Fig. 1b, DPY30 was able to significantly enhance methylation activity of all MLL/SET1 complexes in an NCP-specific manner. Domain mapping confirmed that the dimerization domain (DD, 45–90) of DPY30, which forms a hydrophobic groove that directly interacts with the ASH2L Sdc-DPY30-Interacting domain (SDI, 504–534), was sufficient to stimulate MLL1 activity on the NCP (Fig. 1c).
of ASH2L Linker IDRs (Fig. 3a) identified three highly conserved regions, 247–251, 252–263 and 275–285 (Supplementary Fig. 2a), that were critical for DPY30-dependent regulation (Fig. 3b–d). These results highlight a previously uncharacterized function of ASH2L IDRs in regulating MLL1 activity on the NCP.

We next examined whether IDRs in other complex subunits are important for DPY30-mediated HMT stimulation. The potential IDRs in the MLL1 core complex include the RbBP5 C-terminus (aa. 382–538) and a segment of the SET domain between the WIN motif and the catalytic domain (aa. 3767–3812)30. Sequential C-terminal RbBP5 truncations were tested and none of them abolished DPY30-mediated HMT stimulation (Supplementary Fig. 2b). Notably, larger deletion of RbBP5 C-terminus lowered the overall HMT activities (Supplementary Fig. 2b), consistent with previous studies for yeast homolog Swd1 in the SET1 complex53,54. To test the MLL1 SET IDR, MLL1SETIL (3813–3969) was used so that the MLL1 complex remains active in the absence of the WIN motif or WDR530. Removal of the SET IDR did not affect DPY30-dependent stimulation (Supplementary Fig. 2b). Circumvention of WDR5 in the MLL1SETIL (3813–3969)-containing core complex also indicates that WDR5 is dispensable for DPY30-mediated stimulation. These results suggest that ASH2L IDRs are necessary and sufficient for DPY30-dependent HMT stimulation on the NCP.

DPY30 induces widespread NMR spectra changes in ASH2L IDRs. To evaluate the effects of DPY30 binding on global ASH2L structure and to explore the mechanism by which DPY30 and ASH2L IDRs regulate MLL1 activity, we performed methylTROSY NMR on 13CH3-labeled Ile-Leu-Val (ILV) ASH2L202–534, in the presence of stoichiometric amount of unlabeled RbBP5 peptide (330–363), the minimal region for ASH2L binding (see Methods for details). We identified ~65% of the 100 anticipated peaks in 13CH3-labeled ILV ASH2L (Fig. 4a and Supplementary Fig. 3b and 3c, red peaks). Chemical shift changes of some apo-state peaks were also observed (Fig. 4a). To further characterize these newly appeared peaks, we carried out residue-specific quantification completed using ImageJ108 with %activity calculated relative to wild-type ASH2L-containing complex108.
methyl-assignments by mutagenesis on the ASH2L202–534-DPY30 complex (Supplementary Fig. 4a–d)55. About 60% of total methyl peaks were unambiguously assigned (see Supplementary Table 1), owing to their dispersed chemical shifts. Interestingly, the majority of the DPY30-induced new peaks corresponded to residues in the ASH2L Linker and Loop IDRs (Fig. 4a, blue and orange, respectively). A number of peaks corresponding to residues in the SPRY domain (Fig. 4a, green) were also perturbed (e.g., I274, V287, I300, V322, I488) or newly appeared (e.g., L291, L350). Importantly, deletion of either Linker (blue) or Loop (orange) IDRs in ASH2L (modeled in Supplementary Figs. 5 and 6) abolished DPY30-induced changes in NMR spectra (Supplementary Fig. 6a–c, right). The NMR results suggest that DPY30 mainly affects ASH2L IDRs and the DPY30-dependent NMR changes require all ASH2L IDRs.

Small-angle X-ray scattering (SAXS) of ASH2L and ASH2L/DPY30. The DPY30-dependent changes of ASH2L IDRs in NMR spectra can be due to alterations of inter-or intra-molecular interactions or stabilization of a particular conformation. To gain more insights into these possibilities, we performed SAXS experiment for ASH2L, DPY30, and the ASH2L/DPY30 complex. The molecular weight for ASH2L was estimated to be 65 KDa by the SAXS experiment. Since the combined mass of ASH2L (60.12 KDa) and RbBP5330–363 (4.07 KDa), which was included in all ASH2L SAXS samples (see "Methods"), is ~64 kDa, ASH2L is likely monomeric in solution. This excludes the possibility that DPY30 functions through resolving intermolecular interactions of ASH2L IDRs. Furthermore, SAXS data show that pair distance distribution function of ASH2L had a peak around 30 Å and decreased smoothly (Supplementary Fig. 7a), suggesting that the structural domains in ASH2L were probably not locked in a rigid configuration. As shown in Supplementary Fig. 7a, ASH2L/DPY30 had a similar Dmax (~140 Å) as compared to ASH2L despite a 30% increase in size (Supplementary Fig. 7a). It suggests that ASH2L in the DPY30/ASH2L complex is probably in a more compact conformation. Interestingly, analysis using ensemble-
optimized method (EOM)\textsuperscript{36} identified two distinguishable ASH2L populations in both the \(D_{\text{max}}\) and \(R_g\) plots (Supplementary Fig. 7b), suggesting that ASH2L is likely in a structural equilibrium between two largely different conformations, with one more extended than the other (Supplementary Fig. 7b). We were not able to perform EOM analysis for ASH2L/DPY30 due to method limitation\textsuperscript{36}. Taken together, we speculate that DPY30 binding may shift the structural equilibrium of ASH2L and stabilize ASH2L IDRs in a more compact conformation. This is consistent with the DPY30-dependent appearance of ASH2L NMR peaks with well-dispersed chemical shifts (Fig. 4a).

**Molecular modeling of the DPY30-ASH2L complex.** While it is challenging to determine the exact conformation(s) of the dynamic ASH2L IDRs in the apo-state, we were able to build a structural model to visualize ASH2L IDRs in the DPY30-bound state. The molecular model of the human ASH2L-DPY30 is based on the co-crystal structure of the ySET1 complex subunits Bre2-Sdc1 (PDB code: 6CHG)\textsuperscript{53} as well as crystal structures of the human ASH2L SPRY domain (without Loop IDR, PDB code: 3TOJ)\textsuperscript{47} (Fig. 4b, see “Methods”). When we mapped the residues that showed DPY30-dependent chemical shift in the NMR spectra onto this structural model, the close spatial proximity of these residues was apparent (Fig. 4b). They clustered together in the IDRs (Supplementary Fig. 5a) and SPRY regions (Supplementary Fig. 5b). In this model, ASH2L IDRs, the SPRY domain, and SDI adopt a compact triangular structural arrangement upon interacting with DPY30 (Fig. 4b). ASH2L IDRs form an ordered three-strand \(\beta\)-sheet, comprised of highly conserved residues 247–252 from Linker IDR and residues 416–428 from Loop IDR (Supplementary Fig. 8a, red box). In addition to the \(\beta\)-sheet structure, residues 252–263 and 275–286 of the Linker IDR also adopt a \(\beta\)-sheet-like conformation next to SDI (Supplementary Fig. 8a, blue box), enclosing a binding interface for the \(\alpha\)-helical SDI (orange) and DPY30 (Supplementary Fig. 8b). Although this is only a computational model, many highlighted structural
DPY30/ASH2L IDRs restrict the rotational dynamics of the MLL1 complex on the NCP. Recently, we and others have solved the cryo-EM structure of the MLL1-NCP complex37,38. It reveals the overall architecture of the five component MLL1 core complex with the NCP. In the MLL1-NCP structure, ASH2L binds to the NCP at DNA superhelical loop (SHL) 7 (Fig. 5a), which together with RbBP5 at SHL 1.5, allows MLL1SET to bind above the nucleosome dyad37. To understand the molecular mechanism by which DPY30 regulates MLL1 activity on the NCP, we determined the single-particle cryo-EM structure of the human recombinant MLL1RWSAD complex (4-MLL1), containing four of the five core proteins, i.e. RbBP5 (aa 1–538); WDR5 (aa 22–334); MLL1SET (aa 3762–3969); and ASH2LΔSDI (aa 1–504), bound to the NCP (4-MLL1-NCP). Overall, a total of 1288 K particles were picked from 6242 micrographs collected from 300 keV Titan Krios equipped with the K2 summit direct detector (Supplementary Fig. 9). After several rounds of heterogeneous refinement using cryoSPARC57, we isolated four different subclasses of 4-MLL1-NCP (Class01, 02, 03, and 05). The best behaving particles were further selected from each subset of the 4-MLL1-NCP images after focused refinement and subsequent 3D classification in RELION (Supplementary Fig. 9)58. In the end, we obtained three different subclasses of 4-MLL1-NCP structures (Class 01, 02, and 05, Fig. 5b–d and Supplementary Fig. 9). The overall resolution of these structures ranged from 4.6 Å to 6.9 Å (Supplementary Fig. 10), which were sufficient to dock coordinates of the MLL1 core components and the NCP from our previous MLL1RWSAD-NCP structure (PDB ID: 6PWV [https://doi.org/10.2210/pdb6pwv/pdb])37. In comparison to the MLL1RWSAD-NCP complex (or 5-MLL1-NCP, Fig. 5a)37, the 4-MLL1-NCP complexes displayed much higher dynamics at the ASH2L-NCP

Fig. 4 ASH2L IDRs undergo significant conformational changes upon DPY30 binding. a DPY30 binding induces drastic conformational change in ASH2L. Superimposed methyl-TROSY spectra of [2H, 13CH3-ILV] ASH2L202–534 in the absence (black) or presence (red) of DPY30. The labels indicate assigned residues in the DPY30-bound state. Underlined residues are newly appeared peaks upon DPY30 addition. b Computation model for ASH2L IDRs after DPY30 binding. Underlined residues in (a). are presented as spheres. These residues clustered together into a compact structure in this model. For both a and b, SPRY domain is shown in green, Linker IDR is shown in blue, Loop IDR is shown in orange.
While the majority of the 5-MLL1-NCP complexes anchored on the NCP with RbBP5 and ASH2L at DNA SHL 1.5 and 7, respectively, the 4-MLL1-NCP complex adopted multiple modes of interaction. With RbBP5 anchoring near SHL 1.5, ASH2L binding sites varied from SHL 7 to SHL 4.5 among different subclasses (Fig. 5b, d). Furthermore, local ASH2L binding dynamics on the NCP also increased significantly in the absence of DPY30, as demonstrated by extremely low or complete loss of ASH2L IDR density in a significant subset of the structures (Fig. 5c, d).

The molecular modeling using the iterative template-based fragment assembly refinement (I-TASSER) method showed that ASH2L IDRs make multiple contacts with nucleosomal DNA (Supplementary Fig. 11a). In addition to the conserved basic residues (205-KRK-207) that contribute to overall MLL1 activity on the NCP, DPY30-induced ASH2L changes appear to enable ASH2L residues 419–421, which reside on a short loop between the newly formed three-stranded β-sheet, to provide another contact with DNA (Supplementary Fig. 11a). Consistent with the modeling, K419A/K421A mutation or deletion of 419–421 significantly reduced or abolished DPY30-dependent regulation of MLL1 activity, respectively (Supplementary Fig. 11b). These results are consistent with a model that DPY30 functions through ASH2L IDRs to restrict rotational dynamics of the MLL1 complex on the NCP and promote productive H3K4 methylation (see "Discussion").

DPY30 is essential for establishing de novo H3K4me3 in E14 ESCs. To investigate the function of DPY30 in establishing H3K4me3 in cells, we first examined the correlation of DPY30 binding and H3K4me3 at MLL1 binding sites in E14 ESCs.

Fig. 5 Cryo-EM structure of the 4-MLL1-NCP complexes. a Front views of the MLL1RWSAD-NCP (PDB ID: 6PWV [https://doi.org/10.2210/pdb6pwv/pdb] and EMDB: EMD-20512)37. The 90° top view, on right, shows the relative position of the anchoring points. DNA, yellow; DPY30, blue; ASH2L, orange; WDR5, light green; RbBP5, cyan. The NCP (histone octamer core, orange circle; DNA, black), RbBP5 (cyan circle), and DPY30 (blue circle), and the remaining MLL components were displayed as a gray bar indicating the orientation of the MLL1 complex. b-d Front view of the 4-MLL1-NCP structures. The 90° top view shows its anchoring points on the NCP. The missing EM density of ASH2L IDRs and DPY30 were indicated as black dash-circle in (c–d) and as a dashed line end in the accompanying cartoons. e–f Alternative conformation for the human 5-MLL1-NCP structure and ySET1-NCP as well as their respective top view. ASH2L/Bre2 in both structures interact with the NCP near SHL 7 with slight rotational dynamics at the RbBP5/Swd1-NCP contact points.
at the MLL1 targets (Fig. 6a). A similar close correlation of DPY30 and H3K4me3 was also found at the 2431 ASH2L binding sites, 67% of which colocalized with DPY30 at gene regulatory regions in the E14 ESCs (Fig. 6b and Supplementary Figs. 12b and 13). These results showed that MLL1/ASH2L alone was ineffective for depositing H3K4me3 on chromatin. Instead, DPY30 was required for promoting high levels of H3K4me3 on chromatin.

Next, we tested whether DPY30 plays a causal role in establishing de novo H3K4me3 on chromatin. To this end, we expressed catalytically inactive HA-dCas9 or HA-dCas9-DPY30 in E14 cells and targeted the fusion proteins to randomly selected genomic regions by gRNAs (Fig. 6b, left). The loci were selected from MLL1/ASH2L joint targets that had no prior DPY30 binding (Fig. 6b, right top). Upon HA-dCas9-DPY30 recruitment, there was a significant increase of H3K4me3 at these loci (Fig. 6b, bottom right). In contrast, no increase of H3K4me3 was observed for the no gRNA controls (Fig. 6b) or in cells expressing HA-dCas9 (Supplementary Fig. 12c). These results confirmed that DPY30 is required for de novo establishment of H3K4me3 in cells.

**Discussion**

Using the biochemical, structural, and cellular approaches, we have revealed the mechanism by which DPY30 regulates MLL//SET1 activity on chromatin. We show that DPY30 functions through ASH2L IDR.s and DPY30-induced changes stabilize ASH2L-NCP interactions and restrict the rotational dynamics of the MLL1 complex on the NCP. Consequently, it promotes productive H3K4 methylation, especially at higher methylation
states (i.e., H3K4me3 and H3K4me2). Our study has established a paradigm by which IDRs, the often-ignored segments in chromatin-interacting proteins, contribute to the heterogeneity of the epigenetic landscape in eukaryotic cells.

Previous studies have shown that DPY30 has negligible effects on H3 methylation in vitro16,39,45, yet its deletion leads to global downregulation of H3K4me3 in cells41. Our study shows that DPY30 confers NCP-specific regulation of MLL1 activity by regulating ASH2L-NCP interactions. Combining complementary biophysical, structural, and biochemical experiments with computational modeling, we show that upon DPY30 binding, ASH2L IDRs converge to adopt a compact structural unit at the MLL1-NCP interface, enabling new contacts with the NCP. In support, deletion or mutating ASH2L IDRs greatly impaired DPY30-dependent methyltransferase activity in vitro (Fig. 3 and Supplementary Fig. 1b). The cryo-EM structure of the 4-MLL1-NCP complex shows significant rotational dynamics on the NCP as compared to the 5-MLL1-NCP, 5-MLL3-NCP (Fig. 5e) or ySET1-NCP complexes (Fig. 5)f,37,38,46,62. The 4-MLL1 complex is able to swing across the nucleosome disc with ASH2L binding near SHL4 in a subset of the cryo-EM structures (Fig. 5). Furthermore, ASH2L also exhibits higher local binding dynamics in the absence of DPY30. We envision that increased rotational dynamics of the 4-MLL1 complex or local ASH2L dynamics reduces the probability of the MLL1 SET domain positioning near nucleosome dyad. In this scenario, the MLL1SET domain has to go through multiple spatial arrangements to optimally engage both H3 substrates in the NCP, which negatively affect MLL1 processivity37. By limiting rotational dynamics of the MLL1 complex on the NCP, DPY30 as well as ASH2L IDRs promote productive enzyme-substrate engagement, which has a specific impact on higher methylation states.

Notably, DPY30/ASH2L IDRs regulate all MLL/SET1 family enzymes, regardless of their respective intrinsic activity and processivity (Fig. 1b). We find that despite its selective impact on global H3K4me3 in cells41, DPY30 is able to stimulate H3K4me1 by the MLL3 complex in vitro. The global reduction of H3K4me3, but not H3K4me1 or H3K4me2, after DPY30 deletion/depletion in cells is probably due to compounding effects of relative abundance and activity of different MLL family enzymes as well as the offset of H3K4me1 inhibition by blocking its conversion to higher methylation states. We also would like to point out that DPY30 is able to enhance human SET1 activity on the H2BK120ub-containing NCP (Supplementary Fig. 1d). Thus, it can cooperate with H2BK120ub in H3K4me3 regulation in vivo, which awaits future studies.

It is well established that intrinsically disordered proteins (IDPs), or proteins containing extensive IDRs, have unique biophysical properties53,64. The undefined structures in the solution enable IDRs to adopt many possible conformations and meaningfully engage in versatile protein-protein interactions65–67. As a result, IDRs or IDPs are often found at hubs of protein interaction networks and enable functional diversification and environmental responsiveness during the complex developmental processes66,68. Recent studies also show that IDRs are able to facilitate phase transition and heterochromatin functions in cells69. Our study here provides a paradigm for how IDRs in histone-modifying enzymes may regulate chromatin functions. We show that ASH2L IDRs and their interacting protein DPY30 can exert locus- and context-specific regulation of H3K4me3 in cells. While the exact conformation(s) of apo-state ASH2L IDRs remain to be determined, our study suggests that ASH2L IDRs are probably in a highly dynamic conformational equilibrium and DPY30 binding leads to stabilization of ASH2L IDRs in one of the more structurally organized conformations. Our study also raises the question of whether ASH2L IDRs can be modulated by other proteins beyond DPY30. We envision that proteins that are able to induce perturbations in ASH2L IDRs and/or stabilize ASH2L IDRs could potentially mediate MLL1/SET1-NCP interactions, thereby regulating H3K4 methylation activity on chromatin. Aberrant expression of ASH2L has been reported in a wide spectrum of human tumors, and contributes to disease progression and prognosis24,70–72. Notably, ASH2L cooperates with activating mutations of Ras in cellular transformation33, recruits the oncogene MYC to target genes in conjunction with WDR54,73, and regulates p53 targeting gene expression76. Future studies on ASH2L IDR and IDR interacting proteins will provide insights into the regulation of H3K4me3 heterogeneity in cells, and potentially shed light on human pathogenesis.

Finally, histone-modifying enzyme complexes usually contain multiple IDRs in both catalytic and non-catalytic subunits. Our survey indicates that IDR content can go up to 70–90% for some histone-modifying enzymes (Supplementary Table 3). Furthermore, 60% of lysine HMTs (HKMTs) contain at least 80 residues or more, whereas only 20% of other annotated proteins have IDRs of similar length77. It suggests that IDRs in the histone-modifying enzymes may have especially important regulatory roles, which may constitute a layer of complexity in epigenetic regulations. Inclusion of the IDRs in enzymes or enzyme complexes may be necessary to discover their regulation to the fullest extent.

Methods

Mouse and human ES cell lines. E14tg2a (E14) (ATCC, #30-2002) cell line was used for all cellular experiments. To generate the E14 cell line stably expressing HA-ASH2L, the plasmid expressing ASH2L from the pPiggybac-HA vector as well as plasmids carrying PBase transposase and rTTA element were co-transfected into E14 cells by electroporation. Geneticin was added one day after transfection and selection was carried out for 10 days. Single colonies were picked and screened for stable expression of HA-ASH2L in the presence of Doxycycline.

General protein expression and purification. All MLL1 complex subunits and their mutants were expressed using the pET-28a expression vector with N-terminal 6-histidine and SUMO tag29. To make ASH2L mutants for methyl assignments, codon-optimized ASH2L202–534 DNA (Integrated DNA Technologies) was used as a template for mutagenesis. Each Ile was changed to Leu, and each Leu or Val was changed to Ile. NEBaseChanger web tool (New England Biolabs) was used to design primers for single residue substitution. Mutant plasmids were constructed using Q5 Site-Directed Mutagenesis Kit (NEB, Cat#E0554S). All proteins were expressed in BL21(DE3) E. coli strain in LB media. Cells were grown initially at 37°C till OD600 reached 0.6–0.8 and shifted to 20°C after IPTG was added at a final concentration of 0.2–0.4 mM. Cells were lysed by sonication and lysates were collected after centrifugation at 32,000 × g for 30 min at 4°C. The supernatant was filtered through a 0.45 μm syringe filter and purified through a Ni-NTA metal-affinity column (Qiagen and GoldBio). After extensive washing with 20 mM Tris (pH 8.0), 300–500 mM NaCl, 2 mM β-mercaptoethanol, and 10 mM imidazole (washing buffer), protein was eluted stepwise at 30, 60, 90, 120, 150, 210, and 300 mM imidazole. SUMO protease was added to the pooled fractions during dialysis at 4°C overnight. Ni-NTA purification was repeated to remove 6-histidine tag and other bacterial impurities. Proteins were further purified on a HiLoad 16/60 Superdex 75 pg or 200 pg columns (GE Healthcare Life Sciences). All MLL complex subunits and their mutants are nicely expressed and well-behaved in solution with no noticeable differences in protein stability.

GST-fusion MLL and SET1 proteins. GST-tagged MLL (MLL11,47, MLL22,490, MLL38,65, MLL41,73) and SET1 (SET1A1,47 and SET1B1,84) proteins were expressed using a pGEX-parallel 1 expression vector with N-terminal GST tag and TEV cleavage sequence31. Plasmids were transformed and expressed in BL21(DE3) E. coli in LB media. Cells were grown until OD600 reached 0.6–0.8 when the temperature was reduced to 20°C and, after temperature equilibration, protein expression was induced using 0.4 mM IPTG and grown for 16 h. Cells were harvested and lysed using sonication and the supernatant was collected by centrifugation at 32,000 × g, filtered through a 0.45 μm syringe, and loaded onto a pre-equilibrated Glutathione Sepharose 4B column (GE Healthcare Life Sciences). After several washes with 20 mM Tris HCl (pH 7.5), 300 mM NaCl, 2 mM DTT, 10% v/v glycerol (GST wash buffer), the protein was eluted off the column using GST wash buffer with 10 mM reduced glutathione. Proteins were further purified over a HiLoad 16/60 Superdex 200 pg column (GE Healthcare Life Sciences). The purified SET domains remain soluble and stable for the in vitro assays.
In vitro HMT assay. Mixture of stoichiometric amounts of MLL1 core proteins was used for the in vitro HMT assay. Recombinant mono-nuclease was prepared by chemical dialysis of equal molar histone octamer and 146 bp 601 DNA. The reaction was carried out in 20 μL of the HMT buffer of 20 mM Tris (pH 8.0), 50 mM NaCl, 5 mM Mg²⁺, 1 mM DTT and 10% v/v glycerol. The reaction was initiated by adding 1 μL of 100 μM S-adenosyl-L-methionine and incubated at room temperature for 1 h for the NCP structures or 4 h for the recombinant H3 structure. The 2x SDS-PAGE sample buffer was added to quench the reaction.

Western blotting. The histones were separated on a 10–15% polyacrylamide gel and transferred onto polyvinylidene difluoride membrane (Millipore). The membrane was blocked in blocking solution, consisting of 5% milk in 0.1% Tween 20/TBS (TBST), followed by incubation at 4°C overnight with the primary antibody in blocking membranes. Membranes were washed three times in TBST and incubated with the HRP-conjugated anti-mouse/rabbit secondary antibodies at room temperature for 1 h. The membrane was developed using Pierce™ ECL Western Blotting Substrate (Thermo Fisher Scientific, #32106), and images were captured by ChemiDoc™ Touch Imaging System (Bio-rad).

Antibodies. The primary and secondary antibodies included: Rabbit anti-H3K4me1 (Abcam, cat # ab8895, 1:20,000), Rabbit anti-H3K4me2 (Millipore, cat #07-030, 1:20,000), Rabbit anti-H3K4me3 (Millipore, cat #07-473, 1:10,000), Rabbit anti-Histone H3 (Abcam, cat # ab1791, 1:20,000) and anti-Rabbit IgG Horseradish Peroxidase-linked whole antibody (GE Healthcare, cat # NA934, 1:1000). Anti-HA (clone C29F4) rabbit monoclonal antibody (Cell signaling technology, cat # #3724, 1:1000).

Preparation of ILV U-[2H]3-labeled ASH2L NMR samples. The U-[2H]3Ile-1-[13C]Lys, U-[2H]Val-1-[13C]Leu, U-[2H]Asp-2-[13C]Asp, U-[2H]ASP-3-[13C]ASP, U-[2H]HIS-4-[13C]HIS, and U-[2H]RBP5 (330–363) labeled proteins were produced using a previously developed protocol with modifications. Freshly prepared single colony was inoculated into H2O minimal medium containing 6.5 g/L NaHPO4, 3 g/L KH₂PO₄, 0.5 g/L NaCl, 120 mg/L MgSO₄, 11 mg/L CaCl₂, 10 mg/L thiamine, 30 mg/L kanamycin, 2 g/L D-glucose and 1 g/L NH₄Cl. Cells were cultured at 37°C until OD₅₆₀ reaches 0.25 and harvested to remove H₂O media. Then cells were resuspended in 1L of ASH2L minimal media containing the same salts in H₂O media in which plain glucose was replaced by D-[2H] glucose (CIL, DLM-2062). Cells were cultured at 37°C until OD₅₆₀ reaches 0.7–0.8. The temperature was lowered to 20°C and 70 mL of 1% [3-13C]CH₃COONa, 3.3–H] α-ketobutyrate (Cambridge Isotope Laboratory, CDLM-7318) and 120 mg/L [3-13C]CH₃, 3.4,4,4-H] α-ketovalerate (CIL, CDLM-7317) was added to the culture. After 1 h, IPTG dissolved in D₂O was added to the final concentration of 0.4 mM. Cells were cultured for another 24 h before harvesting. The labeled ASH2L proteins were purified through Ni-NTA column as described above. To prevent potential aggregation of ASH2L at high concentration, 1.5 mM excess of unlabeled RbBP5 (330–363) was added to the concentrated samples. To examine the effect of DYPX, additional 1.2 mM excess of unlabeled dimeric DYPX was added. At this NMR samples were concentrated and buffer exchanged into 25 mM sodium phosphate, pH 6.5, 10 mM NaCl, 0.25 mM d₅-dithiothreitol (CIL, DLM-2622), and 1 mM NaN₃ in 99.99% D₂O (Aldrich Cat#151882).

NMR spectroscopy. NMR experiments were carried out on 800 MHz Bruker Ascend spectrometer equipped with pulsed-field gradient 5 mm inverse triple resonance TCI probe and SampleCASE with 24 sample slots. IconNMR software was used for the automated collection of mutant samples for assignment. All HMQC experiments were acquired at 25°C. Complex points of 2048 and 256 (H, 13C) were used for most of the experiments except for Ile mutants for assignment, for which 128 complex points in 13C dimension were used. The 4H and 13C carrier frequencies were placed at 47 and 17 ppm, respectively. Spectral width was set to 12 and 20 ppm for 1H and 13C dimensions, respectively. A recycle delay of 0.5 s was used with 32–256 scans depending on protein concentration. Residual water was suppressed by the WATERGATE method. 13C WALTZ-16 decoupling was employed during acquisition in the direct dimension. All spectra were processed using the NMRPipe program. Gaussian broad window and sine bell window functions were applied in 1H and 13C dimensions. NMRFAM-Sparky was used to visualize NMR spectra.

Small-angle X-ray scattering. All SAXS data were collected at the 18-ID BioCAT Beamline (Biophysics Collaborative Access Team, Advanced Photon Source, Argonne National Laboratory) using the inline SEC-SAXS configuration, in which a flow cell was connected to a AKTApure FPLC system (GE Healthcare). To prevent potential aggregation, a stoichiometric amount of RbBP5 peptide (330–363) was added to the ASH2L samples as described above. About 200–500 μL of 1 mg/mL ASH2L was injected to a 5 mL column (HiLoad 16/60, GE Healthcare) pre-equilibrated with 20 mM Tris (pH 7.5), 150 mM NaCl and 1 mM DTT. The flow rate was set to 0.7 mL/min during the data collection. The scattering data were collected every 2 s with 1 s exposure during the SEC elution between 5–24 mL. After data reduction, the strongest scattering data around the protein elution peak were selected for sample scattering. Several data points with minimal scattering near the elution peak were chosen for buffer-only scattering. PRIMUS was used for data processing, including averaging scattering data, background subtraction, and calculation of the radius of gyration, Rg, and the Porod Volume. The molecular weight was estimated by dividing Porod Volume by 1.6. The pair distribution function was calculated by GNOM in the GUI version of PRIMUS. For EOM analysis, a pool of 10,000 structures of ASH2L with N-terminal PHD-WH and C-terminal SPRY domains connected by the Linker and Loop IDRs was generated by RANCH. The sequence of RbBP5 (330–363) was not included in the EOM model due to its small size and well-characterized interaction with the rigid SPRY domain, as well as limitation of EOM for multiple polypeptide chains. GAJOE was used to select an ensemble that best fit the experimental data using a generic algorithm.

Molecular modeling of ASH2L IDRs. Human ASH2L protein consists of two domains, PHD-WH domain, and SPRY domain, that have homologous PDB structures, 3S5Z (A-chains), 3TOJ, respectively. The crystal structure of yeast Bre2 determined in the COMPASS complex (PDB: 6CHG) contains the Linker and PHD-WH domain. The cross-linking distribution function was calculated by GNOM and found to be consistent with the rigid SPRY domain57, as well as limitation of EOM for multiple polypeptide chains56. GAJOE was used to select an ensemble that best fit the experimental data using a generic algorithm.

Cryo-EM sample preparation and data collection. The GraFx method was applied to the MLL1RWSA complex to prepare for the cryo-EM grid. In brief, 30–μm-thick ice was incubated with 100 μM MLL1RWSA and 0.5 mM S-adenosyl-L-homocysteine for 30 min at 4°C in the GraFx buffer (50 mM HEPES, pH 7.5, 50 mM NaCl, 1 mM MgCl₂, and 1 mM TCEP). The sample was centrifuged at 10,000 × g at 4°C for 3 h after applying onto a centrifuge tube, which contained a gradient solution of 0–60% glycerol and 0.2–0.6% glutaraldehyde. After centrifugation, the gradient sample was overlayed with the gradient solution by removing glycerol from the GraFx buffer, we performed further buffer exchange using a centrifugal concentrator (Sartorius Vivaspin 500). The sample at 1–mg/ml was applied onto a glow discharge Quantifoil R1/2.13 grid (Electron Microscopy Sciences) at 4°C with 100% humidity. The loaded grid was plunged-frozen in liquid ethane with 10% v/v ethanolamine and 0.2% glutaraldehyde. After 4 s blotting and 30 s waiting using a Mark IV Vitrobot (Thermo Fisher Scientific). The cryo-EM data were collected using Titan Krios (Thermo Fisher Scientific) operating at 300 keV with the K2 Summit direct electron detector. The movie data was recorded in a counting mode at a ×29,000 magnification and the pixel size of 1.01 Å/pixel, with a defocus range between −1.5 to −2.5 μm. A dose rate of 1.28 electrons/Å²/frame with a total 50 frames per 8 s was applied for data collection, resulting in a total dose of 64 electrons per Å². A total of 6,242 movies were collected.

Cryo-EM data processing and model refinement. Micrograph movies were aligned with whole-frame and local drift correction using MotionCor240, and CTF was estimated with CTFFIND4.145. Micrographs with higher than 4.5 Å of the estimated resolution were further selected, which resulted in 613 micrographs. A total of 287,771 particle were picked using WorkWedge. The 3D reconstruction was performed using RELION58, imported into cryoSPARC for 2D classification. After excluding bad particles, a total of 1,194,542 particles were subjected to the first round of ab initio 3D classification into five classes (Supplementary Fig. 4). Two of five classes were subjected to the second round of ab initio 3D classification into five subclasses, and the subsequent heterogeneous refinement was performed. Four of the five subclasses displayed a well-defined map of the MLL and nucleosome complex after the heterogeneous refinement. They were exported for 3D classification. The focused 3D classification was performed at the MLLRWSA region without alignment (35 cycles, T = 4, binary mask: 10 pixels/soft mask: 10 pixels). The CrysAlisPro was excluded because it displayed a structurally heterogeneous and unresolvable EM density even after the focused 3D classification. The best behaving class selected from Class01 (13,086 particles), Class02 (27,730 particles), and...
Class05 (23,236 particles) was subjected to the 3D auto refinement and further post-processed to a resolution of 6.9, 4.6, and 6.0 Å, respectively. Each final cryo-EM map was locally filtered to avoid over-estimation. The resolution of all structures was estimated by RELION with Fourier shell correlation (FSC) at the criteria of 0.143.

For the model building, the rigid-body fitting was performed for each class using Chimera97. The cryo-EM structure of MLL1RWSAD-NCP (PDB ID: 6PWV) was used for the rigid-body fitting for each individual class. For the model refinement, each class was subjected to the real-space refinement using PHENIX30, and model validations were performed by MolProbity100. Statistics for data collection, refinement, and validation were summarized in Supplementary Table 2.

ESC culture and transfection. E14tg2a (E14, AC3T, CRL-1821TM) were grown in the KnockOut™-DMEM medium containing 15% FBS, 2 mM glutamine, 1x non-essential amino acids, 0.1 mM mercaptoethanol and 10^3 U/ml LIF (Millipore, #ESG107), unless otherwise indicated. E14 cells were routinely tested negative for mycoplasma contamination using the LookOut™ Mycoplasma PCR Detection Kit (SIGMA ALDRICH, #MP0035) according to the manufacturer’s instructions. For expressing dCas9 fusion proteins, E14 ESCs were transfected with pcDNA3-dCas9-HA and pcDNA3-dCas9-DYPO-HA plasmids using Fugene 6 (Promega, Cat# E2691) for 2 days and then selected with G418 (400 µg/ml, Gibco, Cat# 10131-035) for 5 days. After selection, the cells were split and transfected with a pool of three ppsRNA-gRNAs for selected genomic loci. The ppsRNA-gRNAs were co-transfected with a repressor (factor 110) that confers puromycin resistance. After 2 days of puromycin selection (1.5 µg/ml, Gibco, Cat# A11138-03), the cells were subject to ChIP using anti-HA antibody (Cell Signaling Technology, cat# 3724) and anti-H3K4me3 antibody (Millipore, Cat# 07-473), respectively. ChIP-qPCRs were performed to detect the enrichment of H3K4me3 and HA in each location.

CUT&RUN. CUT&RUN was performed according to the protocol described previously101. HA-ASH2L E14 and the parental E14 (E14tg2a, ACTT, #CRL-1821TM) were grown in KnockOut™-DMEM medium containing 15% FBS, 2 mM glutamine, 1x non-essential amino acids, 0.1 mM 2-mercaptoethanol and 10^3 Um l^-1 LIF. E14tg2a (E14, AC3T, CRL-1821TM) were subjected to the 3D auto refinement using PHENIX99, and model validations were performed by MolProbity100. Statistics for data collection, refinement, and validation were summarized in Supplementary Table 2.

ChIP-seq data mapping and normalization. ChIP-seq dataset for DPY30 and MLL1 were downloaded from GEO GSE626136 and GEO GSE107406, respectively. Paired-end sequencing reads were trimmed with trim_galore to remove adaptor sequences. We kept reads that were 20 bp or longer after trimming and paired between the mates. All ChIP-seq data were mapped to the mouse mm10 genome by using Bowtie2 (v2.2.4)102 with parameters “-p phred33 –very-sensitive -p 10”. Duplicated reads were removed using SAMTools (v1.5)103. The bigwig files for IP/input ratio were generated from BAM files by using deepTools3102 with command “bamCompare -b CHIP-bam -b Input-bam --ignoreDuplicates --minMappingQuality 30 --normalizeUsing RPMK --binSize 1 --operation ratio” using SamToolMethod None. For each individual class, the bigwig files for visualization were merged using SAMTools and converted to BED format using BEDTools103. Peaks were called from bed files using MACS (v 1.4.2)106 with parameters “-w 0.00001 -g mm”. The input signal was used as the control for peak calling. Heatmap of ChIP-seq signals was visualized using deepTools3.

Cut&Run peak calling and visualization. HA or H3K4me3 CUT&RUN from two independent biological replicates were initially analyzed in parallel. Paired-end sequencing reads were trimmed with trim_galore to remove adaptor sequences. We kept reads that were 20 bp or longer after trimming and paired between the mates. All ChIP-seq data were mapped to the mouse mm10 genome by using Bowtie2 (v2.2.4)102 with parameters “-p phred33 –very-sensitive -p 10”. Duplicated reads were removed using SAMTools (v1.5)103. The bigwig files for IP/input ratio were generated from BAM files by using deepTools3102 with command “bamCompare -b CHIP-bam -b Input-bam --ignoreDuplicates --minMappingQuality 30 --normalizeUsing RPMK --binSize 1 --operation ratio” using SamToolMethod None. For each individual class, the bigwig files for visualization were merged using SAMTools and converted to BED format using BEDTools103. Peaks were called from bed files using MACS (v 1.4.2)106 with parameters “-w -S -p 0.00001 -g mm”. The input signal was used as the control for peak calling. Heatmap of ChIP-seq signals was visualized using deepTools3.

Statistical analysis and reproducibility. Statistical analysis was performed by two-tailed Student’s t-test using GraphPad Prism 7.0 software. Data were presented as the standard error of the mean (SEM). p value of <0.05 was considered statistically significant; “p < 0.05, **p < 0.01, ***p < 0.001. For all in vitro HMT experiments shown Figs. 1–3 as well as Supplementary Figs. 1, 2, and 11, a minimum of three independent experiments were performed and consistent results were obtained.

Data availability. The data that support this study are available from the corresponding author upon reasonable request. ChIP-seq datasets for HA-ASH2L and H3K4me3 generated in this study are accessible at GEO with accession code GSE146933. DPY30 and MLL1 datasets were downloaded from GEO GSE626136 and GEO GSE107406, respectively. Cryo-EM structures for the 4-MLL1-NCP reported in this study are available with accession numbers: Class01 – PDB 6W5I [https://doi.org/10.2210/pdb6w5i/pdb] and EMDB EMD-21545; Class02 – PDB 6W5J [https://doi.org/10.2210/pdb6w5j/pdb] and EMDB EMD-21544. Source data are provided with this paper.

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**Author contributions**

Y.-T.L. conceived, designed, performed most biochemical and NMR experiments and wrote the manuscript. A.A. performed in vitro histone methyltransferase assays and NMR experiments and wrote the manuscript. S.H.P. performed the cryo-EM studies under the supervision of U.-S.C. F.M. performed bioinformatics analyses; W.Z. performed molecular modeling experiment under the supervision of Y.Z. L.X. performed dCas9 transfection and ChIP experiments. L.S. performed CUT&RUN experiments for ASH2L and H3K4me3. Y.D. supervised the overall study and wrote the manuscript. All authors read and approved the final manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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